# The R Environment for Statistical Computing and Graphics 

Reference Index<br>The R Development Core Team

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## Chapter 1

## The base package

## Machine Numerical Characteristics of the Machine

## Description

.Machine is a variable holding information on the numerical characteristics of the machine R is running on, such as the largest double or integer and the machine's precision.

## Usage

.Machine

## Details

The algorithm is based on Cody's (1988) subroutine MACHAR.

## Value

A list with components (for simplicity, the prefix "double" is omitted in the explanations)
double.eps the smallest positive floating-point number x such that $1+\mathrm{x}!=1$. It equals base^ulp.digits if either base is 2 or rounding is 0 ; otherwise, it is (base^ulp.digits) / 2.
double.neg.eps
a small positive floating-point number x such that $1-\mathrm{x}$ != 1 . It equals base^neg.ulp.digits if base is 2 or round is 0 ; otherwise, it is (base^neg.ulp.digits) / 2. As neg.ulp.digits is bounded below by -(digits + 3), neg.eps may not be the smallest number that can alter 1 by subtraction.
double.xmin the smallest non-vanishing normalized floating-point power of the radix, i.e., base^min. exp.
double.xmax the largest finite floating-point number. Typically, it is equal to (1neg.eps) * base^max.exp, but on some machines it is only the second, or perhaps third, largest number, being too small by 1 or 2 units in the last digit of the significand.
double.base the radix for the floating-point representation

```
double.digits the number of base digits in the floating-point significand
double.rounding
the rounding action.
0 if floating-point addition chops;
1 if floating-point addition rounds, but not in the IEEE style;
2 if floating-point addition rounds in the IEEE style;
3 if floating-point addition chops, and there is partial underflow;
4 if floating-point addition rounds, but not in the IEEE style, and there
is partial underflow;
5 if floating-point addition rounds in the IEEE style, and there is partial
underflow
double.guard the number of guard digits for multiplication with truncating arithmetic.
It is 1 if floating-point arithmetic truncates and more than digits base
base digits participate in the post-normalization shift of the floating-point
    significand in multiplication, and 0 otherwise.
double.ulp.digits
    the largest negative integer i such that \(1+\) base^i ! \(=1\), except that it
    is bounded below by -(digits +3 ).
double.neg.ulp.digits
the largest negative integer i such that \(1-\) base^i \(!=1\), except that it
    is bounded below by -(digits +3 ).
double.exponent
                                    the number of bits (decimal places if base is 10) reserved for the repre-
                                    sentation of the exponent (including the bias or sign) of a floating-point
                                    number
double.min.exp
    the largest in magnitude negative integer i such that base - \(i\) is positive
    and normalized.
double.max.exp
    the smallest positive power of base that overflows.
integer.max the largest integer which can be represented.
sizeof.long the number of bytes in a C long type.
sizeof.longlong
the number of bytes in a C long long type. Will be zero if there is no such type.
```


## References

Cody, W. J. (1988) MACHAR: A subroutine to dynamically determine machine parameters. Transactions on Mathematical Software, 14, 4, 303-311.

## See Also

.Platform for details of the platform.

## Examples

```
str(.Machine)
(Meps <- .Machine$double.eps)
## All the following relations must hold :
stopifnot(
```

```
1 + Meps != 1,
1 + .5* Meps == 1,
log2(.Machine$double.xmax) == .Machine$double.max.exp,
log2(.Machine$double.xmin) == .Machine$double.min.exp
)
```

.Platform Platform Specific Variables

## Description

.Platform is a list with some details of the platform under which R was built. This provides means to write OS portable R code.

## Usage

.Platform

## Value

A list with at least the following components:

| OS.type | character, giving the Operating System (family) of the computer. One <br> of the following values is returned: "unix", "mac", or "windows" (in <br> historical order). |
| :--- | :--- |
| file.sep | character, giving the file separator, used on your platform, e.g., "/" on <br> Unix alikes. |
| dynlib.ext | character, giving the file name extension of dynamically loadable <br> libraries, e.g., ".dll" on Windows. <br> character, giving the type of GUI in use, or "unknown" if no GUI can be <br> assumed. <br> character, "big" or "little", giving the endianness of the processor in <br> chdian <br> use. |

## See Also

R.version and Sys.info give more details about the OS. In particular, R.version\$platform is the canonical name of the platform under which $R$ was compiled.
.Machine for details of the arithmetic used, and system for invoking platform-specific system commands.

## Examples

```
## Note: this can be done in a system-independent way by file.info()$isdir
if(.Platform$OS.type == "unix") {
    system.test <- function(...) { system(paste("test", ...)) == 0 }
    dir.exists <- function(dir) sapply(dir, function(d)system.test("-d", d))
    dir.exists(c(R.home(), "/tmp", "~", "/NO"))# > T T T F
}
```

.Script Scripting Language Interface

## Description

Run a script through its interpreter with given arguments.

## Usage

```
.Script(interpreter, script, args, ...)
```


## Arguments

| interpreter | a character string naming the interpreter for the script. |
| :--- | :--- |
| script | a character string with the base file name of the script, which must be <br> located in the 'interpreter' subdirectory of 'R_HOME/share'. |
| args | a character string giving the arguments to pass to the script. |
| $\ldots$ | further arguments to be passed to system when invoking the interpreter <br> on the script. |

## Note

This function is for $R$ internal use only.

## Examples

```
.Script("perl", "maketitle.pl", file.path(.Library, "base", "DESCRIPTION"))
```

```
abbreviate Abbreviate Strings
```


## Description

Abbreviate strings to at least minlength characters, such that they remain unique (if they were).

## Usage

```
abbreviate(names.arg, minlength = 4, use.classes = TRUE,
    dot = FALSE)
```


## Arguments

names.arg a vector of names to be abbreviated.
minlength the minimum length of the abbreviations.
use.classes logical (currently ignored by R).
dot logical; should a dot (".") be appended?

## Details

The algorithm used is similar to that of S. First spaces at the beginning of the word are stripped. Then any other spaces are stripped. Next lower case vowels are removed followed by lower case consonants. Finally if the abbreviation is still longer than minlength upper case letters are stripped.

Letters are always stripped from the end of the word first. If an element of names.arg contains more than one word (words are separated by space) then at least one letter from each word will be retained. If a single string is passed it is abbreviated in the same manner as a vector of strings.
If use.classes is FALSE then the only distinction is to be between letters and space. This has NOT been implemented.

## Value

A character vector containing abbreviations for the strings in its first argument. Duplicates in the original names.arg will be given identical abbreviations. If any non-duplicated elements have the same minlength abbreviations then minlength is incremented by one and new abbreviations are found for those elements only. This process is repeated until all unique elements of names.arg have unique abbreviations.

The character version of names.arg is attached to the returned value as a names argument.

## See Also

substr.

## Examples

```
x <- c("abcd", "efgh", "abce")
abbreviate(x, 2)
data(state)
(st.abb <- abbreviate(state.name, 2))
table(nchar(st.abb))# out of 50, 3 need 4 letters
```


## abline Add a Straight Line to a Plot

## Description

This function adds one or more straight lines through the current plot.

## Usage

```
abline(a, b, untf = FALSE, ...)
abline(h=, untf = FALSE, ...)
abline(v=, untf = FALSE, ...)
abline(coef=, untf = FALSE, ...)
abline(reg=, untf = FALSE, ...)
```


## Arguments

| $\mathrm{a}, \mathrm{b}$ | the intercept and slope. |
| :--- | :--- |
| untf | logical asking to untransform. See Details. |
| h | the y-value for a horizontal line. |
| v | the x-value for a vertical line. |
| coef | a vector of length two giving the intercept and slope. |
| reg | an object with a coef component. See Details. |
| $\ldots$ | graphical parameters. |

## Details

The first form specifies the line in intercept/slope form (alternatively a can be specified on its own and is taken to contain the slope and intercept in vector form).
The $h=$ and $v=$ forms draw horizontal and vertical lines at the specified coordinates.
The coef form specifies the line by a vector containing the slope and intercept.
reg is a regression object which contains reg\$coef. If it is of length 1 then the value is taken to be the slope of a line through the origin, otherwise, the first 2 values are taken to be the intercept and slope.
If untf is true, and one or both axes are log-transformed, then a curve is drawn corresponding to a line in original coordinates, otherwise a line is drawn in the transformed coordinate system. The h and v parameters always refer to original coordinates.
The graphical parameters col and lty can be specified as arguments to abline; see par for details.

## See Also

lines and segments for connected and arbitrary lines given by their endpoints. par.

## Examples

```
data(cars)
z <- lm(dist ~ speed, data = cars)
plot(cars)
abline(z)
```


## Description

These functions compute miscellaneous mathematical functions. The naming follows the standard for computer languages such as C or Fortran.

## Usage

abs ( x )
sqrt(x)

## Arguments

x a numeric vector

## See Also

Arithmetic for simple, log for logarithmic, sin for trigonometric, and Special for special mathematical functions.

## Examples

```
xx <- -9:9
plot(xx, sqrt(abs(xx)), col = "red")
lines(spline(xx, sqrt(abs(xx)), n=101), col = "pink")
```


## Description

Compute all the single terms in the scope argument that can be added to or dropped from the model, fit those models and compute a table of the changes in fit.

## Usage

```
add1(object, scope, ...)
add1.default(object, scope, scale = 0, test = c("none", "Chisq"),
    \(\mathrm{k}=2\), trace = FALSE, ...)
add1.lm(object, scope, scale \(=0\), test = c("none", "Chisq", "F"),
    \(\mathrm{x}=\) NULL, \(\mathrm{k}=2, \ldots\) )
add1.glm(object, scope, scale \(=0\), test \(=c(" n o n e ", ~ " C h i s q ", ~ " F ")\),
        \(\mathrm{x}=\) NULL, \(\mathrm{k}=2, \ldots\) )
drop1(object, scope, ...)
drop1.default(object, scope, scale \(=0\), test \(=c(" n o n e ", ~ " C h i s q ")\),
                \(\mathrm{k}=2\), trace \(=\) FALSE,.. )
drop1.lm(object, scope, scale \(=0\), all.cols = TRUE,
    test=c("none", "Chisq", "F"),k = 2, ...)
drop1.glm(object, scope, scale = 0, test = c("none", "Chisq", "F"),
        \(\mathrm{k}=2, \ldots\) )
```


## Arguments

| object | a fitted model object. |
| :--- | :--- |
| scope | a formula giving the terms to be considered for adding or dropping. |
| scale | an estimate of the residual mean square to be used in computing $C_{p}$. <br> Ignored if 0 or NULL. |
| test | should the results include a test statistic relative to the original model? <br> The F test is only appropriate for lm and aov models or perhaps for glm <br> fits with estimated dispersion. The $\chi^{2}$ test can be an exact test (lm models <br> with known scale) or a likelihood-ratio test or a test of the reduction in <br> scaled deviance depending on the method. |


| k | the penalty constant in AIC $/ C_{p}$. |
| :--- | :--- |
| trace | if TRUE, print out progress reports. |
| x | a model matrix containing columns for the fitted model and all terms in <br> the upper scope. Useful if add1 is to be called repeatedly. |
| all.cols | (Provided for compatibility with S.) Logical to specify whether all columns <br> of the design matrix should be used. If FALSE then non-estimable columns <br> are dropped, but the result is not usually statistically meaningful. |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

For drop1 methods, a missing scope is taken to be all terms in the model. The hierarchy is respected when considering terms to be added or dropped: all main effects contained in a second-order interaction must remain, and so on.

The methods for 1 m and glm are more efficient in that they do not recompute the model matrix and call the fit methods directly.

The default output table gives AIC, defined as minus twice log likelihood plus $2 p$ where $p$ is the rank of the model (the number of effective parameters). This is only defined up to an additive constant (like log-likelihoods). For linear Gaussian models with fixed scale, the constant is chosen to give Mallows' $C_{p}, R S S / s c a l e+2 p-n$. Where $C_{p}$ is used, the column is labelled as Cp rather than AIC.

## Value

An object of class "anova" summarizing the differences in fit between the models.

## Warning

The model fitting must apply the models to the same dataset. Most methods will attempt to use a subset of the data with no missing values for any of the variables if na.action=na.omit, but this may give biased results. Only use these functions with data containing missing values with great care.

## Note

These are not fully equivalent to the functions in S. There is no keep argument, and the methods used are not quite so computationally efficient.

Their authors' definitions of Mallows' $C_{p}$ and Akaike's AIC are used, not those of the authors of the models chapter of S.

## Author(s)

B. D. Ripley

## See Also

```
step, aov, lm, extractAIC.
```


## Examples

```
example(step)#-> swiss
add1(lm1, ~ I(Education^2) + .^2)
drop1(lm1, test="F")
example(glm)
drop1(glm.D93, test="Chisq")
drop1(glm.D93, test="F")
```

```
aggregate Compute Summary Statistics of Data Subsets
```


## Description

Splits the data into subsets, computes summary statistics for each, and returns the result in a convenient form.

```
Usage
aggregate(x, ...)
aggregate.default(x, ...)
aggregate.data.frame(x, by, FUN, ...)
aggregate.ts(x, nfrequency = 1, FUN = sum, ndeltat = 1,
                        ts.eps = getOption("ts.eps"), ...)
```


## Arguments

```
x an R object.
by a list of grouping elements, each as long as the variables in x. Names for
    the grouping variables are provided if they are not given.
FUN a scalar function to compute the summary statistics which can be applied
    to all data subsets.
nfrequency new number of observations per unit of time; must be a divisor of the
    frequency of x.
ndeltat new fraction of the sampling period between successive observations; must
    be a divisor of the sampling interval of x.
ts.eps tolerance used to decide if nfrequency is a sub-multiple of the original
    frequency.
... further arguments passed to or used by methods.
```


## Details

aggregate is a generic functions with methods for data frames and time series.
The default method aggregate.default uses the time series method if x is a time series, and otherwise coerces x to a data frame and calls the data frame method.
aggregate.data.frame is the data frame method. If $x$ is not a data frame, it is coerced to one. Then, each of the variables (columns) in $x$ is split into subsets of cases (rows) of identical combinations of the components of by, and FUN is applied to each such subset with further arguments in ... passed to it. (I.e., tapply (VAR, by, FUN, ..., simplify =

FALSE) is done for each variable VAR in x , conveniently wrapped into one call to lapply().) Empty subsets are removed, and the result is reformatted into a data frame containing the variables in by and x . The ones arising from by contain the unique combinations of grouping values used for determining the subsets, and the ones arising from x the corresponding summary statistics for the subset of the respective variables in x .
aggregate.ts is the time series method. If $x$ is not a time series, it is coerced to one. Then, the variables in $x$ are split into appropriate blocks of length frequency ( $x$ ) / nfrequency, and FUN is applied to each such block, with further (named) arguments in ... passed to it. The result returned is a time series with frequency nfrequency holding the aggregated values.

## Author(s)

Kurt Hornik

## See Also

```
apply, lapply, tapply.
```


## Examples

```
data(state)
## Compute the averages for the variables in 'state.x77', grouped
## according to the region (Northeast, South, North Central, West) that
## each state belongs to.
aggregate(state.x77, list(Region = state.region), mean)
## Compute the averages according to region and the occurrence of more
## than 130 days of frost.
aggregate(state.x77,
    list(Region = state.region,
        Cold = state.x77[,"Frost"] > 130),
        mean)
## (Note that no state in 'South' is THAT cold.)
data(presidents)
## Compute the average annual approval ratings for American presidents.
aggregate(presidents, nf = 1, FUN = mean)
## Give the summer less weight.
aggregate(presidents, nf = 1, FUN = weighted.mean, w = c(1, 1, 0.5, 1))
```

agrep Approximate String Matching (Fuzzy Matching)

## Description

Searches for approximate matches to pattern (the first argument) within the string $x$ (the second argument) using the Levenshtein edit distance.

## Usage

```
agrep(pattern, x, ignore.case = FALSE, value = FALSE, max.distance = 0.1)
```


## Arguments

pattern a non-empty character string to be matched (not a regular expression!)
x
ignore.case if FALSE, the pattern matching is case sensitive and if TRUE, case is ignored during matching.
value if FALSE, a vector containing the (integer) indices of the matches determined is returned and if TRUE, a vector containing the matching elements themselves is returned.
max.distance Maximum distance allowed for a match. Expressed either as integer, or as a fraction of the pattern length (will be replaced by the smallest integer not less than the corresponding fraction), or a list with possible components
all: maximal (overall) distance
insertions: maximum number/fraction of insertions
deletions: maximum number/fraction of deletions substitutions: maximum number/fraction of substitutions

If all is missing, it is set to $10 \%$, the other components default to all. The component names can be abbreviated.

## Details

The Levensthein edit distance is used as measure of approximateness: it is the the total number of insertions, deletions and substitutions required to transform one string into another.

The function is a simple interface to the apse library developed by Jarkko Hietaniemi (also used in the Perl String::Approx module).

## Value

Either a vector giving the indices of the elements that yielded a match, of, if value is TRUE, the matched elements.

## Author(s)

David Meyer 〈David.Meyer@ci.tuwien.ac.at〉 (based on C code by Jarkko Hietaniemi); modifications by Kurt Hornik

## See Also

 grep
## Examples

```
agrep("lasy", "1 lazy 2")
agrep("lasy", "1 lazy 2", max = list(sub = 0))
agrep("laysy", c("1 lazy", "1", "1 LAZY"), max = 2)
agrep("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, value = TRUE)
agrep("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, ignore.case = TRUE)
```


## AIC Akaike Information Criterion

## Description

Generic function calculating the Akaike information criterion for one or several fitted model objects for which a log-likelihood value can be obtained, according to the formula -2 log-likelihood $+k n_{p a r}$, where $n_{\text {par }}$ represents the number of parameters in the fitted model, and $k=2$ for the usual AIC, or $k=\log (n)$ ( $n$ the number of observations) for the so-called BIC or SBC (Schwarz's Bayesian criterion).

## Usage

```
AIC(object, ..., k = 2)
```


## Arguments

| object | a fitted model object, for which there exists a logLik method to ex- <br> tract the corresponding log-likelihood, or an object inheriting from class |
| :--- | :--- |
| logLik. |  |$\quad$| optionally more fitted model objects. |
| :--- |
| k |$\quad$| numeric, the "penalty" per parameter to be used; the default $\mathrm{k}=2$ is the |
| :--- |
| classical AIC. |

## Details

The default method for AIC, AIC.default () entirely relies on the existence of a logLik method computing the log-likelihood for the given class.

When comparing fitted objects, the smaller the AIC, the better the fit.

## Value

If just one object is provided, returns a numeric value with the corresponding AIC (or BIC, or ..., depending on k ); if more than one object are provided, returns a data.frame with rows corresponding to the objects and columns representing the number of parameters in the model (df) and the AIC.

## Author(s)

Jose Pinheiro and Douglas Bates

## References

Sakamoto, Y., Ishiguro, M., and Kitagawa G. (1986). Akaike Information Criterion Statistics. D. Reidel Publishing Company.

## See Also

logLik, AIC.logLik.

## Examples

```
data(swiss)
lm1 <- lm(Fertility ~ . , data = swiss)
AIC(lm1)
stopifnot(all.equal(AIC(lm1),
    AIC(logLik(lm1))))
## a version of BIC or Schwarz' BC :
AIC(lm1, k = log(nrow(swiss)))
```

AIC.logLik

AIC of a logLik Object

## Description

see Description in AIC.

## Usage

AIC(object, ..., k = 2)

## Arguments

object an object inheriting from class "logLik", usually resulting from applying a logLik method to a fitted model object.
... further arguments to be passed to or from methods.
k numeric, the "penalty" per parameter to be used; the default $\mathrm{k}=2$ is the classical AIC.

## Value

a numeric value with the corresponding AIC.

## Author(s)

Jose Pinheiro and Douglas Bates

## References

Sakamoto, Y., Ishiguro, M., and Kitagawa G. (1986). Akaike Information Criterion Statistics. D. Reidel Publishing Company.

## See Also

AIC, logLik.

```
airmiles Commercial Airline Mileage
```


## Description

The revenue passenger miles flown by commercial airlines in the United States for each year from 1937 to 1960.

## Usage

data(airmiles)

## Format

A time-series of 24 observations; yearly, 1937-1960.

## Source

F.A.A. Statistical Handbook of Aviation.

## References

Brown, R. G. (1963) Smoothing, Forecasting and Prediction of Discrete Time Series. Prentice-Hall.

## Examples

```
data(airmiles)
plot(airmiles, main = "airmiles data",
    xlab = "Passenger-miles flown by U.S. commercial airlines", col = 4)
```

airquality New York Air Quality Measurements

## Description

Daily air quality measurements in New York, May to September 1973.

## Usage

data(airquality)

## Format

A data frame with 154 observations on 6 variables.

| $[, 1]$ | Ozone | numeric | Ozone (ppb) |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | Solar.R | numeric | Solar R (lang) |
| $[, 3]$ | Wind | numeric | Wind (mph) |
| $[, 4]$ | Temp | numeric | Temperature (degrees F) |
| $[, 5]$ | Month | numeric | Month (1-12) |
| $[, 6]$ | Day | numeric | Day of month (1-31) |

## Details

Daily readings of the following air quality values for May 1, 1973 (a Tuesday) to September 30, 1973.

- Ozone: Mean ozone in parts per billion from 1300 to 1500 hours at Roosevelt Island
- Solar . R: Solar radiation in Langleys in the frequency band 4000-7700 Angstroms from 0800 to 1200 hours at Central Park
- Wind: Average wind speed in miles per hour at 0700 and 1000 hours at LaGuardia Airport
- Temp: Maximum daily temperature in degrees Fahrenheit at La Guardia Airport.


## Source

The data were obtained from the New York State Department of Conservation (ozone data) and the National Weather Service (meteorological data).

## References

Chambers, J. M., Cleveland, W. S., Kleiner, B. and Tukey, P. A. (1983) Graphical Methods for Data Analysis. Belmont, CA: Wadsworth.

## Examples

```
data(airquality)
pairs(airquality, panel = panel.smooth, main = "airquality data")
```

```
alias Find Aliases (Dependencies) in a Model
```


## Description

Find aliases (linearly dependent terms) in a linear model specified by a formula.

## Usage

```
alias(object, ...)
alias.formula(object, data, ...)
alias.lm(object, complete = TRUE, partial = FALSE,
    partial.pattern = FALSE, ...)
```


## Arguments

| object | A fitted model object, for example from lm or aov, or a formula for <br> alias.formula. |
| :--- | :--- |
| data | Optionally, a data frame to search for the objects in the formula. |
| complete | Should information on complete aliasing be included? |
| partial | $\quad$ Should information on partial aliasing be included? |
| partial.pattern |  |$\quad$| Should partial aliasing be presented in a schematic way? If this is done, |
| :--- |
| the results are presented in a more compact way, usually giving the deciles |
| of the coefficients. |
| further arguments passed to or from other methods. |

## Details

Although the main method is for class "lm", alias is most useful for experimental designs and so is used with fits from aov. Complete aliasing refers to effects in linear models that cannot be estimated independently of the terms which occur earlier in the model and so have their coefficients omitted from the fit. Partial aliasing refers to effects that can be estimated less precisely because of correlations induced by the design.

## Value

A list (of class "listof") containing components
Model Description of the model; usually the formula.
Complete A matrix with columns corresponding to effects that are linearly dependent on the rows; may be of class "mtable" which has its own print method.

Partial The correlations of the estimable effects, with a zero diagonal.

## Note

The aliasing pattern may depend on the contrasts in use: Helmert contrasts are probably most useful.

The defaults are different from those in S.

## Author(s)

B.D. Ripley

## Examples

```
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,55.0,
    62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
                            K=factor(K), yield=yield)
## The next line is optional (for fractions package which gives neater
## results.)
has.VR <- require(MASS, quietly = TRUE)
op <- options(contrasts=c("contr.helmert", "contr.poly"))
npk.aov <- aov(yield ~ block + N*P*K, npk)
alias(npk.aov)
if(has.VR) detach(package:MASS)
options(op)# reset
```


## all

Are All Values True?

## Description

Given a set of logical vectors, are all of the values true?

## Usage

```
all(..., na.rm = FALSE)
```


## Arguments

$\begin{array}{ll}\ldots . & \text { one or more logical vectors. } \\ \text { na.rm } & \text { logical. If true NA values are removed before the result is computed. }\end{array}$

## Value

Given a sequence of logical arguments, a logical value indicating whether or not all of the elements of $x$ are TRUE.
The value returned is TRUE if all the values in x are TRUE, and FALSE if any the values in x are FALSE.
If $x$ consists of a mix of TRUE and NA values, then value is NA.

## See Also

any, the "complement" of all, and stopifnot(*) which is an all(*)"insurance".

## Examples

```
range(x <- sort(round(rnorm(10) - 1.2,1)))
if(all(x < 0)) cat("all x values are negative\n")
```

```
all.equal Test if Two Objects are (Nearly) Equal
```


## Description

all .equal ( $x, y$ ) is a utility to compare $R$ objects $x$ and $y$ testing "near equality". If they are different, comparison is still made to some extent, and a report of the differences is returned. Don't use all.equal directly in if expressions - either use identical or combine the two, as shown in the documentation for identical.

## Usage

all.equal(target, current, ...)
all.equal.numeric(target, current, tolerance= .Machine\$double.eps ^ 0.5, scale=NULL, ...)

## Arguments

| target | R object. |
| :--- | :--- |
| current | other R object, to be compared with target. |
| $\ldots$ | Further arguments for different methods, notably the following two, for <br> numerical comparison: |
| tolerance | numeric $\geq 0$. Differences smaller than tolerance are not considered. <br> scale |
| numeric scalar $>0$ (or NULL). See Details. |  |

## Details

There are several methods available, most of which are dispatched by the default method, see methods("all.equal"). all.equal.list and all.equal.language provide comparison of recursive objects.

Numerical comparisons for scale $=$ NULL (the default) are done by first computing the mean absolute difference of the two numerical vectors. If this is smaller than tolerance or not finite, absolute differences are used, otherwise relative differences scaled by the mean absolute difference.

If scale is positive, absolute comparisons are after scaling (dividing) by scale.
For complex arguments, Mod of difference is used.
attr.all.equal is used for comparing attributes, returning NULL or character.

## Value

Either TRUE or a vector of mode "character" describing the differences between target and current.

Numerical differences are reported by relative error

## See Also

$==$, and all for exact equality testing.

## Examples

```
all.equal(pi, 355/113) # not precise enough (default tol) > relative error
stopifnot(
all.equal(gamma(2:14), cumprod(1:13))) # TRUE, but
all (gamma(2:14) == cumprod(1:13)) # FALSE, since not exactly
all.equal(gamma(2:14), cumprod(1:13), tol=0) # to see difference
all.equal(options(), .Options)
all.equal(options(), as.list(.Options))# TRUE
.Options $ myopt <- TRUE
all.equal(options(), as.list(.Options))
rm(.Options)
```

```
all.names Find All Names in an Expression
```


## Description

Return a character vector containing all the names which occur in an expression or call.

## Usage

```
all.names(expr, functions = TRUE,
    max.names = 200, unique = FALSE)
all.vars(expr, functions = FALSE,
    max.names = 200, unique = TRUE)
```


## Arguments

| expr | an expression or call from which the names are to be extracted. |
| :--- | :--- |
| functions | a logical value indicating whether function names should be included in <br> the result. |
| max.names | the maximum number of names to be returned. |
| unique | a logical value which indicates whether duplicate names should be re- <br> moved from the value. |

## Details

These functions differ only in the default values for their arguments.

## Value

A character vector with the extracted names.

## Examples

all.names $(\operatorname{expression}(\sin (x+y)))$
all.vars (expression $(\sin (x+y)))$

## anova Anova Tables

## Description

Compute analysis of variance (or deviance) tables for one or more fitted model objects.

## Usage

anova(object, ...)

## Arguments

object an object containing the results returned by a model fitting function (e.g. lm or glm).
... additional objects of the same type.

## Value

This (generic) function returns an object of class anova. These objects represent analysis-of-variance and analysis-of-deviance tables. When given a single argument it produces a table which tests whether the model terms are significant.

When given a sequence of objects, anova tests the models against one another in the order specified.

The print method for anova objects prints tables in a "pretty" form.

## Warning

The comparison between two or more models will only be valid if they are fitted to the same dataset. This may be a problem if there are missing values and R's default of na.action = na.omit is used.

## See Also

 coefficients, effects, fitted.values, residuals, summary.```
anova.glm Analysis of Deviance for Generalized Linear Model Fits
```


## Description

Compute an analysis of deviance table for one or more generalized linear model fits.

## Usage

anova(object, ..., dispersion = NULL, test = NULL)

## Arguments

object, ... objects of class glm, typically the result of a call to glm, or a list of objects for the "glmlist" method.
dispersion the dispersion parameter for the fitting family. By default it is obtained from glm.obj.
test a character string, (partially) matching one of "Chisq", "F" or "Cp". See stat.anova.

## Details

Specifying a single object gives a sequential analysis of deviance table for that fit. That is, the reductions in the residual deviance as each term of the formula is added in turn are given in as the rows of a table, plus the residual deviances themselves.
If more than one object is specified, the table has a row for the residual degrees of freedom and deviance for each model. For all but the first model, the change in degrees of freedom and deviance is also given. (This only make statistical sense if the models are nested.) It is conventional to list the models from smallest to largest, but this is up to the user.
The table will optionally contain test statistics (and P values) comparing the reduction in deviance for the row to the residuals. For models with known dispersion (e.g. binomial and Poisson fits) the chi-squared test is most appropriate, and for those with dispersion estimated by moments (e.g. gaussian, quasibinomial and quasipoisson fits) the F test is most appropriate. Mallows' $C_{p}$ statistic is the residual deviance plus twice the estimate of $\sigma^{2}$ times the residual degrees of freedom, which is closely related to AIC (and a multiple of it if the dispersion is known).

## Value

An object of class "anova" inheriting from class "data.frame".

## Warning

The comparison between two or more models by anova or anova.glmlist will only be valid if they are fitted to the same dataset. This may be a problem if there are missing values and R's default of na.action $=$ na.omit is used, and anova.glmlist will detect this with an error.

## See Also

## Examples

```
## --- Continuing the Example from ''?glm'):
anova(glm.D93)
anova(glm.D93, test = "Cp")
anova(glm.D93, test = "Chisq")
```

anova.lm ANOVA for Linear Model Fits

## Description

Compute an analysis of variance table for one or more linear model fits.

## Usage

```
anova(object, ...)
anova.lmlist(object, ..., scale \(=0\), test \(=\) "F")
```


## Arguments

object, ... objects of class lm, usually, a result of a call to lm.
test a character string specifying the test statistic to be used. Can be one of "F", "Chisq" or "Cp", with partial matching allowed, or NULL for no test.
scale numeric. An estimate of the noise variance $\sigma^{2}$. If zero this will be estimated from the largest model considered.

## Details

Specifying a single object gives a sequential analysis of variance table for that fit. That is, the reductions in the residual sum of squares as each term of the formula is added in turn are given in as the rows of a table, plus the residual sum of squares.
The table will contain F statistics (and P values) comparing the mean square for the row to the residual mean square.

If more than one object is specified, the table has a row for the residual degrees of freedom and sum of squares for each model. For all but the first model, the change in degrees of freedom and sum of squares is also given. (This only make statistical sense if the models are nested.) It is conventional to list the models from smallest to largest, but this is up to the user.
Optionally the table can include test statistics. Normally the F statistic is most appropriate, which compares the mean square for a row to the residual sum of squares for the largest model considered. If scale is specified chi-squared tests can be used. Mallows' $C_{p}$ statistic is the residual sum of squares plus twice the estimate of $\sigma^{2}$ times the residual degrees of freedom.

## Value

An object of class "anova" inheriting from class "data.frame".

## Warning

The comparison between two or more models will only be valid if they are fitted to the same dataset. This may be a problem if there are missing values and R's default of na.action = na.omit is used, and anova.lmlist will detect this with an error.

## Note

Versions of R prior to 1.2 .0 based F tests on pairwise comparisons, and this behaviour can still be obtained by a direct call to anovalist.lm.

## See Also

The model fitting function lm.

## Examples

```
## sequential table
data(LifeCycleSavings)
fit <- lm(sr ~ ., data = LifeCycleSavings)
anova(fit)
## same effect via separate models
fit0 <- lm(sr ~ 1, data = LifeCycleSavings)
```

```
fit1 <- update(fit0, . ~ . + pop15)
fit2 <- update(fit1, . ~ . + pop75)
fit3 <- update(fit2, . ~ . + dpi)
fit4 <- update(fit3, . ~ . + ddpi)
anova(fit0, fit1, fit2, fit3, fit4, test="F")
anova(fit4, fit2, fit0, test="F") # unconventional order
```

anscombe Anscombe's Quartet of "Identical" Simple Linear Regressions

## Description

Four $x-y$ datasets which have the same traditional statistical properties (mean, variance, correlation, regression line, etc.), yet are quite different.

## Usage

data(anscombe)

## Format

A data frame with 11 observations on 8 variables.

$$
\begin{aligned}
\mathrm{x} 1==\mathrm{x} 2==\mathrm{x} 3 & \text { the integers } 4: 14, \text { specially arranged } \\
\mathrm{x} 4 & \text { values } 8 \text { and } 19
\end{aligned}
$$

## Source

Tufte, Edward R. (1989) The Visual Display of Quantitative Information, 13-14. Graphics Press.

## References

Anscombe, Francis J. (1973) Graphs in statistical analysis. American Statistician, 27, 17-21.

## Examples

```
data(anscombe)
summary(anscombe)
##-- now some "magic" to do the 4 regressions in a loop:
ff <- y ~ x
for(i in 1:4) {
    ff[2:3] <- lapply(paste(c("y","x"), i, sep=""), as.name)
    ## or ff[[2]] <- as.name(paste("y", i, sep=""))
    ## ff[[3]] <- as.name(paste("x", i, sep=""))
    assign(paste("lm.",i,sep=""), lmi <- lm(ff, data= anscombe))
    print(anova(lmi))
}
## See how close they are (numerically!)
```

```
sapply(objects(pat="lm\.[1-4]$"), function(n) coef(get(n)))
lapply(objects(pat="lm\.[1-4]$"), function(n) summary(get(n))$coef)
## Now, do what you should have done in the first place: PLOTS
op <- par(mfrow=c(2,2), mar=. 1+c(4,4,1,1), oma=c(0,0,2,0))
for(i in 1:4) {
        ff[2:3] <- lapply(paste(c("y","x"), i, sep=""), as.name)
        plot(ff, data =anscombe, col="red", pch=21, bg = "orange", cex = 1.2,
            xlim=c(3,19), ylim=c(3,13))
        abline(get(paste("lm.",i,sep="")), col="blue")
}
mtext("Anscombe's 4 Regression data sets", outer = TRUE, cex=1.5)
par(op)
```

any Are Some Values True?

## Description

Given a set of logical vectors, are any of the values true?

## Usage

any (..., na.rm = FALSE)

## Arguments

... one or more logical vectors.
na.rm logical. If true NA values are removed before the result is computed.

## Value

Given a sequence of logical arguments, a logical value indicating whether or not any of the elements of $x$ are TRUE.
The value returned is TRUE if any the values in x are TRUE, and FALSE if all the values in x are FALSE.

If x consists of a mix of FALSE and NA values, the value is NA.

## See Also

all, the "complement" of any.

## Examples

```
range(x <- sort(round(rnorm(10) - 1.2,1)))
if(any(x < 0)) cat("x contains negative values\n")
```


## aov

## Description

Summarize an analysis of variance model.

## Usage

```
summary(object, intercept = FALSE, split,
    expand.split = TRUE, keep.zero.df = TRUE, ...)
summary(object, ...)
```


## Arguments

| object | An object of class "aov" or "aovlist". |
| :--- | :--- |
| intercept | logical: should intercept terms be included? |
| split | an optional named list, with names corresponding to terms in the model. <br> Each component is itself a list with integer components giving contrasts <br> whose contributions are to be summed. |
| expand.split | logical: should the split apply also to interactions involving the factor? |
| keep.zero.df | logical: should terms with no degrees of freedom be included? |
| $\ldots$ | Arguments to be passed to or from other methods, for summary.aovlist <br> including those for summary.aov. |

## Value

An object of class c("summary.aov", "listof") or "summary.aovlist" respectively.

## Note

The use of expand.split = TRUE is little tested: it is always possible to set it to FALSE and specify exactly all the splits required.

## Author(s)

B. D. Ripley

## See Also

aov, summary, model.tables, TukeyHSD

## Examples

```
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,55.0,
    62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
K=factor(K), yield=yield)
```

```
( npk.aov <- aov(yield ~ block + N*P*K, npk) )
summary(npk.aov)
coefficients(npk.aov)
# Cochran and Cox (1957, p.164)
# 3x3 factorial with ordered factors, each is average of 12.
CC <- data.frame(
    y = c(449, 413, 326, 409, 358, 291, 341, 278, 312)/12,
    P = ordered(gl(3, 3)), N = ordered(gl(3, 1, 9))
)
CC.aov <- aov(y ~ N * P, data = CC , weights = rep(12, 9))
summary(CC.aov)
# Split both main effects into linear and quadratic parts.
summary(CC.aov, split = list(N = list(L = 1, Q = 2), P = list(L = 1, Q = 2)))
# Split only the interaction
summary(CC.aov, split = list("N:P" = list(L.L = 1, Q = 2:4)))
```

aperm Array Transposition

## Description

Transpose an array by permuting its dimensions and optionally resizing it.

## Usage

```
aperm(a, perm, resize = TRUE)
```


## Arguments

a the array to be transposed.
perm the subscript permutation vector, which must be a permutation of the integers $1: n$, where $n$ is the number of dimensions of $a$. The default is to reverse the order of the dimensions.
resize a flag indicating whether the vector should be resized as well as having its elements reordered (default TRUE).

## Value

A transposed version of array a, with subscripts permuted as indicated by the array perm. If resize is TRUE, the array is reshaped as well as having its elements permuted, the dimnames are also permuted; if FALSE then the returned object has the same dimensions as a, and the dimnames are dropped.
The function $t$ provides a faster and more convenient way of transposing matrices.

## Author(s)

Jonathan Rougier, 〈J.C.Rougier@durham.ac.uk $\rangle$ did the faster C implementation.

## See Also

t , to transpose matrices.

## Examples

```
# interchange the first two subscripts on a 3-way array x
x <- array(1:24, 2:4)
xt <- aperm(x, c(2,1,3))
stopifnot(t(xt[,,2]) == x[,,2],
    t(xt[,,3]) == x[, ,3],
    t(xt[, 4] ) == x[, 4])
```

    append Vector Merging
    
## Description

Add elements to a vector.

## Usage

append(x, values, after=length(x))

## Arguments

| x | the vector to be modified. |
| :--- | :--- |
| values | to be included in the modified vector. |
| after | a subscript, after which the values are to be appended. |

## Value

A vector containing the values in x with the elements of values appended after the specified element of x .

## Examples

```
stopifnot(
    append(1:5, 0:1, after=3)
== append(1:3,c(0:1, 4:5)))
```


## apply Apply Functions Over Array Margins

## Description

Returns a vector or array or list of values obtained by applying a function to margins of an array.

## Usage

apply (X, MARGIN, FUN, ...)

## Arguments

X
the array to be used.
MARGIN a vector giving the subscripts which the function will be applied over. 1 indicates rows, 2 indicates columns, $c(1,2)$ indicates rows and columns.
FUN the function to be applied. In the case of functions like,$+ \% * \%$, etc., the function name must be quoted.
... optional arguments to FUN.

## Details

If $X$ is not an array but has a dimension attribute, apply attempts to coerce it to an array via as.matrix if it is two-dimensional (e.g. data frames) or via as.array.

## Value

If each call to FUN returns a vector of length $n$, then apply returns an array of dimension $c(n, \operatorname{dim}(X)[M A R G I N])$ if $n>1$. If $n$ equals 1 , apply returns a vector if MARGIN has length 1 and an array of dimension $\operatorname{dim}(X)$ [MARGIN] otherwise. If $n$ is 0 , the result has length 0 but not necessarily the "correct" dimension.
If the calls to FUN return vectors of different lengths, apply returns a list of length dim (X) [MARGIN].

## See Also

lapply, tapply, and convenience functions sweep and aggregate.

## Examples

```
## Compute row and column sums for a matrix:
x <- cbind(x1 = 3, x2 = c(4:1, 2:5))
dimnames(x)[[1]] <- letters[1:8]
apply(x, 2, mean, trim = .2)
col.sums <- apply(x, 2, sum)
row.sums <- apply(x, 1, sum)
rbind(cbind(x, Rtot = row.sums), Ctot = c(col.sums, sum(col.sums)))
stopifnot( apply(x,2, is.vector)) # not ok in R <= 0.63.2
## Sort the columns of a matrix
```

```
apply(x, 2, sort)
##- function with extra args:
cave <- function(x, c1,c2) c(mean(x[c1]),mean(x[c2]))
apply(x,1, cave, c1="x1", c2=c("x1","x2"))
ma <- matrix(c(1:4, 1, 6:8), nr = 2)
ma
apply(ma, 1, table) #--> a list of length 2
apply(ma, 1, quantile)# 5 x n matrix with rownames
stopifnot(dim(ma) == dim(apply(ma, 1:2, sum)))## wasn't ok before R 0.63.1
```

approxfun Interpolation Functions

## Description

Return a list of points which linearly interpolate given data points, or a function performing the linear (or constant) interpolation.

## Usage

```
approx (x, y, xout, method="linear", n=50,
    yleft, yright, rule = 1, f=0, ties = mean)
approxfun(x, y, method="linear",
    yleft, yright, rule = 1, f=0, ties = mean)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | vectors giving the coordinates of the points to be interpolated. Alternatively a single plotting structure can be specified: see xy.coords. |
| :---: | :---: |
| xout | an optional set of values specifying where interpolation is to take place. |
| method | specifies the interpolation method to be used. Choices are "linear" or "constant". |
| n | If xout is not specified, interpolation takes place at $n$ equally spaced points spanning the interval $[\min (x), \max (x)]$. |
| yleft | the value to be returned when input $x$ values less than $\min (x)$. The default is defined by the value of rule given below. |
| yright | the value to be returned when input $x$ values greater than $\max (x)$. The default is defined by the value of rule given below. |
| rule | an integer describing how interpolation is to take place outside the interval $[\min (x), \max (x)]$. If rule is 1 then NAs are returned for such points and if it is 2 , the value at the closest data extreme is used. |
| f | For method="constant" a number between 0 and 1 inclusive, indicating a compromise between left- and right-continuous step functions. If y0 and y 1 are the values to the left and right of the point then the value is $y 0 *(1-f)+y 1 * f$ so that $f=0$ is right-continuous and $f=1$ is left-continuous. |
| ties | Handling of tied x values. Either a function with a single vector argument returning a single number result or the string "ordered". |

## Details

The inputs can contain missing values which are deleted, so at least two complete ( $\mathrm{x}, \mathrm{y}$ ) pairs are required. If there are duplicated (tied) $x$ values and ties is a function it is applied to the y values for each distinct x value. Useful functions in this context include mean, min, and max. If ties="ordered" the x values are assumed to be already ordered. The first y value will be used for interpolation to the left and the last one for interpolation to the right.

## Value

approx returns a list with components x and y , containing n coordinates which interpolate the given data points according to the method (and rule) desired.
The function approxfun returns a function performing (linear or constant) interpolation of the given data points. For a given set of x values, this function will return the corresponding interpolated values. This is often more useful than approx.

## See Also

spline and splinefun for spline interpolation.

## Examples

```
x <- 1:10
y <- rnorm(10)
par(mfrow = c(2,1))
plot(x, y, main = "approx(.) and approxfun(.)")
points(approx(x, y), col = 2, pch = "*")
points(approx(x, y, method = "constant"), col = 4, pch = "*")
f <- approxfun(x, y)
curve(f(x), 0, 10, col = "green")
points(x, y)
is.function(fc <- approxfun(x, y, method = "const")) # TRUE
curve(fc(x), 0, 10, col = "darkblue", add = TRUE)
## Show treatment of 'ties' :
x <- c(2,2:4,4,4,5,5,7,7,7)
y <- c(1:6, 5:4, 3:1)
approx(x,y, xout=x)$y # warning
(ay <- approx(x,y, xout=x, ties = "ordered")$y)
stopifnot(ay == c(2,2,3,6,6,6,4,4,1,1,1))
approx(x,y, xout=x, ties = min)$y
approx(x,y, xout=x, ties = max)$y
```

apropos Find Objects by (Partial) Name

## Description

apropos returns a character vector giving the names of all objects in the search list matching what.
find is a different user interface to the same task as apropos.

## Usage

```
apropos(what, where = FALSE, mode = "any")
find(what, mode = "any", numeric. = FALSE, simple.words = TRUE)
```


## Arguments

what name of an object, or regular expression to match against
where, numeric.
a logical indicating whether positions in the search list should also be returned
mode character; if not "any", only objects who's mode equals mode are searched.
simple.words logical; if TRUE, the what argument is only searched as whole only word.

## Details

If mode != "any" only those objects which are of mode mode are considered. If where is TRUE, the positions in the search list are returned as the names attribute.
find is a different user interface to the same task as apropos. However, by default (simple.words == TRUE), only full words are searched.

## Author(s)

Kurt Hornik and Martin Maechler (May 1997).

## See Also

objects for listing objects from one place, help.search for searching the help system, search for the search path.

## Examples

```
apropos("lm")
apropos(ls)
apropos("lq")
lm <- 1:pi
find(lm) #> ".GlobalEnv" "package:base"
find(lm, num=TRUE) # numbers with these names
find(lm, num=TRUE, mode="function")# only the second one
rm(lm)
apropos(".", mode="list")
# need a DOUBLE backslash '\\' (in case you don't see it anymore)
apropos("\\[")
# everything
length(apropos("."))
# those starting with 'pr'
apropos("`pr")
# the 1-letter things
apropos("^.$")
```

```
# the 1-2-letter things
apropos("^..?$")
# the 2-to-4 letter things
apropos("`.{2,4}$")
# the 8-and-more letter things
apropos("^.{8,}$")
table(nchar(apropos("`.{8,}$")))
```

args Argument List of a Function

## Description

Displays the argument names and corresponding default values of a function.

## Usage

args (name)

## Arguments

name an interpreted function. If name is a character string then the function with that name is found and used.

## Details

This function is mainly used interactively. For programming, use formals instead.

## Value

A function with identical formal argument list but an empty body if given an interpreted function; NULL in case of a variable or primitive (non-interpreted) function.

## See Also <br> formals, help.

## Examples

```
args(c) # -> NULL (c is a 'primitive' function)
args(plot.default)
```

Arithmetic Arithmetic Operators

## Description

These binary operators perform arithmetic on vector objects.

## Usage

$x+y$
$x-y$
$x$ * $y$
$\mathrm{x} / \mathrm{y}$
x - y
x \% \% y
x \% / \% y

## Details

1 ^ y and y ^ 0 are 1, always. x ^ y should also give the proper "limit" result when either argument is infinite (i.e., +- Inf).
Objects such as arrays or time-series can be operated on this way provided they are conformable.

## Value

They return numeric vectors containing the result of the element by element operations. The elements of shorter vectors are recycled as necessary (with a warning when they are recycled only fractionally). The operators are + for addition, - for subtraction $*$ for multiplication, / for division and ^ for exponentiation.
$\% \%$ indicates $\mathrm{x} \bmod \mathrm{y}$ and $\% / \%$ indicates integer division. It is guaranteed that $\mathrm{x}==$ ( x $\% \% \mathrm{y})+\mathrm{y} *(\mathrm{x} \% / \% \mathrm{y})$ unless $\mathrm{y}==0$ where the result is NA or NaN (depending on the typeof of the arguments).

## See Also

sqrt for miscellaneous and Special for special mathematical functions.
Syntax for operator precedence.

## Examples

```
x <- -1:12
x + 1
2*x + 3
x %% 2 #-- is periodic
x %/% 5
```

```
array Multi-way Arrays
```


## Description

Creates or tests for arrays.

## Usage

```
array(data \(=\) NA, dim \(=\) length(data), dimnames \(=\) NULL)
```

as.array (x)
is.array ( $x$ )

## Arguments

data a vector giving data to fill the array.
dim the dim attribute for the array to be created, that is a vector of length one or more giving the maximal indices in each dimension.
dimnames the names for the dimensions. This is a list with one component for each dimension, either NULL or a character vector of the length given by dim for that dimension. The list can be names, and the names will be used as names for the dimensions.
x an R object.

## Value

array returns an array with the extents specified in dim and naming information in dimnames. The values in data are taken to be those in the array with the leftmost subscript moving fastest. If there are too few elements in data to fill the array, then the elements in data are recycled.
as. $\operatorname{array}()$ coerces its argument to be an array by attaching a dim attribute to it. It also attaches dimnames if x has names. The sole purpose of this is to make it possible to access the dim[names] attribute at a later time.
is.array returns TRUE or FALSE depending on whether its argument is an array (i.e., has a dim attribute) or not.

## See Also

aperm, matrix, dim, dimnames.

## Examples

```
dim(as.array(letters))
array(1:3, c(2,4)) # recycle 1:3 "2 2/3 times"
# [,1] [,2] [,3] [,4]
#[1,] 1 3 2 1
#[2,] 
# funny object:
str(a0 <- array(1:3, 0))
```

```
arrows Add Arrows to a Plot
```


## Description

Draw arrows between pairs of points.

## Usage

```
arrows(x0, y0, x1, y1, length = 0.25, angle = 30, code = 2,
    col = par("fg"), lty = NULL, lwd = par("lwd"), xpd = NULL)
```


## Arguments

| $\mathrm{x} 0, \mathrm{y} 0$ | coordinates of points from which to draw. |
| :--- | :--- |
| $\mathrm{x} 1, \mathrm{y} 1$ | coordinates of points to which to draw. |
| length | length of the edges of the arrow head (in inches). |
| angle | angle from the shaft of the arrow to the edge of the arrow head. <br> code <br> col, lty, lwd, |
|  | integer code, determining kind of arrows to be drawn. |
|  | usual graphical parameters as in par. |

## Details

For each i, an arrow is drawn between the point (x0[i], yO[i]) and the point (x1[i],y1[i]).

If code=2 an arrowhead is drawn at ( x 0 [i],y0[i]) and if code=1 an arrowhead is drawn at ( $\mathrm{x} 1[\mathrm{i}], \mathrm{y} 1[\mathrm{i}]$ ). If code $=3$ a head is drawn at both ends of the arrow. Unless length $=$ 0 , when no head is drawn.
The graphical parameters col and lty can be used to specify a color and line texture for the line segments which make up the arrows (col may be a vector).
The direction of a zero-length arrow is indeterminate, and hence so is the direction of the arrowheads. To allow for rounding error, arrowheads are omitted (with a warning) on any arrow of length less than $1 / 1000$ inch.

## See Also

segments to draw segments.

## Examples

```
x <- runif(12); y <- rnorm(12)
i <- order(x,y); x <- x[i]; y <- y[i]
plot(x,y, main="arrows(.) and segments(.)")
## draw arrows from point to point :
s <- seq(length(x)-1)# one shorter than data
arrows(x[s], y[s], x[s+1], y[s+1], col= 1:3)
s <- s[-length(s)]
segments(x[s], y[s], x[s+2], y[s+2], col= 'pink')
```


## Description

Converts a number or a character string to the corresponding environment on the search path.

## Usage

as.environment (object)

## Arguments

object the object to convert. If it is already an environment, just return it. If it is a number, return the environment corresponding to that position on the search list. If it is a character string, match the string to the names on the search list.

## Value

The corresponding environment object.

## Author(s)

John Chambers

## See Also

environment for creation and manipulation, search.

## Examples

```
as.environment(1) ## the global environment
identical(globalenv(), as.environment(1)) ## is TRUE
as.environment("package:ctest")
```

```
as.function Convert Object to Function
```


## Description

as.function is a generic function which is used to convert objects to functions.
as.function.default works on a list $x$, which should contain the concatenation of a formal argument list and an expression or an object of mode "call" which will become the function body. The function will be defined in a specified environment, by default that of the caller.

## Usage

as.function(x, ...)
as.function.default(x, envir = parent.frame(), ...)

## Arguments

```
x object to convert, a list for the default method.
... additional arguments, depending on object
envir environment in which the function should be defined
```


## Value

The desired function.

## Author(s)

Peter Dalgaard

## See Also

 function; alist which is handy for the construction of argument lists, etc.
## Examples

```
as.function(alist(a=, b=2,a+b))
as.function(alist(a=,b=2,a+b))(3)
```

```
as.POSIX* Date-time Conversion Functions
```


## Description

Functions to manipulate objects of classes "POSIXIt" and "POSIXct" representing calendar dates and times (to the nearest second).

## Usage

```
as.POSIXct(x, tz = "")
as.POSIXlt(x, tz = "")
```


## Arguments

x
tz

An object to be converted.
A timezone specification to be used for the conversion, if one is required. System-specific, but "" is the current timezone, and "GMT" is UTC (Coordinated Universal Time, in French).

## Details

The as.POSIX* functions convert an object to one of the two classes used to represent date/times (calendar dates plus time to the nearest second). They can take convert a wide variety of objects, including objects of the other class and of classes "date" (from package date), "chron" and "dates" (from package chron) to these classes. They can also convert character strings of the formats "2001-02-03" and "2001/02/03" optionally followed by white space and a time in the format " $14: 52$ " or "14:52:03". (Formats such as "01/02/03" are ambiguous but can be converted via a format specification by strptime.)
Logical NAs can be converted to either of the classes, but no other logical vectors can be.

## Value

as.POSIXct and as.POSIXlt return an object of the appropriate class. If $t z$ was specified, as.POSIXlt will give an appropriate "tzone" attribute.

## Note

If you want to extract specific aspects of a time (such as the day of the week) just convert it to class "POSIXIt" and extract the relevant component(s) of the list, or if you want a character representation (such as a named day of the week) use format.POSIXIt or format. POSIXct.
If a timezone is needed and that specified is invalid on your system, what happens is systemspecific but it will probably be ignored.

## See Also

DateTimeClasses for details of the classes; strptime for conversion to and from character representations.

## Examples

```
(z <- Sys.time()) # the current date, as class "POSIXct"
unclass(z) # a large integer
floor(unclass(z)/86400) # the number of days since 1970-01-01
(z <- as.POSIXlt(Sys.time())) # the current date, as class "POSIXlt"
unlist(unclass(z)) # a list shown as a named vector
as.POSIXlt(Sys.time(), "GMT") # the current time in GMT
```


## Description

Change the class of an object to indicate that it should be treated "as is".

## Usage

I (x)

## Arguments

x
an object

## Details

Function I has two main uses.

- In function data.frame. Protecting an object by enclosing it in I() in a call to data.frame inhibits the conversion of character vectors to factors. I can also be used to protect objects which are to be added to a data frame, or converted to a data frame via as.data.frame.
It achieves this by prepending the class "AsIs" to the object's classes. Class "AsIs" has a few of its own methods, including for [, as.data.frame, print and format.
- In function formula. There it is used to inhibit the interpretation of operators such as "+", "-", "*" and "~" as formula operators, so they are used as arithmetical operators. This is interpreted as a symbol by terms.formula.


## Value

A copy of the object with class "AsIs" prepended to the class(es).

See Also<br>data.frame, formula

assign Assign a Value to a Name

## Description

Assign a value to a name in an environment.

## Usage

```
assign(x, value, pos = -1, envir = as.environment(pos),
    inherits = FALSE, immediate = TRUE)
```


## Arguments

x
a variable name (given as a quoted string in the function call).
value a value to be assigned to $x$.
pos where to do the assignment. By default, assigns into the current environment. See the details for other possibilities.
envir the environment to use. See the details section.
inherits should the enclosing frames of the environment be inspected?
immediate an ignored compatibility feature.

## Details

The pos argument can specify the environment in which to assign the object in any of several ways: as an integer (the position in the search list); as the character string name of an element in the search list; or as an environment (including using sys.frame to access the currently active function calls). The envir argument is an alternative way to specify an environment, but is primarily there for back compatibility.
assign does not dispatch assignment methods, so it cannot be used to set elements of vectors, names, attributes, etc.

## Value

This function is invoked for its side effect, which is assigning value to the variable x . If no envir is specified, then the assignment takes place in the currently active environment.

If inherits is TRUE, enclosing environments of the supplied environment are searched until the variable x is encountered. The value is then assigned in the environment in which the variable is encountered. If the symbol is not encountered then assignment takes place in the user's workspace (the global environment).

If inherits is FALSE, assignment takes place in the initial frame of envir.

## See Also

<-, get, exists, environment.

## Examples

```
for(i in 1:6) { #-- Create objects 'r1', 'r2', ... 'r6' --
    nam <- paste("r",i, sep=".")
    assign(nam, 1:i)
}
ls(pat="^r..$")
##-- Global assignment within a function:
myf <- function(x) {
    innerf <- function(x) assign("Global.res", x^2, env = .GlobalEnv)
    innerf(x+1)
}
myf(3)
Global.res # 16
a <- 1:4
assign("a[1]", 2)
a[1] == 2 #FALSE
get("a[1]") == 2 #TRUE
```

assign0ps Assignment Operators

## Description

Assign a value to a name.

## Usage

```
x <- value
x <<- value
value -> x
value ->> x
x = value
```


## Arguments

| x | a variable name (possibly quoted). |
| :--- | :--- |
| value | a value to be assigned to x. |

## Details

There are three different assignment operators: two of them have leftwards and rightwards forms.
The operators <- and = assign into the environment in which they are evaluated. The <can be used anywhere, but the $=$ is only allowed at the top level (that is, in the complete expression typed by the user) or as one of the subexpressions in a braced list of expressions.
The operators <<- and ->> cause a search to made through the environment for an existing definition of the variable being assigned. If such a variable is found then its value is redefined, otherwise assignment takes place globally. Note that their semantics differ from that in the $S$ language, but is useful in conjunction with the scoping rules of $R$.
In all the assignment operator expressions, x can be a name or an expression defining a part of an object to be replaced (e.g., $z[[1]])$. The name does not need to be quoted, though it can be.
The leftwards forms of assignment <- = <<- group right to left, the other from left to right.

## Value

value. Thus one can use a <- b <- c <- 6 .

## See Also

assign, environment.

```
assocplot Association Plots
```


## Description

Produce a Cohen-Friendly association plot indicating deviations from independence of rows and columns in a 2-dimensional contingency table.

## Usage

assocplot(x, col = c("black", "red"), space = 0.3, main $=$ NULL, $x l a b=$ NULL, ylab $=$ NULL)

## Arguments

x
a two-dimensional contingency table in matrix form.
col a character vector of length two giving the colors used for drawing positive and negative Pearson residuals, respectively.
space the amount of space (as a fraction of the average rectangle width and height) left between each rectange.
main overall title for the plot.

| xlab | a label for the x axis. Defaults to the name of the row variable in x if <br> non-NULL. <br> ylab <br> a label for the y axis. Defaults to the column names of the column variable <br> in x if non-NULL. |
| :--- | :--- |

## Details

For a two-way contingency table, the signed contribution to Pearson's $\chi^{2}$ for cell $i, j$ is $d_{i j}=\left(f_{i j}-e_{i j}\right) / \sqrt{e_{i j}}$, where $f_{i j}$ and $e_{i j}$ are the observed and expected counts corresponding to the cell. In the Cohen-Friendly association plot, each cell is represented by a rectangle that has (signed) height proportional to $d_{i j}$ and width proportional to $\sqrt{e_{i j}}$, so that the area of the box is proportional to the difference in observed and expected frequencies. The rectangles in each row are positioned relative to a baseline indicating independence $\left(d_{i j}=0\right)$. If the observed frequency of a cell is greater than the expected one, the box rises above the baseline and is shaded in the color specified by the first element of col, which defaults to black; otherwise, the box falls below the baseline and is shaded in the color specified by the second element of col, which defaults to red.

## References

Cohen, A. (1980), On the graphical display of the significant components in a two-way contingency table. Communications in Statistics—Theory and Methods, A9, 1025-1041.
Friendly, M. (1992), Graphical methods for categorical data. SAS User Group International Conference Proceedings, 17, 190-200. http://www.math.yorku.ca/SCS/sugi/ sugi17-paper.html

## See Also

```
mosaicplot; chisq.test.
```


## Examples

```
data(HairEyeColor)
## Aggregate over sex:
x <- margin.table(HairEyeColor, c(1, 2))
x
assocplot(x, main = "Relation between hair and eye color")
```

attach Attach Set of $R$ Objects to Search Path

## Description

The database is attached to the R search path. This means that the database is searched by $R$ when evaluating a variable, so objects in the database can be accessed by simply giving their names.

## Usage

$\operatorname{attach}($ what, $\mathrm{pos}=2$, name $=$ deparse (substitute(what)))

## Arguments

| what | "database". This may currently be a data.frame or list or a R data file |
| :--- | :--- |
| created with save. |  |

## Details

When evaluating a variable or function name R searches for that name in the databases listed by search. The first name of the appropriate type is used.
By attaching a data frame to the search path it is possible to refer to the variables in the data frame by their names alone, rather than as components of the data frame (eg in the example below, height rather than women\$height).
By default the database is attached in position 2 in the search path, immediately after the user's workspace and before all previously loaded packages and previously attached databases. This can be altered to attach later in the search path with the pos option, but you cannot attach at pos=1.
Note that by default assignment is not performed in an attached database. Attempting to modify a variable or function in an attached database will actually create a modified version in the user's workspace (the R global environment). For this reason attach can lead to confusion.

## Value

The environment is returned invisibly with a "name" attribute.

## See Also

```
library, detach, search, objects, environment.
```


## Examples

```
data(women)
summary(women$height) ## refers to variable 'height' in the dataframe
attach(women)
summary(height) ## The same variable now available by name
height<-height*2.54 ## Don't do this. It creates a new variable
detach("women")
summary(height) ## The new variable created by modifying 'height'
rm(height)
```

attenu The Joyner-Boore Attenuation Data

## Description

This data gives peak accelerations measured at various observation stations for 23 earthquakes in California. The data have been used by various workers to estimate the attenuating affect of distance on ground acceleration.

## Usage

data(attenu)

## Format

A dataframe with 182 observations on 5 variables.

| $[, 1]$ | event | numeric | Event Number |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | mag | numeric | Moment Magnitude |
| $[, 3]$ | station | factor | Station Number |
| $[, 4]$ | dist | numeric | Station-hypocenter distance (km) |
| $[, 5]$ | accel | numeric | Peak acceleration (g) |

## Source

Joyner, W.B., D.M. Boore and R.D. Porcella (1981). Peak horizontal acceleration and velocity from strong-motion records including records from the 1979 Imperial Valley, California earthquake. USGS Open File report 81-365. Menlo Park, Ca.

## References

Boore, D. M. and Joyner, W.B.(1982) The empirical prediction of ground motion, Bull. Seism. Soc. Am., 72, S269-S268.

Bolt, B. A. and Abrahamson, N. A. (1982) New attenuation relations for peak and expected accelerations of strong ground motion, Bull. Seism. Soc. Am., 72, 2307-2321.

Bolt B. A. and Abrahamson, N. A. (1983) Reply to W. B. Joyner \& D. M. Boore's "Comments on: New attenuation relations for peak and expected accelerations for peak and expected accelerations of strong ground motion", Bull. Seism. Soc. Am., 73, 1481-1483.
Brillinger, D. R. and Preisler, H. K. (1984) An exploratory analysis of the Joyner-Boore attenuation data, Bull. Seism. Soc. Am. 74, 1441-1449.

Brillinger, D. R. and Preisler, H. K. (1984) Further analysis of the Joyner-Boore attenuation data. Manuscript.

## Examples

```
data(attenu)
## check the data class of the variables
sapply(attenu, data.class)
summary(attenu)
pairs(attenu, main = "attenu data")
coplot(accel ~ dist | as.factor(event), data = attenu, show = FALSE)
coplot(log(accel) ~ log(dist) | as.factor(event),
    data = attenu, panel = panel.smooth, show.given = FALSE)
```


## Description

From a survey of the clerical employees of a large financial organization, the data are aggregated from the questionnaires of the approximately 35 employees for each of 30 (randomly selected) departments. The numbers give the percent proportion of favourable responses to seven questions in each department.

## Usage

```
data(attitude)
```


## Format

A dataframe with 30 observations on 7 variables. The first column are the short names from the reference, the second one the variable names in the data frame:

| Y | rating | numeric | Overall rating |
| :--- | :--- | :--- | :--- |
| $\mathrm{X}[1]$ | complaints | numeric | Handling of employee complaints |
| $\mathrm{X}[2]$ | privileges | numeric | Does not allow special privileges |
| $\mathrm{X}[3]$ | learning | numeric | Opportunity to learn |
| $\mathrm{X}[4]$ | raises | numeric | Raises based on performance |
| $\mathrm{X}[5]$ | critical | numeric | Too critical |
| $\mathrm{X}[6]$ | advancel | numeric | Advancement |

## Source

Chatterjee, S. and Price, B. (1977) Regression Analysis by Example. New York: Wiley. (Section 3.7, p.68ff of 2nd ed.(1991).)

## Examples

```
data(attitude)
pairs(attitude, main = "attitude data")
summary(attitude)
summary(fm1 <- lm(rating ~ ., data = attitude))
opar <- par(mfrow = c(2, 2), oma = c(0, 0, 1.1, 0),
    mar = c(4.1, 4.1, 2.1, 1.1))
plot(fm1)
summary(fm2 <- lm(rating ~ complaints, data = attitude))
plot(fm2)
par(opar)
```

attr Object Attributes

## Description

Get or set specific attributes of an object.

## Usage

```
attr(x, which)
attr(x, which) <- value
```


## Arguments

```
x an object whose attributes are to be accessed.
which a character string specifying which attribute is to be accessed.
```


## Value

This function provides access to a single object attribute. The simple form above returns the value of the named attribute. The assignment form causes the named attribute to take the value on the right of the assignment symbol.

## See Also

attributes

## Examples

```
# create a 2 by 5 matrix
x <- 1:10
attr(x,"dim") <- c(2, 5)
```


## Description

These functions access an object's attribute list. The first form above returns the an object's attribute list. The assignment forms make the list on the right-hand side of the assignment the object's attribute list (if appropriate).

## Usage

```
attributes(obj)
attributes(obj) <- list
mostattributes(obj) <- list
```


## Arguments

obj an object

## Details

The mostattributes assignment takes special care for the dim, names and dimnames attributes, and assigns them only when that is valid whereas as attributes assignment would give an error in that case.

## See Also

attr.

## Examples

```
x <- cbind(a=1:3, pi=pi) # simple matrix w/ dimnames
str(attributes(x))
## strip an object's attributes:
attributes(x) <- NULL
x # now just a vector of length 6
mostattributes(x) <- list(mycomment = "really special", dim = 3:2,
    dimnames = list(LETTERS[1:3], letters[1:5]), names = paste(1:6))
x # dim(), but not {dim}names
```


## autoload On-demand Loading of Packages

## Description

autoload creates a promise-to-evaluate autoloader and stores it with name name in .AutoloadEnv environment. When R attempts to evaluate name, autoloader is run, the package is loaded and name is re-evaluated in the new package's environment. The result is that $R$ behaves as if file was loaded but it does not occupy memory.

## Usage

autoload(name, package,...)
autoloader (name, package,...)
. AutoloadEnv

## Arguments

| name | string giving the name of an object |
| :--- | :--- |
| package | string giving the name of a package containing the object |
| $\ldots$. | other arguments to library |

## Value

This function is invoked for its side-effect.

## See Also

delay, library

## Examples

```
autoload("line", "eda")
search()
ls("Autoloads")
all(ls("Autoloads") == ls(envir = .AutoloadEnv))
data(cars)
plot(cars)
z<-line(cars)
abline(coef(z))
search()
```

```
detach("package:eda")
search()
z<-line(cars)
search()
```

ave
Group Averages Over Level Combinations of Factors

## Description

Subsets of x[] are averaged, where each subset consist of those observations with the same factor levels.

## Usage

ave (x, ..., FUN = mean)

## Arguments

$\mathrm{x} \quad$ A numeric.
... Grouping variables, typically factors, all of the same length as x .
FUN Function to apply for each factor level combination.

## Value

A numeric vector, say $y$ of length length(x). If ... is $\mathrm{g} 1, \mathrm{~g} 2$, e.g., $\mathrm{y}[\mathrm{i}]$ is equal to FUN ( $\mathrm{x}[\mathrm{j}]$, for all j with $\mathrm{g} 1[\mathrm{j}]==\mathrm{g} 1[\mathrm{i}]$ and $\mathrm{g} 2[\mathrm{j}]==\mathrm{g} 2[\mathrm{i}]$ ).

## See Also

mean, median.

## Examples

```
ave(1:3)# no grouping -> grand mean
data(warpbreaks)
attach(warpbreaks)
ave(breaks, wool)
ave(breaks, tension)
ave(breaks, tension, FUN = function(x)mean(x, trim=.1))
plot(breaks, main =
    "ave( Warpbreaks ) for wool x tension combinations")
lines(ave(breaks, wool, tension ), type='s', col = "blue")
lines(ave(breaks, wool, tension, FUN=median), type='s', col = "green")
legend(40,70, c("mean","median"), lty=1,col=c("blue","green"), bg="gray90")
detach()
```


## Description

Adds an axis to the current plot, allowing the specification of the side, position, labels, and other options.

## Usage

```
axis(side, at = NULL, labels = TRUE, tick = TRUE, line = 0,
    pos = NA, outer = FALSE, font = NA, vfont = NULL,
    lty = "solid", lwd = 1, col = NULL, ...)
```


## Arguments

side an integer specifying which side of the plot the axis is to be drawn on. The axis is placed as follows: $1=$ below, $2=$ left, $3=$ above and $4=$ right.
at the points at which tick-marks are to be drawn. Non-finite (infinite, NaN or NA) values are omitted. By default, when NULL, tickmark locations are computed, see Details below.
labels this can either be a logical value specifying whether (numerical) annotations are to be made at the tickmarks, or a vector of character strings to be placed at the tickpoints.
tick a logical value specifying whether tickmarks should be drawn
line the number of lines into the margin which the axis will be drawn. This overrides the value of the graphical parameter mgp [3]. The relative placing of tickmarks and tick labels is unchanged.
pos the coordinate at which the axis line is to be drawn. this overrides the value of both line and mgp [3].
outer a logical value indicating whether the axis should be drawn in the outer plot margin, rather than the standard plot margin.
font font for text.
vfont vector font for text.
lty, lwd line type, width for the axis line and the tick marks.
col color for the axis line and the tick marks. The default NULL means to use par("fg").
... other graphical parameters may also be passed as arguments to this function, e.g., las for vertical/horizontal label orientation, see par(las=.).

## Details

The axis line is drawn from the lowest to the highest value of at, but will be clipped at the plot region. Only ticks which are drawn from points within the plot region (up to a tolerance for rounding error) are plotted, but the ticks and their labels may well extend outside the plot region.
When at $=$ NULL, pretty tick mark locations are computed internally, from par("usr","lab"), and par("xlog") (or ylog respectively).

## Value

This function is invoked for its side effect, which is to add an axis to an already existing plot.

## Examples

```
plot(1:4, rnorm(4), axes=FALSE)
axis(1, 1:4, LETTERS[1:4])
axis(2)
box() #- to make it look "as usual"
plot(1:7, rnorm(7), main = "axis() examples",
    type = "s", xaxt="n", frame = FALSE, col = "red")
axis(1, 1:7, LETTERS[1:7], col.axis = "blue")
# unusual options:
axis(4, col = "violet", col.axis="dark violet",lwd = 2)
axis(3, col = "gold", lty = 2, lwd = 0.5)
```

```
axis.POSIXct Date-time Plotting Functions
```


## Description

Functions to manipulate objects of classes "POSIXIt" and "POSIXct" representing calendar dates and times (to the nearest second).

## Usage

```
plot.POSIXct(x, y, xlab = "", ...)
plot.POSIXlt(x, y, xlab = "", ...)
axis.POSIXct(side, x, at, format, ...)
```


## Arguments

| x, at | A date-time object. |
| :--- | :--- |
| y | numeric values to be plotted against x. |
| xlab | a character string giving the label for the x axis. |
| side | See axis. |
| format | See strptime. |
| $\ldots$ | Further arguments to be passed from or to other methods, typically graph- <br> ical parameters. |

## Details

The functions plot against an x -axis of date-times. axis.POSIXct works quite hard to choose suitable time units (years, months, days, hours, minutes or seconds) and a sensible output format, but this can be overridden by supplying a format specification.
If at is supplied for axis.POSIXct it specifies the locations of the ticks and labels: if x is specified a suitable grid of labels is chosen.

## See Also

DateTimeClasses for details of the classes.

## Examples

```
if(require(MASS, quietly = TRUE)) {
data(beav1)
attach(beav1)
time <- strptime(paste(1990, day, time %/% 100, time %% 100),
    "%Y %j %H %M")
plot(time, temp, type="l") # axis at 4-hour intervals.
# now label every hour on the time axis
plot(time, temp, type="l", xaxt="n")
r <- as.POSIXct(round(range(time), "hours"))
axis.POSIXct(1, at=seq(r[1], r[2], by="hour"), format="%H")
rm(time)
detach(beav1)
detach(package:MASS)
}
plot(.leap.seconds, 1:22, type="n", yaxt="n",
    xlab="leap seconds", ylab="", bty="n")
rug(.leap.seconds)
```


## Description

Compute tickmark locations, the same way as R does internally. This is only non-trivial when $\log$ coordinates are active. By default, gives the at values which axis(side) would use.

## Usage

axTicks(side, axp $=$ NULL, usr $=$ NULL, log = NULL)

## Arguments

side $\quad$ integer in $1: 4$, as for axis.
$\operatorname{axp} \quad$ numeric vector of length three, defaulting to $\operatorname{par}(" Z a x p ")$ where " Z " is " x " or " y " depending on the side argument.
usr numeric vector of length four, defaulting to par("usr") giving horizontal (' $x$ ') and vertical (' $y$ ') user coordinate limits.
log logical indicating if $\log$ coordinates are active; defaults to par("Zlog") where ' $Z$ ' is as for the axp argument above.

## Details

The axp, usr, and log arguments must be consistent as their default values (the par (..) results) are. Note that the meaning of axp alters very much when log is TRUE, see the documentation on $\operatorname{par}$ (xaxp=.).
axTicks() can be regarded as an R implementation of the C function CreateAtVector() in '..../src/main/graphics.c' which is called by axis(side,$*$ ) when no argument at is specified.

## Value

numeric vector of coordinate values at which axis tickmarks can be drawn. By default, when only the first argument is specified, these values should be identical to those that axis (side) would use or has used.

## See Also

axis,par.

## Examples

```
plot(1:7, 10*21:27)
axTicks(1)
axTicks(2)
stopifnot(identical(axTicks(1), axTicks(3)),
    identical(axTicks(2), axTicks(4)))
## Show how axTicks() and axis() correspond :
op <- par(mfrow = c(3,1))
for(x in 9999*c(1,2,8)) {
    plot(x,9, log = "x")
    cat(formatC(par("xaxp"),wid=5),";",T <- axTicks(1),"\n")
    rug(T, col="red")
}
par(op)
```

backsolve Solve an Upper or Lower Triangular System

## Description

Solves a system of linear equations where the coefficient matrix is upper or lower triangular.

## Usage

```
    backsolve(r, x, k= ncol(r), upper.tri = TRUE, transpose = FALSE)
```



## Arguments

| $\mathrm{r}, \mathrm{l}$ | an upper (or lower) triangular matrix giving the coefficients for the system |
| :--- | :--- |
| to be solved. Values below (above) the diagonal are ignored. |  |
| x | a matrix whose columns give "right-hand sides" for the equations. |
| k | The number of columns of r and rows of x to use. |

upper.tri logical; if TRUE (default), the upper triangular part of r is used. Otherwise, the lower one.
transpose logical; if TRUE, solve $r^{\prime} * y=x$ for $y$, i.e., $\mathrm{t}(\mathrm{r}) \% * \% \mathrm{y}==\mathrm{x}$.

## Value

The solution of the triangular system. The result will be a vector if x is a vector and a matrix if x is a matrix.

## References

Dongarra, J. J., Bunch,J. R., Moler, C. B. and Stewart, G. W. (1978) LINPACK Users Guide. Philadelphia: SIAM Publications.

## See Also

```
chol, qr, solve.
```


## Examples

```
## upper triangular matrix 'r':
r <- rbind(c(1,2,3),
    c (0,1,1),
    c(0,0,2))
( y <- backsolve(r, x <- c(8,4,2)) ) # -1 3 1
r %*% y # == x = ( }8,4,2
( y2 <- backsolve(r, x, transpose = TRUE)) # 8 -12 -5
all(t(r) %*% y2 == x)# exactly on Linux (Pentium)
all(y == backsolve(t(r), x, upper = FALSE, transpose = TRUE))
all(y2 == backsolve(t(r), x, upper = FALSE, transpose = FALSE))
```


## bandwidth Bandwidth Selectors for Kernel Density Estimation

## Description

Bandwidth selectors for gaussian windows in density.

## Usage

```
bw.nrd0(x)
bw.nrd(x)
bw.ucv(x, nb = 1000, lower, upper)
bw.bcv(x, nb = 1000, lower, upper)
bw.SJ(x, nb=1000, lower, upper, method=c("ste", "dpi"))
```


## Arguments

```
x A data vector.
nb number of bins to use.
lower, upper Range over which to minimize. The default is almost always satisfactory.
method Either "ste" ("solve-the-equation") or "dpi" ("direct plug-in").
```


## Details

bw.nrd0 implements a rule-of-thumb for choosing the bandwidth of a Gaussian kernel density estimator. It defaults to 0.9 times the minimum of the standard deviation and the interquartile range divided by 1.34 times the sample size to the negative one-fifth power (= Silverman's "rule of thumb", Silverman(1986, page 48, eqn (3.31)) unless the quartiles coincide when a positive result will be guaranteed.
bw.nrd is the more common variation given by Scott (1992), using factor 1.06.
bw.ucv and bw.bcv implement unbiased and biased cross-validation respectively.
bw. SJ implements the methods of Sheather \& Jones (1991) to select the bandwidth using pilot estimation of derivatives.

## Value

A bandwidth on a scale suitable for the bw argument of density.

## References

Scott, D. W. (1992) Multivariate Density Estimation: Theory, Practice, and Visualization. Wiley.

Sheather, S. J. and Jones, M. C. (1991) A reliable data-based bandwidth selection method for kernel density estimation. Journal of the Royal Statistical Society series B 53, 683-690.

Silverman, B. W. (1986) Density Estimation. London: Chapman and Hall.
Venables, W. N. and Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. Springer.

## See Also

density.
bandwith.nrd, ucv, bcv and width.SJ in MASS, which are all scaled to the width argument of density and so give answers four times as large.

## Examples

```
data(precip)
plot(density(precip, n=1000))
rug(precip)
lines(density(precip, bw="nrd"), col = 2)
lines(density(precip, bw="ucv"), col = 3)
lines(density(precip, bw="bcv"), col = 4)
lines(density(precip, bw="SJ-ste"), col = 5)
lines(density(precip, bw="SJ-dpi"), col = 6)
legend(55, 0.035
legend = c("nrdO", "nrd", "ucv", "bcv", "SJ-ste", "SJ-dpi"),
col = 1:6, lty = 1)
```

```
barplot Bar Plots
```


## Description

Creates a bar plot with vertical or horizontal bars.

## Usage

```
barplot(height, width = 1, space = NULL,
    names.arg \(=\) NULL, legend.text \(=\) NULL, beside \(=\) FALSE,
    horiz = FALSE, density \(=\) NULL, angle \(=45\),
    col = heat.colors(NR), border = par("fg"),
    main \(=\) NULL, sub \(=\) NULL, \(x l a b=\) NULL, ylab \(=\) NULL,
    xlim = NULL, ylim = NULL, xpd = TRUE,
    axes \(=\) TRUE, axisnames \(=\) TRUE,
    cex.axis = par("cex.axis"), cex.names = par("cex.axis"),
    inside \(=\) TRUE, plot \(=\) TRUE, axis.lty \(=0\), ...)
```


## Arguments

| height | either a vector or matrix of values describing the bars which make up the <br> plot. If height is a vector, the plot consists of a sequence of rectangular <br> bars with heights given by the values in the vector. If height is a matrix <br> and beside is FALSE then each bar of the plot corresponds to a column <br> of height, with the values in the column giving the heights of stacked <br> "sub-bars" making up the bar. If height is a matrix and beside is TRUE, <br> then the values in each column are juxtaposed rather than stacked. <br> optional vector of bar widths. Re-cycled to length the number of bars <br> drawn. Specifying a single value will no visible effect unless xlim is spec- <br> ified. <br> the amount of space (as a fraction of the average bar width) left before |
| :--- | :--- |
| each bar. May be given as a single number or one number per bar. If |  |
| space |  |
| height is a matrix and beside is TRUE, space may be specified by two |  |
| numbers, where the first is the space between bars in the same group, |  |
| and the second the space between the groups. If not given explicitly, it |  |
| defaults to c (0,1) if height is a matrix and beside is TRUE, and to 0.2 |  |
| otherwise. |  |


| horiz | a logical value. If FALSE, the bars are drawn vertically with the first bar |
| :--- | :--- |
| to the left. If TRUE, the bars are drawn horizontally with the first at the |  |
| bottom. |  |

## Details

This is a generic function, it currently only has a default method. A formula interface may be added eventually.

## Value

A numeric vector (or matrix, when beside $=$ TRUE), say mp, giving the coordinates of all the bar midpoints drawn, useful for adding to the graph.
If beside is true, use colMeans (mp) for the midpoints of each group of bars, see example.

## Note

Prior to R 1.6.0, barplot behaved as if axis.lty $=1$, unintentionally.

See Also

```
plot(..., type="h"), dotchart, hist.
```


## Examples

```
tN <- table(Ni <- rpois(100, lambda=5))
r <- barplot(tN, col='gray')
#- type = "h" plotting *is* 'bar'plot
lines(r, tN, type='h', col='red', lwd=2)
barplot(tN, space = 1.5, axisnames=FALSE,
    sub = "barplot(..., space= 1.5, axisnames = FALSE)")
data(VADeaths, package = "base")
barplot(VADeaths, plot = FALSE)
barplot(VADeaths, plot = FALSE, beside = TRUE)
mp <- barplot(VADeaths) # default
tot <- colMeans(VADeaths)
text(mp, tot + 3, format(tot), xpd = TRUE, col = "blue")
barplot(VADeaths, beside = TRUE,
    col = c("lightblue", "mistyrose", "lightcyan",
    "lavender", "cornsilk"),
    legend = rownames(VADeaths), ylim = c(0, 100))
title(main = "Death Rates in Virginia", font.main = 4)
hh <- t(VADeaths)[, 5:1]
mybarcol <- "gray20"
mp <- barplot(hh, beside = TRUE,
    col = c("lightblue", "mistyrose",
        "lightcyan", "lavender"),
    legend = colnames(VADeaths), ylim= c(0,100),
    main = "Death Rates in Virginia", font.main = 4,
    sub = "Faked upper 2*sigma error bars", col.sub = mybarcol,
    cex.names = 1.5)
segments(mp, hh, mp, hh + 2*sqrt(1000*hh/100), col = mybarcol, lwd = 1.5)
stopifnot(dim(mp) == dim(hh))# corresponding matrices
mtext(side = 1, at = colMeans(mp), line = -2,
    text = paste("Mean", formatC(colMeans(hh))), col = "red")
# Bar shading example
barplot(VADeaths, angle = 15+10*1:5, density = 20, col = "black",
    legend = rownames(VADeaths))
title(main = list("Death Rates in Virginia", font = 4))
# border :
barplot(VADeaths, border = "dark blue")
```


## BATCH $\quad$ Batch Execution of $R$

## Description

Run R non-interactively with input from a given file and place output (stdout/stderr) to another file.

## Usage

R CMD BATCH [options] infile [outfile]

## Arguments

| infile | the name of a file with $R$ code to be executed. |
| :--- | :--- |
| options | a list of $R$ command line options, e.g., for setting the amount of memory <br> available and controlling the load/save process. If infile starts with a <br> -, use -- as the final option. |
| outfile | the name of a file to which to write output. If not given, the name used <br> is the one of infile, with a possible ' $\cdot R^{\prime}$ extension stripped, and '.Rout' <br> appended. |

## Details

By default, the input commands are printed along with the output. To suppress this behavior, add options (echo = FALSE) at the beginning of infile.
The infile can have end of line marked by LF or CRLF (but not just CR), and files with a missing EOL mark are processed correctly.
Using R CMD BATCH sets the GUI to "none", so none of x11, jpeg and png are available.

## Note

Unlike Splus BATCH, this does not run the R process in the background. In most shells, R CMD BATCH [options] infile [outfile] \& will do so.

## Bessel Bessel Functions

## Description

Bessel Functions of integer and fractional order, of first and second kind, $J_{\nu}$ and $Y_{\nu}$, and Modified Bessel functions (of first and third kind), $I_{\nu}$ and $K_{\nu}$.
gammaCody is the $(\Gamma)$ function as from the Specfun package and originally used in the Bessel code.

## Usage

```
besselI(x, nu, expon.scaled = FALSE)
besselK(x, nu, expon.scaled = FALSE)
besselJ(x, nu)
besselY(x, nu)
gammaCody(x)
```


## Arguments

$\mathrm{x} \quad$ numeric,$\geq 0$.
nu numeric; The order (maybe fractional!) of the corresponding Bessel function.
expon.scaled logical; if TRUE, the results are exponentially scaled in order to avoid overflow $\left(I_{\nu}\right)$ or underflow $\left(K_{\nu}\right)$, respectively.

## Details

The underlying C code stems from Netlib (http://www.netlib.org/specfun/r[ijky] besl).
If expon.scaled $=$ TRUE, $e^{-x} I_{\nu}(x)$, or $e^{x} K_{\nu}(x)$ are returned.
gammaCody may be somewhat faster but less precise and/or robust than R's standard gamma. It is here for experimental purpose mainly, and may be defunct very soon.

For $\nu<0$, formulae 9.1.2 and 9.6.2 from the reference below are applied (which is probably suboptimal), unless for bessel K which is symmetric in nu.

## Value

Numeric vector of the same length of $x$ with the (scaled, if expon.scale=TRUE) values of the corresponding Bessel function.

## Author(s)

Original Fortran code: W. J. Cody, Argonne National Laboratory Translation to C and adaption to R: Martin Maechler 〈maechler@stat.math.ethz.ch.)

## References

Abramowitz, M. and Stegun, I. A. (1972) Handbook of Mathematical Functions. Dover, New York; Chapter 9: Bessel Functions of Integer Order.

## See Also

Other special mathematical functions, as the gamma, $\Gamma(x)$, and beta, $B(x)$.

## Examples

```
nus <- c(0:5,10,20)
x <- seq(0,4, len= 501)
plot(x,x, ylim = c(0,6), ylab="",type='n', main = "Bessel Functions I_nu(x)")
for(nu in nus) lines(x,besselI(x,nu=nu), col = nu+2)
legend(0,6, leg=paste("nu=",nus), col = nus+2, lwd=1)
x <- seq(0,40,len=801); yl <- c(-.8,.8)
plot(x,x, ylim = yl, ylab="",type='n', main = "Bessel Functions J_nu(x)")
for(nu in nus) lines(x,besselJ(x,nu=nu), col = nu+2)
legend(32,-.18, leg=paste("nu=",nus), col = nus+2, lwd=1)
## Negative nu's :
xx <- 2:7
nu <- seq(-10,9, len = 2001)
op <- par(lab = c(16,5,7))
matplot(nu, t(outer(xx,nu, besselI)), type = 'l', ylim = c(-50,200),
    main = expression(paste("Bessel ",I[nu](x)," for fixed ", x,
                                    ", as ",f(nu))),
        xlab = expression(nu))
abline(v=0, col = "light gray", lty = 3)
legend(5,200, leg = paste("x=",xx), col=seq(xx), lty=seq(xx))
par(op)
x0 <- 2^(-20:10)
```

```
plot(x0,x0^-8, log='xy', ylab="",type='n',
    main = "Bessel Functions J_nu(x) near O\n log - log scale")
for(nu in sort(c(nus,nus+.5))) lines(x0,besselJ(x0,nu=nu), col = nu+2)
legend(3,1e50, leg=paste("nu=", paste(nus,nus+.5, sep=",")), col=nus+2, lwd=1)
plot(x0,x0^-8, log='xy', ylab="",type='n',
    main = "Bessel Functions K_nu(x) near O\n log - log scale")
for(nu in sort(c(nus,nus+.5))) lines(x0,besselK(x0,nu=nu), col = nu+2)
legend(3,1e50, leg=paste("nu=", paste(nus,nus+.5, sep=",")), col=nus+2, lwd=1)
x <- x[x > 0]
plot(x,x, ylim=c(1e-18,1e11),log="y", ylab="",type='n',
    main = "Bessel Functions K_nu(x)")
for(nu in nus) lines(x,besselK(x,nu=nu), col = nu+2)
legend(0,1e-5, leg=paste("nu=",nus), col = nus+2, lwd=1)
## Check the Scaling :
for(nu in nus)
    print(all(abs(1- besselK(x,nu)*exp( x) / besselK(x,nu,expo=TRUE)) < 2e-15))
for(nu in nus)
    print(all(abs(1- besselI(x,nu)*exp(-x) / besselI(x,nu,expo=TRUE)) < 1e-15))
yl <- c(-1.6, .6)
plot(x,x, ylim = yl, ylab="",type='n', main = "Bessel Functions Y_nu(x)")
for(nu in nus) {xx <- x[x > .6*nu]; lines(xx,besselY(xx,nu=nu), col = nu+2)}
legend(25,-.5, leg=paste("nu=",nus), col = nus+2, lwd=1)
```


## Beta The Beta Distribution

## Description

Density, distribution function, quantile function and random generation for the Beta distribution with parameters shape1 and shape2 (and optional non-centrality parameter ncp).

## Usage

```
dbeta(x, shape1, shape2, ncp=0, log = FALSE)
pbeta(q, shape1, shape2, ncp=0, lower.tail = TRUE, log.p = FALSE)
qbeta(p, shape1, shape2, lower.tail = TRUE, log.p = FALSE)
rbeta(n, shape1, shape2)
```


## Arguments

```
x, q vector of quantiles.
p vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the
    number required.
shape1, shape2
    positive parameters of the Beta distribution.
```

```
ncp non-centrality parameter.
log, log.p logical; if TRUE, probabilities p are given as log(p).
lower.tail logical; if TRUE (default), probabilities are P[X\leqx], otherwise, P[X>
    x].
```


## Details

The Beta distribution with parameters shape $1=a$ and shape $2=b$ has density

$$
f(x)=\frac{\Gamma(a+b)}{\Gamma(a) \Gamma(b)} x^{a}(1-x)^{b}
$$

for $a>0, b>0$ and $0<x<1$.

## Value

dbeta gives the density, pbeta the distribution function, qbeta the quantile function, and rbeta generates random deviates.

## See Also

beta for the Beta function, and dgamma for the Gamma distribution.

## Examples

```
x <- seq(0, 1, length=21)
dbeta(x, 1, 1)
pbeta(x, 1, 1)
```


## bindenv Binding and Environment Adjustments

## Description

These functions represent an experimental interface for adjustments to environments and bindings within environments. They allow for locking environments as well as individual bindings, and for linking a variable to a function.

## Usage

```
lockEnvironment(env, bindings = FALSE)
environmentIsLocked(env)
lockBinding(sym, env)
bindingIsLocked(sym, env)
makeActiveBinding(sym, fun, env)
bindingIsActive(sym, env)
```


## Arguments

| env | an environment. |
| :--- | :--- |
| bindings | logical specifying whether bindings should be locked. |
| sym | a name object or character string |
| fun | a function taking zero or one arguments |

## Details

The function lockEnvironment locks its environment argument, which must be a proper environment, not NULL. Locking the NULL (base) environment may be supported later. Locking the environment prevents adding or removing variable bindings from the environment. Changing the value of a variable is still possible unless the binding has been locked.
lockBinding locks individual bindings in the specified environment. The value of a locked binding cannot be changed. Locked bindings may be removed from an environment unless the environment is locked
makeActiveBinding installs fun so that getting the value of sym calls fun with no arguments, and assigning to sym calls fun with one argument, the value to be assigned. This allows things like C variables linked to R variables and variables linked to data bases to be implemented. It may also be useful for making thread-safe versions of some system globals.

## Author(s)

Luke Tierney

## Examples

```
# locking environments
e<-new.env()
assign("x",1, env=e)
get("x",env=e)
lockEnvironment(e)
get("x",env=e)
assign("x",2, env=e)
try(assign("y",2, env=e)) # error
# locking bindings
e<-new.env()
assign("x",1, env=e)
get("x",env=e)
lockBinding("x", e)
try(assign("x",2, env=e)) # error
# active bindings
f<-local({
    x <- 1
    function(v) {
            if (missing(v))
                cat("get\n")
            else {
                cat("set\n")
                x <<- v
            }
            x
    }
})
makeActiveBinding("fred", f, .GlobalEnv)
bindingIsActive("fred", .GlobalEnv)
fred
fred<-2
fred
```


## Binomial The Binomial Distribution

## Description

Density, distribution function, quantile function and random generation for the binomial distribution with parameters size and prob.

## Usage

dbinom(x, size, prob, log = FALSE)
pbinom(q, size, prob, lower.tail = TRUE, log.p = FALSE)
qbinom(p, size, prob, lower.tail = TRUE, log.p = FALSE)
rbinom(n, size, prob)

## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n number of observations. If length $(\mathrm{n})>1$, the length is taken to be the number required.
size number of trials.
prob probability of success on each trial.
$\log , \log . \mathrm{p} \quad \operatorname{logical}$; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The binomial distribution with size $=n$ and prob $=p$ has density

$$
p(x)=\binom{n}{x} p^{x}(1-p)^{n-x}
$$

for $x=0, \ldots, n$.
If an element of $\mathbf{x}$ is not integer, the result of dbinom is zero, with a warning. $p(x)$ is computed using Loader's algorithm, see the reference below.
The quantile is defined as the smallest value $x$ such that $F(x) \geq p$, where $F$ is the distribution function.

## Value

dbinom gives the density, pbinom gives the distribution function, qbinom gives the quantile function and rbinom generates random deviates.
If size is not an integer, NaN is returned.

## References

Catherine Loader (2000). Fast and Accurate Computation of Binomial Probabilities; manuscript available from http://cm.bell-labs.com/cm/ms/departments/sia/ catherine/dbinom

## See Also

 dnbinom for the negative binomial, and dpois for the Poisson distribution.
## Examples

```
# Compute P(45 < X < 55) for X Binomial(100,0.5)
sum(dbinom(46:54, 100, 0.5))
## Using "log = TRUE" for an extended range :
n <- 2000
plot (0:n, dbinom(0:n, n, pi/10, log=TRUE), type='l',
    main = "dbinom(*, log=TRUE) is better than log(dbinom(*))")
lines(0:n, log(dbinom(0:n, n, pi/10)), col='red', lwd=2)
mtext("dbinom(k, log=TRUE)", adj=0)
mtext("extended range", adj=0, line = -1, font=4)
mtext("log(dbinom(k))", col="red", adj=1)
```

```
birthday Probability of coincidences
```


## Description

Computes approximate answers to a generalised "birthday paradox" problem. pbirthday computes the probability of a coincidence and qbirthday computes the number of observations needed to have a specified probability of coincidence.

## Usage

```
qbirthday(prob = 0.5, classes = 365, coincident = 2)
pbirthday(n, classes = 365, coincident = 2)
```


## Arguments

| classes | How many distinct categories the people could fall into |
| :--- | :--- |
| prob | The desired probability of coincidence |
| n | The number of people |
| coincident | The number of people to fall in the same category |

## Details

The birthday paradox is that a very small number of people, 23 , suffices to have a $50-50$ chance that two of them have the same birthday. This function generalises the calculation to probabilities other than 0.5 , numbers of coincident events other than 2 , and numbers of classes other than 365 .

This formula is approximate, as the example below shows. For coincident=2 the exact computation is straightforward and may be preferable.

## Value

| qbirthday | Number of people needed for a probability prob that k of them have the <br> same one out of classes equiprobable labels. |
| :--- | :--- |
| pbirthday | Probability of the specified coincidence |

body

## References

Diaconis P, Mosteller F. "Methods for studying coincidences" JASA 84:853-861

## Examples

\#\# the standard version
qbirthday()
\#\# same 4-digit PIN number
qbirthday (classes=10^4)
\#\# 0.9 probability of three coincident birthdays
qbirthday (coincident=3, prob=0.9)
\#\# Chance of 4 coincident birthdays in 150 people
pbirthday (150, coincident=4)
\#\# Accuracy compared to exact calculation
x1<- sapply(10:100, pbirthday)
x2<-1-sapply(10:100, function(n) $\operatorname{prod}((365:(365-n+1)) / r e p(365, n)))$
$\operatorname{par}(m f r o w=c(2,2))$
plot(x1,x2,xlab="approximate", ylab="exact")
abline ( 0,1 )
plot(x1,x1-x2,xlab="approximate", ylab="error")
abline ( $\mathrm{h}=0$ )
plot( $\mathrm{x} 1, \mathrm{x} 2, \log =" \mathrm{xy} ", x \operatorname{lab}=$ "approximate", yl lab="exact")
abline ( 0,1 )
plot(1-x1,1-x2,log="xy",xlab="approximate",ylab="exact")
abline $(0,1)$
body Access to and Manipulation of the Body of a Function

## Description

Get or set the body of a function.

## Usage

body(fun = sys.function(sys.parent()))
body(fun) <- list

## Arguments

fun a function object or a character string naming the function to be manipulated. If not specified, the function calling body is used.
list a list of $R$ expressions.

## Value

body returns the body of the function specified.
The assignment form sets the body of a function to the list on the right hand side.

## See Also

alist, args, function.

## Examples

```
body(body)
f <- function(x) x^5
body(f) <- expression(5`x)
f(3) # = 125
str(body(f))
```

box
Draw a Box around a Plot

## Description

This function draws a box around the current plot in the given color and linetype. The bty parameter determines the type of box drawn. See par for details.

## Usage

```
box(which="plot", lty="solid", ...)
```


## Arguments

which character, one of "plot", "figure", "inner" and "outer".
lty line type of the box.
... further graphical parameters, such as bty, col, or lwd, see par.

## See Also

rect for drawing of arbitrary rectangles.

## Examples

```
plot(1:7,abs(rnorm(7)), type='h', axes = FALSE)
axis(1, labels = letters[1:7])
box(lty='137', col = 'red')
```

```
boxplot Box Plots
```


## Description

Produce box-and-whisker plot(s) of the given (grouped) values.

## Usage

```
boxplot(x, ..., range = 1.5, width = NULL, varwidth = FALSE,
    notch = FALSE, outline = TRUE, names, boxwex = 0.8, plot = TRUE,
    border = par("fg"), col = NULL, log = "", pars = NULL,
    horizontal = FALSE, add = FALSE, at = NULL)
```


## Arguments

| x , | for specifying data from which the boxplots are to be produced as well as for giving graphical parameters. The named arguments in this (more precisely, in list (x, ...)) are treated as graphical parameters in addition to the ones given by argument pars. The other arguments specify the data, either as separate vectors, each corresponding to a component boxplot, or as a single list containing such vectors. NAs are allowed in the data. |
| :---: | :---: |
| range | this determines how far the plot whiskers extend out from the box. If range is positive, the whiskers extend to the most extreme data point which is no more than range times the interquartile range from the box. A value of zero causes the whiskers to extend to the data extremes. |
| width | a vector giving the relative widths of the boxes making up the plot. |
| varwidth | if varwidth is TRUE, the boxes are drawn with widths proportional to the square-roots of the number of observations in the groups. |
| notch | if notch is TRUE, a notch is drawn in each side of the boxes. If the notches of two plots do not overlap then the medians are significantly different at the 5 percent level. |
| outline | if outline is not true, the boxplot lines are not drawn. |
| names | group labels which will be printed under each boxplot. |
| boxwex | a scale factor to be applied to all boxes. When there are only a few groups, the appearance of the plot can be improved by making the boxes narrower. |
| plot | if TRUE (the default) then a boxplot is produced. If not, the summaries which the boxplots are based on are returned. |
| border | an optional vector of colors for the outlines of the boxplots. The values in border are recycled if the length of border is less than the number of plots. |
| col | if col is non-null it is assumed to contain colors to be used to col the bodies of the box plots. |
| $\log$ | character indicating if x or y or both coordinates should be plotted in log scale. |
| pars | graphical parameters can also be passed as arguments to boxplot. |
| horizontal | logical indicating if the boxplots should be horizontal; default FALSE means vertical boxes. |
| add | logical, if true add boxplot to current plot. |
| at | numeric vector giving the locations where the boxplots should be drawn, particularly when add = TRUE; defaults to $1: \mathrm{n}$ where n is the number of boxes. |

## Details

The generic function boxplot currently has a default method (boxplot.default) and a formula interface (boxplot.formula).

## Value

List with the following components:
stats a matrix, each column contains the extreme of the lower whisker, the lower hinge, the median, the upper hinge and the extreme of the upper whisker for one group/plot.
n
conf
out the values of any data points which lie beyond the extremes of the whiskers.
group a vector of the same length as out whose elements indicate which group the outlier belongs to
names a vector of names for the groups

## See Also

boxplot.formula for the formula interface; boxplot.stats which does the computation, bxp for the plotting; and stripchart for an alternative (with small data sets).

## Examples

```
## boxplot on a formula:
data(InsectSprays)
boxplot(count ~ spray, data = InsectSprays, col = "lightgray")
# *add* notches (somewhat funny here):
boxplot(count ~ spray, data = InsectSprays,
    notch = TRUE, add = TRUE, col = "blue")
data(OrchardSprays)
boxplot(decrease ~ treatment, data = OrchardSprays,
        log = "y", col="bisque")
rb <- boxplot(decrease ~ treatment, data = OrchardSprays, col="bisque")
title("Comparing boxplot()s and non-robust mean +/- SD")
mn.t <- tapply(OrchardSprays$decrease, OrchardSprays$treatment, mean)
sd.t <- tapply(OrchardSprays$decrease, OrchardSprays$treatment, sd)
xi <- 0.3 + seq(rb$n)
points(xi, mn.t, col = "orange", pch = 18)
arrows(xi, mn.t - sd.t, xi, mn.t + sd.t,
    code = 3, col = "pink", angle = 75, length = .1)
## boxplot on a matrix:
mat <- cbind(Uni05 = (1:100)/21, Norm = rnorm(100),
    T5 = rt(100, df = 5), Gam2 = rgamma(100, shape = 2))
boxplot(data.frame(mat), main = "boxplot(data.frame(mat), main = ...)")
par(las=1)# all axis labels horizontal
boxplot(data.frame(mat), main = "boxplot(*, horizontal = TRUE)",
        horizontal = TRUE)
## Using 'at = ' and adding boxplots -- example idea by Roger Bivand :
data(ToothGrowth)
```

```
boxplot(len ~ dose, data = ToothGrowth,
    boxwex = 0.25, at = 1:3 - 0.2,
    subset= supp == "VC", col="yellow",
    main="Guinea Pigs' Tooth Growth",
    xlab="Vitamin C dose mg",
    ylab="tooth length", ylim=c(0,35))
boxplot(len ~ dose, data = ToothGrowth, add = TRUE,
    boxwex = 0.25, at = 1:3 + 0.2,
    subset= supp == "OJ", col="orange")
legend(2, 9, c("Ascorbic acid", "Orange juice"),
    fill = c("yellow", "orange"))
```

boxplot.formula Formula Notation for Box Plots

## Description

Produce box-and-whisker plot(s) of the given (grouped) values using formula notation.

## Usage

```
boxplot(formula, data = NULL, ..., subset)
```


## Arguments

| formula | a formula, such as $y \sim \mathrm{x}$. |
| :--- | :--- |
| data | a data.frame (or list) from which the variables in formula should be taken. |
| $\ldots$ | arguments to the default boxplot method and graphical parameters may <br> also be passed as arguments, see par. |
| subset | an optional vector specifying a subset of observations to be used for plot- <br> ting. |

## Details

This is a method of the generic function boxplot. It operates by setting up the data from the formula specification, and then calling boxplot. default.

## See Also

boxplot.default

## Examples

```
data(OrchardSprays)
boxplot(decrease ~ treatment, data = OrchardSprays,
    log = "y", col="bisque")
```

```
boxplot.stats Box Plot Statistics
```


## Description

This function is typically called by boxplot to gather the statistics necessary for producing box plots, but may be invoked separately.

## Usage

boxplot.stats(x, coef = 1.5, do.conf=TRUE, do.out=TRUE)

## Arguments

x
coef this determines how far the plot "whiskers" extend out from the box. If coef is positive, the whiskers extend to the most extreme data point which is no more than coef times the length of the box away from the box. A value of zero causes the whiskers to extend to the data extremes (and no outliers be returned).
do.conf, do.out
logicals; if FALSE, the conf or out component respectively will be empty in the result.

## Details

The two "hinges" are versions of the first and third quartile, i.e. close to quantile ( x , $\mathrm{c}(1,3) / 4$ ). The hinges equal the quartiles for odd $n$ (where n <- length $(\mathrm{x})$ ) and differ for even $n$. Where the quartiles only equal observations for $\mathrm{n} \% \% 4==1(n \equiv 1 \bmod 4)$, the hinges do so additionally for $\mathrm{n} \% \% 4=2(n \equiv 2 \bmod 4)$, and are in the middle of two observations otherwise.

## Value

List with named components as follows:
stats a vector of length 5, containing the extreme of the lower whisker, the lower "hinge", the median, the upper "hinge" and the extreme of the upper whisker.
n
conf the lower and upper extremes of the "notch" (if (do.conf)).
out the values of any data points which lie beyond the extremes of the whiskers (if(do.out)).

Note that \$stats and \$conf are sorted in increasing order, unlike S, and that \$n and \$out include any +- Inf values.

## See Also

fivenum, boxplot, bxp.

## Examples

```
x <- c(1:100, 1000)
str(b1 <- boxplot.stats(x))
str(b2 <- boxplot.stats(x, do.conf=FALSE, do.out=FALSE))
stopifnot(b1 $ stats == b2 $ stats) # do.out=F is still robust
str(boxplot.stats(x, coef = 3, do.conf=FALSE))
## no outlier treatment:
str(boxplot.stats(x, coef = 0))
str(boxplot.stats(c(x, NA))) # slight change : n + 1
str(r <- boxplot.stats(c(x, -1:1/0)))
stopifnot(r$out == c(1000, -Inf, Inf))
```


## Description

The browseEnv function opens a browser with list of objects currently in sys.frame() environment.

## Usage

```
browseEnv(envir = .GlobalEnv, pattern, excludepatt = "^last\\.warning",
                    html = .Platform$OS.type != "mac",
    expanded = TRUE, properties = NULL,
    main = NULL, debugMe = FALSE)
```


## Arguments

| envir | an environment the objects of which are to be browsed. |
| :--- | :--- |
| pattern | a regular expression for object subselection is passed to the internal ls() <br> call. |
| excludepatt | a regular expression for dropping objects with matching names. <br> is used on non Macintosh machines to display the workspace on a HTML <br> page in your favorite browser. |
| html | whether to show one level of recursion. It can be useful to switch it to <br> FALSE if your workspace is large. This option is ignored if html is set to <br> FALSE. |
| properties | a named list of global properties (of the objects chosen) to be showed in the <br> browser; when NULL (as per default), user, date, and machine information |
| is used. |  |
| main | a title string to be used in the browser; when NULL (as per default) a title <br> is constructed. |
| debugMe | logical switch; if true, some diagnostic output is produced. |

## Details

Very experimental code. Only allows one level of recursion into object structures. The HTML version is not dynamic.

It can be generalized. See sources ('.../library/base/R/databrowser.R') for details.
wsbrowser() is currently just an internally used function; its argument list will certainly change.

Most probably, this should rather work through using the 'tkWidget' package (from www. Bioconductor.org) but the 'tcltk' package is not yet available for the Carbon version for MacOS.

## See Also

```
str, ls.
```


## Examples

```
if(interactive()) {
    ## create some interesting objects :
    ofa <- ordered(4:1)
    ex1 <- expression(1+ 0:9)
    ex3 <- expression(u,v, 1+ 0:9)
    example(factor, echo = FALSE)
    example(table, echo = FALSE)
    example(ftable, echo = FALSE)
    example(lm, echo = FALSE)
    example(str, echo = FALSE)
    ## and browse them:
    browseEnv()
    ## a (simple) function's environment:
    af12 <- approxfun(1:2, 1:2, method = "const")
    browseEnv(envir = environment(af12))
}
```

browser Environment Browser

## Description

Interrupt the execution of an expression and allow the inspection of the environment where browser was called from.

## Usage

browser ()

## Details

A call to browser causes a pause in the execution of the current expression and runs a copy of the R interpreter which has access to variables local to the environment where the call took place.

Local variables can be listed with ls, and manipulated with $R$ expressions typed to this sub-interpreter. The interpreter copy is exited by typing c. Execution then resumes at the statement following the call to browser.

Typing n causes the step-through-debugger, to start and it is possible to step through the remainder of the function one line at a time.

Typing $Q$ quits the current execution and returns you to the top-level prompt.

## See Also

debug, and traceback for the stack on error.

```
browseURL Load URL into a WWW Browser
```


## Description

Load a given URL into a WWW browser.

## Usage

browseURL(url, browser = getOption("browser"))

## Arguments

url a non-empty character string giving the URL to be loaded.
browser a non-empty character string giving the name of the program to be used as hypertext browser. It should be in the PATH, or a full path specified.

## Details

If browser supports remote control and R knows how to perform it, the URL is opened in any already running browser or a new one if necessary. This mechanism currently is available for browsers which support the "-remote openURL(...)" interface (which includes Netscape 4.x, 6.2.x (but not $6.0 / 1$ ), Opera $5 / 6$ and Mozilla $>=0.9 .5$ ), Galeon, KDE konqueror (via kfmclient) and the GNOME interface to Mozilla. Netscape 7.0 behaves slightly differently, and you will need to open it first.

Because "-remote" will use any browser displaying on the X server (whatever machine it is running on), the remote control mechanism is only used if DISPLAY points to the local host. This may not allow displaying more than one URL at a time.

## Description

Invokes an editor to write a bug report and optionally mail it to the automated r-bugs repository at 〈r-bugs@r-project.org〉. Some standard information on the current version and configuration of R are included automatically.

## Usage

```
bug.report(subject = "", ccaddress = Sys.getenv("USER"),
    method = getOption("mailer"), address = "r-bugs@r-project.org",
    file = "R.bug.report")
```


## Arguments

| subject | Subject of the email. Please do not use single quotes (') in the subject! <br> File separate bug reports for multiple bugs |
| :--- | :--- |
| ccaddress | Optional email address for copies (default is current user). Use ccaddress <br> = FALSE for no copies. |
| method | Submission method, one of "mailx", "gnudoit", "none", or "ess". <br> address |
| file | Recipient's email address. |
|  | File to use for setting up the email (or storing it when method is "none" <br> or sending mail fails). |

## Details

Currently direct submission of bug reports works only on Unix systems. If the submission method is "mailx", then the default editor is used to write the bug report. Which editor is used can be controlled using options, type getOption("editor") to see what editor is currently defined. Please use the help pages of the respective editor for details of usage. After saving the bug report (in the temporary file opened) and exiting the editor the report is mailed using a Unix command line mail utility such as mailx. A copy of the mail is sent to the current user.

If method is "gnudoit", then an emacs mail buffer is opened and used for sending the email.

If method is "none" or NULL (which is the default on Windows systems), then only an editor is opened to help writing the bug report. The report can then be copied to your favorite email program and be sent to the r-bugs list.

If method is "ess" the body of the mail is simply sent to stdout.

## Value

Nothing useful.

## When is there a bug?

If $R$ executes an illegal instruction, or dies with an operating system error message that indicates a problem in the program (as opposed to something like "disk full"), then it is certainly a bug.
Taking forever to complete a command can be a bug, but you must make certain that it was really R's fault. Some commands simply take a long time. If the input was such that you KNOW it should have been processed quickly, report a bug. If you don't know whether the command should take a long time, find out by looking in the manual or by asking for assistance.
If a command you are familiar with causes an $R$ error message in a case where its usual definition ought to be reasonable, it is probably a bug. If a command does the wrong thing, that is a bug. But be sure you know for certain what it ought to have done. If you aren't familiar with the command, or don't know for certain how the command is supposed to work, then it might actually be working right. Rather than jumping to conclusions, show the problem to someone who knows for certain.

Finally, a command's intended definition may not be best for statistical analysis. This is a very important sort of problem, but it is also a matter of judgment. Also, it is easy to come to such a conclusion out of ignorance of some of the existing features. It is probably best not to complain about such a problem until you have checked the documentation in the usual ways, feel confident that you understand it, and know for certain that what you want is not available. The mailing list $\mathbf{r}$-devel@r-project.org is a better place for discussions of this sort than the bug list.

If you are not sure what the command is supposed to do after a careful reading of the manual this indicates a bug in the manual. The manual's job is to make everything clear. It is just as important to report documentation bugs as program bugs.

If the online argument list of a function disagrees with the manual, one of them must be wrong, so report the bug.

## How to report a bug

When you decide that there is a bug, it is important to report it and to report it in a way which is useful. What is most useful is an exact description of what commands you type, from when you start R until the problem happens. Always include the version of R, machine, and operating system that you are using; type 'version' in $R$ to print this. To help us keep track of which bugs have been fixed and which are still open please send a separate report for each bug.

The most important principle in reporting a bug is to report FACTS, not hypotheses or categorizations. It is always easier to report the facts, but people seem to prefer to strain to posit explanations and report them instead. If the explanations are based on guesses about how R is implemented, they will be useless; we will have to try to figure out what the facts must have been to lead to such speculations. Sometimes this is impossible. But in any case, it is unnecessary work for us.

For example, suppose that on a data set which you know to be quite large the command data.frame(x, y, z, monday, tuesday) never returns. Do not report that data.frame() fails for large data sets. Perhaps it fails when a variable name is a day of the week. If this is so then when we got your report we would try out the data.frame() command on a large data set, probably with no day of the week variable name, and not see any problem. There is no way in the world that we could guess that we should try a day of the week variable name.

Or perhaps the command fails because the last command you used was a [ method that had a bug causing R's internal data structures to be corrupted and making the data.frame() command fail from then on. This is why we need to know what other commands you have typed (or read from your startup file).

It is very useful to try and find simple examples that produce apparently the same bug, and somewhat useful to find simple examples that might be expected to produce the bug but actually do not. If you want to debug the problem and find exactly what caused it, that is wonderful. You should still report the facts as well as any explanations or solutions.

Invoking R with the --vanilla option may help in isolating a bug. This ensures that the site profile and saved data files are not read.

On some systems a bug report can be generated using the bug.report() function. This automatically includes the version information and sends the bug to the correct address. Alternatively the bug report can be emailed to 〈r-bugs@r-project.org〉 or submitted to the Web page at http://bugs.r-project.org.

## Author(s)

This help page is adapted from the Emacs manual

## See Also

R FAQ
builtins Returns the names of all built-in objects

## Description

Return the names of all the built-in objects. These are fetched directly from the symbol table of the R interpreter.

## Usage

builtins(internal = FALSE)

## Arguments

internal a logical indicating whether only "internal" functions (which can be called via . Internal) should be returned.
bxp Box Plots from Summaries

## Description

bxp draws box plots based on the given summaries in $z$. It is usually called from within boxplot, but can be invoked directly.

```
Usage
    bxp(z, notch = FALSE, width = NULL, varwidth = FALSE, outline = TRUE,
        notch.frac = 0.5, boxwex = 0.8, border = par("fg"), col = NULL,
        log = "", pars = NULL, frame.plot = axes, horizontal = FALSE,
        add = FALSE, at = NULL, show.names=NULL, ...)
```


## Arguments

z
notch
width a vector giving the relative widths of the boxes making up the plot.
varwidth if varwidth is TRUE, the boxes are drawn with widths proportional to the square-roots of the number of observations in the groups.
outline if outline is not true, the boxplot lines are not drawn.
boxwex a scale factor to be applied to all boxes. When there are only a few groups, the appearance of the plot can be improved by making the boxes narrower.
notch.frac numeric in $(0,1)$. When notch=TRUE, the fraction of the box width that the notches should use.
border character, the color of the box borders. Is recycled for multiple boxes.
col character; the color within the box. Is recycled for multiple boxes
log character, indicating if any axis should be drawn in logarithmic scale, as in plot. default.
frame.plot logical, indicating if a "frame" (box) should be drawn; defaults to TRUE, unless axes = FALSE is specified.
horizontal logical indicating if the boxplots should be horizontal; default FALSE means vertical boxes.
add logical, if true $a d d$ boxplot to current plot.
at numeric vector giving the locations where the boxplots should be drawn, particularly when add $=$ TRUE; defaults to $1: \mathrm{n}$ where n is the number of boxes.
show. names Set to TRUE or FALSE to override the defaults on whether an x-axis label is printed for each group.
pars,... Graphical parameters can be passed as arguments to this function, either as a list (pars) or normally (. . .).

## Value

An invisible vector, actually identical to the at argument, with the coordinates ("x" if horizontal is false, "y" otherwise) of box centers, useful for adding to the plot.

## Examples

```
set.seed(753)
str(bx.p <- boxplot(split(rt(100, 4), gl(5,20))))
op <- par(mfrow= c(2,2))
bxp(bx.p, xaxt = "n")
bxp(bx.p, notch = TRUE, axes = FALSE, pch = 4)
bxp(bx.p, notch = TRUE, col= "lightblue", frame= FALSE, outl= FALSE,
    main = "bxp(*, frame= FALSE, outl= FALSE)")
bxp(bx.p, notch = TRUE, col= "lightblue", border="red", ylim = c(-4,4),
    pch = 22, bg = "green", log = "x", main = "... log='x', ylim=*")
par(op)
op <- par(mfrow= c(1,2))
data(PlantGrowth)
## single group -- no label
boxplot(weight~ group,data=PlantGrowth, subset=group=="ctrl")
bx<-boxplot(weight~group,data=PlantGrowth,subset=group=="ctrl",plot=FALSE)
## with label
bxp(bx,show.names=TRUE)
par(op)
```

by

Apply a Function to a Data Frame split by Factors

## Description

Function by is an object-oriented wrapper for tapply applied to data frames.

## Usage

by(data, INDICES, FUN, ...)

## Arguments

| data | an $R$ object, normally a data frame, possibly a matrix. |
| :--- | :--- |
| INDICES | a factor or a list of factors, each of length nrow $(x)$. |
| FUN | a function to be applied to data frame subsets of x. |
| $\ldots$ | further arguments to FUN. |

## Details

A data frame is split by row into data frames subsetted by the values of one or more factors, and function FUN is applied to each subset in term.
Object data will be coerced to a data frame by default.

## Value

A list of class "by", giving the results for each subset.

## See Also

```
tapply
```


## Examples

```
data(warpbreaks)
attach(warpbreaks)
by(warpbreaks[, 1:2], tension, summary)
by(warpbreaks[, 1], list(wool=wool, tension=tension), summary)
by(warpbreaks, tension, function(x) lm(breaks ~ wool, data=x))
## now suppose we want to extract the coefficients by group
tmp <- by(warpbreaks, tension, function(x) lm(breaks ~ wool, data=x))
sapply(tmp, coef)
detach("warpbreaks")
```

C
Sets Contrasts for a Factor

## Description

Sets the "contrasts" attribute for the factor.

## Usage

C(object, contr, how.many, ...)

## Arguments

| object | a factor or ordered factor |
| :--- | :--- |
| contr | which contrasts to use. Can be a matrix with one row for each level of the <br> factor or a suitable function like contr. poly or a character string giving <br> the name of the function |
| how.many | the number of contrasts to set, by default one less than nlevels (object). <br> $\ldots$. |
|  | Additional arguments for the function contr. |

## Details

For compatibility with S, contr can be treatment, helmert, sum or poly (without quotes) as shorthand for contr.treatment and so on.

## Value

The factor object with the "contrasts" attribute set.

## Author(s)

B.D. Ripley

## See Also

contrasts, contr.sum, etc.

## Examples

```
## reset contrasts to defaults
options(contrasts=c("contr.treatment", "contr.poly"))
data(warpbreaks)
attach(warpbreaks)
tens <- C(tension, poly, 1)
attributes(tens)
detach()
## tension SHOULD be an ordered factor, but as it is not we can use
aov(breaks ~ wool + tens + tension, data=warpbreaks)
## show the use of ... The default contrast is contr.treatment here
summary(lm(breaks ~ wool + C(tension, base=2), data=warpbreaks))
data(esoph) # following on from help(esoph)
model3 <- glm(cbind(ncases, ncontrols) ~ agegp + C(tobgp, , 1) +
    C(alcgp, , 1), data = esoph, family = binomial())
summary(model3)
```

c
Combine Values into a Vector or List

## Description

This is a generic function which combines its arguments.
The default method combines its arguments to form a vector. All arguments are coerced to a common type which is the type of the returned value.

## Usage

$c(. . ., r e c u r s i v e=F A L S E)$

## Arguments

... objects to be concatenated.
recursive logical. If recursive=TRUE, the function recursively descends through lists combining all their elements into a vector.

See Also
unlist and as.vector to produce attribute-free vectors.

## Examples

```
c(1,7:9)
c(1:5, 10.5, "next")
## append to a list:
ll <- list(A = 1, c="C")
## do *not*
c(ll, d = 1:3) # which is == c(ll, as.list (c(d=1:3))
## but rather
```

```
c(ll, d = list(1:3))\# c() combining two lists
    \(c(\) list \((A=c(B=1))\), recursive=TRUE)
    c(options(), recursive=TRUE)
    \(c(\) list \((A=c(B=1, C=2), B=c(E=7))\), recursive=TRUE)
```

    call Function Calls
    
## Description

Create or test for objects of mode "call".

## Usage

```
call(name, ...)
is.call(x)
as.call(x)
```


## Arguments

```
name a character string naming the function to be called.
... arguments to be part of the call.
x an arbitrary R object.
```


## Value

call returns an unevaluated function call, that is, an unevaluated expression which consists of the named function applied to the given arguments (name must be a quoted string which gives the name of a function to be called).
is.call is used to determine whether $x$ is a call (i.e., of mode "call").
Objects of mode "list" can be coerced to mode "call". The first element of the list becomes the function part of the call, so should be a function or the name of one (as a symbol; a quoted string will not do).

## See Also

do.call for calling a function by name and argument list; Recall for recursive calling of functions; further is.language, expression, function.

## Examples

```
is.call(call) #-> FALSE: Functions are NOT calls
# set up a function call to round with argument 10.5
cl <- call("round", 10.5)
is.call(cl)# TRUE
cl
# such a call can also be evaluated.
eval(cl)# [1] 10
```

capabilities Report Capabilities of this Build of $R$

## Description

Report on the optional features which have been compiled into this build of $R$.

## Usage

```
capabilities(what = NULL)
```


## Arguments

what character vector or NULL, specifying required components. NULL implies that all are required.

## Value

A named logical vector. Current components are
jpeg Is the jpeg function operational?
png Is the png function operational?
tcltk Is the tcltk package operational?
X11 (Unix) Are X11 and the data editor available?
GNOME (Unix) Is the GNOME GUI in use and are GTK and GNOME graphics devices available?
libz Is gzfile available? From R 1.5.0 this will always be true.
http/ftp Are url and the internal method for download.file available?
sockets Are make.socket and related functions available?
libxml Is there support for integrating libxml with the $R$ event loop?
cledit Is command-line editing available in the current R session? This is false in non-interactive sessions.
IEEE754 Does this platform have IEEE 754 arithmetic? Note that this is more correctly known by the international standard IEC 60559.
bzip2 Is bzfile available?
PCRE Is the Perl-Compatible Regular Expression library available? This is needed for the perl $=$ TRUE option to grep are related function.

## See Also

```
    .Platform
```


## Examples

```
capabilities()
if(!capabilities("http/ftp"))
    warning("internal download.file() is not available")
## See also the examples for 'connections'.
```


## cars Stopping Distances of Cars

## Description

The data give the speed of cars and the distances taken to stop. Note that the data were recorded in the 1920s.

## Usage

data(cars)

## Format

A data frame with 50 observations on 2 variables.

| $[, 1]$ | speed | numeric | Speed (mph) |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | dist | numeric | Stopping distance (ft) |

## Source

Ezekiel, M. (1930) Methods of Correlation Analysis. Wiley.

## References

McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(cars)
plot(cars, xlab = "Speed (mph)", ylab = "Stopping distance (ft)",
    las = 1)
lines(lowess(cars$speed, cars$dist, f = 2/3, iter = 3), col = "red")
title(main = "cars data")
plot(cars, xlab = "Speed (mph)", ylab = "Stopping distance (ft)",
    las = 1, log = "xy")
title(main = "cars data (logarithmic scales)")
lines(lowess(cars$speed, cars$dist, f = 2/3, iter = 3), col = "red")
summary(fm1 <- lm(log(dist) ~ log(speed), data = cars))
opar <- par(mfrow = c(2, 2), oma = c(0, 0, 1.1, 0),
            mar = c(4.1, 4.1, 2.1, 1.1))
plot(fm1)
par(opar)
## An example of polynomial regression
plot(cars, xlab = "Speed (mph)", ylab = "Stopping distance (ft)",
    las = 1, xlim = c(0, 25))
d <- seq(0, 25, len = 200)
for(degree in 1:4) {
    fm <- lm(dist ~ poly(speed, degree), data = cars)
    assign(paste("cars", degree, sep="."), fm)
    lines(d, predict(fm, data.frame(speed=d)), col = degree)
}
anova(cars.1, cars.2, cars.3, cars.4)
```


## Description

Simple utilities returning (non-missing) case names, and (non-eliminated) variable names.

## Usage

```
case.names(object, ...)
case.names.lm(object, full = FALSE, ...)
variable.names(object, ...)
variable.names.lm(object, full = FALSE, ...)
```


## Arguments

```
object an R object, typically a fitted model.
full logical; if TRUE, all names (including zero weights,...) are returned.
... further arguments passed to or from other methods.
```


## Value

A character vector.

## See Also

$$
\mathrm{lm}
$$

## Examples

```
x <- 1:20
y <- x + (x/4 - 2) ^3 + rnorm(20, s=3)
names(y) <- paste("0",x,sep=".")
ww <- rep(1,20); ww[13] <- 0
summary(lmxy <- lm(y ~ x + I (x^2)+I(x^3) + I((x-10)^2),
                            weights = ww), cor = TRUE)
variable.names(lmxy)
variable.names(lmxy, full= TRUE)# includes the last
case.names(lmxy)
case.names(lmxy, full = TRUE)# includes the 0-weight case
```

cat Concatenate and Print

## Description

Prints the arguments, coercing them if necessary to character mode first.

## Usage

```
cat(... , file = "", sep = " ", fill = FALSE, labels = NULL,
    append = FALSE)
```


## Arguments

| $\ldots$. | R objects which are coerced to character strings, concatenated, and <br> printed, with the remaining arguments controlling the output. |
| :--- | :--- |
| file | A connection, or a character string naming the file to print to. If "" (the <br> default), cat prints to the standard output connection, the console unless <br> redirected by sink. If it is " $\mid$ cmd", the output is piped to the command <br> given by 'cmd', by opening a pipe connection. |
| sep | character string to insert between the objects to print. |
| fill | a logical or numeric controlling how the output is broken into successive <br> lines. If FALSE (default), only newlines created explicitly by <br> n are printed. Otherwise, the output is broken into lines with print width <br> equal to the option width if fill is TRUE, or the value of fill if this is <br> numeric. |
| nabels | character vector of labels for the lines printed. Ignored if fill is FALSE. |
| append | logical. Only used if the argument file is the name of file (and not a con- <br> nection or " I cmd"). If TRUE output will be appended to file; otherwise, <br> it will overwrite the contents of file. |

## Details

cat converts its arguments to character strings, concatenates them, separating them by the given sep= string, and then prints them.
No linefeeds are printed unless explicitly requested by " n " or if generated by filling (if argument fill is TRUE or numeric.)
cat is useful for producing output in user-defined functions.

## Value

None (invisible NULL).

## See Also

print, format, and paste which concatenates into a string.

## Examples

```
## print an informative message
cat("iteration = ", iter <- iter + 1, "\n")
## 'fill' and label lines:
cat(paste(letters, 100* 1:26), fill = TRUE,
    labels = paste("{",1:10,"}:",sep=""))
```

Cauchy The Cauchy Distribution

## Description

Density, distribution function, quantile function and random generation for the Cauchy distribution with location parameter location and scale parameter scale.

## Usage

```
dcauchy(x, location = 0, scale = 1, log = FALSE)
pcauchy(q, location = 0, scale = 1, lower.tail = TRUE, log.p = FALSE)
qcauchy(p, location = 0, scale = 1, lower.tail = TRUE, log.p = FALSE)
rcauchy(n, location = 0, scale = 1)
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n number of observations. If length (n) > 1, the length is taken to be the number required.
location, scale
location and scale parameters.
$\log , \log \cdot \mathrm{p} \quad \operatorname{logical}$; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

If location or scale are not specified, they assume the default values of 0 and 1 respectively.
The Cauchy distribution with location $l$ and scale $s$ has density

$$
f(x)=\frac{1}{\pi s}\left(1+\left(\frac{x-l}{s}\right)^{2}\right)^{-1}
$$

for all $x$.

## Value

dcauchy, pcauchy, and qcauchy are respectively the density, distribution function and quantile function of the Cauchy distribution. rcauchy generates random deviates from the Cauchy.

## See Also

$d t$ for the t distribution which generalizes dcauchy $(*, l=0, \mathrm{~s}=1)$.

## Examples

```
all.equal(dcauchy(-1:4), 1 / (pi*(1 + (-1:4) ^2)))
```

cbind $\quad$ Combine $R$ Objects by Rows or Columns

## Description

Take a sequence of vector, matrix or data frames arguments and combine by columns or rows, respectively. There may be methods for other R classes.

## Usage

cbind(..., deparse.level = 1)
rbind(..., deparse.level = 1)

## Arguments

... vectors or matrices. These can be given as named arguments.
deparse.level integer controlling the construction of labels; currently, 1 is the only possible value.

## Details

The functions cbind and rbind are generic, with methods for data frames. The data frame method will be used if an argument is a data frame and the rest are vectors or matrices. There can be other methods; in particular, there is one for time series objects.
If there are several matrix arguments, they must all have the same number of columns (or rows) and this will be the number of columns (or rows) of the result. If all the arguments are vectors, the number of columns (rows) in the result is equal to the length of the longest vector. Values in shorter arguments are recycled to achieve this length (with a warning if they are recycled only fractionally).
When the arguments consist of a mix of matrices and vectors the number of columns (rows) of the result is determined by the number of columns (rows) of the matrix arguments. Any vectors have their values recycled or subsetted to achieve this length.
For cbind (rbind), vectors of zero length are ignored unless the result would have zero rows (columns), for $S$ compatibility. (Zero-extent matrices do not occur in $S$ and are not ignored in R.)

## Value

A matrix or data frame combining the . . . arguments column-wise or row-wise.
For cbind (rbind) the column (row) names are taken from the names of the arguments, or where those are not supplied by deparsing the expressions given (if that gives a sensible name). The names will depend on whether data frames are included: see the examples.

## Note

The method dispatching is not done via UseMethod(), but by C-internal dispatching. Therefore, there is no need for, e.g., rbind.default.
The dispatch algorithm is described in the source file ( ${ }^{〔} . . / \mathrm{src} /$ main/bind.c') as

1. For each argument we get the list of possible class memberships from the class attribute.
2. We inspect each class in turn to see if there is an an applicable method.
3. If we find an applicable method we make sure that it is identical to any method determined for prior arguments. If it is identical, we proceed, otherwise we immediately drop through to the default code.

If you want to combine other objects with data frames, it may be necessary to coerce them to data frames first.

## See Also

c to combine vectors (and lists) as vectors, data.frame to combine vectors and matrices as a data frame.

## Examples

```
cbind(1, 1:7) # the '1' (= shorter vector) is recycled
cbind(1:7, diag(3))# vector is subset -> warning
cbind(0, rbind(1, 1:3))
cbind(I=0, X=rbind(a=1, b=1:3)) # use some names
xx <- data.frame(I=rep (0,2))
cbind(xx, X=rbind(a=1, b=1:3)) # named differently
cbind(0, matrix(1, nrow=0, ncol=4))#> Warning (making sense)
dim(cbind(0, matrix(1, nrow=2, ncol=0)))#-> 2 x 1
```

char.expand
Expand a String with Respect to a Target Table

## Description

Seeks a unique match of its first argument among the elements of its second. If successful, it returns this element; otherwise, it performs an action specified by the third argument.

## Usage

```
char.expand(input, target, nomatch = stop("no match"))
```


## Arguments

input a character string to be expanded.
target a character vector with the values to be matched against.
nomatch an $R$ expression to be evaluated in case expansion was not possible.

## Details

This function is particularly useful when abbreviations are allowed in function arguments, and need to be uniquely expanded with respect to a target table of possible values.

## See Also

charmatch and pmatch for performing partial string matching.

## Examples

```
locPars <- c("mean", "median", "mode")
char.expand("me", locPars, warning("Could not expand!"))
char.expand("mo", locPars)
```

```
character Character Vectors
```


## Description

Create or test for objects of type "character".

## Usage

```
character(length = 0)
as.character(x, ...)
is.character(x)
```


## Arguments

| length | desired length. |
| :--- | :--- |
| x | object to be coerced or tested. |
| $\ldots$ | further arguments passed to or from other methods. |

## Value

character creates a character vector of the specified length. The elements of the vector are all equal to " ".
as.character attempts to coerce its argument to character type.
is.character returns TRUE or FALSE depending on whether its argument is of character type or not.

## Note

as. character truncates components of language objects to 500 characters (was about 70 before 1.3.1).

## See Also

paste, substr and strsplit for character concatenation and splitting, chartr for character translation and casefolding (e.g. upper to lower case) and sub, grep etc for string matching and substitutions. Note that help.search (keyword = "character") gives even more links. deparse, which is normally preferable to as.character for language objects.

## Examples

```
form <- y ~ a + b + c
as.character(form) ## length 3
deparse(form) ## like the input
```

```
charmatch
Partial String Matching
```


## Description

charmatch seeks matches for the elements of its first argument among those of its second.

## Usage

charmatch (x, table, nomatch = NA)

## Arguments

| x | the values to be matched. |
| :--- | :--- |
| table | the values to be matched against. |
| nomatch | the value returned at non-matching positions. |

## Details

Exact matches are preferred to partial matches (those where the value to be matched has an exact match to the initial part of the target, but the target is longer).

If there is a single exact match or no exact match and a unique partial match then the index of the matching value is returned; if multiple exact or multiple partial matches are found then 0 is returned and if no match is found then NA is returned.

## Author(s)

This function is based on a C function written by Terry Therneau.

## See Also

```
pmatch, match.
```

grep or regexpr for more general (regexp) matching of strings.

## Examples

```
charmatch("", "") # returns 1
charmatch("m", c("mean", "median", "mode")) # returns 0
charmatch("med", c("mean", "median", "mode")) # returns 2
```

chartr
Character Translation and Casefolding

## Description

Translate characters in character vectors, in particular from upper to lower case or vice versa.

## Usage

```
chartr(old, new, x)
tolower(x)
toupper(x)
casefold(x, upper = FALSE)
```


## Arguments

$\mathrm{x} \quad \mathrm{a}$ character vector.
old a character string specifying the characters to be translated.
new a character string specifying the translations.
upper logical: translate to upper or lower case?.

## Details

chartr translates each character in x that is specified in old to the corresponding character specified in new. Ranges are supported in the specifications, but character classes and repreated characters are not. If old contains more characters than new, an error is signaled; if it contains fewer characters, the extra characters at the end of new are ignored.
tolower and toupper convert upper-case characters in a character vector to lower-case, or vice versa. Non-alphabetic characters are left unchanged.
casefold is a wrapper for tolower and toupper provided for compatibility with S-PLUS.

## See Also

sub and gsub for other substitutions in strings.

## Examples

```
x <- "MiXeD cAsE 123"
chartr("iXs", "why", x)
chartr("a-cX", "D-Fw", x)
tolower(x)
toupper(x)
```

check.options Set Options with Consistency Checks

## Description

Utility function for setting options with some consistency checks. The attributes of the new settings in new are checked for consistency with the model (often default) list in name.opt.

## Usage

```
check.options(new, name.opt, reset = FALSE, assign.opt = FALSE,
    envir = .GlobalEnv, check.attributes = c("mode", "length"),
    override.check = FALSE)
```


## Arguments

| new | a named list |
| :--- | :--- |
| name.opt | character with the name of R object containing the "model" (default) list. |
| reset | logical; if TRUE, reset the options from name. opt. If there is more than <br> one R object with name name.opt, remove the first one in the search() <br> path. |
| assign.opt | logical; if TRUE, assign the $\ldots$ <br> envir <br> check.attributes environment used for get and assign. |
| character containing the attributes which check. options should check. |  |

## Value

A list of components with the same names as the one called name.opt. The values of the components are changed from the new list, as long as these pass the checks (when these are not overridden according to override. check).

## Author(s)

Martin Maechler

## See Also

ps.options which uses check.options.

## Examples

```
L1 <- list(a=1:3, b=pi, ch="CH")
str(L2 <- check.options(list(a=0:2), name.opt = "L1"))
str(check.options(NULL, reset = TRUE, name.opt = "L1"))
```

chickwts Chicken Weights by Feed Type

## Description

An experiment was conducted to measure and compare the effectiveness of various feed supplements on the growth rate of chickens.

## Usage

data(chickwts)

## Format

A data frame with 71 observations on 2 variables.
weight a numeric variable giving the chick weight.
feed a factor giving the feed type.

## Details

Newly hatched chicks were randomly allocated into six groups, and each group was given a different feed supplement. Their weights in grams after six weeks are given along with feed types.

## Source

Anonymous (1948) Biometrika, 35, p. 214 .

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(chickwts)
boxplot(weight ~ feed, data = chickwts, col = "lightgray",
    varwidth = TRUE, notch = TRUE, main = "chickwt data",
    ylab = "Weight at six weeks (gm)")
anova(fm1 <- lm(weight ~ feed, data = chickwts))
opar <- par(mfrow = c(2, 2), oma = c(0, 0, 1.1, 0),
    mar = c(4.1, 4.1, 2.1, 1.1))
plot(fm1)
par(opar)
```

Chisquare The (non-central) Chi-Squared Distribution

## Description

Density, distribution function, quantile function and random generation for the chi-squared $\left(\chi^{2}\right)$ distribution with df degrees of freedom and optional non-centrality parameter ncp.

## Usage

```
dchisq(x, df, ncp=0, log = FALSE)
pchisq(q, df, ncp=0, lower.tail = TRUE, log.p = FALSE)
qchisq(p, df, ncp=0, lower.tail = TRUE, log.p = FALSE)
rchisq(n, df, ncp=0)
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the number required.
df degrees of freedom (positive, but can be non-integer).
ncp non-centrality parameter. For rchisq, $n c p=0$ is the only possible value.
$\log$, log.p logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The chi-squared distribution with $\mathrm{df}=n$ degrees of freedom has density

$$
f_{n}(x)=\frac{1}{2^{n / 2} \Gamma(n / 2)} x^{n / 2-1} e^{-x / 2}
$$

for $x>0$. The mean and variance are $n$ and $2 n$.
The non-central chi-squared distribution with $\mathrm{df}=n$ degrees of freedom and non-centrality parameter ncp $=\lambda$ has density

$$
f(x)=e^{-\lambda / 2} \sum_{r=0}^{\infty} \frac{(\lambda / 2)^{r}}{r!} f_{n+2 r}(x)
$$

for $x \geq 0$. It is the distribution of the sum of squares of $n$ normals each with variance one, $\lambda$ being the sum of squares of the normal means.

## Value

dchisq gives the density, pchisq gives the distribution function, qchisq gives the quantile function, and rchisq generates random deviates.

## See Also

dgamma for the Gamma distribution which generalizes the chi-squared one.

## Examples

```
dchisq(1, df=1:3)
pchisq(1, df= 3)
pchisq(1, df= 3, ncp = 0:4)# includes the above
x <- 1:10
## Chi-squared(df = 2) is a special exponential distribution
all.equal(dchisq(x, df=2), dexp(x, 1/2))
all.equal(pchisq(x, df=2), pexp(x, 1/2))
```

```
chol The Choleski Decomposition
```


## Description

Compute the Choleski factorization of a real symmetric positive-definite square matrix.

## Usage

```
chol(x, pivot = FALSE)
```

La.chol(x)

## Arguments

$\mathrm{x} \quad$ a real symmetric, positive-definite matrix
pivot Should pivoting be used?

## Details

chol provides an interface to the LINPACK routine DCHDC. La. chol provides an interface to the LAPACK routine DPOTRF.

Note that only the upper triangular part of x is used, so that $R^{\prime} R=x$ when x is symmetric.
If pivot = FALSE and x is not non-negative definite an error occurs. If x is positive semidefinite (i.e. some zero eigenvalues) an error will also occur, as a numerical tolerance is used.
If pivot $=$ TRUE, then the Choleski decomposition of a positive semi-definite x can be computed. The rank of $x$ is returned as attr ( $Q$, "rank"), subject to numerical errors. The pivot is returned as attr(Q, "pivot"). It is no longer the case that $t(Q) \% * \%$ Q equals x. However, setting pivot <- attr(Q, "pivot") and oo <- order (pivot), it is true that $t(Q[, \infty]) \% * \% ~ Q[, \infty]$ equals $x$, or, alternatively, $t(Q) \% * \% Q$ equals $x[p i v o t$, pivot]. See the examples.

## Value

The upper triangular factor of the Choleski decomposition, i.e., the matrix $R$ such that $R^{\prime} R=x$ (see example).
If pivoting is used, then two additional attributes "pivot" and "rank" are also returned.

## Warning

The code does not check for symmetry
If pivot $=$ TRUE and $x$ is not non-negative definite then there will be no error message but a meaningless result will occur. So only use pivot $=$ TRUE when $x$ is non-negative definite by construction.

## References

Dongarra, J. J., Bunch, J. R., Moler, C. B. and Stewart, G. W. (1978) LINPACK Users Guide. Philadelphia: SIAM Publications.
Anderson. E. and ten others (1999) LAPACK Users' Guide. Third Edition. SIAM.
Available on-line at http://www.netlib.org/lapack/lug/lapack_lug.html.

## See Also

chol2inv for its inverse (without pivoting), backsolve for solving linear systems with upper triangular left sides. qr, svd for related matrix factorizations.

## Examples

```
( m <- matrix(c(5,1,1,3),2,2) )
( cm <- chol(m) )
t(cm) %*% cm #-- = 'm'
all(abs(m - t(cm) %*% cm) < 100* .Machine$double.eps) # TRUE
( Lcm <- La.chol(m) )
crossprod(Lcm)
all(abs(m - crossprod(Lcm)) < 100* .Machine$double.eps) # TRUE
x <- matrix(c(1:5, (1:5)^2), 5, 2)
m <- crossprod(x)
Q <- chol(m)
all.equal(t(Q) %*% Q, m) # TRUE
Q <- chol(m, pivot = TRUE)
pivot <- attr(Q, "pivot")
oo <- order(pivot)
all.equal(t(Q[, o০]) %*% Q[, oo], m) # TRUE
all.equal(t(Q) %*% Q, m[pivot, pivot]) # TRUE
# now for something positive semi-definite
x <- cbind(x, x[, 1]+3*x[, 2])
m <- crossprod(x)
qr(m)$rank # is 2, as it should be
# chol() may fail, depending on numerical rounding:
# chol() unlike qr() does not use a tolerance.
test <- try(Q <- chol(m))
(Q <- chol(m, pivot = TRUE)) # NB wrong rank here ... see Warning section.
pivot <- attr(Q, "pivot")
\circo <- order(pivot)
all.equal(t(Q[, oo]) %*% Q[, oo], m) # TRUE
all.equal(t(Q) %*% Q, m[pivot, pivot]) # TRUE
```


## Description

Invert a symmetric, positive definite square matrix from its Choleski decomposition.

## Usage

chol2inv( $x$, size $=n c o l(x))$
La.chol2inv(x, size $=n c o l(x))$

## Arguments

$\mathrm{x} \quad$ a matrix. The first nc columns of the upper triangle contain the Choleski decomposition of the matrix to be inverted.
size the number of columns of x containing the Choleski decomposition.

## Details

chol provides an interface to the LINPACK routine DPODI. La.chol provides an interface to the LAPACK routine DPOTRI.

## Value

The inverse of the decomposed matrix.

## References

Dongarra, J. J., Bunch, J. R., Moler, C. B. and Stewart, G. W. (1978) LINPACK Users Guide. Philadelphia: SIAM Publications.

Anderson. E. and ten others (1999) LAPACK Users' Guide. Third Edition. SIAM. Available on-line at http://www.netlib.org/lapack/lug/lapack_lug.html.

## See Also

chol, solve.

## Examples

```
cma <- chol(ma <- cbind(1, 1:3, c(1,3,7)))
t(cma) %*% cma # = ma
all.equal(diag(3), ma %*% chol2inv(cma))
all.equal(diag(3), ma %*% La.chol2inv(cma))
```

chull Compute Convex Hull of a Set of Points

## Description

Computes the subset of points which lie on the convex hull of the set of points specified.

## Usage

chull $(x, y=N U L L)$

## Arguments

$\mathrm{x}, \mathrm{y} \quad$ coordinate vectors of points. This can be specified as two vectors x and y , a 2-column matrix x , a list x with components x and y

## Details

xy. coords is used to interpret the specification of the points. The algorithm is that given by Eddy (1977).
'Peeling' as used in the S function chull can be implemented by calling chull recursively.

## Value

An integer vector giving the indices of the points lying on the convex hull, in clockwise order.

## Author(s)

B. D. Ripley

## References

Eddy, W. F. (1977) A new convex hull algorithm for planar sets. ACM Transactions on Mathematical Software, 3, 398-403.

Eddy, W. F. (1977) Algorithm 523. CONVEX, A new convex hull algorithm for planar sets[Z]. ACM Transactions on Mathematical Software, 3, 411-412.

## See Also

xy.coords,polygon

## Examples

```
X <- matrix(rnorm(2000), ncol=2)
plot(X, cex=0.5)
hpts <- chull(X)
hpts <- c(hpts, hpts[1])
lines(X[hpts, ])
```

```
class Object Classes
```


## Description

$R$ possesses a simple generic function mechanism which can be used for an object-oriented style of programming. Method despatch takes place based on the class of the first argument to the generic function.

## Usage

class ( x )
class(x) <- names
unclass (x)
inherits(x, what, which = FALSE)

## Arguments

$x$ an objects
what a character vector naming classes.
which logical affecting return value: see Details.

## Details

An R "object" is a data object which has a class attribute. A class attribute is a character vector giving the names of the classes which the object "inherits" from. When a generic function fun is applied to an object with class attribute c("first", "second"), the system searches for a function called fun.first and, if it finds it, applies it to the object. If no such function is found, a function called fun.second is tried. If no class name produces a suitable function, the function fun.default is used.

The function class prints the vector of names of classes an object inherits from. Correspondingly, class<- sets the classes an object inherits from.
unclass returns (a copy of) its argument with its class information removed.
inherits indicates whether its first argument inherits from any of the classes specified in the what argument. If which is TRUE then an integer vector of the same length as what is returned. Each element indicates the position in the class(x) matched by the element of what; zero indicates no match. If which is FALSE then TRUE is returned by inherits if any of the names in what match with any class.

## See Also

UseMethod, NextMethod.

## Examples

```
x <- 10
inherits(x, "a") #FALSE
class(x)<-c("a", "b")
inherits(x,"a") #TRUE
inherits(x, "a", TRUE) # 1
inherits(x, c("a", "b", "c"), TRUE) # 1 2 0
```

```
    close.socket Close a Socket
```


## Description

Closes the socket and frees the space in the file descriptor table. The port may not be freed immediately.

## Usage

close.socket(socket, ...)

## Arguments

```
socket A socket object
... further arguments passed to or from other methods.
```


## Value

logical indicating success or failure

## Author(s)

Thomas Lumley

## See Also

make.socket, read.socket
co2 Mauna Loa Atmospheric CO2 Concentration

## Description

Atmospheric concentrations of $\mathrm{CO}_{2}$ are expressed in parts per million (ppm) and reported in the preliminary 1997 SIO manometric mole fraction scale.

## Usage

data(co2)

## Format

A time series of 468 observations; monthly from 1959 to 1997.

## Details

The values for February, March and April of 1964 were missing and have been obtained by interpolating linearly between the values for January and May of 1964.

## Source

Keeling, C. D. and Whorf, T. P., Scripps Institution of Oceanography (SIO), University of California, La Jolla, California USA 92093-0220.
ftp://cdiac.esd.ornl.gov/pub/maunaloa-co2/maunaloa.co2.

## References

Cleveland, W. S. (1993) Visualizing Data. New Jersey: Summit Press.

## Examples

```
data(co2)
plot(co2, ylab = expression("Atmospheric concentration of CO"[2]),
    las = 1)
title(main = "co2 data set")
```

```
codes Factor Codes
```


## Description

This (generic) function returns a numeric coding of a factor. It can also be used to assign to a factor using the coded form.

## Usage

codes (x, ...)
codes (x) <- value

## Arguments

$\mathrm{x} \quad$ an object from which to extract or set the codes.
... further arguments passed to or from other methods.

## Value

For an ordered factor, it returns the internal coding ( 1 for the lowest group, 2 for the second lowest, etc.).
For an unordered factor, an alphabetical ordering of the levels is assumed, i.e., the level that is coded 1 is the one whose name is sorted first according to the prevailing collating sequence. Warning: the sort order may well depend on the locale, and should not be assumed to be ASCII.

## Note

Normally codes is not the appropriate function to use with an unordered factor. Use unclass or as.numeric to extract the codes used in the internal representation of the factor, as these do not assume that the codes are sorted.
The behaviour for unordered factors is dubious, but compatible with $S$ version 3 . To get the internal coding of a factor, use as.integer. Note in particular that the codes may not be the same in different language locales because of collating differences.

## See Also

```
factor, levels, nlevels.
```


## Examples

```
codes(rep(factor(c(20,10)),3))
x <- gl(3,5)
codes(x)[3] <- 2
x
data(esoph)
( ag <- esoph$alcgp[12:1] )
codes(ag)
codes(factor(1:10)) # BEWARE!
```

coefficients Extract Model Coefficients

## Description

coef is a generic function which extracts model coefficients from objects returned by modeling functions. coefficients is an alias for it.

## Usage

```
coef(object, ...)
coefficients(object, ...)
```


## Arguments

object an object for which the extraction of model coefficients is meaningful.
... other arguments.

## Details

All object classes which are returned by model fitting functions should provide a coef method. (Note that the method is coef and not coefficients.)

## Value

Coefficients extracted from the model object object.

## See Also

fitted.values and residuals for related methods; glm, lm for model fitting.

## Examples

```
x <- 1:5; coef(lm(c(1:3,7,6) ~ x ))
```


## Description

Returns a matrix of integers indicating their column number in the matrix.

## Usage

$\operatorname{col}(x$, as.factor=FALSE)

## Arguments

x
a matrix.
as.factor a logical value indicating whether the value should be returned as a factor rather than as numeric.

## Value

An integer matrix with the same dimensions as $x$ and whose $i j$-th element is equal to $j$.

## See Also

row to get rows.

## Examples

```
# extract an off-diagonal of a matrix
ma <- matrix(1:12, 3, 4)
ma[row(ma) == col(ma) + 1]
# create an identity 5-by-5 matrix
x <- matrix(0, nr = 5, nc = 5)
x[row(x) == col(x)] <- 1
```

```
col2rgb
```

Color to RGB Conversion

## Description

"Any R color" to RGB (red/green/blue) conversion.

## Usage

col2rgb (col)

## Arguments

col
vector of any of the three kind of R colors, i.e., either a color name (an element of colors()), a hexadecimal string of the form "\#rrggbb", or an integer i meaning palette() [i].

## Details

For integer colors, 0 is shorthand for the current par("bg"), and NA means "nothing" which effectively does not draw the corresponding item.
For character colors, "NA" is equivalent to NA above.

## Value

an integer matrix with three rows and number of columns the length (and names if any) as col.

## Author(s)

Martin Maechler

## See Also

```
rgb, colors, palette, etc.
```


## Examples

```
col2rgb("peachpuff")
col2rgb(c(blu = "royalblue", reddish = "tomato")) # names kept
col2rgb(1:8)# the ones from the palette() :
pp <- palette(); names(pp) <- pp # add & use names :
stopifnot(col2rgb(1:8) ==
    print(col2rgb(pp)))
col2rgb(paste("gold", 1:4, sep=""))
stopifnot(col2rgb("#08a0ff") == c(8, 160, 255))
## all three kind of colors mixed :
col2rgb(c(red="red", palette= 1:3, hex="#abcdef"))
##-- NON-INTRODUCTORY examples --
grC <- col2rgb(paste("gray",0:100,sep=""))
stopifnot(grC["red",] == grC["green",],
    grC["red",] == grC["blue",],
    grC["red", 1:4] == c(0,3,5,8))
table(print(diff(grC["red",])))# '2' or '3': almost equidistant
## The 'named' grays are in between {"slate gray" is not gray, strictly}
col2rgb(c(g66="gray66", darkg= "dark gray", g67="gray67",
    g74="gray74", gray = "gray", g75="gray75",
    g82="gray82", light="light gray", g83="gray83"))
crgb <- col2rgb(cc <- colors())
colnames(crgb) <- cc
t(crgb)## The whole table
ccodes <- c(256^(2:0) %*% crgb)## = internal codes
## How many names are 'aliases' of each other:
table(tcc <- table(ccodes))
length(uc <- unique(sort(ccodes))) # 502
## All the multiply named colors:
mult <- uc[tcc >= 2]
```

```
cl <- lapply(mult, function(m) cc[ccodes == m])
names(cl) <- apply(col2rgb(sapply(cl, function(x)x[1])),
    2, function(n)paste(n, collapse=","))
str(cl)
    if(require(xgobi)) { ## Look at the color cube dynamically :
        tc <- t(crgb[, !duplicated(ccodes)])
        table(is.gray <- tc[,1] == tc[,2] & tc[,2] == tc[,3])# (397, 105)
        xgobi(tc, color = c("gold", "gray")[1 + is.gray])
    }
```

colors Color Names

## Description

Returns the built-in color names which R knows about.

## Usage

colors()

## Details

These color names can be used with a col= specification in graphics functions.
An even wider variety of colors can be created with primitives rgb and hsv or the derived rainbow, heat.colors, etc.

## Value

A character vector containing all the built-in color names.

## See Also

palette for setting the "palette" of colors for $\operatorname{par}(\mathrm{col}=<n u m>)$; rgb, hsv, gray; rainbow for a nice example; and heat.colors, topo.colors for images.
col2rgb for translating to RGB numbers and extended examples.

## Examples

```
str(colors())
```

```
colSums Form Row and Column Sums and Means
```


## Description

Form row and column sums and means for numeric arrays.

## Usage

colSums (x, na.rm = FALSE, dims = 1)
rowSums ( $x$, na.rm = FALSE, dims = 1)
colMeans ( $x$, na.rm = FALSE, dims = 1)
rowMeans(x, na.rm = FALSE, dims = 1)

## Arguments

$\mathrm{x} \quad$ an array of two or more dimensions, containing numeric, complex, integer or logical values, or a numeric data frame.
na.rm logical. Should missing values (including NaN) be omitted from the calculations?
dims Which dimensions are regarded as "rows" or "columns" to sum over. For col*, the sum or mean is over dimensions dims+1, ...; for row* it is over dimensions 1:dims.

## Details

These functions are equivalent to use of apply with FUN = mean or FUN = sum with appropriate margins, but are a lot faster. As they are written for speed, they blur over some of the subtleties of NaN and NA. If na.rm = FALSE and either NaN or NA appears in a sum, the result will be one of NaN or NA, but which might be platform-dependent.

## Value

A numeric or complex array of suitable size, or a vector if the result is one-dimensional. The dimnames (or names for a vector result) are taken from the original array.

If there are no non-missing values in a range to be summed over, the component of the output is set to NA.

## See Also

apply, rowsum

## Examples

```
## Compute row and column sums for a matrix:
x <- cbind(x1 = 3, x2 = c(4:1, 2:5))
rowSums(x); colSums(x)
dimnames(x)[[1]] <- letters[1:8]
rowSums(x); colSums(x); rowMeans(x); colMeans(x)
x[] <- as.integer(x)
rowSums(x); colSums(x)
x[] <- x < 3
```

```
rowSums(x); colSums(x)
x <- cbind(x1 = 3, x2 = c(4:1, 2:5))
x[3, ] <- NA; x[4, 2] <- NA
rowSums(x); colSums(x); rowMeans(x); colMeans(x)
rowSums(x, na.rm = TRUE); colSums(x, na.rm = TRUE)
rowMeans(x, na.rm = TRUE); colMeans(x, na.rm = TRUE)
## an array
data(UCBAdmissions)
dim(UCBAdmissions)
rowSums(UCBAdmissions); rowSums(UCBAdmissions, dims = 2)
colSums(UCBAdmissions); colSums(UCBAdmissions, dims = 2)
## complex case
x <- cbind(x1 = 3 + 2i, x2 = c(4:1, 2:5) - 5i)
x[3, ] <- NA; x[4, 2] <- NA
rowSums(x); colSums(x); rowMeans(x); colMeans(x)
rowSums(x, na.rm = TRUE); colSums(x, na.rm = TRUE)
rowMeans(x, na.rm = TRUE); colMeans(x, na.rm = TRUE)
```

commandArgs Extract Command Line Arguments

## Description

Provides access to a copy of the command line arguments supplied when this R session was invoked.

```
Usage
    commandArgs()
```


## Details

These arguments are captured before the standard R command line processing takes place. This means that they are the unmodified values. If it were useful, we could provide support an argument which indicated whether we want the unprocessed or processed values.

## Value

A character vector containing the name of the executable and the user-supplied command line arguments. The first element is the name of the executable by which $R$ was invoked. As far as I am aware, the exact form of this element is platform dependent. It may be the fully qualified name, or simply the last component (or basename) of the application.

## See Also

BATCH

## Examples

```
commandArgs()
## Spawn a copy of this application as it was invoked.
## system(paste(commandArgs(), collapse=" "))
```

```
comment Query or Set a 'Comment' Attribute
```


## Description

These functions set and query a comment attribute for any R objects. This is typically useful for data.frames or model fits.

Contrary to other attributes, the comment is not printed (by print or print.default).

## Usage

```
comment ( x )
comment(x) <- value
```


## Arguments

| x | any R object |
| :--- | :--- |
| value | a character vector |

## See Also

attributes and attr for "normal" attributes.

## Examples

```
x <- matrix(1:12, 3,4)
comment(x) <- c("This is my very important data from experiment #0234",
    "Jun 5, 1998")
x
comment (x)
```


## Comparison

 Relational Operators
## Description

Binary operators which allow the comparison of values in vectors.

## Usage

$$
\begin{aligned}
& x<y \\
& x>y \\
& x<=y \\
& x>=y \\
& x==y \\
& x \quad!=y
\end{aligned}
$$

## Details

Comparison of strings in character vectors is lexicographic within the strings using the collating sequence of the locale in use: see locales. The collating sequence of locales such as en_US is normally different from C (which should use ASCII) and can be surprising.

## Value

A vector of logicals indicating the result of the element by element comparison. The elements of shorter vectors are recycled as necessary.

Objects such as arrays or time-series can be compared this way provided they are conformable.

## Note

Don't use $==$ and $!=$ for tests, such as in if expressions, where you must get a single TRUE or FALSE. Unless you are absolutely sure that nothing unusual can happen, you should use the identical function instead.

For numerical values, remember $==$ and $!=$ do not allow for the finite representation of fractions, nor for rounding error. Using all.equal with identical is almost always preferable. See the examples.

## See Also

Syntax for operator precedence.

## Examples

```
x <- rnorm(20)
x < 1
x[x > 0]
x1<- 0.5-0.3
x2 <- 0.3-0.1
x1 == x2 # FALSE on most machines
identical(all.equal(x1, x2), TRUE) # TRUE everywhere
```


## Description

Compile given source files so that they can subsequently be collected into a shared library using R CMD SHLIB and be loaded into R using dyn.load().

## Usage

R CMD COMPILE [options] srcfiles

## Arguments

srcfiles A list of the names of source files to be compiled. Currently, C, C++ and FORTRAN are supported; the corresponding files should have the extensions '.c', '.cc' (or '.cpp' or '.C'), and '. $f$ ', respectively.
options A list of compile-relevant settings, such as special values for CFLAGS or FFLAGS, or for obtaining information about usage and version of the utility.

## Details

Note that Ratfor is not supported. If you have Ratfor source code, you need to convert it to FORTRAN. On many Solaris systems mixing Ratfor and FORTRAN code will work.

## See Also

SHLIB, dyn.load

```
complete.cases Find Complete Cases
```


## Description

Return a logical vector indicating which cases are complete, i.e., have no missing values.

## Usage

complete.cases(...)

## Arguments

... a sequence of vectors, matrices and data frames.

## Value

A logical vector specifying which observations/rows have no missing values across the entire sequence.

## See Also

```
is.na, na.omit, na.fail.
```


## Examples

```
data(airquality)
x <- airquality[, -1] # x is a regression design matrix
y <- airquality[, 1] # y is the corresponding response
stopifnot(complete.cases(y) != is.na(y))
ok <- complete.cases(x,y)
sum(!ok) # how many are not "ok" ?
x <- x[ok,]
y <- y [ok]
```


## Description

Basic functions which support complex arithmetic in $R$.

## Usage

```
complex(length.out = 0, real = numeric(), imaginary = numeric(),
    modulus = 1, argument = 0)
as.complex(x, ...)
is.complex(x)
Re(x)
Im(x)
Mod(x)
Arg(x)
Conj(x)
```


## Arguments

| length.out | numeric. Desired length of the output vector, inputs being recycled as <br> needed. |
| :--- | :--- |
| real | numeric vector. |
| imaginary | numeric vector. |
| modulus | numeric vector. |
| argument | numeric vector. |
| x | an object, probably of mode complex. |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

Complex vectors can be created with complex. The vector can be specified either by giving its length, its real and imaginary parts, or modulus and argument. (Giving just the length generates a vector of complex zeroes.)
Note that is.complex and is.numeric are never both TRUE.
The functions Re, Im, Mod, Arg and Conj have their usual interpretation as returning the real part, imaginary part, modulus, argument and complex conjugate for complex values. Modulus and argument are also called the polar coordinates. If $z=x+i y$ with real $x$ and $y, \operatorname{Mod}(z)=\sqrt{x^{2}+y^{2}}$, and for $\phi=\operatorname{Arg}(z), x=\cos (\phi)$ and $y=\sin (\phi)$.
In addition, the elementary trigonometric, logarithmic and exponential functions are available for complex values.

## Examples

```
( z <- 0i ~ (-3:3) )
stopifnot(Re(z) == 0 ~ (-3:3))
matrix(1i^ (-6:5), nr=4)#- all columns are the same
0 ~ 1i # a complex NaN
## create a complex normal vector
z <- complex(real = rnorm(100), imag = rnorm(100))
## or also (less efficiently):
z2 <- 1:2 + 1i*(8:9)
all(Mod ( 1 - sin(z) / ( (exp(1i*z)-exp(-1i*z))/(2*1i) ))
    < 100*.Machine$double.eps)
## The Arg(.) is an angle:
zz <- (rep(1:4,len=9) + 1i*(9:1))/10
zz.shift <- complex(modulus = Mod(zz), argument= Arg(zz) + pi)
plot(zz, xlim=c(-1,1), ylim=c(-1,1), col="red", asp = 1,
    main = expression(paste("Rotation by "," ", pi == 180^o)))
abline(h=0,v=0, col="blue", lty=3)
points(zz.shift, col="orange")
```

conflicts Search for Masked Objects on the Search Path

## Description

conflicts reports on objects that exist with the same name in two or more places on the search path, usually because an object in the user's workspace or a package is masking a system object of the same name. This helps discover unintentional masking.

## Usage

conflicts(where=search(), detail=FALSE)

## Arguments

where A subset of the search path, by default the whole search path.
detail If TRUE, give the masked or masking functions for all members of the search path.

## Value

If detail=FALSE, a character vector of masked objects. If detail=TRUE, a list of character vectors giving the masked or masking objects in that member of the search path. Empty vectors are omitted.

Author(s)
B.D. Ripley

## Examples

```
lm <- 1:3
conflicts(, TRUE)
## gives something like
# $.GlobalEnv
# [1] "lm"
#
# $package:base
# [1] "lm"
## Remove things from your "workspace" that mask others:
remove(list = conflicts(detail=TRUE)$.GlobalEnv)
```

connections Functions to Manipulate Connections

## Description

Functions to create, open and close connections.

## Usage

```
file(description = "", open = "", blocking = TRUE,
    encoding = getOption("encoding"))
pipe(description, open = "", encoding = getOption("encoding"))
fifo(description = "", open = "", blocking = FALSE,
    encoding = getOption("encoding"))
gzfile(description, open = "", encoding = getOption("encoding"),
            compression = 6)
unz(description, filename, open = "", encoding = getOption("encoding"))
bzfile(description, open = "", encoding = getOption("encoding"))
url(description, open = "", blocking = TRUE,
    encoding = getOption("encoding"))
socketConnection(host = "localhost", port, server = FALSE,
    blocking = FALSE, open = "a+",
    encoding = getOption("encoding"))
open(con, open \(=\) " r ", blocking \(=\) TRUE, ...)
close(con, type = "rw", ...)
isOpen(con, rw = "")
isIncomplete(con)
```


## Arguments

description character. A description of the connection. For file and pipe this is a path to the file to be opened. For url it is a complete URL, including schemes (http://, ftp:// or file://). file also accepts complete URLs.
filename a filename within a zip file.
con a connection.

| host | character. Host name for port. |
| :--- | :--- |
| port | integer. The TCP port number. |
| server | logical. Should the socket be a client or a server? <br> open |
| character. A description of how to open the connection (if at all). See |  |
| blocking | Details for possible values. <br> logical. See 'Blocking' section below. |
| encoding | An integer vector of length 256. |
| compression | integer in 0-9. The amount of compression to be applied when writing, <br> from none to maximal. The default is a good space/time compromise. |
| type | character. Currently ignored. <br> where |
| integer. A file position (relative to the origin specified by origin), or NA. |  |
| rw | character. Empty or "read" or "write", partial matches allowed. |
| .. | arguments passed to or from other methods. |

## Details

The first eight functions create connections. By default the connection is not opened (except for socketConnection), but may be opened by setting a non-empty value of argument open. gzfile applies to files compressed by 'gzip', and bzfile to those compressed by 'bzip2': such connections can only be binary.
unz reads (only) single files within zip files, in binary mode. The description is the full path, with '.zip' extension if required.

All platforms support (gz)file connections and url("file://") connections. The other types may be partially implemented or not implemented at all. (They do work on most Unix platforms, and all but fifo on Windows.)

Proxies can be specified for url connections: see download.file.
open, close and seek are generic functions: the following applies to the methods relevant to connections.
open opens a connection. In general functions using connections will open them if they are not open, but then close them again, so to leave a connection open call open explicitly.
close closes and destroys a connection.
Possible values for the mode open to open a connection are
"r" or "rt" Open for reading in text mode.
"w" or "wt" Open for writing in text mode.
"a" or "at" Open for appending in text mode.
"rb" Open for reading in binary mode.
"wb" Open for writing in binary mode.
"ab" Open for appending in binary mode.
" $\mathrm{r}+\mathrm{"}$, " $\mathrm{r}+\mathrm{b} "$ Open for reading and writing.
" $\mathrm{w}+\mathrm{"}, \mathrm{w}+\mathrm{b}$ " Open for reading and writing.
" $\mathrm{r}+\mathrm{"}, \mathrm{lr} \mathrm{r} \mathrm{b}$ " Open for reading and writing, truncating file initially.
" $\mathrm{a}+\mathrm{"}, \mathrm{a} \mathrm{a}+\mathrm{b}$ " Open for reading and appending.

Not all modes are applicable to all connections: for example URLs can only be opened for reading. Only file and socket connections can be opened for reading and writing/appending. For many connections there is little or no difference between text and binary modes, but there is for file-like connections on Windows, and pushBack is text-oriented and is only allowed on connections open for reading in text mode.

If for a file connection the description is "", the file is immediately opened in " $\mathrm{w}+$ " mode and unlinked from the file system. This provides a temporary file to write to and then read from.
The encoding vector is used to map the input from a file or pipe to the platform's native character set. Supplied examples are native.enc as well as MacRoman, WinAnsi and ISOLatin1, whose actual encoding is platform-dependent. Missing characters are mapped to a space in these encodings.

## Value

file, pipe, fifo, url, gzfile and socketConnection return a connection object which inherits from class "connection" and has a first more specific class.
isOpen returns a logical value, whether the connection is currently open.
isIncomplete returns a logical value, whether last read attempt was blocked, or for an output text connection whether there is unflushed output.

## Blocking

The default condition for all but fifo and socket connections is to be in blocking mode. In that mode, functions do not return to the R evaluator until they are complete. In nonblocking mode, operations return as soon as possible, so on input they will return with whatever input is available (possibly none) and for output they will return whether or not the write succeeded.

The function readLines behaves differently in respect of incomplete last lines in the two modes: see its help page.

Even when a connection is in blocking mode, attempts are made to ensure that it does not block the event loop and hence the operation of GUI parts of R. These do not always succeed, and the whole process will be blocked during a DNS lookup on Unix, for example.

Most blocking operations on URLs and sockets are subject to the timeout set by options("timeout"). Note that this is a timeout for no response at all, not for the whole operation.

## Fifos

Fifos default to non-blocking. That follows Svr4 and it probably most natural, but it does have some implications. In particular, opening a non-blocking fifo connection for writing (only) will fail unless some other process is reading on the fifo.

Opening a fifo for both reading and writing (in any mode: one can only append to fifos) connects both sides of the fifo to the $R$ process, and provides an similar facility to file().

## Note

R's connections are modelled on those in S version 4 (see Chambers, 1998). However R goes well beyond the Svr4 model, for example in output text connections and URL, gzfile, bzfile and socket connections.

The default mode in R is " r " except for socket connections. This differs from Svr4, where it is the equivalent of " $\mathrm{r}+$ ", known as "*".

## References

Chambers, J. M. (1998) Programming with Data. A Guide to the S Language. Springer.

## See Also

textConnection, seek, readLines, showConnections, pushBack. capabilities to see if gzfile, url, fifo and socketConnection are supported by this build of $R$.

## Examples

```
zz <- file("ex.data", "w") # open an output file connection
cat("TITLE extra line", "2 3 5 7", "", "11 13 17", file = zz, sep = "\n")
cat("One more line\n", file = zz)
close(zz)
readLines("ex.data")
unlink("ex.data")
zz <- gzfile("ex.gz", "w") # compressed file
cat("TITLE extra line", "2 3 5 7", "", "11 13 17", file = zz, sep = "\n")
close(zz)
readLines(gzfile("ex.gz"))
unlink("ex.gz")
if(capabilities("bzip2")) {
    zz <- bzfile("ex.bz2", "w") # bzip2-ed file
    cat("TITLE extra line", "2 3 5 7", "", "11 13 17", file = zz, sep = "\n")
    close(zz)
    print(readLines(bzfile("ex.bz2")))
    unlink("ex.bz2")
}
## An example of a file open for reading and writing
Tfile <- file("test1", "w+")
c(isOpen(Tfile, "r"), isOpen(Tfile, "w")) # both TRUE
cat("abc\ndef\n", file=Tfile)
readLines(Tfile)
seek(Tfile, 0, rw="r") # reset to beginning
readLines(Tfile)
cat("ghi\n", file=Tfile)
readLines(Tfile)
close(Tfile)
unlink("test1")
## We can do the same thing with an anonymous file.
Tfile <- file()
cat("abc\ndef\n", file=Tfile)
readLines(Tfile)
close(Tfile)
if(capabilities("fifo")) {
    zz <- fifo("foo", "w+")
    writeLines("abc", zz)
```

```
    print(readLines(zz))
    close(zz)
    unlink("foo")
}
## Unix examples of use of pipes
# read listing of current directory
readLines(pipe("ls -1"))
# remove trailing commas. Suppose
% cat data2
450, 390, 467, 654, 30, 542, 334, 432, 421,
357, 497, 493, 550, 549, 467, 575, 578, 342,
446, 547, 534, 495, 979, 479
# Then read this by
scan(pipe("sed -e s/,$// data2"), sep=",")
# convert decimal point to comma in output
zz <- pipe(paste("sed s/\\./,/ >", "outfile"), "w")
cat(format(round(rnorm(100), 4)), sep = "\n", file = zz)
close(zz)
file.show("outfile", delete.file=TRUE)
## example for Unix machine running a finger daemon
con <- socketConnection(port = 79, blocking = TRUE)
writeLines(paste(system("whoami", intern=TRUE), "\r", sep=""), con)
gsub(" *$", "", readLines(con))
close(con)
## two R processes communicating via non-blocking sockets
# R process 1
con1 <- socketConnection(port = 6011, server=TRUE)
writeLines(LETTERS, con1)
close(con1)
# R process 2
con2 <- socketConnection(Sys.info()["nodename"], port = 6011)
# as non-blocking, may need to loop for input
readLines(con2)
while(isIncomplete(con2)) {Sys.sleep(1); readLines(con2)}
close(con2)
```


## Constants Built-in Constants

## Description

Constants built into R .

## Usage

LETTERS
letters

```
month.abb
month.name
pi
```


## Details

R has a limited number of built-in constants (there is also a rather larger library of data sets which can be loaded with the function data).
The following constants are available:

- LETTERS: the 26 upper-case letters of the Roman alphabet;
- letters: the 26 lower-case letters of the Roman alphabet;
- month.abb: the three-letter abbreviations for the English month names;
- month. name: the English names for the months of the year;
- pi: the ratio of the circumference of a circle to its diameter.


## See Also

data.

## Examples

```
stopifnot(
    nchar(letters) == 1,
    month.abb == substr(month.name, 1, 3)
)
eps <- .Machine$double.eps
all.equal(pi, 4*atan(1), tol= 2*eps)
# John Machin (1705) computed 100 decimals of pi :
all.equal(pi/4, 4*atan(1/5) - atan(1/239), 4*eps)
```

contour Display Contours

## Description

Create a contour plot, or add contour lines to an existing plot.

## Usage

```
contour (x, ....)
contour (x \(=\operatorname{seq}(0,1\), len \(=\operatorname{nrow}(z)), y=\operatorname{seq}(0,1\), len \(=n c o l(z))\),
            z,
            nlevels = 10, levels = pretty(zlim, nlevels), labels = NULL,
            xlim = range(x, finite = TRUE),
            ylim = range(y, finite = TRUE),
            zlim = range(z, finite = TRUE),
            labcex = 0.6, drawlabels = TRUE, method = "flattest",
            vfont = c("sans serif", "plain"),
            axes = TRUE, frame.plot = axes,
            col = par("fg"), lty = par("lty"), lwd = par("lwd"),
            add \(=\) FALSE, ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | locations of grid lines at which the values in $\mathbf{z}$ are measured. These must be in ascending order. By default, equally spaced values from 0 to 1 are used. If x is a list, its components $\mathrm{x} \$ \mathrm{x}$ and $\mathrm{x} \$ \mathrm{y}$ are used for x and y , respectively. If the list has component $z$ this is used for $z$. |
| :---: | :---: |
| z | a matrix containing the values to be plotted (NAs are allowed). Note that x can be used instead of z for convenience. |
| nlevels | number of contour levels desired iff levels is not supplied. |
| levels | numeric vector of levels at which to draw contour lines. |
| labels | a vector giving the labels for the contour lines. If NULL then the levels are used as labels. |
| labcex | cex for contour labelling. |
| drawlabels | logical. Contours are labelled if TRUE. |
| method | character string specifying where the labels will be located. Possible values are "simple", "edge" and "flattest" (the default). See the Details section. |
| vfont | if a character vector of length 2 is specified, then Hershey vector fonts are used for the contour labels. The first element of the vector selects a typeface and the second element selects a fontindex (see text for more information). |
| xlim, ylim, zlim |  |
|  | x -, y - and z -limits for the plot. |
| axes, frame.plot |  |
|  | logical indicating whether axes or a box should be drawn, see plot. default. |
| col | color for the lines drawn. |
| lty | line type for the lines drawn. |
| lwd | line width for the lines drawn. |
| add | logical. If TRUE, add to a current plot. |
|  | additional graphical parameters (see par) and the arguments to title may also be supplied. |

## Details

contour is a generic function with only a default method in base $R$.
There is currently no documentation about the algorithm. The source code is in '\$R_HOME/src/main/plot3d.c'.
The methods for positioning the labels on contours are "simple" (draw at the edge of the plot, overlaying the contour line), "edge" (draw at the edge of the plot, embedded in the contour line, with no labels overlapping) and "flattest" (draw on the flattest section of the contour, embedded in the contour line, with no labels overlapping). The second and third may not draw a label on every contour line.
For information about vector fonts, see the help for text and Hershey.

## See Also

filled.contour for "color-filled" contours, image and the graphics demo which can be invoked as demo(graphics).

## Examples

```
x <- -6:16
op <- par(mfrow = c(2, 2))
contour(outer(x, x), method = "edge", vfont = c("sans serif", "plain"))
z <- outer(x, sqrt(abs(x)), FUN = "/")
## Should not be necessary:
z[!is.finite(z)] <- NA
image(x, x, z)
contour(x, x, z, col = "pink", add = TRUE, method = "edge",
    vfont = c("sans serif", "plain"))
contour(x, x, z, ylim = c(1, 6), method = "simple", labcex = 1)
contour(x, x, z, ylim = c(-6, 6), nlev = 20, lty = 2, method = "simple")
par(op)
## Persian Rug Art:
x <- y <- seq(-4*pi, 4*pi, len = 27)
r <- sqrt(outer(x^2, y^2, "+"))
opar <- par(mfrow = c(2, 2), mar = rep(0, 4))
for(f in pi^(0:3))
    contour(cos(r^2)*exp(-r/f),
                drawlabels = FALSE, axes = FALSE, frame = TRUE)
data("volcano")
rx <- range(x <- 10*1:nrow(volcano))
ry <- range(y <- 10*1:ncol(volcano))
ry <- ry + c(-1,1) * (diff(rx) - diff(ry))/2
tcol <- terrain.colors(12)
par(opar); opar <- par(pty = "s", bg = "lightcyan")
plot(x = 0, y = 0,type = "n", xlim = rx, ylim = ry, xlab = "", ylab = "")
u <- par("usr")
rect(u[1], u[3], u[2], u[4], col = tcol[8], border = "red")
contour(x, y, volcano, col = tcol[2], lty = "solid", add = TRUE,
    vfont = c("sans serif", "plain"))
title("A Topographic Map of Maunga Whau", font = 4)
abline(h = 200*0:4, v = 200*0:4, col = "lightgray", lty = 2, lwd = 0.1)
par(opar)
```

```
contrast
Contrast Matrices
```


## Description

Return a matrix of contrasts.

## Usage

```
contr.helmert(n, contrasts = TRUE)
contr.poly(n, contrasts = TRUE)
contr.sum(n, contrasts = TRUE)
contr.treatment(n, base = 1, contrasts = TRUE)
```


## Arguments

n
contrasts
base
a vector of levels for a factor, or the number of levels.
a logical indicating whether contrasts should be computed.
an integer specifying which group is considered the baseline group. Ignored if contrasts is FALSE.

## Details

These functions are used for creating contrast matrices for use in fitting analysis of variance and regression models. The columns of the resulting matrices contain contrasts which can be used for coding a factor with n levels. The returned value contains the computed contrasts. If the argument contrasts is FALSE then a square indicator matrix is returned.

Note that as from R version 0.62 .2 , contr.poly returns contrasts based on orthogonal (rather than raw) polynomials.

## Value

A matrix with n rows and k columns, with $\mathrm{k}=\mathrm{n}-1$ if contrasts is TRUE and $\mathrm{k}=\mathrm{n}$ if contrasts is FALSE.

## See Also

contrasts, C, and aov, glm, lm.

## Examples

```
(cH <- contr.helmert(4))
apply(cH, 2,sum) # column sums are 0!
crossprod(cH) # diagonal -- columns are orthogonal
contr.helmert(4, contrasts = FALSE) # just the 4 x 4 identity matrix
(cT <- contr.treatment(5))
all(crossprod(cT) == diag(4)) # TRUE: even orthonormal
(cP <- contr.poly(3)) # Linear and Quadratic
zapsmall(crossprod(cP), dig=15) # orthonormal up to fuzz
```

```
contrasts Get and Set Contrast Matrices
```


## Description

Set and view the contrasts associated with a factor.

## Usage

```
contrasts(x, contrasts = TRUE)
contrasts(x, how.many) <- ctr
```


## Arguments

x
contrasts logical. See Details.
how.many How many contrasts should be made. Defaults to one less than the number of levels of x . This need not be the same as the number of columns of ctr.
ctr either a matrix whose columns give coefficients for contrasts in the levels of x , or the (quoted) name of a function which computes such matrices.

## Details

If contrasts are not set for a factor the default functions from options("contrasts") are used.

The argument contrasts is ignored if x has a matrix contrasts attribute set. Otherwise if contrasts $=$ TRUE it is passed to a contrasts function such as contr.treatment and if contrasts = FALSE an identity matrix is returned.

## Note

Prior to R version 1.2.0, contrasts (, FALSE) called a contrasts function with contrasts $=$ FALSE. This normally gave the same result, but not for contr.poly, the default for ordered factors.

## See Also

C, contr.helmert, contr.poly, contr.sum, contr.treatment; glm, aov, lm.

## Examples

```
example(factor)
(fff <- factor(ff))
contrasts(fff) # treatment contrasts by default
contrasts(C(fff, sum))
contrasts(fff, contrasts = FALSE) # the 5x5 identity matrix
contrasts(fff) <- contr.sum(5); contrasts(fff) # set sum contrasts
contrasts(fff, 2) <- contr.sum(5); contrasts(fff) # set 2 contrasts
# supply 2 contrasts, compute 2 more to make full set of 4.
contrasts(fff) <- contr.sum(5)[,1:2]; contrasts(fff)
```

```
contributors }\quadR\mathrm{ Project Contributors
```


## Description

The R Who-is-who, describing who made significant contributions to the development of R .

## Usage

contributors()

## Control Control Flow

## Description

These are the basic control-flow constructs of the R language. They function in much the same way as control statements in any algol-like language.

## Usage

```
if(cond) expr
if(cond) cons.expr else alt.expr
for(var in seq) expr
while(cond) expr
repeat expr
break
next
```


## Details

Note that expr and cons.expr, etc, in the Usage section above means an expression in a formal sense. This is either a simple expression or a so called compound expression, usually of the form \{ expr1 ; expr2 \}.

Note that it is a common mistake to forget putting braces ( $\{$. $\}$ ) around your statements, e.g. after if (..) or for(....). For that reason, one (somewhat extreme) attitude of defensive programming uses braces always, e.g. for if clauses.

The index seq in a for loop is evaluated at the start of the loop; changing it subsequently does not affect the loop.

## See Also

Syntax for the basic R syntax and operators, Paren for parentheses and braces; further, ifelse, switch.

## Examples

```
for(i in 1:5) print(1:i)
for(n in c(2,5,10,20,50)) {
    x <- rnorm(n)
    cat(n,":", sum(x^2),"\n")
}
```

```
convolve Fast Convolution
```


## Description

Use the Fast Fourier Transform to compute the several kinds of convolutions of two sequences.

## Usage

```
convolve(x, y, conj = TRUE, type = c("circular", "open", "filter"))
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ numeric sequences of the same length to be convolved.
conj logical; if TRUE, take the complex conjugate before back-transforming (default, and used for usual convolution).
type character; one of "circular", "open", "filter" (beginning of word is ok). For circular, the two sequences are treated as circular, i.e., periodic.
For open and filter, the sequences are padded with 0s (from left and right) first; "filter" returns the middle sub-vector of "open", namely, the result of running a weighted mean of $x$ with weights $y$.

## Details

The Fast Fourier Transform, fft, is used for efficiency.
The input sequences x and y must have the same length if circular is true.
Note that the usual definition of convolution of two sequences $x$ and $y$ is given by convolve(x, rev(y), type = "o").

## Value

If $\mathrm{r}<-\operatorname{convol}^{2}(\mathrm{x}, \mathrm{y}$, type $=$ "open") and $\mathrm{n}<-$ length $(\mathrm{x}), \mathrm{m}<-$ length $(\mathrm{y})$, then

$$
r_{k}=\sum_{i} x_{k-m+i} y_{i}
$$

where the sum is over all valid indices $i$, for $k=1, \ldots, n+m-1$
If type == "circular", $n=m$ is required, and the above is true for $i, k=1, \ldots, n$ when $x_{j}:=x_{n+j}$ for $j<1$.

## References

Brillinger, D. R. (1981) Time Series: Data Analysis and Theory, Second Edition. San Francisco: Holden-Day.

## See Also

$f f t$, nextn, and particularly filter (from the 'ts' package) which may be more appropriate.

## Examples

```
x <- c(0,0,0,100,0,0,0)
y <- c(0,0,1, 2 ,1,0,0)/4
zapsmall(convolve(x,y)) # *NOT* what you first thought.
zapsmall(convolve(x, y[3:5], type="f")) # rather
x <- rnorm(50)
y <- rnorm(50)
# Circular convolution *has* this symmetry:
all.equal(convolve(x,y, conj = FALSE),
    rev(convolve(rev(y),x)))
n <- length(x <- -20:24)
y <- (x-10)^2/1000 + rnorm(x)/8
Han <- function(y) # Hanning
    convolve(y, c(1,2,1)/4, type = "filter")
plot(x,y, main="Using convolve(.) for Hanning filters")
lines(x[-c(1 , n) ], Han(y), col="red")
lines(x[-c(1:2, (n-1):n)], Han(Han(y)), lwd=2, col="dark blue")
```

coplot Conditioning Plots

## Description

This function produces two variants of the conditioning plots discussed in the reference below.

## Usage

```
coplot(formula, data, given.values, panel = points, rows, columns,
        show.given = TRUE, col = par("fg"), pch = par("pch"),
        bar.bg = c(num = gray(0.8), fac = gray(0.95)),
        xlab = c(x.name, paste("Given :", a.name)),
        ylab = c(y.name, paste("Given :", b.name)),
        subscripts = FALSE,
        axlabels = function(f) abbreviate(levels(f)),
        number = 6, overlap = 0.5, xlim, ylim, ...)
co.intervals(x, number = 6, overlap = 0.5)
```


## Arguments

formula a formula describing the form of conditioning plot. A formula of the form y $\sim \mathrm{x} \mid \mathrm{a}$ indicates that plots of y versus x should be produced conditional on the variable $a$. A formula of the form $y \sim x \mid a * b$ indicates that plots of y versus x should be produced conditional on the two variables a and b .
All three or four variables may be either numeric or factors. When x or y are factors, the result is almost as if as.numeric() was applied, whereas for factor a or b , the conditioning (and its graphics if show. given is true) are adapted.
$\left.\begin{array}{ll}\text { data } & \begin{array}{l}\text { a data frame containing values for any variables in the formula. By default } \\ \text { the environment where coplot was called from is used. }\end{array} \\ \text { given.values } \\ \text { a value or list of two values which determine how the conditioning on a } \\ \text { and b is to take place. } \\ \text { When there is no b (i.e., conditioning only on a), usually this is a matrix } \\ \text { with two columns each row of which gives an interval, to be conditioned } \\ \text { on, but is can also be a single vector of numbers or a set of factor levels } \\ \text { (if the variable being conditioned on is a factor). In this case (no b), the } \\ \text { result of co.intervals can be used directly as given.values argument. } \\ \text { a function (x, y, col, pch, ...) which gives the action to be carried } \\ \text { out in each panel of the display. The default is points. }\end{array}\right\}$

## Details

In the case of a single conditioning variable a, when both rows and columns are unspecified, a "close to square" layout is chosen with columns >= rows.
In the case of multiple rows, the order of the panel plots is from the bottom and from the left (corresponding to increasing a, typically).

## Value

co.intervals(., number, .) returns a (number $\times 2$ ) matrix, say ci, where ci[k,] is the range of x values for the k -th interval.

## References

Cleveland, W. S. (1993) Visualizing Data. New Jersey: Summit Press.

## See Also

## Examples

```
## Tonga Trench Earthquakes
data(quakes)
coplot(lat ~ long | depth, data = quakes)
given.depth <- co.intervals(quakes$depth, number=4, overlap=.1)
coplot(lat ~ long | depth, data = quakes, given.v=given.depth, rows=1)
## Conditioning on 2 variables:
ll.dm <- lat ~ long | depth * mag
coplot(ll.dm, data = quakes)
coplot(ll.dm, data = quakes, number=c(4,7), show.given=c(TRUE,FALSE))
coplot(ll.dm, data = quakes, number=c(3,7),
    overlap=c(-.5,.1)) # negative overlap DROPS values
data(warpbreaks)
## given two factors
coplot(breaks ~ 1:54 | wool * tension, data = warpbreaks, show.given = 0:1)
coplot(breaks ~ 1:54 | wool * tension, data = warpbreaks,
    col = "red", bg = "pink", pch = 21,
    bar.bg = c(fac = "light blue"))
## Example with empty panels:
data(state)
attach(data.frame(state.x77))#> don't need 'data' arg. below
coplot(Life.Exp ~ Income | Illiteracy * state.region, number = 3,
    panel = function(x, y, ...) panel.smooth(x, y, span = .8, ...))
## y ~ factor -- not really sensical, but 'show off':
coplot(Life.Exp ~ state.region | Income * state.division,
        panel = panel.smooth)
detach() # data.frame(state.x77)
```


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cor $\quad$ Correlation, Variance and Covariance (Matrices)

## Description

var, cov and cor compute the variance of x and the covariance or correlation of x and y if these are vectors. If x and y are matrices then the covariances (or correlations) between the columns of x and the columns of y are computed.

## Usage

```
var(x, y = NULL, na.rm = FALSE, use)
cor(x, y = NULL, use = "all.obs")
cov(x, y = NULL, use = "all.obs")
```


## Arguments

$x \quad$ a numeric vector, matrix or data frame.
y NULL (default) or a vector, matrix or data frame with compatible dimensions to x . The default is equivalent to $\mathrm{y}=\mathrm{x}$ (but more efficient).
use an optional character string giving a method for computing covariances in the presence of missing values. This must be (an abbreviation of) one of the strings "all.obs", "complete.obs" or "pairwise.complete.obs".
na.rm logical. Should missing values be removed?

## Details

For cov and cor one must either give a matrix or data frame for x or give both x and y .
var just another interface to cov, where na.rm is used to determine the default for use when that is unspecified. If na.rm is TRUE then the complete observations (rows) are used (use = "complete") to compute the variance. Otherwise (use = "all"), var will give an error if there are missing values.
If use is "all.obs", then the presence of missing observations will produce an error. If use is "complete.obs" then missing values are handled by casewise deletion. Finally, if use has the value "pairwise.complete.obs" then the correlation between each pair of variables is computed using all complete pairs of observations on those variables. This can result in covariance or correlation matrices which are not positive semidefinite.
The denominator $n-1$ is used which gives an unbiased estimator of the (co)variance for i.i.d. observations. These functions return NA when there is only one observation (whereas S-plus has been returning NaN ), and from R 1.2.3 fail if x has length zero.

## Value

For $r<-\operatorname{cor}(*$, use $=$ "all.obs" $)$, it is now guaranteed that all $(r<=1)$.

## See Also

 cov.wt for weighted covariance computation, sd for standard deviation (vectors).
## Examples

```
var(1:10)# 9.166667
var(1:5,1:5)# 2.5
## Two simple vectors
cor(1:10,2:11)# == 1
## Correlation Matrix of Multivariate sample:
data(longley)
(Cl <- cor(longley))
## Graphical Correlation Matrix:
symnum(Cl) # highly correlated
##--- Missing value treatment:
data(swiss)
C1 <- cov(swiss)
range(eigen(C1, only=TRUE)$val) # 6.19 1921
swiss[1,2] <- swiss[7,3] <- swiss[25,5] <- NA # create 3 "missing"
    C2 <- cov(swiss) # Error: missing obs...
C2 <- cov(swiss, use = "complete")
range(eigen(C2, only=TRUE)$val) # 6.46 1930
C3 <- cov(swiss, use = "pairwise")
range(eigen(C3, only=TRUE)$val) # 6.19 1938
```

```
count.fields Count the Number of Fields per Line
```


## Description

count.fields counts the number of fields, as separated by sep, in each of the lines of file read.

## Usage

```
count.fields(file, sep = "", quote = "\"'", skip = 0,
blank.lines.skip = TRUE, comment.char = "\#")
```


## Arguments

| file | a character string naming an ASCII data file, or a connection, which will <br> be opened if necessary, and if so closed at the end of the function call. |
| :--- | :--- |
| sep | the field separator character. Values on each line of the file are sepa- <br> rated by this character. By default, arbitrary amounts of whitespace can <br> separate fields. |
| quote | the set of quoting characters |

skip the number of lines of the data file to skip before beginning to read data. blank.lines.skip
logical: if TRUE blank lines in the input are ignored.
comment.char character: a character vector of length one containing a single character or an empty string.

## Details

This used to be used by read.table and can still be useful in discovering problems in reading a file by that function.

For the handling of comments, see scan.

## Value

A vector with the numbers of fields found.

## See Also

read.table

## Examples

```
cat("NAME", "1:John", "2:Paul", file = "foo", sep = "\n")
count.fields("foo", sep = ":")
unlink("foo")
```

```
cov.wt Weighted Covariance Matrices
```


## Description

Returns a list containing estimates of the weighted covariance matrix and the mean of the data, and optionally of the (weighted) correlation matrix.

## Usage

cov.wt(x, wt $=\operatorname{rep}(1 / \operatorname{nrow}(x), \operatorname{nrow}(x))$, cor $=$ FALSE, center $=$ TRUE)

## Arguments

$x \quad$ a matrix or data frame. As usual, rows are observations and columns are variables.
wt a non-negative and non-zero vector of weights for each observation. Its length must equal the number of rows of x .
cor A logical indicating whether the estimated correlation weighted matrix will be returned as well.
center Either a logical or a numeric vector specifying the centers to be used when computing covariances. If TRUE, the (weighted) mean of each variable is used, if FALSE, zero is used. If center is numeric, its length must equal the number of columns of $x$.

## Details

The covariance matrix is divided by one minus the sum of squares of the weights, so if the weights are the default $(1 / n)$ the conventional unbiased estimate of the covariance matrix with divisor $(n-1)$ is obtained. This differs from the behaviour in S-PLUS.

## Value

A list containing the following named components:
cov the estimated (weighted) covariance matrix
center an estimate for the center (mean) of the data.
n.obs the number of observations (rows) in x .
wt the weights used in the estimation. Only returned if given as an argument.
cor the estimated correlation matrix. Only returned if cor is TRUE.

## See Also

cov and var.

```
crossprod Matrix Crossproduct
```


## Description

Given matrices x and y as arguments, crossprod returns their matrix cross-product. This is formally equivalent to, but faster than, the call $t(x) \% * \% y$.

## Usage

crossprod (x, y = NULL)

## Arguments

$\mathrm{x}, \mathrm{y} \quad$ matrices: $\mathrm{y}=\mathrm{NULL}$ is taken to be the same matrix as x .

## See Also

$\% * \%$ and outer product $\% \%$.

## Examples

```
(z <- crossprod(1:4)) # = sum(1 + 2^2 + 3^2 + 4^2)
drop(z) # scalar
```


## Description

Returns a vector whose elements are the cumulative sums, products, minima or maxima of the elements of the argument.

## Usage

```
cumsum (x)
cumprod (x)
cummax ( x )
cummin( x )
```


## Arguments

x
a numeric object.

## Details

An NA value in x causes the corresponding and following elements of the return value to be NA.

## Examples

```
cumsum(1:10)
cumprod(1:10)
cummin(c(3:1, 2:0, 4:2))
cummax(c(3:1, 2:0, 4:2))
```

curve

## Description

Draws a curve corresponding to the given function or expression (in x ) over the interval [from,to].

## Usage

```
curve(expr, from, to, n = 101, add = FALSE, type = "l",
            ylab = NULL, log = NULL, xlim = NULL, ...)
plot(x, from = 0, to = 1, xlim = NULL, ...)
```


## Arguments

expr an expression written as a function of x , or alternatively a function which will be plotted.
x
a 'vectorizing' numeric $R$ function.
from, to the range over which the function will be plotted.
n
integer; the number of $x$ values at which to evaluate.
add logical; if TRUE add to already existing plot.
xlim numeric of length 2 ; if specified, it serves as default for c (from, to).
type, ylab, log,...
graphical parameters can also be specified as arguments. plot.function passes all these to curve.

## Details

The evaluation of expr is at n points equally spaced over the range [from, to], possibly adapted to $\log$ scale. The points determined in this way are then joined with straight lines. $\mathrm{x}(\mathrm{t})$ or expr (with x inside) must return a numeric of the same length as the argument t or x .

If add = TRUE, $c$ (from,to) default to xlim which defaults to the current x-limits. Further, log is taken from the current plot when add is true.

This used to be a quick hack which now seems to serve a useful purpose, but can give bad results for functions which are not smooth.

For "expensive" expressions, you should use smarter tools.

## See Also

splinefun for spline interpolation, lines.

## Examples

```
op <- par(mfrow=c(2,2))
curve(x^3-3*x, -2, 2)
curve(x^2-2, add = TRUE, col = "violet")
plot(cos, xlim = c(-pi,3*pi), n = 1001, col = "blue")
chippy <- function(x) sin(cos(x)*exp(-x/2))
curve(chippy, -8, 7, n=2001)
curve(chippy, -8, -5)
for(ll in c("","x","y","xy"))
    curve(log(1+x), 1,100, log=ll, sub=paste("log= '",1l,"'",sep=""))
par(op)
```


## Description

cut divides the range of x into intervals and codes the values in x according to which interval they fall. The leftmost interval corresponds to level one, the next leftmost to level two and so on.

## Usage

```
cut(x, ...)
cut.default(x, breaks, labels = NULL,
include.lowest \(=\) FALSE, right \(=\) TRUE, dig.lab \(=3, \ldots\) )
```


## Arguments

$\mathrm{x} \quad$ a numeric vector which is to be converted to a factor by cutting.
breaks either a vector of cut points or number giving the number of intervals which x is to be cut into.
labels labels for the levels of the resulting category. By default, labels are constructed using " (a,b]" interval notation. If labels = FALSE, simple integer codes are returned instead of a factor.
include.lowest
logical, indicating if an ' $x[i]$ ' equal to the lowest (or highest, for right $=$ FALSE) 'breaks' value should be included.
right logical, indicating if the intervals should closed on the right (and open on the left) or vice versa.
dig.lab integer which is used when labels are not given. It determines the number of digits used in formatting the break numbers.
... further arguments passed to or from other methods.

## Details

If a labels parameter is specified, its values are used to name the factor levels. If none is specified, the factor level labels are constructed as " (b1, b2] ", " (b2, b3]" etc. for right=TRUE and as "[b1, b2)", ...if right=FALSE. In this case, dig.lab indicates how many digits should be used in formatting the numbers b1, b2, ...

## Value

A factor is returned, unless labels = FALSE which results in the mere integer level codes.

## Note

Instead of table(cut(x, br)), hist(x, br, plot = FALSE) is more efficient and less memory hungry.

## See Also

split for splitting a variable according to a group factor; factor, tabulate, table.

## Examples

```
Z <- rnorm(10000)
table(cut(Z, br = -6:6))
system.time(print(sum(table(cut(Z, br = -6:6, labels=FALSE)))))
system.time(print(sum( hist (Z, br = -6:6, plot=FALSE)$counts)))
cut(rep(1,5),4)#-- dummy
tx0 <- c(9, 4, 6, 5, 3, 10, 5, 3, 5)
x <- rep(0:8, tx0)
stopifnot(table(x) == tx0)
table( cut(x, b = 8))
table( cut(x, br = 3*(-2:5)))
table( cut(x, br = 3*(-2:5), right = FALSE))
##--- some values OUTSIDE the breaks :
table(cx <- cut(x, br = 2*(0:4)))
table(cxl <- cut(x, br = 2*(0:4), right = FALSE))
which(is.na(cx)); x[is.na(cx)] #-- the first 9 values 0
which(is.na(cxl)); x[is.na(cxl)] #-- the last 5 values 8
## Label construction:
y <- rnorm(100)
table(cut(y, breaks = pi/3*(-3:3)))
table(cut(y, breaks = pi/3*(-3:3), dig.lab=4))
table(cut(y, breaks = 1*(-3:3), dig.lab=4))# extra digits don't "harm" here
table(cut(y, breaks = 1*(-3:3), right = FALSE))#- the same, since no exact INT!
```

cut.POSIXt $\quad$ Convert a Date-Time Object to a Factor

## Description

Method for cut applied to date-time objects.

## Usage

cut(x, breaks, labels=NULL, start.on.monday=TRUE, ...)

## Arguments

x
an object inheriting from class "POSIXt".
breaks a vector of cut points or number giving the number of intervals which x is to be cut into or an interval specification, one of "secs", "mins", "hours", "days", "weeks", "months" or "years".
labels labels for the levels of the resulting category. By default, labels are constructed using " (a,b]" interval notation. If labels = FALSE, simple integer codes are returned instead of a factor.
start.on.monday
logical. If breaks = "weeks", should the week start on Mondays or Sundays?
... arguments to be passed to or from other methods.

## Value

A factor is returned, unless labels $=$ FALSE which returns the integer level codes.

## See Also

seq.POSIXt, cut

## Examples

```
## random dates in a 10-week period
cut(ISOdate(2001, 1, 1) + 70*86400*runif(100), "weeks")
```

```
data Data Sets
```


## Description

Loads specified data sets, or list the available data sets.

## Usage

```
data(..., list = character(0), package = .packages(),
    lib.loc = NULL, verbose = getOption("verbose"))
```


## Arguments

... a sequence of names or character strings.
list a character vector.
package a name or character vector giving the packages to look into for data sets. By default, all packages in the search path are used, then the 'data' directory (if present) of the current working directory.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. If the default is used, the loaded packages are searched before the libraries.
verbose a logical. If TRUE, additional diagnostics are printed.

## Details

Currently, four formats of data files are supported:

1. files ending '.RData' or '.rda' are load()ed.
2. files ending '. $R$ ' or '. $r$ ' are source()d in, with the $R$ working directory changed temporarily to the directory containing the respective file.
3. files ending '.tab' or '.txt' are read using read.table(..., header $=$ TRUE), and hence result in a data frame.
4. files ending '.csv’ are read using read.table(..., header = TRUE, sep = ";"), and also result in a data frame.

The data sets to be loaded can be specified as a sequence of names or character strings, or as the character vector list, or as both.
If no data sets are specified, data lists the available data sets. It looks for a file '00Index' in a 'data' directory of each specified package, and uses these files to prepare a listing. If there is a 'data' area but no index a warning is given: such packages are incomplete. The information about available data sets is returned in an object of class "packageIQR". The structure of this class is experimental. In earlier versions of $R$, an empty character vector was returned along with listing available data sets.
If lib.loc is not specified, the datasets are searched for amongst those packages already loaded, followed by the 'data' directory (if any) of the current working directory and then packages in the specified libraries. If lib.loc is specified, packages are searched for in the specified libraries, even if they are already loaded from another library.
To just look in the 'data' directory of the current working directory, set package $=$ NULL.

## Value

a character vector of all data sets specified, or information about all available data sets in an object of class "packageIQR" if none were specified.

## Note

The data files can be many small files. On some file systems it is desirable to save space, and the files in the 'data' directory of an installed package can be zipped up as a zip archive 'Rdata.zip'. You will need to provide a single-column file 'filelist' of file names in that directory.
One can take advantage of the search order and the fact that a '. $R$ ' file will change directory. If raw data are stored in 'mydata.txt' then one can set up 'mydata.R' to read 'mydata.txt' and pre-process it, e.g. using transform. For instance one can convert numeric vectors to factors with the appropriate labels. Thus, the '. $R$ ' file can effectively contain a metadata specification for the plaintext formats.

## See Also

help for obtaining documentation on data sets, save for creating the first ('.rda') kind of data, typically the most efficient one.

## Examples

```
data() # list all available data sets
data(package = base) # list the data sets in the base package
data(USArrests, "VADeaths") # load the data sets 'USArrests' and 'VADeaths'
help(USArrests) # give information on data set 'USArrests'
```

data.class Object Classes

## Description

Determine the class of an arbitrary $R$ object.

## Usage

```
data.class(x)
```


## Arguments

$x \quad$ an R object.

## Value

character string giving the "class" of x.
The "class" is the (first element) of the class attribute if this is non-NULL, or inferred from the object's dim attribute if this is non-NULL, or mode( x ).
Simply speaking, data.class(x) returns what is typically useful for method dispatching. (Or, what the basic creator functions already and maybe eventually all will attach as a class attribute.)

## Note

For compatibility reasons, there is one exception to the rule above: When x is integer, the result of data.class( x ) is "numeric" even when x is classed.

## See Also

class

## Examples

```
x <- LETTERS
data.class(factor(x)) # has a class attribute
data.class(matrix(x, nc = 13)) # has a dim attribute
data.class(list(x)) # the same as mode(x)
data.class(x) # the same as mode(x)
stopifnot(data.class(1:2) == "numeric")# compatibility "rule"
```

```
data.frame Data Frames
```


## Description

These functions create or manipulate data frames, tightly coupled collections of variables which share many of the properties of matrices and of lists, used as the fundamental data structure by most of R's modeling software.

## Usage

```
data.frame(..., row.names = NULL, check.rows = FALSE,
    check.names = TRUE)
as.data.frame(x, row.names = NULL, optional = FALSE)
is.data.frame(x)
row.names(x)
row.names(x) <- names
```


## Arguments

```
... these arguments are of either the form value or tag=value. Component
                                    names are created based on the tag (if present) or the deparsed argument
                                    itself.
row.names NULL or a character vector giving the row names for the data frame. Miss-
    ing values are not allowed.
check.rows if TRUE then the rows are checked for consistency of length and names.
check.names logical. If TRUE then the names of the variables in the data frame are
    checked to ensure that they are syntactically valid variable names. If
    necessary they are adjusted (by make.names) so that they are.
    optional logical. If TRUE, setting row names is optional.
    x object of class data.frame.
```


## Details

A data frame is a list of variables of the same length with unique row names, given class "data.frame".
Character variables passed to data.frame are converted to factor columns unless protected by I. It also applies to adding columns to a data frame.

If a list or data frame or matrix is passed to data.frame it is as if each column had been passed as a separate argument, with the exception of matrices of class model.matrix.
Objects passed to data.frame should have the same number of rows, but atomic vectors and factors will be recycled a whole number of times if necessary.

## Value

For data.frame(.) a data frame, a matrix-like stucture whose columns may be of differing types (numeric, factor and character).
as.data.frame is generic function with many methods. It attempts to coerce its argument to be a data frame.
is.data.frame returns TRUE if its argument is a data frame and FALSE otherwise.
row.names can be used to set and retrieve the row names of a data frame, similarly to rownames for arrays (and it is a generic function that calls rownames for an array argument.

## Note

In versions of R prior to 1.4.0 (and in S3 but not S4) logical columns were converted to factors.

## See Also

I, print.data.frame, read.table, Math.data.frame etc, about Group methods for data.frames; make.names.

## Examples

```
L3 <- LETTERS[1:3]
str(d <- data.frame(cbind(x=1, y=1:10), fac=sample(L3, 10, repl=TRUE)))
## The same with automatic column names:
```

```
str( data.frame(cbind( 1, 1:10), sample(L3, 10, repl=TRUE)))
is.data.frame(d)
## do not convert to factor, using I() :
str(cbind(d, char = I(letters[1:10])), vec.len = 10)
stopifnot(1:10 == row.names(d))# {coercion}
(dO <- d[, FALSE]) # NULL dataframe with 10 rows
(d.0 <- d[FALSE, ]) # <0 rows> dataframe (3 cols)
(d00 <- dO[FALSE,]) # NULL dataframe with 0 rows
```

```
data.matrix Data Frame to Numeric Matrix
```


## Description

Return the matrix obtained by converting all the variables in a data frame to numeric mode and then binding them together as the columns of a matrix. Factors and ordered factors are replaced by their codes.

## Usage

data.matrix(frame)

## Arguments

frame a data frame whose components are logical vectors, factors or numeric vectors.

## See Also

as.matrix, codes, data.frame, matrix.

```
dataentry Spreadsheet Interface for Entering Data
```


## Description

A spreadsheet-like editor for entering or editing data.

## Usage

```
data.entry(..., Modes = NULL, Names = NULL)
dataentry(data, modes)
de(..., Modes = list(), Names = NULL)
```


## Arguments

| $\ldots$. | A list of variables: currently these should be numeric or character vectors |
| :--- | :--- |
| or list containing such vectors. |  |
| Modes | The modes to be used for the variables. |
| Names | The names to be used for the variables. |
| data | A list of numeric and/or character vectors. |
| modes | A list of length up to that of data giving the modes of (some of) the <br> variables. list() is allowed. |

## Details

The data entry editor is only available on some platforms and GUIs. Where available it provides a means to visually edit a matrix or a collection of variables (including a data frame) as described in the "Notes" section.
data.entry has side effects, any changes made in the spreadsheet are reflected in the variables. The functions de, de.ncols, de.setup and de.restore are designed to help achieve these side effects. If the user passes in a matrix, X say, then the matrix is broken into columns before dataentry is called. Then on return the columns are collected and glued back together and the result assigned to the variable X. If you don't want this behaviour use dataentry directly.
The primitive function is dataentry. It takes a list of vectors of possibly different lengths and modes (the second argument) and opens a spreadsheet with these variables being the columns. The columns of the dataentry window are returned as vectors in a list when the spreadsheet is closed.
de.ncols counts the number of columns which are supplied as arguments to data.entry. It attempts to count columns in lists, matrices and vectors. de.setup sets things up so that on return the columns can be regrouped and reassigned to the correct name. This is handled by de.restore.

## Value

de and dataentry return the edited value of their arguments. data.entry invisibly returns a vector of variable names but its main value is its side effect of assigning new version of those variables in the user's workspace.

## Note

The details of interface to the data grid may differ by platform and GUI. The following description applies to the X11-based implementation under Unix.

You can navigate around the grid using the cursor keys or by clicking with the (left) mouse button on any cell. The active cell is highlighted by thickening the surrounding rectangle. Moving to the right or down will scroll the grid as needed: there is no constraint to the rows or columns currently in use.
The are alternative ways to navigate using the keys. Return and (keypad) Enter and LineFeed all move down. Tab moves right and Shift-Tab move left. Home moves to the top left.
PageDown or Control-F moves down a page, and PageUp or Control-B up by a page. End will show the last used column and the last few rows used (in any column).

Using any other key starts an editing process on the currently selected cell: moving away from that cell enters the edited value whereas Esc cancels the edit and restores the previous
value. When the editing process starts the cell is cleared. In numerical columns (the default) only letters making up a valid number (including -.eE) are accepted, and entering an invalid edited value (such as blank) enters NA in that cell. The last entered value can be deleted using the BackSpace or Del(ete) key. Only a limited number of characters (currently 29) can be entered in a cell, and if necessary only the start or end of the string will be displayed, with the omissions indicated by > or <. (The start is shown except when editing.)
Entering a value in a cell further down a column than the last used cell extends the variable and fills the gap (if any) by NAs (not shown on screen).
The column names can only be selected by clicking in them. This gives a popup menu to select the column type (currently Real (numeric) or Character) or to change the name. Changing the type converts the current contents of the column (and converting from Character to Real may generate NAs.) If changing the name is selected the header cell becomes editable (and is cleared). As with all cells, the value is entered by moving away from the cell by clicking elsewhere or by any of the keys for moving down (only).

New columns are created by entering values in them (and not by just assigning a new name). The mode of the column is auto-detected from the first value entered: if this is a valid number it gives a numeric column. Unused columns are ignored, so adding data in var5 to a three-column grid adds one extra variable, not two.

The Copy button copies the currently selected cell: paste copies the last copied value to the current cell, and right-clicking selects a cell and copies in the value. Initially the value is blank, and attempts to paste a blank value will have no effect.
Control-L will refresh the display, recalculating field widths to fit the current entries.
In the default mode the column widths are chosen to fit the contents of each column, with a default of 10 characters for empty columns. you can specify fixed column widths by setting option de.cellwidth to the required fixed width (in characters). (set it to zero to return to variable widths). The displayed width of any field is limited to 600 pixels (and by the window width).

## See Also

vi, edit: edit uses dataentry to edit data frames.

## Examples

```
# call data entry with variables x and y
data.entry(x,y)
```


## date System Date and Time

## Description

Returns a character string of the current system date and time.

## Usage

date()

## Value

The string has the form "Fri Aug 20 11:11:00 1999", i.e. length 24, since it relies on POSIX' ctime ensuring the above fixed format. Timezone and Daylight Saving Time are taken account of, but not indicated in the result.

## Examples

```
(d <- date())
```

nchar $(\mathrm{d})==24$

```
DateTimeClasses Date-Time Classes
```


## Description

Description of the classes "POSIXlt" and "POSIXct" representing calendar dates and times (to the nearest second).

## Usage

```
print.POSIXct(x, ...)
print.POSIXlt(x, ...)
summary.POSIXct(object, digits = 15, ...)
summary.POSIXlt(object, digits = 15, ...)
time + number
time - number
time1 lop time2
```


## Arguments

\(\left.\begin{array}{ll}\mathrm{x}, object \& An object to be printed or summarized. <br>
digits \& Number of significant digits for the computations: should be high enough <br>

to represent the least important time unit exactly.\end{array}\right]\)| Further arguments to be passed from or to other methods. |  |
| :--- | :--- |
| $\ldots$ |  |
| time, time1, | time2 |
| number | date-time objects. |
| lop numeric object. |  |$\quad$| One of $==,!=,<,<=,>$ or $>=$. |
| :--- |

## Details

There are two basic classes of date/times. Class "POSIXct" represents the (signed) number of seconds since the beginning of 1970 as a numeric vector. Class "POSIXlt" is a named list of vectors representing
sec $0-61$ : seconds
min 0-59: minutes
hour 0-23: hours
mday 1-31: day of the month
mon $0-11$ : months after the first of the year.
year Years since 1900.
wday $0-6$ day of the week, starting on Sunday.
yday $0-365$ : day of the year.
isdst Daylight savings time flag. Positive if in force, zero if not, negative if unknown.
The classes correspond to the ANSI C constructs of "calendar time" (the time_t data type) and "local time" (or broken-down time, the struct tm data type), from which they also inherit their names.
"POSIXct" is more convenient for including in data frames, and "POSIXlt" is closer to human-readable forms. A virtual class "POSIXt" inherits from both of the classes: it is used to allow operations such as subtraction to mix the two classes.

Logical comparisons and limited arithmetic are available for both classes. One can add or subtract a number of seconds or a difftime object from a date-time object, but not add two date-time objects. Subtraction of two date-time objects is equivalent to using difftime. Be aware that "POSIXIt" objects will be interpreted as being in the current timezone for these operations, unless a timezone has been specified.
"POSIXlt" objects will often have an attribute "tzone", a character vector of length 3 giving the timezone name from the "TZ" environment variable and the names of the base timezone and the alternate (daylight-saving) timezone. Sometimes this may just be of length one, giving the timezone name.
Unfortunately, the conversion is complicated by the operation of time zones and leap seconds ( 22 days have been 86401 seconds long so far: the times of the extra seconds are in the object .leap.seconds). The details of this are entrusted to the OS services where possible. This will usually cover the period 1970-2037, and on Unix machines back to 1902 (when time zones were in their infancy). Outside those ranges we use our own C code. This uses the offset from GMT in use in the timezone in 2000, and uses the alternate (daylight-saving) timezone only if isdst is positive.
It seems that some systems use leap seconds but most do not. This is detected and corrected for at build time, so all "POSIXct" times used by R do not include leap seconds. (Conceivably this could be wrong if the system has changed since build time, just possibly by changing locales.)
Using c on "POSIXlt" objects converts them to the current time zone.

## Warning

Some Unix-like systems (especially Linux ones) do not have "TZ" set, yet have internal code that expects it (as does POSIX). We have tried to work around this, but if you get unexpected results try setting "TZ".

## See Also

as.POSIXct and as.POSIXlt for conversion between the classes.
strptime for conversion to and from character representations.
Sys.time for clock time as a "POSIXct" object.
difftime for time intervals.
cut.POSIXt, seq.POSIXt, round.POSIXt and trunc.POSIXt for methods for these classes. weekdays.POSIXt for convenience extraction functions.

## Examples

```
(z <- Sys.time()) # the current date, as class "POSIXct"
Sys.time() - 3600 # an hour ago
as.POSIXlt(Sys.time(), "GMT") # the current time in GMT
format(.leap.seconds) # all 22 leapseconds in your timezone
```

dcf

## Description

Reads or writes an R object from/to a file in Debian Control File format.

## Usage

```
read.dcf(file, fields=NULL)
write.dcf(x, file = "", append = FALSE,
            indent \(=0.1\) * getOption("width"),
            width \(=0.9 *\) getOption("width"))
```


## Arguments

file either a character string naming a file or a connection. " " indicates output to the console.
fields Fields to read from the DCF file. Default is to read all fields.
$\mathrm{x} \quad$ the object to be written, typically a data frame. If not, it is attempted to coerce x to a data frame.
append logical. If TRUE, the output is appended to the file. If FALSE, any existing file of the name is destroyed.
indent a positive integer specifying the indentation for continuation lines in output entries.
width a positive integer giving the target column for wrapping lines in the output.

## Details

DCF is a simple format for storing databases in plain text files that can easily be directly read and written by humans. DCF is used in various places to store R system information, like descriptions and contents of packages.
The DCF rules as implemented in $R$ are:

1. A database consists of one or more records, each with one or more named fields. Not every record must contain each field.
2. Regular lines start with a non-whitespace character.
3. Regular lines are of form tag:value, i.e., have a name tag and a value for the field, seperated by : (only the first : counts). The value can be empty (=whitespace only).
4. Lines starting with whitespace are continuation lines (to the preceding field) if at least one character in the line is non-whitespace.
5. Records are seperated by one or more empty (=whitespace only) lines.
read.dcf returns a character matrix with one line per record and one column per field. Leading and trailing whitespace of field values is ignored. If a tag name is specified, but the corresponding value is empty, then an empty string of length 0 is returned. If the tag name of a fields is never used in a record, then NA is returned.

See Also write.table.

## Examples

```
## Create a reduced version of the 'CONTENTS' file in package 'eda'
x <- read.dcf(file = system.file("CONTENTS", package = "eda"),
    fields = c("Entry", "Description"))
write.dcf(x)
```

debug Debug a function

## Description

Set or unset the debugging flag on a function.

## Usage

debug(fun)
undebug (fun)

## Arguments

fun any interpreted $R$ function.

## Details

When a function flagged for debugging is entered, normal execution is suspended and the body of function is executed one statement at a time. A new browser context is initiated for each step (and the previous one destroyed). Currently you can only debug functions that have bodies enclosed in braces. This is a bug and will be fixed soon. You take the next step by typing carriage return, $n$ or next. You can see the values of variables by typing their names. Typing c or cont causes the debugger to continue to the end of the function. You can debug new functions before you step in to them from inside the debugger. Typing $\mathbf{Q}$ quits the current execution and returns you to the top-level prompt. Typing where causes the debugger to print out the current stack trace (all functions that are active). If you have variables with names that are identical to the controls (eg. c or n ) then you need to use print(c) and print(n) to evaluate them.

## See Also

browser, traceback to see the stack after an Error: ... message; recover for another debugging approach.

```
debugger Post-Mortem Debugging
```


## Description

Functions to dump the evaluation environments (frames) and to examine dumped frames.

## Usage

dump.frames(dumpto = "last.dump", to.file = FALSE)
debugger (dump = last. dump)

## Arguments

| dumpto | a character string. The name of the object or file to dump to. |
| :--- | :--- |
| to.file | logical. Should the dump be to an $R$ object or to a file? |
| dump | An R dump object created by dump.frames. |

## Details

To use post-mortem debugging, set the option error to be a call to dump.frames. By default this dumps to an R object "last.dump" in the workspace, but it can be set to dump to a file (as dump of the object produced by a call to save). The dumped object contain the call stack, the active environments and the last error message as returned by geterrmessage.
When dumping to file, dumpto gives the name of the dumped object and the file name has .rda appended.

A dump object of class "dump.frames" can be examined by calling debugger. This will give the error message and a list of environments from which to select repeatedly. When an environment is selected, it is copied and the browser called from within the copy.
If dump.frames is installed as the error handler, execution will continue even in noninteractive sessions. See the examples for how to dump and then quit.

## Value

None.

## Note

Functions such as sys.parent and environment applied to closures will not work correctly inside debugger.

Of course post-mortem debugging will not work if R is too damaged to produce and save the dump, for example if it has run out of workspace.

## Author(s)

B. D. Ripley

## See Also

options for setting error options; recover is an interactive debugger working similarly to debugger but directly after the error occurs.

## Examples

```
options(error=quote(dump.frames("testdump", TRUE)))
f <- function() {
    g <- function() stop("test dump.frames")
    g()
}
f() # will generate a dump on file "testdump.rda"
options(error=NULL)
## possibly in another R session
load("testdump.rda")
debugger(testdump)
Available environments had calls:
1: f()
2: g()
3: stop("test dump.frames")
Enter an environment number, or 0 to exit
Selection: 1
Browsing in the environment with call:
f()
Called from: debugger.look(ind)
Browse[1]> ls()
[1] "g"
Browse[1]> g
function() stop("test dump.frames")
<environment: 759818>
Browse[1]>
Available environments had calls:
1: f()
2: g()
3: stop("test dump.frames")
Enter an environment number, or 0 to exit
Selection: 0
## A possible setting for non-interactive sessions
options(error=quote({dump.frames(to.file=TRUE); q()}))
```

Defunct Defunct Functions

## Description

The functions or variables listed here are no longer part of R as they are not needed (any more).

```
Usage
    .Defunct()
Version()
provide(package)
    .Provided
category(...)
dnchisq(.)
pnchisq(.)
qnchisq(.)
rnchisq(.)
print.anova.glm(.)
print.anova.lm(.)
print.tabular(.)
print.plot(.)
save.plot(.)
system.test(.)
dotplot(...)
stripplot(...)
getenv(...)
read.table.url(url, method,...)
scan.url(url, file = tempfile(), method, ...)
source.url(url, file = tempfile(), method, ...)
httpclient(url, port=80, error.is.fatal=TRUE, check.MIME.type=TRUE,
                                    file=tempfile(), drop.ctrl.z=TRUE)
parse.dcf(text = NULL, file = "", fields = NULL, versionfix = FALSE)
.Alias(expr)
reshapeWide(x, i = reshape.i, j = reshape.j, val = reshape.v,
    jnames = levels(j))
reshapeLong(x,jvars, ilev = row.names(x),
                                    jlev = names(x)[jvars], iname = "reshape.i",
    jname = "reshape.j", vname = "reshape.v")
piechart(x, labels = names(x), edges = 200, radius = 0.8,
    density = NULL, angle = 45, col = NULL, main = NULL, ...)
print.ordered(.)
.Dyn.libs
.lib.loc
```


## Details

.Defunct is the function to which defunct functions are set.
category has been an old-S function before there were factors; should be replaced by factor throughout!
The *chisq() functions now take an optional non-centrality argument, so the $*$ nchisq() functions are no longer needed.

The new function dev.print() should now be used for saving plots to a file or printing them.
provide and its object . Provided have been removed. They were never used for their intended purpose, to allow one package to subsume another.
dotplot and stripplot have been renamed to dotchart and stripchart, respectively.
getenv has been replaced by Sys.getenv.
*. url are replaced by calling read.table, scan or source on a url connection.
httpclient was used by the deprecated "socket" method of download.file.
parse.dcf has been replaced by read.dcf, which is much faster, but has a slightly different interface.
.Alias provided an unreliable way to create duplicate references to the same object. There is no direct replacement. Where multiple references to a single object are required for semantic reasons consider using environments or external pointers. There are some notes on http://developer.r-project.org.
reshape*, which were experimental, are replaced by reshape. This has a different syntax and allows multiple time-varying variables.
piechart is the old name for pie, but clashed with usage in Trellis.
.Dyn.libs and .lib.loc were internal variables used for storing and manipulating the information about packages with dynloaded shared libs, and the known R library trees. These are now dynamic variables which one can get or set using .dynLibs and .libPaths, respectively.

```
See Also
Deprecated
```


## delay Delay Evaluation

## Description

delay creates a promise to evaluate the given expression in the specified environment if its value is requested. This provides direct access to lazy evaluation mechanism used by R for the evaluation of (interpreted) functions.

## Usage

delay (x, env=.GlobalEnv)

## Arguments

```
x an expression.
env an evaluation environment
```


## Details

This is an experimental feature and its addition is purely for evaluation purposes.

## Value

A promise to evaluate the expression. The value which is returned by delay can be assigned without forcing its evaluation, but any further accesses will cause evaluation.

## Examples

```
x <- delay({
    for(i in 1:7)
        cat("yippee!\n")
    1 0
})
x^2#- yippee
x^2#- simple number
```

```
delete.response Modify Terms Objects
```


## Description

delete.response returns a terms object for the same model but with no response variable. drop.terms removes variables from the right-hand side of the model.
reformulate creates a formula from a character vector.

## Usage

```
delete.response(termobj)
reformulate(termlabels, response = NULL)
drop.terms(termobj, dropx = NULL, keep.response = FALSE)
```


## Arguments

termobj A terms object
termlabels character vector giving the right-hand side of a model formula.
response character string, symbol or call giving the left-hand side of a model formula.
dropx vector of positions of variables to drop from the right-hand side of the model.
keep.response Keep the response in the resulting object?

## Value

delete.response and drop.terms return a terms object.
reformulate returns a formula.

## See Also

## Examples

```
ff <- y ~ z + x + w
tt <- terms(ff)
tt
delete.response(tt)
drop.terms(tt, 2:3, keep.response = TRUE)
reformulate(attr(tt, "term.labels"))
## keep LHS :
reformulate("x*w", ff[[2]])
fS <- surv(ft, case) ~ a + b
reformulate(c("a", "b*f"), fS[[2]])
stopifnot(identical( ~ var, reformulate("var")),
    identical(~ a + b + c, reformulate(letters[1:3])),
    identical( y ~ a + b, reformulate(letters[1:2], "y"))
    )
```

demo Demonstrations of $R$ functionality

## Description

demo is a user-friendly interface to running some demonstration R scripts. demo() gives the list of available topics.

## Usage

```
demo(topic, device = getOption("device"),
    package = .packages(), lib.loc = NULL,
    character.only = FALSE, verbose = getOption("verbose"))
```


## Arguments

topic the topic which should be demonstrated. If omitted, the list of available topics is displayed.
device the graphics device to be used.
package a name or character vector giving the packages to look into for data sets. By default, all packages in the search path are used.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. If the default is used, the loaded packages are searched before the libraries.
character.only
logical; if TRUE, use topic as character string instead of name.
verbose a logical. If TRUE, additional diagnostics are printed.

## Details

If no topics are given, demo lists the available demos. The corresponding information is returned in an object of class "packageIQR". The structure of this class is experimental. In earlier versions of R , an empty character vector was returned along with listing available demos.

## See Also

 source which is called by demo.
## Examples

```
demo() # for attached packages
## All available demos:
demo(package = .packages(all.available = TRUE))
demo(lm.glm)
ch <- "scoping"
demo(ch, character = TRUE)
```

density

Kernel Density Estimation

## Description

The function density computes kernel density estimates with the given kernel and bandwidth.

## Usage

```
density(x, bw = "nrd0", adjust = 1,
    kernel = c("gaussian", "epanechnikov", "rectangular", "triangular",
            "biweight", "cosine", "optcosine"),
    window = kernel, width,
    give.Rkern = FALSE,
    n = 512, from, to, cut = 3, na.rm = FALSE)
```


## Arguments

$\mathrm{x} \quad$ the data from which the estimate is to be computed.
bw the smoothing bandwidth to be used. The kernels are scaled such that this is the standard deviation of the smoothing kernel. (Note this differs from the reference books cited below, and from S-PLUS.)
bw can also be a character string giving a rule to choose the bandwidth. See bw.nrd.
The specified (or computed) value of bw is multiplied by adjust.
adjust the bandwidth used is actually adjust*bw. This makes it easy to specify values like "half the default" bandwidth.
kernel, window
a character string giving the smoothing kernel to be used. This must be one of "gaussian", "rectangular", "triangular", "epanechnikov", "biweight", "cosine" or "optcosine", with default "gaussian", and may be abbreviated to a unique prefix (single letter).
"cosine" is smoother than "optcosine", which is the usual "cosine" kernel in the literature and almost MSE-efficient. However, "cosine" is the version used by $S$.

| width | this exists for compatibility with S ; if given, and bw is not, will set bw to width if this is a character string, or to a kernel-dependent multiple of width if this is numeric. |
| :---: | :---: |
| give.Rkern | logical; if true, no density is estimated, and the "canonical bandwidth" of the chosen kernel is returned instead. |
| n | the number of equally spaced points at which the density is to be estimated. When $\mathrm{n}>512$, it is rounded up to the next power of 2 for efficiency reasons (fft). |
| from, to | the left and right-most points of the grid at which the density is to be estimated. |
| cut | by default, the values of left and right are cut bandwidths beyond the extremes of the data. This allows the estimated density to drop to approximately zero at the extremes. |
| na.rm | logical; if TRUE, missing values are removed from x. If FALSE any missing values cause an error. |

## Details

The algorithm used in density disperses the mass of the empirical distribution function over a regular grid of at least 512 points and then uses the fast Fourier transform to convolve this approximation with a discretized version of the kernel and then uses linear approximation to evaluate the density at the specified points.
The statistical properties of a kernel are determined by $\sigma_{K}^{2}=\int t^{2} K(t) d t$ which is always $=1$ for our kernels (and hence the bandwidth bw is the standard deviation of the kernel) and $R(K)=\int K^{2}(t) d t$.
MSE-equivalent bandwidths (for different kernels) are proportional to $\sigma_{K} R(K)$ which is scale invariant and for our kernels equal to $R(K)$. This value is returned when give. Rkern $=$ TRUE. See the examples for using exact equivalent bandwidths.
Infinite values in x are assumed to correspond to a point mass at $+/-\operatorname{Inf}$ and the density estimate is of the sub-density on ( $-\operatorname{Inf},+\operatorname{Inf}$ ).

## Value

If give. Rkern is true, the number $R(K)$, otherwise an object with class "density" whose underlying structure is a list containing the following components.
$\mathrm{x} \quad$ the n coordinates of the points where the density is estimated.
$y \quad$ the estimated density values.
bw the bandwidth used.
N the sample size after elimination of missing values.
call the call which produced the result.
data.name the deparsed name of the x argument.
has.na logical, for compatibility (always FALSE).

## References

Scott, D. W. (1992) Multivariate Density Estimation. Theory, Practice and Visualization. New York: Wiley.

Sheather, S. J. and Jones M. C. (1991) A reliable data-based bandwidth selection method for kernel density estimation. J. Roy. Statist. Soc. B, 683-690.

Silverman, B. W. (1986) Density Estimation. London: Chapman and Hall.
Venables, W. N. and Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. New York: Springer.

See Also
bw.nrd, plot.density, hist.

## Examples

```
plot(density(c(-20,rep(0,98),20)), xlim = c(-4,4))# IQR = 0
# The Old Faithful geyser data
data(faithful)
d <- density(faithful$eruptions, bw = "sj")
d
plot(d)
plot(d, type = "n")
polygon(d, col = "wheat")
## Missing values:
x <- xx <- faithful$eruptions
x[i.out <- sample(length(x), 10)] <- NA
doR <- density(x, bw = 0.15, na.rm = TRUE)
lines(doR, col = "blue")
points(xx[i.out], rep(0.01, 10))
(kernels <- eval(formals(density)$kernel))
## show the kernels in the R parametrization
plot (density(0, bw = 1), xlab = "",
    main="R's density() kernels with bw = 1")
for(i in 2:length(kernels))
    lines(density(0, bw = 1, kern = kernels[i]), col = i)
legend(1.5,.4, legend = kernels, col = seq(kernels),
        lty = 1, cex = .8, y.int = 1)
## show the kernels in the S parametrization
plot(density(0, from=-1.2, to=1.2, width=2, kern="gaussian"), type="l",
    ylim = c(0, 1), xlab="", main="R's density() kernels with width = 1")
for(i in 2:length(kernels))
    lines(density(0, width=2, kern = kernels[i]), col = i)
legend(0.6, 1.0, legend = kernels, col = seq(kernels), lty = 1)
(RKs <- cbind(sapply(kernels, function(k)density(kern = k, give.Rkern = TRUE))))
100*round(RKs["epanechnikov",]/RKs, 4) ## Efficiencies
data(precip)
bw <- bw.SJ(precip) ## sensible automatic choice
plot(density(precip, bw = bw, n = 2^13),
    main = "same sd bandwidths, 7 different kernels")
for(i in 2:length(kernels))
    lines(density(precip, bw = bw, kern = kernels[i], n = 2^13), col = i)
## Bandwidth Adjustment for "Exactly Equivalent Kernels"
h.f <- sapply(kernels, function(k)density(kern = k, give.Rkern = TRUE))
```

```
(h.f <- (h.f["gaussian"] / h.f)^ .2)
## -> 1, 1.01, .995, 1.007,... close to 1 => adjustment barely visible..
plot(density(precip, bw = bw, n = 2^13),
    main = "equivalent bandwidths, 7 different kernels")
for(i in 2:length(kernels))
    lines(density(precip, bw = bw, adjust = h.f[i], kern = kernels[i],
        n = 2^13), col = i)
legend(55, 0.035, legend = kernels, col = seq(kernels), lty = 1)
```

    deparse Expression Deparsing
    
## Description

Turn unevaluated expressions into character strings.

## Usage

```
deparse(expr, width.cutoff = 60)
```


## Arguments

expr any $R$ expression.
width.cutoff integer in [20,500] determining the cutoff at which line-breaking is tried.

## Details

This function turns unevaluated expressions (where "expression" is taken in a wider sense than the strict concept of a vector of mode "expression" used in expression) into character strings (a kind of inverse parse).
A typical use of this is to create informative labels for data sets and plots. The example shows a simple use of this facility. It uses the functions deparse and substitute to create labels for a plot which are character string versions of the actual arguments to the function myplot.

## See Also

substitute, parse, expression.

## Examples

```
deparse(args(lm))
deparse(args(lm), width = 500)
myplot <-
function(x, y)
    plot(x, y, xlab=deparse(substitute(x)),
        ylab=deparse(substitute(y)))
```


## Deprecated Deprecated Functions

## Description

These functions are provided for compatibility with older versions of R only, and may be defunct as soon as of the next release.

## Usage

.Deprecated (new)

## Details

.Deprecated ("<new name>") is called from deprecated functions. The original help page for these functions is often available at help("oldName-deprecated") (note the quotes).
Machine() and Platform() are functions returning the variables .Machine and .Platform respectively.
restart() should be replaced by try(), in preparation for an exception-based implementation. If you use restart() in a way that cannot be replaced with try() then ask for help on r-devel.

See Also

Defunct,

## deriv Symbolic and Algorithmic Derivatives of Simple Expressions

## Description

Compute derivatives of simple expressions, symbolically.

## Usage

D (expr, name)
deriv(expr, namevec, function.arg, tag = ".expr", hessian = FALSE)
deriv3(expr, namevec, function.arg, tag = ".expr", hessian = TRUE)

## Arguments

expr expression or call to be differentiated.
name, namevec character vector, giving the variable names (only one for $D()$ ) with respect to which derivatives will be computed.
function.arg If specified, a character vector of arguments for a function return, or a function (with empty body) or TRUE, the latter indicating that a function with argument names namevec should be used.
tag character; the prefix to be used for the locally created variables in result.
hessian a logical value indicating whether the second derivatives should be calculated and incorporated in the return value.

## Details

D is modelled after its S namesake for taking simple symbolic derivatives.
deriv is a generic function with a default and a formula method. It returns a call for computing the expr and its (partial) derivatives, simultaneously. It uses so-called "algorithmic derivatives". If function.arg is a function, its arguments can have default values, see the fx example below.
Currently, deriv.formula just calls deriv.default after extracting the expression to the right of ${ }^{\sim}$.
deriv3 and its methods are equivalent to deriv and its methods except that hessian defaults to TRUE for deriv3.

## Value

D returns a call and therefore can easily be iterated for higher derivatives.
deriv and deriv3 normally return an expression object whose evaluation returns the function values with a "gradient" attribute containing the gradient matrix. If hessian is TRUE the evaluation also returns a "hessian" attribute containing the Hessian array.
If function.arg is specified, deriv and deriv3 return a function with those arguments rather than an expression.

## References

Griewank, A. and Corliss, G. F. (1991) Automatic Differentiation of Algorithms: Theory, Implementation, and Application. SIAM proceedings, Philadelphia.

## See Also

nlm and optim for numeric minimization which could make use of derivatives, nls in package nls.

## Examples

```
## formula argument :
dx2x <- deriv(~ x^2, "x") ; dx2x
expression({
    .value <- x^2
    .grad <- array(0, c(length(.value), 1), list(NULL, c("x")))
    .grad[, "x"] <- 2 * x
    attr(.value, "gradient") <- .grad
    .value
})
mode(dx2x)
x <- -1:2
eval(dx2x)
## Something 'tougher':
trig.exp <- expression(sin(cos(x + y^2)))
( D.sc <- D(trig.exp, "x") )
all.equal(D(trig.exp[[1]], "x"), D.sc)
( dxy <- deriv(trig.exp, c("x", "y")) )
y <- 1
eval(dxy)
```

```
eval(D.sc)
stopifnot(eval(D.sc) ==
    attr(eval(dxy),"gradient")[,"x"])
## function returned:
deriv((ff <- y ~ sin(cos(x) * y)), c("x","y"), func = TRUE)
stopifnot(all.equal(deriv(ff, c("x","y"), func = TRUE ),
                                    deriv(ff, c("x","y"), func = function(x,y){ } )))
## function with defaulted arguments:
(fx <- deriv(y ~ b0 + b1 * 2^ (-x/th), c("b0", "b1", "th"),
        function(b0, b1, th, x = 1:7){} ) )
fx(2,3,4)
## Higher derivatives
deriv3(y ~ b0 + b1 * 2^(-x/th), c("b0", "b1", "th"),
        c("b0", "b1", "th", "x") )
## Higher derivatives:
DD <- function(expr,name, order = 1) {
    if(order < 1) stop("'order' must be >= 1")
    if(order == 1) D(expr,name)
    else DD(D(expr, name), name, order - 1)
}
DD(expression(sin(x^2)), "x", 3)
## showing the limits of the internal "simplify()" :
-sin(x^2) * (2 * x) * 2 + ((cos(x^2) * (2 * x) * (2 * x) + sin(x^2) *
    2) * (2 * x) + sin(x^2) * (2 * x) * 2)
```

det
Calculate the Determinant of a Matrix

## Description

det calculates the determinant of a matrix either by QR decomposition or from the eigenvalues, see qr and eigen.

## Usage

$\operatorname{det}(x, \operatorname{method}=c(" q r ", \quad$ eigenvalues"), tol $=1 \mathrm{e}-07)$

## Arguments

| x | numeric matrix. |
| :--- | :--- |
| method | "qr" (default) or "eigenvalues". |

tol tolerance, used only for method "qr".

## Value

The determinant, or zero if qr determines the matrix to be numerically singular.

## Note

Often, computing the determinant is not what you should be doing to solve a given problem. The "qr" method is much faster for large matrices.

## See Also

eigen, qr, svd

## Examples

```
(x <- matrix(1:4, ncol=2))
det(x)
det(x, method="eigenvalues")
det(print(cbind(1,1:3,c(2,0,1))))
```

```
detach Detach Objects from the Search Path
```


## Description

Detach a database, i.e., remove it from the search() path of available R objects. Usually, this either a data.frame which has been attached or a package which was required previously.

## Usage

detach (name, pos = 2)

## Arguments

| name | The object to detach. Defaults to search () [pos]. This can be a name |
| :--- | :--- |
| or a character string but not a character vector. |  |
| pos | Index position in search() of database to detach. When name is numeric, |
| pos = name is used. |  |

## Value

The attached database is returned invisibly, either as data.frame or as list.

## Note

You cannot detach either the workspace (position 1) or the base package (the last item in the search list).

## See Also

attach, library, search, objects.

## Examples

```
require(eda)#package
detach(package:eda)
## could equally well use detach("package:eda")
## but NOT pkg <- "package:eda"; detach(pkg)
## Instead, use
library(eda)
pkg <- "package:eda"
detach(pos = match(pkg, search()))
library(mva)
detach(2)# 'pos' used for 'name'
```

dev.xxx Control Multiple Devices

## Description

These functions provide control over multiple graphics devices.
Only one device is the active device. This is the device in which all graphics operations occur.

Devices are associated with a name (e.g., "X11" or "postscript") and a number; the "null device" is always device 1 .
dev.off shuts down the specified (by default the current) device. graphics.off() shuts down all open graphics devices.
dev.set makes the specified device the active device.
A list of the names of the open devices is stored in .Devices. The name of the active device is stored in .Device.

## Usage

```
dev.cur()
dev.list()
dev.next(which = dev.cur())
dev.prev(which = dev.cur())
dev.off(which = dev.cur())
dev.set(which = dev.next())
graphics.off()
```


## Arguments

which An integer specifying a device number

## Value

dev.cur returns the number and name of the active device, or 1 , the null device, if none is active.
dev.list returns the numbers of all open devices, except device 1, the null device. This is a numeric vector with a names attribute giving the names, or NULL is there is no open device.
dev.next and dev.prev return the number and name of the next / previous device in the list of devices. The list is regarded as a circular list, and "null device" will be included only if there are no open devices.
dev. off returns the name and number of the new active device (after the specified device has been shut down).
dev. set returns the name and number of the new active device.

## See Also

Devices, such as postscript, etc; layout and its links for setting up plotting regions on the current device.

## Examples

```
## Unix-specific example
x11()
plot(1:10)
x11()
plot(rnorm(10))
dev.set(dev.prev())
abline(0,1)# through the 1:10 points
dev.set(dev.next())
abline(h=0, col="gray")# for the residual plot
dev.set(dev.prev())
dev.off(); dev.off()#- close the two X devices
```

dev2
Copy Graphics Between Multiple Devices

## Description

dev. copy copies the graphics contents of the current device to the device specified by which or to a new device which has been created by the function specified by device (it is an error to specify both which and device).
dev.print copies the graphics contents of the current device to a new device which has been created by the function specified by device and then shuts the new device.
dev.copy2eps is similar to dev.print but produces an EPSF output file, in portrait orientation (horizontal = FALSE)
dev.control allows the user to control the recording of graphics operations in a device. If displaylist is "inhibit" then recording is turned off.

```
Usage
    dev.copy(device, ..., which=dev.next())
dev.print(device=postscript, ...)
dev.copy2eps(...)
dev.control(displaylist)
```


## Arguments

device A device function (e.g., x11, postscript, ...)
... Arguments to the device function above. For dev.print, this includes which and by default any postscript arguments.
which A device number specifying the device to copy to
displaylist A character string: the only valid value is "inhibit".

## Details

For dev.copy2eps, width and height are taken from the current device unless otherwise specified. If just one of width and height is specified, the other is adjusted to preserve the aspect ratio of the device being copied. The default file name is Rplot.eps.
The default for dev.print is to produce and print a postscript copy, if options("printcmd") is set suitably.
dev.print is most useful for producing a postscript print (its default) when the following applies. Unless file is specified, the plot will be printed. Unless width, height and pointsize are specified the plot dimensions will be taken from the current device, shrunk if necessary to fit on the paper. (pointsize is rescaled if the plot is shrunk.) If horizontal is not specified and the plot can be printed at full size by switching its value this is done instead of shrinking the plot region.

If dev.print is used with a specified device (even postscript) it sets the width and height in the same way as dev.copy2eps.

## Value

dev. copy returns the name and number of the device which has been copied to.
dev.print and dev.copy2eps return the name and number of the device which has been copied from.

## Note

Most devices (including all screen devices) have a display list which records all of the graphics operations that occur in the device. dev. copy copies graphics contents by copying the display list from one device to another device. Also, automatic redrawing of graphics contents following the resizing of a device depends on the contents of the display list.
After the command dev. control("inhibit"), graphics operations are not recorded in the display list so that dev.copy and dev.print will not copy anything and the contents of a device will not be redrawn automatically if the device is resized.
The recording of graphics operations is relatively expensive in terms of memory so the command dev.control("inhibit") can be useful if memory usage is an issue.

## See Also

dev.cur and other dev.xxx functions

## Examples

```
x11()
plot(rnorm(10), main="Plot 1")
dev.copy(device=x11)
```

```
mtext("Copy 1", 3)
dev.print(width=6, height=6, horizontal=FALSE) # prints it
dev.off(dev.prev())
dev.off()
```

dev2bitmap Graphics Device for Bitmap Files via GhostScript

## Description

bitmap generates a graphics file. dev2bitmap copies the current graphics device to a file in a graphics format.

## Usage

```
bitmap(file, type = "png256", height = 6, width = 6, res = 72,
    pointsize, ...)
dev2bitmap(file, type = "png256", height = 6, width = 6, res = 72,
                    pointsize, ...)
```


## Arguments

| file | The output file name, with an appropriate extension. |
| :--- | :--- |
| type | The type of bitmap. the default is "png256". |
| height | The plot height, in inches. |
| width | The plot width, in inches. |
| res | Resolution, in dots per inch. <br> pointsize |
| The pointsize to be used for text: defaults to something reasonable given <br> the width and height |  |
| $\ldots$ | Other parameters passed to postscript. |

## Details

dev2bitmap works by copying the current device to a postscript device, and postprocessing the output file using ghostscript. bitmap works in the same way using a postscript device and postprocessing the output as "printing".
You will need a recent version of ghostscript ( 5.10 and later have been tested): the full path to the executable can be set by the environment variable "R_GSCMD".

The types available will depend on the version of ghostscript, but are likely to include "pcxmono", "pcxgray", "pcx16", "pcx256", "pcx24b", "pcxcmyk", "pbm", "pbmraw", "pgm", "pgmraw", "pgnm", "pgnmraw", "pnm", "pnmraw", "ppm", "ppmraw", "pkm", "pkmraw", "tiffcrle", "tiffg3", "tiffg32d", "tiffg4", "tifflzw", "tiffpack", "tiff12nc", "tiff24nc", "psmono", "psgray", "psrgb", "bit", "bitrgb", "bitcmyk", "pngmono", "pnggray", "png16", "png256", "png16m", "jpeg", "jpeggray", "pdfwrite".
Note: despite the name of the functions they can produce PDF via type = "pdfwrite", and the PDF produced is not bitmapped.
For dev2bitmap if just one of width and height is specified, the other is chosen to preserve aspect ratio of the device being copied.

## Value

None.

## Author(s)

B. D. Ripley

## See Also

postscript, png and jpeg and on Windows bmp.
pdf generate PDF directly.

```
deviance Model Deviance
```


## Description

Returns the deviance of a fitted model object.

## Usage

```
deviance(object, ...)
deviance.lm (object, ...)
deviance.glm(object, ...)
deviance.mlm(object, ...)
deviance.default(object, ...)
```


## Arguments

| object | an object for which the deviance is desired. |
| :--- | :--- |
| $\ldots$ | additional optional argument. |

## Details

This is a generic function which can be used to extract deviances for fitted models. Consult the individual modeling functions for details on how to use this function.

There is no default method for this function.

## Value

The value of the deviance extracted from the object object.

## See Also

df.residual, extractAIC, glm, lm.

Devices List of Graphical Devices

## Description

The following graphics devices are currently available:

- postscript Writes PostScript graphics commands to a file
- pdf Write PDF graphics commands to a file
- pictex Writes LaTeX/PicTeX graphics commands to a file
- xfig Device for XFIG graphics file format
- bitmap bitmap pseudo-device via GhostScript (if available).

The following devices will be available if R was compiled to use them and started with the appropriate --gui argument:

- X11 The graphics driver for the X11 Window system
- png PNG bitmap device
- jpeg JPEG bitmap device
- GTK, GNOME Graphics drivers for the GNOME GUI.

None of these are available under R CMD BATCH.

## Usage

```
X11(...)
postscript(...)
pdf(...)
pictex(...)
png(...)
jpeg(...)
GTK(...)
GNOME(...)
xfig(...)
bitmap(...)
dev.interactive()
```


## Details

If no device is open, using a high-level graphics function will cause a device to be opened. Which device is given by options("device") which is initially set as the most appropriate for each platform: a screen device in interactive use and postscript otherwise.

## Value

dev.interactive() returns a logical, TRUE iff an interactive (screen) device is in use.

## See Also

The individual help files for further information on any of the devices listed here;
dev.cur, dev.print, graphics.off, image, dev2bitmap. capabilities to see if X11, jpeg and png are available.

## Examples

```
## open the default screen device on this platform if no device is
## open
if(dev.cur() == 1) get(getOption("device"))()
```

```
df.residual Residual Degrees-of-Freedom
```


## Description

Returns the residual degrees-of-freedom extracted from a fitted model object.

## Usage

df.residual(object, ...)

## Arguments

> object an object for which the degrees-of-freedom are desired.
> $\ldots \quad$ additional optional arguments.

## Details

This is a generic function which can be used to extract residual degrees-of-freedom for fitted models. Consult the individual modeling functions for details details on how to use this function.

The default method just extracts the df.residual component.

## Value

The value of the residual degrees-of-freedom extracted from the object x .

## See Also

deviance, glm, lm.

## diag Matrix Diagonals

## Description

Extract or replace the diagonal of a matrix, or construct a diagonal matrix.

## Usage

diag(x = 1, nrow, ncol)
diag(x) <- value

## Arguments

$x \quad$ a matrix, vector or 1D array.
nrow, ncol Optional dimensions for the result.

## Value

If x is a matrix then $\operatorname{diag}(\mathrm{x})$ returns the diagonal of x . The resulting vector will have names if the matrix $x$ has matching column and row names.
If $x$ is a vector (or 1D array) of length two or more, then $\operatorname{diag}(x)$ returns a diagonal matrix whose diagonal is x .

If x is a vector of length one then $\operatorname{diag}(\mathrm{x})$ returns an identity matrix of order the nearest integer to $x$. The dimension of the returned matrix can be specified by nrow and ncol (the default is square).

The assignment form sets the diagonal of the matrix x to the given value(s).

## Note

Using diag ( x ) can have unexpected effects if x is a vector that could be of length one. Use $\operatorname{diag}(x$, nrow $=$ length $(x))$ for consistent behaviour.

## See Also

## Examples

```
dim(diag(3))
diag(10,3,4) # guess what?
all(diag(1:3) == {m <- matrix(0,3,3); diag(m) <- 1:3; m})
diag(var(M <- cbind(X=1:5, Y=rnorm(5))))#-> vector with names "X" and "Y"
rownames(M) <- c(colnames(M),rep("",3));
M; diag(M) # named as well
```


## diff

Lagged Differences

## Description

Returns suitably lagged and iterated differences.

## Usage

diff(x, ...)
diff.default(x, lag=1, differences=1, ...)

## Arguments

$x \quad$ a numeric vector or matrix containing the values to be differenced.
lag an integer indicating which lag to use.
differences an integer indicating the order of the difference.
... further arguments to be passed to or from methods.

## Details

diff is a generic function with a default method and one for class ts objects. NA's propagate.

## Value

If x is a vector of length n and differences $=1$, then the computed result is equal to the successive differences $x[(1+l a g): n]-x[1:(n-l a g)]$.
If difference is larger than one this algorithm is applied recursively to $x$. Note that the returned value is a vector which is shorter than x .

If x is a matrix then the difference operations are carried out on each column separately.

## See Also

diff.ts.

## Examples

```
diff(1:10, 2)
diff(1:10, 2, 2)
x <- cumsum(cumsum(1:10))
stopifnot(diff(x, lag = 2) == x[(1+2):10] - x[1:(10 - 2)],
    diff(x, lag = 2) == (3:10)^2,
    diff(diff(x)) == diff(x, differences = 2))
```

```
difftime Time Intervals
```


## Description

Create, print and round time intervals.

## Usage

```
time1 - time2
difftime(time1, time2, tz = "",
    units = c("auto", "secs", "mins", "hours", "days", "weeks"))
round(x, digits = 0)
```


## Arguments

```
time1, time2 date-time objects.
```

tz A timezone specification to be used for the conversion. System-specific, but "" is the current time zone, and "GMT" is UTC.
units character. Units in which the results are desired. Can be abbreviated.
x
An object inheriting from class "difftime".
digits integer. Number of significant digits to retain.

## Details

Function difftime takes a difference of two date/time objects (of either class) and returns an object of class "difftime" with an attribute indicating the units. There is a round method for objects of this class.

If units = "auto", a suitable set of units is chosen, the largest possible (excluding "weeks") in which all the absolute differences are greater than one.

Subtraction of two date-time objects gives an object of this class, by calling difftime with units="auto".

## See Also

DateTimeClasses.

## Examples

```
(z <- Sys.time() - 3600)
Sys.time() - z # just over 3600 seconds.
## time interval between releases of 1.2.2 and 1.2.3.
ISOdate(2001, 4, 26) - ISOdate(2001, 2, 26)
```

```
dim Dimensions of an Object
```


## Description

Retrieve or set the dimension of an object.

## Usage

```
dim(x)
dim(x) <- values
```


## Arguments

$x \quad$ an $R$ object, for example a matrix, array or data frame.

## Details

The functions dim and dim<- are generic.
For an array (and hence in particular, for a matrix) they retrieve or set the dim attribute of the object. It is always integer or NULL.
dim has a method for data.frames, which returns the length of the row.names attribute of x and the length of x (the numbers of "rows" and "columns").

## See Also

ncol, nrow and dimnames.

## Examples

```
x <- 1:12 ; dim(x) <- c(3,4)
x
# simple versions of nrow and ncol could be defined as follows
nrow0 <- function(x) dim(x)[1]
ncol0 <- function(x) dim(x)[2]
```

dimnames Dimnames of an Object

## Description

Retrieve or set the dimnames of an object.

## Usage

```
dimnames(x)
dimnames(x) <- nlist
```


## Arguments

$\begin{array}{ll}\mathrm{x} & \text { an } \mathrm{R} \text { object, for example a matrix, array or data frame. } \\ \mathrm{nlist} & \begin{array}{l}\text { a list of the length } \operatorname{dim}(\mathrm{x}) \text { whose components are either null or character } \\ \text { vectors the length of the appropriate dimension of } \mathrm{x} .\end{array}\end{array}$

## Details

The functions dimnames and dimnames<- are generic.
For an array (and hence in particular, for a matrix), they retrieve or set the dimnames attribute (see attributes) of the object. The list nlist can have names, and these will be used to label the dimensions of the array where appropriate.
Both have methods for data frames. The dimnames of a data frame are its row.names attribute and its names.

## See Also

```
rownames, colnames; array, matrix, data.frame.
```


## Examples

```
## simple versions of rownames and colnames
## could be defined as follows
rownames0 <- function(x) dimnames(x) [[1]]
colnames0 <- function(x) dimnames(x)[[2]]
```

discoveries Numbers of Important Discoveries

## Description

The numbers of "great" inventions and scientific discoveries in each year from 1860 to 1959.

## Usage

```
data(discoveries)
```


## Format

A time series of 100 values.

## Source

The World Almanac and Book of Facts, 1975 Edition, pages 315-318.

## References

McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(discoveries)
plot(discoveries, ylab = "Number of important discoveries",
    las = 1)
title(main = "discoveries data set")
```

```
do.call
Execute a Function Call
```


## Description

do. call executes a function call from the name of the function and a list of arguments to be passed to it.

## Usage

do.call(what, args)

## Arguments

| what | a character string naming the function to be called. |
| :--- | :--- |
| args | a list of arguments to the function call. The names attribute of args gives |
| the argument names. |  |

## Value

The result of the (evaluated) function call.

## See Also

call which creates an unevaluated call.

## Examples

```
do.call("complex", list(imag = 1:3))
```

dotchart Cleveland Dot Plots

## Description

Draw a Cleveland dot plot.

Usage

```
dotchart(x, labels = NULL, groups = NULL, gdata = NULL,
    cex = par("cex"), pch = 21, gpch = 21, bg = par("bg"),
    color = par("fg"), gcolor = par("fg"), lcolor = "gray",
    xlim = range(x[is.finite(x)]),
    main = NULL, xlab = NULL, ylab = NULL, ...)
```


## Arguments

X
either a vector or matrix of numeric values (NAs are allowed). If $x$ is a matrix the overall plot consists of juxtaposed dotplots for each row.
labels a vector of labels for each point. For vectors the default is to use names (x) and for matrices the row labels dimnames (x) [[1] ].
groups an optional factor indicating how the elements of x are grouped. If x is a matrix, groups will default to the columns of x .
gdata data values for the groups. This is typically a summary such as the median or mean of each group.
cex the character size to be used. Setting cex to a value smaller than one can be a useful way of avoiding label overlap.
pch the plotting character or symbol to be used.
gpch the plotting character or symbol to be used for group values.
bg the background color to be used.
color the color(s) to be used for points an labels.
gcolor the single color to be used for group labels and values.
lcolor the color(s) to be used for the horizontal lines.
xlim horizontal range for the plot, see plot.window, e.g.
main overall title for the plot, see title.
xlab, ylab
axis annotations as in title.
graphical parameters can also be specified as arguments.

## Value

This function is invoked for its side effect, which is to produce two variants of dotplots as described in Cleveland (1985).

Dot plots are a reasonable substitute for bar plots.

## References

Cleveland, W. S. (1985) The Elements of Graphing Data. Monterey, CA: Wadsworth.

## Examples

```
data(VADeaths)
dotchart(VADeaths, main = "Death Rates in Virginia - 1940")
op <- par(xaxs="i")# 0 -- 100%
dotchart(t(VADeaths), xlim = c(0,100),
    main = "Death Rates in Virginia - 1940")
par(op)
```

double
Double Precision Vectors

## Description

Create, coerce to or test for a double-precision vector.

## Usage

```
double(length = 0)
as.double(x, ...)
is.double(x)
single(length = 0)
as.single(x, ...)
```


## Arguments

| length | desired length. |
| :--- | :--- |
| x | object to be coerced or tested. |
| $\ldots$ | further arguments passed to or from other methods. |

## Value

double creates a double precision vector of the specified length. The elements of the vector are all equal to 0 .
as. double attempts to coerce its argument to be of double type.
is. double returns TRUE or FALSE depending on whether its argument is of double type or not.

## Note

R has no single precision data type. All real numbers are stored in double precision format. The functions as.single and single are identical to as.double and double except they set the attribute Csingle that is used in the .C and .Fortran interface, and they are intended only to be used in that context.

## See Also

integer.

## Examples

```
is.double(1)
all(double(3) == 0)
```

download.file Download File from the Internet

## Description

This function can be used to download a file from the Internet.

## Usage

```
download.file(url, destfile, method, quiet = FALSE, mode="w",
                        cacheOK = TRUE)
```


## Arguments

| url | A character string naming the URL of a resource to be downloaded. |
| :--- | :--- |
| destfile | A character string with the name where the downloaded file is saved. |
| Tilde-expansion is performed. |  |
| method | Method to be used for downloading files. Currently download methods <br> "internal", "wget" and "lynx" are available. The default is to choose <br> the first of these which will be "internal". The method can also be set <br> through the option "download.file.method": see options(). |
| quiet | If TRUE, suppress status messages (if any). <br> character. The mode with which to write the file. Useful values are "w", |
|  | "wb" (binary), "a" (append) and "ab". Only used for the "internal" <br> method. |
| cache0K | logical. Is a server-side cached value acceptable? Implemented for the <br> "internal" and "wget" methods. |

## Details

The function download.file can be used to download a single file as described by url from the internet and store it in destfile. The url must start with a scheme such as "http://", "ftp://" or "file://".
cacheOK = FALSE is useful for "http://" URLs, and will attempt to get a copy directly from the site rather than from an intermediate cache. (Not all platforms support it.) It is used by CRAN. packages.
The remaining details apply to method "internal" only.
The timeout for many parts of the transder can be set by the option timeout which defaults to 60 seconds.

The level of detail provided during transfer can be set by the quiet argument and the internet.info option. The details depend on the platform and scheme, but setting internet.info to 0 gives all available details, including all server responses. Using 2 (the default) gives only serious messages, and 3 or more suppresses all messages.
Method "wget" can be used with proxy firewalls which require user/password authentication if proper values are stored in the configuration file for wget.

## Setting Proxies

This applies to the internal code only.
Proxies can be specified via environment variables. Setting "no_proxy" stops any proxy being tried. Otherwise the setting of "http_proxy" or "ftp_proxy" (or failing that, the all upper-case version) is consulted and if non-empty used as a proxy site. For FTP transfers, the username and password on the proxy can be specified by "ftp_proxy_user" and "ftp_proxy_password". The form of "http_proxy" should be "http://proxy.dom.com/" or "http://proxy.dom.com:8080/" where the port defaults to 80 and the trailing slash may be omitted. For "ftp_proxy" use the form "ftp://proxy.dom.com:3128/" where the default port is 21 . These environment variables must be set before the download code is first used: they cannot be altered later by calling Sys.putenv.

## Note

Methods "wget" and "lynx" are for historical compatibility. They will block all other activity on the R process.

For methods "wget" and "lynx" a system call is made to the tool given by method, and the respective program must be installed on your system and be in the search path for executables.

## See Also

options to set the timeout and internet.info options.
url for a finer-grained way to read data from URLs.
url.show, CRAN.packages, download.packages for applications

```
dput
Write an Internal Object to a File
```


## Description

Writes an ASCII text representation of an R object to a file or connection, or uses one to recreate the object.

## Usage

dput(x, file = "")
$\operatorname{dget}(f i l e)$

## Arguments

x
file either a character string naming a file or a connection. " " indicates output to the console.

## Details

dput opens file and deparses the object x into that file. The object name is not written (contrary to dump). If x is a function the associated environment is stripped. Hence scoping information can be lost.

Using dget, the object can be recreated (with the limitations mentioned above).
dput will warn if fewer characters were written to a file than expected, which may indicate a full or corrupt file system.

## See Also

deparse, dump, write.

## Examples

```
## Write an ASCII version of mean to the file "foo"
dput(mean, "foo")
## And read it back into 'bar'
bar <- dget("foo")
unlink("foo")
```

drop Drop Redundant Extent Information

## Description

Delete the dimensions of an array which have only one level.

## Usage

drop(x)

## Arguments

$\mathrm{x} \quad$ an array (including a matrix).

## Value

If $x$ is an object with a dim attribute (e.g., a matrix or array), then drop returns an object like $x$, but with any extents of length one removed. Any accompanying dimnames attribute is adjusted and returned with x .
Array subsetting ([) performs this reduction unless used with drop = FALSE, but sometimes it is useful to invoke drop directly.

## See Also

drop1 which is used for dropping terms in models.

## Examples

```
dim(drop(array(1:12, dim=c(1,3,1,1,2,1,2))))# = 3 2 2
drop(1:3 %*% 2:4)# scalar product
```


## Description

This extracts coefficients in terms of the original levels of the coefficients rather than the coded variables.

## Usage

```
dummy.coef(object, ...)
dummy.coef.lm(object, use.na = FALSE, ...)
dummy.coef.aovlist(object, use.na = FALSE, ...)
```


## Arguments

object a linear model fit.
use.na logical flag for coefficients in a singular model. If use.na is true, undetermined coefficients will be missing; if false they will get one possible value.
... arguments passed to or from other methods.

## Details

A fitted linear model has coefficients for the contrasts of the factor terms, usually one less in number than the number of levels. This function re-expresses the coefficients in the original coding; as the coefficients will have been fitted in the reduced basis, any implied constraints (e.g. zero sum for contr.helmert or contr.sum will be respected. There will be little point in using dummy.coef for contr.treatment contrasts, as the missing coefficients are by definition zero.
The method used has some limitations, and will give incomplete results for terms such as $\operatorname{poly}(\mathrm{x}, 2))$. However, it is adequate for its main purpose, aov models.

## Value

A list giving for each term the values of the coefficients. For a multistratum aov model, such a list for each stratum.

## Warning

This function is intended for human inspection of the output: it should not be used for calculations. Use coded variables for all calculations.

The results differ from $S$ for singular values, where $S$ can be incorrect.

## Author(s)

B.D. Ripley

## See Also

aov, model.tables

## Examples

```
options(contrasts=c("contr.helmert", "contr.poly"))
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
    K=factor(K), yield=yield)
npk.aov <- aov(yield ~ block + N*P*K, npk)
dummy.coef(npk.aov)
npk.aovE <- aov(yield ~ N*P*K + Error(block), npk)
dummy.coef (npk.aovE)
```

dump Text Representations of $R$ Objects

## Description

This function takes a vector of names of $R$ objects and produces text representations of the objects on a file or connection. A dump file can be sourced into another R (or S) session.

## Usage

dump(list, file="dumpdata.R", append=FALSE)

## Arguments

| list | character. The names of one or more R objects to be dumped. |
| :--- | :--- |
| file | either a character string naming a file or a connection. " " indicates output <br> to the console. |
| append | if TRUE, output will be appended to file; otherwise, it will overwrite the <br> contents of file. |

## Details

At present the implementation of dump is very incomplete and it really only works for functions and simple vectors.
dump will warn if fewer characters were written to a file than expected, which may indicate a full or corrupt file system.
The function save is designed to be used for transporting $R$ data between machines.

## See Also

dput, dget,write.

## Examples

```
x <- 1; y <- 1:10
dump(ls(patt='^[xyz]'), "xyz.Rdmped")
unlink("xyz.Rdmped")
```

duplicated Determine Duplicate Elements

## Description

Determines which elements of a vector of data frame are duplicates of elements with smaller subscripts, and returns a logical vector indicating which elements (rows) are duplicates.

## Usage

duplicated(x, incomparables = FALSE, ...)
duplicated.array(x, incomparables = FALSE, MARGIN = 1, ...)

## Arguments

x
incomparables
an atomic vector or a data frame or an array.
incomparables possible value, meaning that all values can be compared.
... arguments for particular methods.
MARGIN the array margin to be held fixed: see apply.

## Details

This is a generic function with methods for vectors, data frames and arrays (including matrices).

The data frame method works by pasting together a character representation of the rows separated by
$r$, so may be imperfect if the data frame has characters with embedded carriage returns or columns which do not reliably map to characters.

The array method calculates for each element of the sub-array specified by MARGIN if the remaining dimensions are identical to those for an earlier element (in row-major order). This would most commonly be used to find duplicated rows (the default) or columns (with MARGIN = 2).

## See Also

unique.

## Examples

```
x <- c(9:20, 1:5, 3:7, 0:8)
## extract unique elements
(xu <- x[!duplicated(x)])
stopifnot(xu == unique(x), # but unique(x) is more efficient
    0:20 == sort(x[!duplicated(x)]))
data(iris)
stopifnot(duplicated(iris)[143] == TRUE)
data(iris3)
duplicated(iris3, MARGIN = c(1, 3))
```

dyn.load Foreign Function Interface

## Description

Load or unload shared libraries, and test whether a C function or Fortran subroutine is available.

## Usage

```
dyn.load(x, local = TRUE, now = TRUE)
dyn.unload(x)
is.loaded(symbol, PACKAGE="")
symbol.C(name)
symbol.For(name)
```


## Arguments

| x |  |
| :--- | :--- |
| local | a character string giving the pathname to a shared library or DLL. |
| a logical value controlling whether the symbols in the shared library are |  |
| stored in their own local table and not shared across shared libraries, or |  |
| added to the global symbol table. Whether this has any effect is system- |  |
| dependent. |  |
| now |  |
| a logical controlling whether all symbols are resolved (and relocated) im- |  |
| mediately the library is loaded or deferred until they are used. This |  |
| control is useful for developers testing whether a library is complete and |  |
| has all the necessary symbols and for users to ignore missing symbols. |  |
| Whether this has any effect is system-dependent. |  |

## Details

See 'See Also' and the Writing $R$ Extensions manual for how to create a suitable shared library. Note that unlike some versions of S-PLUS, dyn.load does not load an object (.o) file but a shared library or DLL.
Unfortunately a very few platforms (Compaq Tru64) do not handle the PACKAGE argument correctly, and may incorrectly find symbols linked into R.
The additional arguments to dyn.load mirror the different aspects of the mode argument to the dlopen() routine on UNIX systems. They are available so that users can exercise greater control over the loading process for an individual library. In general, the defaults values are appropriate and one should override them only if there is good reason and you understand the implications.
The local argument allows one to control whether the symbols in the DLL being attached are visible to other DLLs. While maintaining the symbols in their own namespace is good practice, the ability to share symbols across related "chapters" is useful in many cases. Additionally, on certain platforms and versions of an operating system, certain libraries must have their symbols loaded globally to successfully resolve all symbols.

One should be careful of the potential side-effect of using lazy loading via the 'now' argument as FALSE. If a routine is called that has a missing symbol, the process will terminate immediately and unsaved session variables will be lost. The intended use is for library developers to call specify a value TRUE to check that all symbols are actually resolved and for regular users to all with FALSE so that missing symbols can be ignored and the available ones can be called.
The initial motivation for adding these was to avoid such termination in the _init () routines of the Java virtual machine library. However, symbols loaded locally may not be (read probably) available to other DLLs. Those added to the global table are available to all other elements of the application and so can be shared across two different DLLs.
Some systems do not provide (explicit) support for local/global and lazy/eager symbol resolution. This can be the source of subtle bugs. One can arrange to have warning messages emitted when unsupported options are used. This is done by setting either of the options verbose or warn to be non-zero via the options function. Currently, we know of only 2 platforms that do not provide a value for local load (RTLD_LOCAL). These are IRIX6.4 and unpatched versions of Solaris 2.5.1.

There is a short discussion of these additional arguments with some example code available at http://cm.bell-labs.com/stat/duncan/R/dynload.

## Value

The function dyn.load is used for its side effect which links the specified shared library to the executing R image. Calls to .C, .Fortran and .External can then be used to execute compiled C functions or Fortran subroutines contained in the library.
The function dyn.unload unlinks the shared library.
Functions symbol.C and symbol. For map function or subroutine names to the symbol name in the compiled code: is.loaded checks if the symbol name is loaded and hence available for use in .C or .Fortran.

## Note

The creation of shared libraries and the runtime linking of them into executing programs is very platform dependent. In recent years there has been some simplification in the process because the C subroutine call dlopen has become the standard for doing this under UNIX.

Under UNIX dyn.load uses the dlopen mechanism and should work on all platforms which support it. On Windows it uses the standard mechanisms for loading 32-bit DLLs.

The original code for loading DLLs in UNIX was provided by Heiner Schwarte. The compatibility code for HP-UX was provided by Luke Tierney.

## See Also

library.dynam to be used inside a package's .First.lib initialization. SHLIB for how to create suitable shared libraries.
.C, .Fortran, .External, .Call.

## Examples

```
is.loaded(symbol.For("hcass2")) #-> probably FALSE
library(mva)
is.loaded(symbol.For("hcass2")) #-> TRUE
```


## Description

Invoke a text editor on an R object.

## Usage

```
edit(name = NULL, file = "", editor = getOption("editor"), ...)
vi(name = NULL, file = "")
emacs(name = NULL, file = "")
pico(name = NULL, file = "")
xemacs(name = NULL, file = "")
xedit(name = NULL, file = "")
```


## Arguments

name a named object that you want to edit. If name is missing then the file specified by file is opened for editing.
file a string naming the file to write the edited version to.
editor a string naming the text editor you want to use. On Unix the default is set from the environment variables EDITOR or VISUAL if either is set, otherwise vi is used. On Windows it defaults to notepad.
... further arguments to be passed to or from methods.

## Details

edit invokes the text editor specified by editor with the object name to be edited. It is a generic function, currently with a default method and one for data frames.
data.entry can be used to edit data, and is used by edit to edit matrices and data frames on systems for which data.entry is available.

It is important to realize that edit does not change the object called name. Instead, a copy of name is made and it is that copy which is changed. Should you want the changes to apply to the object name you must assign the result of edit to name. (Try fix if you want to make permanent changes to an object.)
In the form edit (name), edit deparses name into a temporary file and invokes the editor editor on this file. Quitting from the editor causes file to be parsed and that value returned. Should an error occur in parsing, possibly due to incorrect syntax, no value is returned. Calling edit(), with no arguments, will result in the temporary file being reopened for further editing.

## Note

The functions vi, emacs, pico, xemacs, xedit rely on the corresponding editor being available and being on the path. This is system-dependent.

## See Also

```
edit.data.frame, data.entry, fix.
```


## Examples

```
# use xedit on the function mean and assign the changes
mean <- edit(mean, editor = "xedit")
# use vi on mean and write the result to file mean.out
vi(mean, file = "mean.out")
```

```
edit.data.frame Edit Data Frames and Matrices
```


## Description

Use data editor on data frame or matrix contents.

```
Usage
    edit(name, factor.mode = c("character", "numeric"),
    edit.row.names = any(row.names(name) != 1:nrow(name)),
    ...)
edit(name, edit.row.names = any(rownames(name) != 1:nrow(name))), ...)
```


## Arguments

```
name A data frame or matrix.
factor.mode How to handle factors (as integers or using character levels) in a data
frame.
edit.row.names
logical. Show the row names be displayed as a separate editable column?
... further arguments passed to or from other methods.
```


## Details

At present, this only works on simple data frames containing numeric or character vectors and factors. Factors are represented in the spreadsheet as either numeric vectors (which is more suitable for data entry) or character vectors (better for browsing). After editing, vectors are padded with NA to have the same length and factor attributes are restored. The set of factor levels can not be changed by editing in numeric mode; invalid levels are changed to NA and a warning is issued. If new factor levels are introduced in character mode, they are added at the end of the list of levels in the order in which they encountered.

It is possible to use the data-editor's facilities to select the mode of columns to swap between numerical and factor columns in a data frame. Changing any column in a numerical matrix to character will cause the result to be coerced to a character matrix.

## Value

The edited data frame.

## Note

fix(dataframe) works for in-place editing by calling this function.
If the data editor is not available, a dump of the object is presented for editing using the default method of edit.

## Author(s)

Peter Dalgaard

```
See Also
data.entry, edit
```


## Examples

```
data(InsectSprays)
edit(InsectSprays)
edit(InsectSprays, factor.mode="numeric")
```


## Description

Computes the efficiencies of fixed-effect terms in an analysis of variance model with multiple strata.

## Usage

eff.aovlist(aovlist)

## Arguments

aovlist The result of a call to aov with a Error term.

## Details

Fixed-effect terms in an analysis of variance model with multiple strata may be estimable in more than one stratum, in which case there is less than complete information in each. The efficiency is the fraction of the maximum possible precision (inverse variance) obtainable by estimating in just that stratum.

This is used to pick strata in which to estimate terms in model.tables.aovlist and elsewhere.

## Value

A matrix giving for each non-pure-error stratum (row) the efficiencies for each fixed-effect term in the model.

## Author(s)

B.D. Ripley

## See Also

aov, model.tables.aovlist, se.contrast.aovlist

## Examples

```
## for balanced designs all efficiencies are zero or one.
## so as a statistically meaningless test:
options(contrasts=c("contr.helmert", "contr.poly"))
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
    K=factor(K), yield=yield)
npk.aovE <- aov(yield ~ N*P*K + Error(block), npk)
eff.aovlist(npk.aovE)
```


## effects Effects from Fitted Model

## Description

Returns (orthogonal) effects from a fitted model, usually a linear model. This is a generic function, but currently only has a method for objects inheriting from class "lm".

## Usage

```
effects(object, ...)
effects.lm(object, set.sign=FALSE, ...)
```


## Arguments

object an R object; typically, the result of a model fitting function such as 1 m .
set.sign logical. If TRUE, the sign of the effects corresponding to coefficients in the model will be set to agree with the signs of the corresponding coefficients, otherwise the sign is arbitrary.
... arguments passed to or from other methods.

## Details

For a linear model fitted by lm or aov, the effects are the uncorrelated single-degree-offreedom values obtained by projecting the data onto the successive orthogonal subspaces generated by the QR decomposition during the fitting process. The first $r$ (the rank of the model) are associated with coefficients and the remainder span the space of residuals (but are not associated with particular residuals).

## Value

A (named) numeric vector of the same length as residuals, or a matrix if there were multiple responses in the fitted model, in either case of class "coef".

The first $r$ rows are labelled by the corresponding coefficients, and the remaining rows are unlabelled. Note that in rank-deficient models the "corresponding" coefficients will be in a different order if pivoting occurred.

## See Also

```
coef
```


## Examples

```
y <- c(1:3,7,5)
x <- c(1:3,6:7)
( ee <- effects(lm(y ~ x)) )
c(round(ee - effects(lm(y+10 ~ I(x-3.8))),3))# just the first is different
```

eigen Spectral Decomposition of a Matrix

## Description

Function eigen computes eigenvalues and eigenvectors by providing an interface to the EISPACK routines RS, RG, CH and CG.

Function La.eigen uses the LAPACK routines DSYEV/DSYEVR, DGEEV, ZHEEV and ZGEEV.

## Usage

```
eigen(x, symmetric, only.values = FALSE)
La.eigen(x, symmetric, only.values = FALSE,
    method = c("dsyevr", "dsyev"))
```


## Arguments

| x | a matrix whose spectral decomposition is to be computed. |
| :--- | :--- |
| symmetric | if TRUE, the matrix is assumed to be symmetric (or Hermitian if complex) <br> and only its lower triangle is used. If symmetric is not specified, the <br> matrix is inspected for symmetry. |
| only.values | if TRUE, only the eigenvalues are computed and returned, otherwise both <br> eigenvalues and eigenvectors are returned. |
| method | The LAPACK routine to use in the real symmetric case. |

## Details

If symmetric is unspecified, the code attempts to determine if the matrix is symmetric up to plausible numerical inaccuracies. It is faster and surer to set the value yourself.

La.eigen is preferred to eigen for new projects, but its eigenvectors may differ in sign and (in the asymmetric case) in normalization. (They may also differ between methods and between platforms.)

The LAPACK routine DSYEVR is usually substantially faster than DSYEV: see http:// www.cs.berkeley.edu/~demmel/DOE2000/Report0100.html. Most benefits are seen with an optimized BLAS system.
Computing the eigenvectors is the slow part for large matrices.
Using method="dsyevr" requires IEEE 754 arithmetic. Should this not be supported on your platform, method="dsyev" is used, with a warning.

## Value

The spectral decomposition of x is returned as components of a list.
values a vector containing the $p$ eigenvalues of x , sorted in decreasing order, according to Mod(values) if they are complex.
vectors a $p \times p$ matrix whose columns contain the eigenvectors of x , or NULL if only.values is TRUE.
For eigen (, symmetric = FALSE) the choice of length of the eigenvectors is not defined by LINPACK. In all other cases the vectors are normalized to unit length.
Recall that the eigenvectors are only defined up to a constant: even when the length is specified they are still only defined up to a scalar of modulus one (the sign for real matrices).

## References

Smith, B. T, Boyle, J. M., Dongarra, J. J., Garbow, B. S., Ikebe,Y., Klema, V., and Moler, C. B. (1976). Matrix Eigensystems Routines - EISPACK Guide. Springer-Verlag Lecture Notes in Computer Science.
Anderson. E. and ten others (1999) LAPACK Users' Guide. Third Edition. SIAM.
Available on-line at http://www.netlib.org/lapack/lug/lapack_lug.html.

## See Also

svd, a generalization of eigen; qr, and chol for related decompositions.
To compute the determinant of a matrix, the qr decomposition is much more efficient: det. capabilities to test for IEEE 754 arithmetic.

## Examples

```
eigen(cbind(c(1, -1),c(-1,1)))
eigen(cbind(c(1,-1),c(-1,1)), symmetric = FALSE)# same (different algorithm).
eigen(cbind(1,c(1,-1)), only.values = TRUE)
eigen(cbind(-1,2:1)) # complex values
eigen(print(cbind(c(0,1i), c(-1i,0))))# Hermite ==> real Eigen values
## 3 x 3:
eigen(cbind( 1,3:1,1:3))
eigen(cbind(-1,c(1:2,0),0:2)) # complex values
Meps <- .Machine$double.eps
set.seed(321, kind = "default") # force a particular seed
m <- matrix(round(rnorm(25),3), 5,5)
sm <- m + t(m) #- symmetric matrix
em <- eigen(sm); V <- em$vect
print(lam <- em$values) # ordered DEcreasingly
stopifnot(
    abs(sm %*% V - V %*% diag(lam)) < 60*Meps,
    abs(sm - V %*% diag(lam) %*% t(V)) < 60*Meps)
##------- Symmetric = FALSE: -- different to above : ---
em <- eigen(sm, symmetric = FALSE); V2 <- em$vect
print(lam2 <- em$values) # ordered decreasingly in ABSolute value !
    # and V2 is not normalized (where V is):
print(i <- rev(order(lam2)))
stopifnot(abs(lam - lam2[i]) < 60 * Meps)
zapsmall(Diag <- t(V2) %*% V2) # orthogonal, but not normalized
```

```
print(norm2V <- colSums(V2 * V2))
stopifnot( abs(1- norm2V / diag(Diag)) < 60*Meps)
V2n <- sweep(V2,2, STATS= sqrt(norm2V), FUN="/")## V2n are now Normalized EV
apply(V2n * V2n, 2, sum)
##[1] 1 1 1 1 1
## Both are now TRUE:
stopifnot(abs(sm %*% V2n - V2n %*% diag(lam2)) < 60*Meps,
    abs(sm - V2n %*% diag(lam2) %*% t(V2n)) < 60*Meps)
## Re-ordered as with symmetric:
sV <- V2n[,i]
slam <- lam2[i]
all(abs(sm %*% sV - sV %*% diag(slam)) < 60*Meps)
all(abs(sm - sV %*% diag(slam) %*% t(sV)) < 60*Meps)
## sV *is* now equal to V -- up to sign (+-) and rounding errors
all(abs(c(1 - abs(sV / V))) < 1000*Meps) # TRUE (P ~ 0.95)
```

environment Environment Access

## Description

Get, set, test for and create environments.

## Usage

```
environment(fun = NULL)
environment(fun) <- value
is.environment(obj)
.GlobalEnv
globalenv()
new.env(hash=FALSE, parent=parent.frame())
parent.env(env)
parent.env(env) <- value
```


## Arguments

| fun | a function, a formula, or NULL, which is the default. |
| :--- | :--- |
| value | an environment to associate with the function |
| obj | an arbitrary R object. |
| hash | a logical, if TRUE the environment will be hashed |
| parent | an environment to be used as the parent of the environment created. |
| env | an environment |

## Details

The global environment .GlobalEnv is the first item on the search path, more often known as the user's workspace. It can also be accessed by globalenv().

The variable .BaseNamespaceEnv is part of some experimental support for name space management.

The assignment function parent.env<- is extremely dangerous as it can be used to destructively change environments in ways that violate assumptions made by the internal C code. It may be removed in the near future.

## Value

If fun is a function or a formula then environment (fun) returns the environment associated with that function or formula. If fun is NULL then the current evaluation environment is returned.

The assignment form sets the environment of the function or formula fun to the value given.
is.environment (obj) returns TRUE iff obj is an environment.
new.env returns a new (empty) environment enclosed in the parent's environment, by default.
parent.env returns the parent environment of its argument.
parent.env<- sets the parent environment of its first argument.

## See Also

The envir argument of eval.

## Examples

```
##-- all three give the same:
environment()
environment(environment)
.GlobalEnv
ls(envir=environment(approxfun(1:2,1:2, method="const")))
is.environment(.GlobalEnv)# TRUE
e1 <- new.env(TRUE, NULL)
e2 <- new.env(FALSE, NULL)
assign("a", 3, env=e2)
parent.env(e1) <- e2
get("a", env=e1)
```

esoph Smoking, Alcohol and (O)esophageal Cancer

## Description

Data from a case-control study of (o)esophageal cancer in Ile-et-Vilaine, France.

## Usage

data(esoph)

## Format

data frame with records for 88 age/alcohol/tobacco combinations.

| [,1] | "agegp" | Age group | 125-34 years |
| :---: | :---: | :---: | :---: |
|  |  |  | 2 35-44 |
|  |  |  | 3 45-54 |
|  |  |  | 4 55-64 |
|  |  |  | $565-74$ |
|  |  |  | $675+$ |
| [,2] | "alcgp" | Alcohol consumption | $10-39 \mathrm{gm} /$ day |
|  |  |  | 2 40-79 |
|  |  |  | 3 80-119 |
|  |  |  | $4120+$ |
| [,3] | "tobgp" | Tobacco consumption | $10-9 \mathrm{gm} /$ day |
|  |  |  | 2 10-19 |
|  |  |  | 3 20-29 |
|  |  |  | $430+$ |
| [,4] | "ncases" | Number of cases |  |
| [,5] | "ncontrols" | Number of controls |  |

## Author(s)

Thomas Lumley

## Source

Breslow, N. E. and Day, N. E. (1980) Statistical Methods in Cancer Research. 1: The Analysis of Case-Control Studies. IARC Lyon / Oxford University Press.

## Examples

```
data(esoph)
summary(esoph)
## effects of alcohol, tobacco and interaction, age-adjusted
model1 <- glm(cbind(ncases, ncontrols) ~ agegp + tobgp * alcgp,
    data = esoph, family = binomial())
anova(model1)
## Try a linear effect of alcohol and tobacco
model2 <- glm(cbind(ncases, ncontrols) ~ agegp + codes(tobgp)
    + codes(alcgp),
    data = esoph, family = binomial())
```

```
summary(model2)
## Re-arrange data for a mosaic plot
ttt <- table(esoph$agegp, esoph$alcgp, esoph$tobgp)
ttt[ttt == 1] <- esoph$ncases
tt1 <- table(esoph$agegp, esoph$alcgp, esoph$tobgp)
tt1[tt1 == 1] <- esoph$ncontrols
tt <- array(c(ttt, tt1), c(dim(ttt),2),
    c(dimnames(ttt), list(c("Cancer", "control"))))
mosaicplot(tt, main = "esoph data set", color = TRUE)
```

```
euro Euro Conversion Rates
```


## Description

Conversion rates between the various Euro currencies.

## Usage

```
data(euro)
```


## Format

euro is a named vector of length 11, euro.cross a named matrix of size 11 by 11 .

## Details

The data set euro contains the value of 1 Euro in all currencies participating in the European monetary union (Austrian Schilling ATS, Belgian Franc BEF, German Mark DEM, Spanish Peseta ESP, Finnish Markka FIM, French Franc FRF, Irish Punt IEP, Italian Lira ITL, Luxembourg Franc LUF, Dutch Guilder NLG and Portugese Escudo PTE). These conversion rates were fixed by the European Union on December 31, 1998. To convert old prices to Euro prices, divide by the respective rate and round to 2 digits.
The data set euro.cross contains conversion rates between the various Euro currencies, i.e., the result of outer (1 / euro, euro).

## Examples

```
data(euro)
cbind(euro)
## These relations hold:
euro == signif(euro,6) # [6 digit precision in Euro's definition]
all(euro.cross == outer(1/euro, euro))
## Convert 20 Euro to Belgian Franc
20 * euro["BEF"]
## Convert 20 Austrian Schilling to Euro
20 / euro["ATS"]
## Convert 20 Spanish Pesetas to Italian Lira
20 * euro.cross["ESP", "ITL"]
dotchart(euro,
```

```
    main = "euro data: 1 Euro in currency unit")
dotchart(1/euro,
    main = "euro data: 1 currency unit in Euros")
dotchart(log(euro, 10),
    main = "euro data: log10(1 Euro in currency unit)")
```

```
eurodist Distances Between Cities in Europe
```


## Description

The data give the road distances (in km) between 21 cities in Europe. The data are taken from a table in "The Cambridge Encyclopaedia".

## Usage

data(eurodist)

## Format

A dist object based on 21 objects. (You must have the mva package loaded to have the methods for this kind of object available).

## Source

Crystal, D. Ed. (1990) The Cambridge Encyclopaedia. Cambridge: Cambridge University Press,

```
eval Evaluate an (Unevaluated) Expression
```


## Description

Evaluate an R expression in a specified environment.

## Usage

```
eval(expr, envir = parent.frame(),
    enclos = if(is.list(envir) || is.pairlist(envir)) parent.frame())
evalq(expr, envir, enclos)
eval.parent(expr, n = 1)
local(expr, envir = new.env())
```


## Arguments

```
expr object of mode expression orcall or an "unevaluated expression".
envir the environment in which expr is to be evaluated. May also be a list, a
    data frame, or an integer as in sys.call.
enclos Relevant when envir is a list or a data frame. Specifies the enclosure,
    i.e., where R looks for objects not found in envir.
n
    parent generations to go back
```


## Details

eval evaluates the expression expr argument in the environment specified by envir and returns the computed value. If envir is not specified, then sys.frame(sys.frame()), the environment where the call to eval was made is used.
The evalq form is equivalent to eval (quote (expr), ...).
As eval evaluates its first argument before passing it to the evaluator, it allows you to assign complicated expressions to symbols and then evaluate them. evalq avoids this.

```
eval.parent(expr, n) is a shorthand for eval(expr, parent.frame(n)).
```

local evaluates an expression in a local environment. It is equivalent to evalq except the its default argument creates a new, empty environment. This is useful to create anonymous recursive functions and as a kind of limited namespace feature since variables defined in the environment are not visible from the outside.

## Note

Due to the difference in scoping rules, there are some differences between $R$ and $S$ in this area. In particular, the default enclosure in S is the global environment.
When evaluating expressions in dataframes that has been passed as argument to a function, the relevant enclosure is often the caller's environment, i.e., one needs eval(x, data, parent.frame()).

## See Also

expression, quote, sys.frame, parent.frame, environment.

## Examples

```
eval(2 - 2 - 3)
mEx <- expression(2^2^3); mEx; 1 + eval(mEx)
eval({ xx <- pi; xx^2}) ; xx
a <- 3 ; aa <- 4 ; evalq(evalq(a+b+aa, list(a=1)), list(b=5)) # == 10
a <- 3 ; aa <- 4 ; evalq(evalq(a+b+aa, -1), list(b=5)) # == 12
ev <- function() {
    e1 <- parent.frame()
    ## Evaluate a in e1
    aa <- eval(expression(a),e1)
    ## evaluate the expression bound to a in e1
    a <- expression(x+y)
    list(aa = aa, eval = eval(a, e1))
}
tst.ev <- function(a = 7) { x <- pi; y <- 1; ev() }
tst.ev()#-> aa : 7, eval : 4.14
##
## Uses of local()
##
# Mutual recursives.
# gg gets value of last assignment, an anonymous version of f.
gg <- local({
```

```
    k <- function(y)f(y)
    f <- function(x) if(x) x*k(x-1) else 1
})
gg(10)
sapply(1:5, gg)
# Nesting locals. a is private storage accessible to k
gg <- local({
    k <- local({
                a <- 1
                function(y){print(a <<- a+1);f(y)}
    })
    f <- function(x) if(x) x*k(x-1) else 1
})
sapply(1:5, gg)
ls(envir=environment(gg))
ls(envir=environment(get("k", envir=environment(gg))))
```

example Run an Examples Section from the Online Help

## Description

Run all the R code from the Examples part of R's online help topic topic with two possible exceptions, dontrun and testonly, see Details below.

## Usage

```
example(topic, package = .packages(), lib.loc = NULL,
    echo = TRUE, verbose = getOption("verbose"),
    prompt.echo = paste(abbreviate(topic, 6),"> ", sep=""))
```


## Arguments

| topic | name or character: The online help topic the examples of which should <br> be run. |
| :--- | :--- |
| package | a character vector with package names. By default, all packages in the <br> search path are used. |
| lib.loc | a character vector of directory names of R libraries, or NULL. The default <br> value of NULL corresponds to all libraries currently known. If the default <br> is used, the loaded packages are searched before the libraries. |
| echo | logical; if TRUE, show the R input when sourcing. |
| verbose | logical; if TRUE, show even more when running example code. |
| prompt.echo $\quad$ character; gives the prompt to be used if echo = TRUE. |  |

## Details

If lib.loc is not specified, the packages are searched for amongst those already loaded, then in the specified libraries. If lib.loc is specified, they are searched for only in the specified libraries, even if they are already loaded from another library.
An attempt is made to load the package before running the examples, but this will not replace a package loaded from another location.

As detailed in the manual Writing R Extensions, the author of the help page can markup parts of the examples for two exception rules
dontrun encloses code that should not be run.
testonly encloses code that is invisible on help pages, but will be run both by the package checking tools, and the example() function.

## Value

(the value of the last evaluated expression).

## Note

The examples can be many small files. On some file systems it is desirable to save space, and the files in the 'R-ex' directory of an installed package can be zipped up as a zip archive 'Rex.zip'.

## Author(s)

Martin Maechler and others

## See Also

demo

## Examples

```
example(dbinom)
## force use of the standard package eda:
example("smooth", package="eda", lib.loc=.Library)
```

```
exists Is an Object Defined?
```


## Description

Search for an R object of the given name on the search path.

## Usage

```
exists(x, where = -1, envir = parent.frame(),
    frame = NULL, mode = "any", inherits = TRUE)
```


## Arguments

mode the mode of object sought.
inherits
x
where where to look for the object (see the details section); if omitted, the function will search, as if the name of the object appeared in unquoted in an expression.
envir an alternative way to specify an environment to look in, but it's usually simpler to just use the where argument.
frame a frame in the calling list. Equivalent to giving where as sys.frame(frame).
a variable name (given as a character string). an

## Details

The where argument can specify the environment in which to look for the object in any of several ways: as an integer (the position in the search list); as the character string name of an element in the search list; or as an environment (including using sys.frame to access the currently active function calls). The envir argument is an alternative way to specify an environment, but is primarily there for back compatibility.

This function looks to see if the name $x$ has a value bound to it. If inherits is TRUE and a value is not found for x , then the parent frames of the environment are searched until the name x is encountered. Warning: This is the default behaviour for R but not for S .

If mode is specified then only objects of that mode are sought. The function returns TRUE if the variable is encountered and FALSE if not.

The mode includes collections such as "numeric" and "function": any member of the collection will suffice.

## Value

Logical, true if and only if the object is found on the search path.

## See Also

get.

## Examples

```
## Define a substitute function if necessary:
if(!exists("some.fun", mode="function"))
    some.fun <- function(x) { cat("some.fun(x)\n"); x }
search()
exists("ls", 2) # true even though ls is in pos=3
exists("ls", 2, inherits=FALSE) # false
```

expand.grid
Create a Data Frame from All Combinations of Factors

## Description

Create a data frame from all combinations of the supplied vectors or factors. See the description of the return value for precise details of the way this is done.

## Usage

expand.grid(...)

## Arguments

... Vectors, factors or a list containing these.

## Value

A data frame containing one row for each combination of the supplied factors. The first factors vary fastest. The columns are labelled by the factors if these are supplied as named arguments or named components of a list.

## Author(s)

B.D. Ripley

## Examples

```
expand.grid(height = seq}(60, 80, 5), weight = seq(100, 300, 50)
```

    sex = c("Male","Female"))
    expand.model.frame $\quad$ Add new variables to a model frame

## Description

Evaluates new variables as if they had been part of the formula of the specified model. This ensures that the same na.action and subset arguments are applied and allows e.g. x to be recovered for a model using $\sin (x)$ as a predictor.

## Usage

expand.model.frame(model, extras, envir=environment(formula(model)), na. expand $=$ FALSE)

## Arguments

| model | a fitted model |
| :--- | :--- |
| extras | one-sided formula or vector of character strings describing new variables <br> to be added |
| envir | an environment to evaluate things in |
| na.expand | logical; see below |

## Details

If na.expand=FALSE then NA values in the extra variables will be passed to the na.action function used in model. This may result in a shorter data frame (with na.omit) or an error (with na.fail). If na.expand=TRUE the returned data frame will have precisely the same rows as model.frame(model), but the columns corresponding to the extra variables may contain NA.

## Value

A data frame.

## See Also

```
model.frame,predict
```


## Examples

```
data(trees)
model <- lm(log(Volume) ~ log(Girth) + log(Height), data=trees)
expand.model.frame(model, ~ Girth) # prints data.frame like
dd <- data.frame(x=1:5, y=rnorm(5), z=c(1,2,NA,4,5))
model <- glm(y ~ x, data=dd, subset=1:4, na.action=na.omit)
expand.model.frame(model, "z", na.expand=FALSE) # = default
expand.model.frame(model, "z", na.expand=TRUE)
```

Exponential The Exponential Distribution

## Description

Density, distribution function, quantile function and random generation for the exponential distribution with rate rate (i.e., mean $1 /$ rate).

## Usage

```
dexp(x, rate = 1, log = FALSE)
pexp(q, rate = 1, lower.tail = TRUE, log.p = FALSE)
qexp(p, rate = 1, lower.tail = TRUE, log.p = FALSE)
rexp(n, rate = 1)
```


## Arguments

```
x, q vector of quantiles.
p vector of probabilities.
n number of observations. If length (n) > 1, the length is taken to be the
number required.
rate vector of rates.
log, log.p logical; if TRUE, probabilities p are given as log(p).
lower.tail logical; if TRUE (default), probabilities are P[X\leqx], otherwise, P[X>
x].
```


## Details

If rate is not specified, it assumes the default value of 1.
The exponential distribution with rate $\lambda$ has density

$$
f(x)=\lambda e^{-\lambda x}
$$

for $x \geq 0$.

## Value

dexp gives the density, pexp gives the distribution function, qexp gives the quantile function, and rexp generates random deviates.

## Note

The cumulative hazard $H(t)=-\log (1-F(t))$ is $-\operatorname{pexp}(\mathrm{t}, \mathrm{r}$, lower $=\mathrm{FALSE}, \log =$ TRUE).

## See Also

$\exp$ for the exponential function, dgamma for the gamma distribution and dweibull for the Weibull distribution, both of which generalize the exponential.

## Examples

```
dexp(1) - exp(-1) #-> 0
r <- rexp(100)
all(abs(1 - dexp(1, r) / (r*exp(-r))) < 1e-14)
```

expression Unevaluated Expressions

## Description

Creates or tests for objects of mode "expression".

## Usage

expression(...)
is.expression(x)
as.expression(x, ...)

## Arguments

... valid R expressions.
x an arbitrary R object.

## Value

expression returns a vector of mode "expression" containing its arguments as unevaluated "calls".
is.expression returns TRUE if expr is an expression object and FALSE otherwise.
as.expression attempts to coerce its argument into an expression object.

## See Also

 call, eval, function. Further, text and legend for plotting math expressions.
## Examples

```
length(ex1 <- expression(1+ 0:9))# 1
ex1
eval(ex1)# 1:10
length(ex3 <- expression(u,v, 1+ 0:9))# 3
mode(ex3 [3]) # expression
mode(ex3[[3]])# call
rm(ex3)
```

Extract Extract or Replace Parts of an Object

## Description

Operators act on vectors, arrays, dataframes and lists to extract or replace subsets.

## Usage

x[i]
x[i, j, ...]
x[i, j, ... , drop=TRUE]
x[[i]]
$\mathrm{x}[\mathrm{i}, \mathrm{j}, \ldots]]$
x\$name
.subset (x, ...)
.subset2(x, ...)

## Arguments

[^1]
## Details

If one of these expressions appears on the left side of an assignment then that part of x is set to the value of the right hand side of the assignment.

These operators are generic. You can write methods to handle subsetting of specific classes of data.
The [ [ operator requires all relevant subscripts be supplied. With the [ operator a comma separated blank indicates that all entries in that dimension are selected.

When [.data.frame is used for subsetting rows of a data.frame, it returns a dataframe with unique (and non-missing)row names, if necessary transforming the names using make.names ( * , unique $=$ TRUE). See the swiss example below.
When operating on a list, the [ [ operator gives the specified element of the list while the [ operator returns a list with the specified element(s) in it.

The operators $\$$ and $\$<-$ do not evaluate their second argument. It is translated to a string and that string is used to locate the correct component of the first argument.

The functions .subset and .subset2 are essentially equivalent to the [ and [[ operators, except that methods dispatch does not take place. This is to avoid expensive unclassing in order to apply the default method to an object. They should not normally be invoked by end users.

## See Also

list, array, matrix.
Syntax for operator precedence.

## Examples

```
x <- 1:12; m <- matrix(1:6,nr=2); li <- list(pi=pi, e = exp(1))
x[10] # the tenth element of x
m[1,] # the first row of matrix m
m[1, , drop = FALSE] # is a 1-row matrix
li[[1]] # the first element of list li
y <- list (1,2,a=4,5)
y[c(3,4)] # a list containing elements 3 and 4 of y
y$a # the element of y named a
data(swiss)
swiss[ c(1, 1:2), ] # duplicate row, unique row names
```

```
extractAIC Extract AIC from a Fitted Model
```


## Description

Computes the (generalized) Akaike Information Criterion for a fitted parametric model.

## Usage

```
extractAIC (fit, scale, k = 2, ...)
extractAIC.lm (fit, scale = 0, k = 2, ...)
extractAIC.glm(fit, scale = 0, k = 2, ...)
extractAIC.aov(fit, scale = 0, k = 2, ...)
extractAIC.coxph (fit, scale, k = 2, ...)
extractAIC.negbin (fit, scale, k = 2, ...)
extractAIC.survreg(fit, scale, k = 2, ...)
```


## Arguments

| fit | fitted model, usually the result of a fitter like lm. |
| :--- | :--- |
| scale | optional numeric specifying the scale parameter of the model, see scale <br> in step. |
| k | numeric specifying the "weight" of the equivalent degrees of freedom <br> $(\equiv$ (三df) part in the AIC formula. |
| $\ldots$ | further arguments (currently unused in base R). |

## Details

The criterion used is

$$
A I C=-2 \log L+k \times \mathrm{edf},
$$

where $L$ is the likelihood and edf the equivalent degrees of freedom (i.e., the number of parameters for usual parametric models) of fit.

For generalized linear models (i.e., for 1 m , aov, and glm ), $-2 \log L$ is the deviance, as computed by deviance(fit), plus a constant.
$\mathrm{k}=2$ corresponds to the traditional AIC, using $\mathrm{k}=\log (\mathrm{n})$ provides the BIC (Bayes IC) instead.

For further information, particularly about scale, see step.

## Value

A numeric vector of length 2, giving
edf the "equivalent degrees of freedom" of the fitted model fit.
AIC the (generalized) Akaike Information Criterion for fit.

## Note

These functions are used in add1, drop1 and step and that may be their main use.

## Author(s)

B. D. Ripley

## References

Venables, W. N. and Ripley, B. D. (1997) Modern Applied Statistics with S-PLUS. New York: Springer (2nd ed).

```
See Also
```



## Examples

```
example(glm)
extractAIC(glm.D93)#>> 5 15.129
```


## Extremes Maxima and Minima

## Description

Returns the (parallel) maxima and minima of the input values.

## Usage

```
max(..., na.rm=FALSE)
min(..., na.rm=FALSE)
pmax(..., na.rm=FALSE)
pmin(..., na.rm=FALSE)
```


## Arguments

| $\ldots$. | numeric arguments. |
| :--- | :--- |
| na.rm | a logical indicating whether missing values should be removed. |

## Value

max and min return the maximum or minimum of all the values present in their arguments, as integer if all are integer, or as double otherwise.
The minimum and maximum of an empty set are $+\operatorname{Inf}$ and $-\operatorname{Inf}$ (in this order!) which ensures transitivity, e.g., $\min (x 1, \min (x 2))==\min (x 1, x 2)$. In $R$ versions before 1.5, $\min (i n t e g e r(0))==$.Machine\$integer.max, and analogously for max, preserving argument type, whereas from R version 1.5.0, $\max (\mathrm{x})==-\operatorname{Inf}$ and $\min (\mathrm{x})==+\operatorname{Inf}$ whenever length $(x)=0$ (after removing missing values if requested).

If na.rm is FALSE an NA value in any of the arguments will cause a value of NA to be returned, otherwise NA values are ignored.
pmax and pmin take several vectors as arguments and return a single vector giving the parallel maxima (or minima) of the vectors. The first element of the result is the maximum (minimum) of the first elements of all the arguments, the second element of the result is the maximum (minimum) of the second elements of all the arguments and so on. Shorter vectors are recycled if necessary. If na.rm is FALSE, NA values in the input vectors will produce NA values in the output. If na.rm is TRUE, NA values are ignored. attributes (such as names or dim) are transferred from the first argument (if applicable).

## See Also

range (both min and max) and which.min (which.max) for the arg min, i.e., the location where an extreme value occurs.

## Examples

```
min(5:1,pi)
pmin(5:1, pi)
x <- sort(rnorm(100)); cH <- 1.35
pmin(cH, quantile(x)) # no names
pmin(quantile(x), cH) # has names
plot(x, pmin(cH, pmax(-cH, x)), type='b', main= "Huber's function")
```

factor Factors

## Description

The function factor is used to encode a vector as a factor (the names category and enumerated type are also used for factors). If ordered is TRUE, the factor levels are assumed to be ordered. For compatibility with $S$ there is also a function ordered.
is.factor, is.ordered, as.factor and as.ordered are the membership and coercion functions for these classes.

## Usage

```
factor(x, levels = sort(unique(x), na.last = TRUE), labels = levels,
    exclude = NA, ordered = is.ordered(x))
ordered(x, ...)
is.factor(x)
is.ordered(x)
as.factor(x)
as.ordered(x)
```


## Arguments

$\mathrm{x} \quad$ a vector of data, usually taking a small number of distinct values
levels an optional vector of the values that x might have taken. The default is the set of values taken by x , sorted into increasing order.
labels either an optional vector of labels for the levels (in the same order as levels after removing those in exclude), or a character string of length 1.
exclude a vector of values to be excluded when forming the set of levels. This should be of the same type as $x$, and will be coerced if necessary.
ordered logical flag to determine if the levels should be regarded as ordered (in the order given).
... (in ordered (.)): any of the above, apart from ordered itself.

## Details

The type of the vector x is not restricted.
Ordered factors differ from factors only in their class, but methods and the model-fitting functions treat the two classes quite differently.

The encoding of the vector happens as follows. First all the values in exclude are removed from levels. If $\mathrm{x}[\mathrm{i}]$ equals levels[j], then the $i$-th element of the result is j . If no match is found for $\mathrm{x}[\mathrm{i}]$ in levels, then the i-th element of the result is set to NA.

Normally the 'levels' used as an attribute of the result are the reduced set of levels after removing those in exclude, but this can be altered by supplying labels. This should either be a set of new labels for the levels, or a character string, in which case the levels are that character string with a sequence number appended.
factor ( x ) applied to a factor is a no-operation unless there are unused levels: in that case, a factor with the reduced level set is returned. If exclude is used it should also be a factor with the same level set as x or a set of codes for the levels to be excluded.

The codes of a factor may contain NA. For a numeric $x$, set exclude=NULL to make NA an extra level ("NA"), by default the last level.
If "NA" is a level, the way to set a code to be missing is to use is.na on the left-hand-side of an assignment. Under those circumstances missing values are printed as <NA>.

## Value

factor returns an object of class "factor" which has a set of numeric codes the length of x with a "levels" attribute of mode character. If ordered is true (or ordered is used) the result has class c("ordered", "factor").
is.factor returns TRUE or FALSE depending on whether its argument is of type factor or not. Correspondingly, is.ordered returns TRUE when its argument is ordered and FALSE otherwise.
as.factor coerces its argument to a factor. It is an abbreviated form of factor.
as.ordered( $x$ ) returns $x$ if this is ordered, and ordered( $x$ ) otherwise.

## Warning

The interpretation of a factor depends on both the codes and the "levels" attribute. Be careful only to compare factors with the same set of levels (in the same order). In particular, as.numeric applied to a factor is meaningless, and may happen by implicit coercion.
The levels of a factor are by default sorted, but the sort order may well depend on the locale at the time of creation, and should not be assumed to be ASCII.

## See Also

gl for construction of "balanced" factors and C for factors with specified contrasts. levels and nlevels for accessing the levels, and codes to get integer codes.

## Examples

```
ff <- factor(substring("statistics", 1:10, 1:10), levels=letters)
ff
codes(ff)
factor(ff)# drops the levels that do not occur
factor(factor(letters[7:10])[2:3]) # exercise indexing and reduction
```

```
factor(letters[1:20], label="letter")
class(ordered(4:1))# "ordered", inheriting from "factor"
## suppose you want "NA" as a level, and to allowing missing values.
(x <- factor(c(1, 2, "NA"), exclude = ""))
is.na(x)[2] <- TRUE
x # [1] 1 <NA> NA, <NA> used because NA is a level.
is.na(x)
# [1] FALSE TRUE FALSE
```

```
factor.scope
```

Compute Allowed Changes in Adding to or Dropping from a Formula

## Description

add.scope and drop.scope compute those terms that can be individually added to or dropped from a model while respecting the hierarchy of terms.

## Usage

```
add.scope(terms1, terms2)
drop.scope(terms1, terms2)
factor.scope(factor, scope)
```


## Arguments

terms 1 the terms or formula for the base model.
terms2 the terms or formula for the upper (add.scope) or lower (drop.scope) scope. If missing for drop.scope it is taken to be the null formula, so all terms (except any intercept) are candidates to be dropped.
factor the "factor" attribute of the terms of the base object.
scope a list with one or both components drop and add giving the "factor" attribute of the lower and upper scopes respectively.

## Details

factor.scope is not intended to be called directly by users.

## Value

For add.scope and drop.scope a character vector of terms labels. For factor.scope, a list with components drop and add, character vectors of terms labels.

## Author(s)

B.D. Ripley

## See Also

add1, drop1, aov, lm

## Examples

```
add.scope( ~ a + b + c + a:b, ~ (a + b + c) ^ 3)
# [1] "a:c" "b:c"
drop.scope( ~ a + b + c + a:b)
# [1] "c" "a:b"
```

faithful Old Faithful Geyser Data

## Description

The 'faithful' data frame has 272 rows and 2 columns; the waiting time between eruptions and the duration of the eruption for the Old Faithful geyser in Yellowstone National Park, Wyoming, USA.

## Usage

data(faithful)

## Format

A data frame with 272 observations on 2 variables.

| $[, 1]$ | eruptions | numeric | Eruption time in mins |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | waiting | numeric | Waiting time to next eruption |

## Details

A closer look at faithful\$eruptions reveals that these are heavily rounded times originally in seconds, where multiples of 5 are more frequent than expected under non-human measurement. For a "better" version of the eruptions times, see the example below.

There are many versions of this dataset around: Azzalini and Bowman (1990) use a more complete version.

## Source

W. Härdle.

## References

Härdle, W. (1991) Smoothing Techniques with Implementation in $S$. New York: Springer.
Azzalini, A. and Bowman, A. W. (1990). A look at some data on the Old Faithful geyser. Applied Statistics 39, 357-365.

## See Also

geyser in package MASS for the Azzalini-Bowman version.

## Examples

```
data(faithful)
f.tit <- "faithful data: Eruptions of Old Faithful"
```

```
ne60 <- round(e60 <- 60 * faithful$eruptions)
all.equal(e60, ne60) # relative diff. ~ 1/10000
table(zapsmall(abs(e60 - ne60))) # 0, 0.02 or 0.04
faithful$better.eruptions <- ne60 / 60
te <- table(ne60)
te[te >= 4] # (too) many multiples of 5 !
plot(names(te), te, type="h", main = f.tit, xlab = "Eruption time (sec)")
plot(faithful[, -3], main = f.tit,
    xlab = "Eruption time (min)",
    ylab = "Waiting time to next eruption (min)")
lines(lowess(faithful$eruptions, faithful$waiting, f = 2/3, iter = 3),
        col = "red")
```

family Family Objects for Models

## Description

Family objects provide a convenient way to specify the details of the models used by functions such as glm. See the documentation for glm for the details on how such model fitting takes place.

## Usage

```
family(object, ...)
    binomial(link = "logit")
    gaussian(link ="identity")
    Gamma(link = "inverse")
    inverse.gaussian(link = "1/mu^2")
    poisson(link = "log")
    quasi(link = "identity", variance = "constant")
    quasibinomial(link = "logit")
    quasipoisson(link = "log")
```


## Arguments

| link | a specification for the model link function. The binomial family admits |
| :--- | :--- |
| the links "logit", "probit", "log", and "cloglog" (complementary log- |  |
| log); the Gamma family the links "identity", "inverse", and "log"; the |  |
| poisson family the links "identity", "log", and "sqrt"; the quasi |  |
| family the links "logit", "probit", "cloglog", "identity", "inverse", |  |
| "log", "1/mu^2" and "sqrt". The function power can also be used to |  |
| create a power link function for the quasi family. |  |
| The other families have only one permissible link function: "identity" |  |
| for the gaussian family, and "1/mu^2" for the inverse.gaussian family. |  |
| variance | for all families, other than quasi, the variance function is determined by |
|  | the family. The quasi family will accept the specifications "constant", |
|  | "mu(1-mu)", "mu", "mu^2" and "mu^3" for the variance function. |

object the function family accesses the family objects which are stored within objects created by modelling functions (e.g. glm).
... further arguments passed to methods.

## Details

The quasibinomial and quasipoisson families differ from the binomial and poisson families only in that the dispersion parameter is not fixed at one, so they can "model" overdispersion. For the binomial case see McCullagh and Nelder (1989, pp. 124-8). Although they show that there is (under some restrictions) a model with variance proportional to mean as in the quasi-binomial model, note that glm does not compute maximum-likelihood estimates in that model. The behaviour of S-PLUS is closer to the quasi- variants.

## References

McCullagh P. and Nelder, J. A. (1989) Generalized Linear Models. London: Chapman and Hall.
Dobson, A. J. (1983) An Introduction to Statistical Modelling. London: Chapman and Hall. Cox, D. R. and Snell, E. J. (1981). Applied Statistics; Principles and Examples. London: Chapman and Hall.

## See Also

glm, power.

## Examples

```
nf <- gaussian()# Normal family
nf
str(nf)# internal STRucture
gf <- Gamma()
gf
str(gf)
gf$linkinv
all(1:10 == gf$linkfun(gf$linkinv(1:10)))# is TRUE
gf$variance(-3:4) #- == (.)^2
## quasipoisson. compare with example(glm)
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
d.AD <- data.frame(treatment, outcome, counts)
glm.qD93 <- glm(counts ~ outcome + treatment, family=quasipoisson())
glm.qD93
anova(glm.qD93, test="F")
summary(glm.qD93)
## for Poisson results use
anova(glm.qD93, dispersion = 1, test="Chisq")
summary(glm.qD93, dispersion = 1)
## tests of quasi
x <- rnorm(100)
y <- rpois(100, exp(1+x))
glm(y ~x, family=quasi(var="mu", link="log"))
```

```
# which is the same as
glm(y ~x, family=poisson)
glm(y ~x, family=quasi(var="mu^2", link="log"))
glm(y ~x, family=quasi(var="mu^3", link="log")) # should fail
y <- rbinom(100, 1, plogis(x))
# needs to set a starting value for the next fit
glm(y ~x, family=quasi(var="mu(1-mu)", link="logit"), start=c(0,1))
```

FDist The F Distribution

## Description

Density, distribution function, quantile function and random generation for the F distribution with df1 and df2 degrees of freedom (and optional non-centrality parameter ncp).

## Usage

$\mathrm{df}(\mathrm{x}, \mathrm{df} 1, \mathrm{df} 2, \log =\mathrm{FALSE})$
pf(q, df1, df2, ncp=0, lower.tail = TRUE, log.p = FALSE)
$\mathrm{qf}(\mathrm{p}, \mathrm{df} 1, \mathrm{df} 2, \quad$ lower.tail $=$ TRUE, log. $\mathrm{p}=\mathrm{FALSE})$
rf( $n, \mathrm{df} 1, \mathrm{df} 2$ )

## Arguments

```
x, q vector of quantiles.
p vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the
    number required.
df1, df2 degrees of freedom.
ncp non-centrality parameter.
log, log.p logical; if TRUE, probabilities p are given as log(p).
lower.tail logical; if TRUE (default), probabilities are P[X\leqx], otherwise, P[X>
    x].
```


## Details

The F distribution with $\mathrm{df} 1=n_{1}$ and df2 $=n_{2}$ degrees of freedom has density

$$
f(x)=\frac{\Gamma\left(n_{1} / 2+n_{2} / 2\right)}{\Gamma\left(n_{1} / 2\right) \Gamma\left(n_{2} / 2\right)}\left(\frac{n_{1}}{n_{2}}\right)^{n_{1} / 2} x^{n_{1} / 2-1}\left(1+\frac{n_{1} x}{n_{2}}\right)^{-\left(n_{1}+n_{2}\right) / 2}
$$

for $x>0$.
It is the distribution of the ratio of the mean squares of $n_{1}$ and $n_{2}$ independent standard normals, and hence of the ratio of two independent chi-squared variates each divided by its degrees of freedom. Since the ratio of a normal and the root mean-square of $m$ independent normals has a Student's $t_{m}$ distribution, the square of a $t_{m}$ variate has a F distribution on 1 and $m$ degrees of freedom.
The non-central F distribution is again the ratio of mean squares of independent normals of unit variance, but those in the numerator are allowed to have non-zero means and ncp is the sum of squares of the means. See Chisquare for further details on non-central distributions.

## Value

df gives the density, pf gives the distribution function qf gives the quantile function, and rf generates random deviates.

## See Also

dchisq for chi-squared and dt for Student's t distributions.

## Examples

```
## the density of the square of a t_m is 2*dt (x,m)/(2*x)
# check this is the same as the density of F_{1,m}
x <- seq(0.001, 5, len=100)
all.equal (df (x^2, 1, 5), dt (x, 5)/x)
## Identity: qf(2*p - 1, 1, df)) == qt(p, df)^2) for p >= 1/2
p <- seq(1/2, .99, length=50); df <- 10
rel.err <- function(x,y) ifelse(x==y,0, abs(x-y)/mean(abs(c(x,y))))
quantile(rel.err(qf(2*p - 1, df1=1, df2=df), qt(p, df) ^2), .90)# ~}=7e-
```

fft Fast Discrete Fourier Transform

## Description

Performs the Fast Fourier Transform of an array.

```
Usage
    fft(z, inverse = FALSE)
    mvfft(z, inverse = FALSE)
```


## Arguments

| $z$ | a real or complex array containing the values to be transformed. |
| :--- | :--- |
| inverse | if TRUE, the unnormalized inverse transform is computed (the inverse has |
|  | $a+$ in the exponent of $e$, but here, we do not divide by $1 /$ length $(\mathrm{x}))$. |

## Value

When $z$ is a vector, the value computed and returned by $f f t$ is the unnormalized univariate Fourier transform of the sequence of values in $\mathbf{z}$. When z contains an array, fft computes and returns the multivariate (spatial) transform. If inverse is TRUE, the (unnormalized) inverse Fourier transform is returned, i.e., if $y<-\operatorname{fft}(z)$, then $z$ is $f f t(y$, inverse $=$ TRUE) / length(y).
By contrast, mvfft takes a real or complex matrix as argument, and returns a similar shaped matrix, but with each column replaced by its discrete Fourier transform. This is useful for analyzing vector-valued series.
The FFT is fastest when the length of of the series being transformed is highly composite (i.e. has many factors). If this is not the case, the transform may take a long time to compute and will use a large amount of memory.

## References

Singleton, R. C. (1979) Mixed Radix Fast Fourier Transforms, in Programs for Digital Signal Processing, IEEE Digital Signal Processing Committee eds. IEEE Press.

## See Also

convolve, nextn.

## Examples

```
x <- 1:4
fft(x)
all(fft(fft(x), inverse = TRUE)/(x*length(x)) == 1+0i)
eps <- 1e-11 ## In general, not exactly, but still:
for(N in 1:130) {
    cat("N=",formatC(N,wid=3),": ")
    x <- rnorm(N)
    if(N %% 5 == 0) {
        m5 <- matrix(x,ncol=5)
        cat("mvfft:",all(apply(m5,2,fft) == mvfft(m5)),"")
    }
    dd <- Mod(1 - (f2 <- fft(fft(x), inverse=TRUE)/(x*length(x))))
    cat(if(all(dd < eps))paste(" all < ", formatC(eps)) else
        paste("NO: range=",paste(formatC(range(dd)),collapse=",")),"\n")
}
plot(fft(c(9:0,0:13, numeric(301))), type = "l")
periodogram <- function(x, mean.x = mean(x)) { # simple periodogram
    n <- length(x)
    x <- unclass(x) - mean.x
    Mod(fft(x))[2:(n%/% + + 1)]^2 / (2*pi*n) # drop I(0)
}
data(sunspots)
plot(10*log10(periodogram(sunspots)), type = "b", col = "blue")
```

file.access Ascertain File Accessibility

## Description

Utility function to access information about files on the user's file systems.

## Usage

```
file.access(names, mode = 0)
```


## Arguments

$$
\begin{array}{ll}
\text { names } & \text { character vector containing file names. } \\
\text { mode } & \text { integer specifying access mode required. }
\end{array}
$$

## Details

The mode value can be the exclusive or of the following values
0 test for existence.
1 test for execute permission.
2 test for write permission.
4 test for read permission.
Permission will be computed for real user ID and real group ID (rather than the effective IDs).

## Value

An integer vector with values 0 for success and -1 for failure.

## Note

This is intended as a replacement for the S-PLUS function access, a wrapper for the C function of the same name, which explains the return value encoding. Note that the return value is false for success.

## Author(s)

B. D. Ripley

## See Also

file.info

## Examples

```
fa <- file.access(dir("."))
table(fa) # count successes & failures
```

file.choose
Choose a File Interactively

## Description

Choose a file interactively.

## Usage

file.choose(new=FALSE)

## Arguments

new Logical: choose the style of dialog box presented to the user: at present only new $=$ FALSE is used.

## Value

A character vector of length one giving the file path.

```
file.info Extract File Information
```


## Description

Utility function to extract information about files on the user's file systems.

## Usage

```
file.info(...)
```


## Arguments

... character vectors containing file names.

## Details

What is meant by "file access" and hence the last access time is system-dependent.
On most systems symbolic links are followed, so information is given about the file to which the link points rather than about the link.

## Value

A data frame with row names the file names and columns
size integer: File size in bytes.
isdir logical: Is the file a directory?
mode integer of class "octmode". The file permissions, printed in octal, for example 644.
mtime, ctime, atime
integer of class "POSIXct": file modification, creation and last access times.
uid integer: the user ID of the file's owner.
gid integer: the group ID of the file's group.
uname character: uid interpreted as a user name.
grname character: gid interpreted as a group name.
Unknown user and group names will be NA.
Entries for non-existent or non-readable files will be NA. The uid

## Note

This function will only be operational on systems with the stat system call, but that seems very widely available.

Author(s)
B. D. Ripley

See Also
files, file.access, list.files, and DateTimeClasses for the date formats.

## Examples

```
ncol(finf <- file.info(dir()))# at least six
finf # the whole list
## Those that are more than 100 days old :
finf[difftime(Sys.time(), finf[,"mtime"], units="days") > 100 , 1:4]
file.info("no-such-file-exists")
```

file.path
Construct Path to File

## Description

Construct the path to a file from components in a platform-independent way.

## Usage

```
file.path(..., fsep=.Platform$file.sep)
```


## Arguments

| $\ldots$. | Character vectors |
| :--- | :--- |
| fsep | The path separator to use |

Value
A character vector of length one.

```
file.show Display One or More Files
```


## Description

Display one or more files.

## Usage

```
file.show(..., header, title="R Information",
        delete.file=FALSE, pager=getOption("pager"))
```


## Arguments

> ... one or more character vectors containing the names of the files to be displayed.
> header character vector (of the same length as the number of files specified in ...) giving a header for each file being displayed. Defaults to empty strings.
> title an overall title for the display. If a single separate window is used for the display, title will be used as the window title. If multiple windows are used, their titles should combine the title and the file-specific header.
> delete.file should the files be deleted after display? Used for temporary files.
> pager the pager to be used.

## Details

This function provides the core of the R help system, but it can be used for other purposes as well.

## Note

How the pager is implemented is highly system dependent.
The basic Unix version concatenates the files (using the headers) to a temporary file, and displays it in the pager selected by the pager argument, which is a character vector specifying a system command to run on the set of files.

Most GUI systems will use a separate pager window for each file, and let the user leave it up while R continues running. The selection of such pagers could either be done using "magic" pager names being intercepted by lower-level code (such as "internal" and "console" on Windows), or by letting pager be an R function which will be called with the same arguments as file.show and take care of interfacing to the GUI.

Not all implementations will honour delete.file.

## Author(s)

Ross Ihaka, Brian Ripley.

## See Also

```
files, list.files, help.
```


## Examples

```
file.show(file.path(R.home(), "COPYRIGHTS"))
```

```
files File Manipulation
```


## Description

These functions provide a low-level interface to the computer's file system.

## Usage

```
file.create(...)
file.exists(...)
file.remove(...)
file.rename(from, to)
file.append(file1, file2)
file.copy(from, to, overwrite = FALSE)
dir.create(path)
basename(path)
dirname(path)
path.expand(path)
```


## Arguments

..., file1, file2, from, to, path
character vectors, containing file names.
overwrite logical; should the destination files be overwritten?

## Details

The . . . arguments are concatenated to form one character string: you can specify the files separately or as one vector.
file.create creates files with the given names if they do not already exist and truncates them if they do. It returns a logical vector indicating the success or failure of the operation for each file.
file.exists returns a logical vector indicating whether the files named by its argument exist.
file.remove attempts to remove the files named in its argument. It returns a logical vector indicating whether or not it succeeded in removing each file.
file.rename attempts to rename a file. It returns a logical value indicating whether the operation succeeded.
file. append attempts to append the files named by its second argument to those named by its first. The R subscript recycling rule is used to align names given in vectors of different lengths.
file.copy works in a similar way to file.append but with the arguments in the natural order for copying. Copying to existing destination files is skipped unless overwrite = TRUE. The to argument can specify a single existing directory.
dir.create creates the last element of the path. It returns a logical, true for success.
basename removes all of the path up to the last path separator (if any).
dirname returns the part of the path up to (but excluding) the last path separator, or "." if there is no path separator. Tilde expansion is done: see the description for path.expand below.
In both basename and dirname trailing file separators are removed before dissecting the path, and for dirname any trailing file separators are removed from the result.
path.expand expands path(s) by replacing a leading tilde by the user's home directory (if defined on that platform). On some Unix versions, a leading ~user will expand to the home directory of user, but not on Unix versions without readline installed.

## Author(s)

Ross Ihaka, Brian Ripley

## See Also

file.info, file.access, file.path, file.show, list.files, unlink.

## Examples

```
cat("file A\n", file="A")
cat("file B\n", file="B")
file.append("A", "B")
file.create("A")
file.append("A", rep("B", 10))
if(interactive()) file.show("A")
file.copy("A", "C")
dir.create("tmp")
file.copy(c("A", "B"), "tmp")
unlink("tmp", recursive=TRUE)
file.remove("A", "B", "C")
basename(file.path("","p1","p2","p3","filename"))
dirname(file.path("","p1","p2","p3","filename"))
path.expand("~/foo")
```

filled.contour Level (Contour) Plots

## Description

This function produces a contour plot with the areas between the contours filled in solid color (Cleveland calls this a level plot). A key showing how the colors map to z values is shown to the right of the plot.

## Usage

```
filled.contour(x = seq(0, 1, len = nrow(z)),
    y = seq(0, 1, len = ncol(z)),
    z,
    xlim = range(x, finite=TRUE),
    ylim = range(y, finite=TRUE),
    zlim = range(z, finite=TRUE),
```

```
levels = pretty(zlim, nlevels), nlevels = 20,
color.palette = cm.colors,
col = color.palette(length(levels) - 1),
plot.title, plot.axes, key.title, key.axes,
asp = NA, xaxs = "i", yaxs = "i", las = 1, axes = TRUE,
...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | locations of grid lines at which the values in $z$ are measured. These must be in ascending order. By default, equally spaced values from 0 to 1 are used. If x is a list, its components $\mathrm{x} \$ \mathrm{x}$ and $\mathrm{x} \$ \mathrm{y}$ are used for x and y , respectively. If the list has component $\mathbf{z}$ this is used for $\mathbf{z}$. |
| :---: | :---: |
| z | a matrix containing the values to be plotted (NAs are allowed). Note that $x$ can be used instead of $z$ for convenience. |
| xlim | x limits for the plot. |
| ylim | $y$ limits for the plot. |
| zlim | z limits for the plot. |
| levels | a set of levels which are used to partition the range of $z$. Must be strictly increasing (and finite). Areas with z values between consecutive levels are painted with the same color. |
| nlevels | if levels is not specified, the range of $z$, values is divided into approximately this many levels. |
| color.palette | a color palette function to be used to assign colors in the plot. |
| col | an explicit set of colors to be used in the plot. This argument overrides any palette function specification. |
| plot.title | statements which add titles the main plot. |
| plot.axes | statements which draw axes on the main plot. This overrides the default axes. |
| key.title | statements which add titles for the plot key. |
| key.axes | statements which draw axes on the plot key. This overrides the default axis. |
| asp | the $y / x$ aspect ratio, see plot.window. |
| xaxs | the x axis style. The default is to use internal labeling. |
| yaxs | the y axis style. The default is to use internal labeling. |
| las | the style of labeling to be used. The default is to use horizontal labeling. |

## Note

This function currently uses the layout function and so is restricted to a full page display. In future it is likely to be replaced by a genuine levelplot function which will work in multipanel displays.
The ouput produced by filled.contour is actually a combination of two plots; one is the filled contour and one is the legend. Two separate coordinate systems are set up for these two plots, but they are only used internally - once the function has returned these coordinate systems are lost. If you want to annotate the main contour plot, for example to add points, you can specify graphics commands in the plot.axes argument. An example is given below.

## Author(s)

Ross Ihaka.

## References

Cleveland, W. S. (1993) Visualizing Data. Summit, New Jersey: Hobart.

## See Also

```
contour, image, palette.
```


## Examples

```
data(volcano)
filled.contour(volcano, color = terrain.colors, asp = 1)# simple
x <- 10*1:nrow(volcano)
y <- 10*1:ncol(volcano)
filled.contour(x, y, volcano, color = terrain.colors,
    plot.title = title(main = "The Topography of Maunga Whau",
    xlab = "Meters North", ylab = "Meters West"),
    plot.axes = { axis(1, seq(100, 800, by = 100))
                axis(2, seq(100, 600, by = 100)) },
    key.title = title(main="Height\n(meters)"),
    key.axes = axis(4, seq(90, 190, by = 10)))# maybe also asp=1
mtext(paste("filled.contour(.) from", R.version.string),
        side = 1, line = 4, adj = 1, cex = .66)
# Annotating a filled contour plot
a <- expand.grid(1:20, 1:20)
b <- matrix(a[,1] + a[,2], 20)
filled.contour(x = 1:20, y = 1:20, z = b,
    plot.axes={ axis(1); axis(2); points(10,10) })
```


## findInterval Find Interval Numbers or Indices

## Description

Find the indices of x in vec, where vec must be sorted (non-decreasingly); i.e., if i <findinterval ( $\mathrm{x}, \mathrm{v}$ ), we have $v_{i_{j}} \leq x_{j}<v_{i_{j}+1}$ where $v_{0}:=-\infty, v_{N+1}:=+\infty$, and $\mathrm{N}<-$ length (vec). At the two boundaries, the returned index may differ by 1 , depending on the optional arguments rightmost.closed and all.inside.

## Usage

```
findInterval(x, vec, rightmost.closed = FALSE, all.inside = FALSE)
```


## Arguments

```
x numeric.
vec numeric, sorted (weakly) increasingly, of length N, say.
rightmost.closed
                            logical; if true, the rightmost interval, vec[N-1] .. vec[N] is treated
                    as closed, see below.
all.inside logical; if true, the returned indices are coerced into {1,\ldots,N-1}, i.e. 0
    is mapped to 1 and N to N-1.
```


## Details

The function findInterval finds the index of one vector x in another, vec, where the latter must be non-decreasing. Where this is trivial, equivalent to apply ( outer ( x , vec, " $>=$ "), 1, sum), as a matter of fact, the internal algorithm uses interval search ensuring $O(n \log N)$ complexity where n <- length(x) (and N <- length (vec)). For (almost) sorted x, it will be even faster, basically $O(n)$.
This is the same computation as for the empirical distribution function, and indeed, findInterval (t, sort(X)) is identical to $n F_{n}\left(t ; X_{1}, \ldots, X_{n}\right)$ where $F_{n}$ is the empirical distribution function of $X_{1}, \ldots, X_{n}$.
When rightmost.closed $=$ TRUE, the result for $\mathrm{x}[\mathrm{j}]=\mathrm{vec}[\mathrm{N}](=\max (v e c))$, is $\mathrm{N}-1$ as for all other values in the last interval.

## Value

vector of length length( x ) with values in $0: \mathrm{N}$ where N <- length(vec), or values coerced to 1: ( $\mathrm{N}-1$ ) iff all.inside $=$ TRUE (equivalently coercing all x values inside the intervals).

## Author(s)

Martin Maechler

## See Also

approx(*, method = "constant") which is a generalization of findInterval(), ecdf for computing the empirical distribution function which is (up to a factor of $n$ ) also basically the same as findInterval(.).

## Examples

```
N <- 100
X <- sort(round(rt(N, df=2), 2))
tt <- c(-100, seq(-2,2, len=201), +100)
it <- findInterval(tt, X)
tt[it < 1 | it >= N] # only first and last are outside range(X)
## See that this is N * Fn(.) :
tt <- c(tt,X)
eps <- 100 * .Machine$double.eps
require(stepfun)
stopifnot( it [c(1,203)] == c(0, 100),
    all.equal(N * ecdf(X)(tt),
    findInterval(tt, X), tol = eps),
    findInterval(tt,X) == apply( outer(tt, X, ">="), 1, sum)
)
```

```
fitted.values Extract Model Fitted Values
```


## Description

fitted is a generic function which extracts fitted values from objects returned by modeling functions. fitted.values is an alias for it.

All object classes which are returned by model fitting functions should provide a fitted method. (Note that the generic is fitted and not fitted.values.)
Methods can make use of napredict methods to compensate for the omission of missing values. The default, lm and glm methods do.

## Usage

fitted(object, ...)
fitted.values(object, ...)

## Arguments

object an object for which the extraction of model fitted values is meaningful.
... other arguments.

## Value

Fitted values extracted from the object x.

## See Also

coefficients, glm, lm, residuals.

## fivenum Tukey Five-Number Summaries

## Description

Returns Tukey's five number summary (minimum, lower-hinge, median, upper-hinge, maximum) for the input data.

## Usage

fivenum(x, na.rm = TRUE)

## Arguments

$x \quad$ numeric, maybe including NAs and $+/$-Infs.
na.rm logical; if TRUE, all NA and NaNs are dropped, before the statistics are computed.

## Value

A numeric vector of length 5 containing the summary information. See boxplot.stats for more details.

## See Also

IQR, boxplot.stats, median, quantile, range.

## Examples

```
fivenum(c(rnorm(100),-1:1/0))
```

fix Fix an Object

## Description

fix invokes the editor specified in options("editor") on $x$ and then assigns the new (edited) version of $x$ in the user's workspace.

## Usage

fix(x, ...)

## Arguments

x
an R object.
... arguments to pass to editor.

## See Also <br> edit, edit.data.frame

## Examples

```
## Assume 'my.fun' is a user defined function :
fix(my.fun)
## now my.fun is changed
## Also,
fix(my.data.frame) # calls up data editor
fix(my.data.frame, factor.mode="char") # use of ...
```


## Foreign Foreign Function Interface

## Description

Functions to make calls to compiled code that has been loaded into R.

## Usage

```
                        .C(name, ..., NAOK = FALSE, DUP = TRUE, PACKAGE)
                .Fortran(name, ..., NAOK = FALSE, DUP = TRUE, PACKAGE)
                .External(name, ..., PACKAGE)
            .Call(name, ..., PACKAGE)
.External.graphics(name, ..., PACKAGE)
    .Call.graphics(name, ..., PACKAGE)
```


## Arguments

name a character string giving the name of a C function or Fortran subroutine.
... arguments to be passed to the foreign function.
NAOK if TRUE then any NA or NaN or Inf values in the arguments are passed on to the foreign function. If FALSE, the presence of NA or NaN or Inf values is regarded as an error.
DUP if TRUE then arguments are "duplicated" before their address is passed to C or Fortran.

PACKAGE if supplied, confine the search for the name to the DLL given by this argument (plus the conventional extension, .so, .sl, .dll,...). This is intended to add safety for packages, which can ensure by using this argument that no other package can override their external symbols. Use PACKAGE="base" for symbols linked in to R.

## Details

The functions . C and .Fortran can be used to make calls to C and Fortran code.
.External and .External.graphics can be used to call compiled code that uses R objects in the same way as internal R functions.
. Call and . Call.graphics can be used call compiled code which makes use of internal R objects. The arguments are passed to the C code as a sequence of R objects. It is included to provide compatibility with S version 4.
For details about how to write code to use with .Call and .External, see the chapter on "System and foreign language interfaces" in "Writing R Extensions" in the 'doc/manual' subdirectory of the R source tree).

## Value

The functions . C and .Fortran return a list similar to the . . . list of arguments passed in, but reflecting any changes made by the C or Fortran code.
.External, Call, .External.graphics, and .Call.graphics return an R object.
These calls are typically made in conjunction with dyn.load which links DLLs to R.

The .graphics versions of . Call and .External are used when calling code which makes low-level graphics calls. They take additional steps to ensure that the device driver display lists are updated correctly.

## Argument types

The mapping of the types of R arguments to C or Fortran arguments in .C or .Fortran is

| R | C | Fortran |
| :--- | :--- | :--- |
| integer | int $*$ | integer |
| numeric | double ${ }^{*}$ | double precision |
| - or - | float $*$ | real |
| complex | Rcomplex * | double complex |
| logical | int ${ }^{*}$ | integer |
| character | char $* *$ | [see below] |
| list | SEXP $*$ | not allowed |
| other | SEXP | not allowed |

Numeric vectors in R will be passed as type double * to C (and as double precision to Fortran) unless (i) .C or .Fortran is used, (ii) DUP is false and (iii) the argument has attribute Csingle set to TRUE (use as.single or single). This mechanism is only intended to be used to facilitate the interfacing of existing C and Fortran code.
The C type Rcomplex is defined in 'Complex.h' as a typedef struct \{double r; double i;\}. Fortran type double complex is an extension to the Fortran standard, and the availability of a mapping of complex to Fortran may be compiler dependent.

Note: The C types corresponding to integer and logical are int, not long as in S.
The first character string of a character vector is passed as a C character array to Fortran: that string may be usable as character*255 if its true length is passed separately. Only up to 255 characters of the string are passed back. (How well this works, or even if it works at all, depends on the C and Fortran compilers and the platform.)
Missing (NA) string values are passed to . C as the string "NA". As the C char type can represent all possible bit patterns there appears to be no way to distinguish missing strings from the string "NA". If this distinction is important use .Call.
Functions, expressions, environments and other language elements are passed as the internal R pointer type SEXP. This type is defined in 'Rinternals.h' or the arguments can be declared as generic pointers, void *. Lists are passed as C arrays of SEXP and can be declared as void * or SEXP *.
$R$ functions can be invoked using call_S or call_R and can be passed lists or the simple types as arguments.

## Header files for external code

Writing code for use with .External and . Call will use internal R structures. If possible use just those defined in 'Rinternals.h' and/or the macros in 'Rdefines.h', as other header files are not installed and are even more likely to be changed.

## Note

$D U P=F A L S E$ is dangerous.
There are two dangers with using DUP=FALSE.

The first is that if you pass a local variable to .C/.Fortran with DUP=FALSE, your compiled code can alter the local variable and not just the copy in the return list. Worse, if you pass a local variable that is a formal parameter of the calling function, you may be able to change not only the local variable but the variable one level up. This will be very hard to trace.
The second is that lists are passed as a single R SEXP with DUP=FALSE, not as an array of SEXP. This means the accessor macros in 'Rinternals.h' are needed to get at the list elements and the lists cannot be passed to call_S/call_R. New code using R objects should be written using . Call or .External, so this is now only a minor issue.
(Prior to R version 1.2.0 there has a third danger, that objects could be moved in memory by the garbage collector. The current garbage collector never moves objects.)
It is safe and useful to set DUP=FALSE if you do not change any of the variables that might be affected, e.g.,

```
.C("Cfunction", input=x, output=numeric(10)).
```

In this case the output variable did not exist before the call so it cannot cause trouble. If the input variable is not changed in the C code of Cfunction you are safe.

Neither . Call nor .External copy their arguments. You should treat arguments you receive through these interfaces as read-only.

```
See Also
dyn.load.
```

Formaldehyde Determination of Formaldehyde

## Description

These data are from a chemical experiment to prepare a standard curve for the determination of formaldehyde by the addition of chromatropic acid and concentrated sulpuric acid and the reading of the resulting purple color on a spectophotometer.

## Usage

data(Formaldehyde)

## Format

A data frame with 6 observations on 2 variables.

| $[, 1]$ | carb | numeric | Carbohydrate (ml) |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | optden | numeric | Optical Density |

## Source

Bennett, N. A. and N. L. Franklin (1954) Statistical Analysis in Chemistry and the Chemical Industry. New York: Wiley.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(Formaldehyde)
plot(optden ~ carb, data = Formaldehyde,
    xlab = "Carbohydrate (ml)", ylab = "Optical Density",
    main = "Formaldehyde data", col = 4, las = 1)
abline(fm1 <- lm(optden ~ carb, data = Formaldehyde))
summary(fm1)
opar <- par(mfrow = c(2,2), oma = c(0, 0, 1.1, 0))
plot(fm1)
par(opar)
```

formals Access to and Manipulation of the Formal Arguments

## Description

Get or set the formal arguments of a function.

## Usage

```
formals(fun = sys.function(sys.parent()))
formals(fun) <- list
```


## Arguments

fun a function object or a character string naming the function to be manipulated. If not specified, the function calling body is used.
list a list of R expressions.

## Value

formals returns the formal argument list of the function specified.
The assignment form sets the formals of a function to the list on the right hand side.

## See Also

args for a "human-readable" version, alist, body, function.

## Examples

```
length(formals(lm)) # the number of formal arguments
names(formals(boxplot)) # formal arguments names
f <- function(x)a+b
formals(f) <- alist(a=,b=3) # function(a,b=3)a+b
f(2) # result = 5
```


## format

Encode in a Common Format

## Description

Format an R object for pretty printing: format.pval is intended for formatting p-values.

## Usage

```
format(x, ...)
format.AsIs(x, width = 12, ...)
format.data.frame(x, ..., justify = "none")
format.default(x, trim = FALSE, digits = getOption("digits"),
        nsmall = 0, justify = c("left", "right", "none"),
        big.mark = "", big.interval = 3,
        small.mark = "", small.interval = 5,
    decimal.mark = ".", ...)
format.factor(x, ...)
format.pval(pv, digits = max(1, getOption("digits") - 2),
    eps = .Machine$double.eps, na.form = "NA")
prettyNum(x, big.mark = "", big.interval = 3,
            small.mark = "", small.interval = 5,
        decimal.mark = ".", ...)
```


## Arguments

| x | any R object (conceptually); typically numeric. |
| :---: | :---: |
| trim | logical; if TRUE, leading blanks are trimmed off the strings. |
| digits | how many significant digits are to be used for numeric x . This is suggestion: enough decimal places will be used so that the smallest (in magnitude) number has this many significant digits. |
| nsmall | number of digits which will always appear to the right of the decimal point in formatting real/complex numbers in non-scientific formats. Allowed values 0 <= nsmall <= 20 . |
| justify | should character vector be left-justified, right-justified or left alone. When justifying, the field width is that of the longest string. |
| big.mark | character; if not empty used as mark between every big.interval decimals before (hence big) the decimal point. |
| big.interval | see big.mark above; defaults to 3 . |
| small.mark | character; if not empty used as mark between every small.interval decimals after (hence small) the decimal point. |
| small.interval |  |
|  | see small.mark above; defaults to 5 . |
| decimal.mark | the character used to indicate the numeric decimal point. |
| pv | a numeric vector. |
| eps | a numerical tolerance: see Details. |
| na.form | character representation of NAs. |
| width | the returned vector has elements of at most width. |
|  | her arguments passed to or from other methods. |

## Details

These functions convert their first argument to a vector (or array) of character strings which have a common format (as is done by print), fulfilling length(format*(x, *)) == length $(x)$. The trimming with trim $=$ TRUE is useful when the strings are to be used for plot axis annotation.
format.AsIs deals with columns of complicated objects that have been extracted from a data frame.
format.pval is mainly an auxiliary function for print.summary.lm etc., and does separate formatting for fixed, floating point and very small values; those less than eps are formatted as "< [eps]" (where "[eps]" stands for format(eps, digits).
The function formatC provides a rather more flexible formatting facility for numbers, but does not provide a common format for several numbers, nor it is platform-independent.
format.data.frame formats the data frame column by column, applying the appropriate method of format for each column.
prettyNum is the utility function for prettifying $x$. If $x$ is not a character, format (x[i], ...) is applied to each element, and then it is left unchanged if all the other arguments are at their defaults. Note that prettyNum (x) may behave unexpectedly if $x$ is a character not resulting from something like format (<number>).

## Note

Currently format drops trailing zeroes, so format(6.001, digits=2) gives "6" and format (c(6.0, 13.1), digits=2) gives c(" 6", "13").
Character(s) " in input strings $x$ are escaped to $\backslash$ ".

## See Also

format.info indicates how something would be formatted; formatC, paste, as.character.

## Examples

```
format(1:10)
zz <- data.frame("(row names)"= c("aaaaa", "b"), check.names=FALSE)
format(zz)
format(zz, justify="left")
## use of nsmall
format(13.7)
format(13.7, nsmall=3)
r <- c("76491283764.97430", "29.12345678901", "-7.1234", "-100.1","1123")
## American:
prettyNum(r, big.mark = ",")
## Some Europeans:
prettyNum(r, big.mark = "'", decimal.mark = ",")
(dd <- sapply(1:10, function(i)paste((9:0)[1:i],collapse="")))
prettyNum(dd, big.mark="'")
pN <- pnorm(1:7, lower=FALSE)
cbind(format (pN, small.mark = " ", digits = 15))
```

```
cbind(formatC(pN, small.mark = " ", digits = 17, format = "f"))
## handling of quotes
zz <- data.frame(a=I("abc"), b=I("def\"gh"))
format(zz)
p <- c(47,13,2,.1,.023,.0045, 1e-100)/1000
format.pval(p)
format.pval(p / 0.9)
format.pval(p / 0.9, dig=3)
```

```
format.info format(.) Information
```


## Description

Information is returned on how format(x, digits $=$ options("digits")) would be formatted.

## Usage

format.info(x)

## Arguments

$\mathrm{x} \quad$ (numeric) vector; potential argument of format (x, ...).

## Value

An integer vector of length 3 , say r.
$r$ [1] width (number of characters) used for format ( x )
$r$ [2] number of digits after decimal point.
$r$ [3] in 0:2; if $\geq 1$, exponential representation would be used, with exponent length of $r[3]+1$.

## Note

The result depends on the value of options("digits").

## See Also

format, formatC.

## Examples

```
dd <- options("digits") ; options(digits = 7) #-- for the following
format.info(123) # 3 0 0
format.info(pi) # 8 6 0
format.info(1e8) # 5 0 1 - exponential "1e+08"
format.info(1e222)#6 0 2 - exponential "1e+222"
x <- pi*10^c(-10,-2,0:2,8,20)
names(x) <- formatC(x,w=1,dig=3,format="g")
```

```
cbind(sapply(x,format))
t(sapply(x, format.info))
# Reset old options:
options(dd)
```

formatC Formatting Using C-style Formats

## Description

Formatting numbers individually and flexibly, using C style format specifications. format. char is a helper function for formatC.

## Usage

```
formatC(x, digits = NULL, width = NULL,
            format = NULL, flag = "", mode = NULL,
            big.mark = "", big.interval = 3,
            small.mark = "", small.interval = 5,
        decimal.mark = ".")
format.char(x, width = NULL, flag = "-")
```


## Arguments

x
digits the desired number of digits after the decimal point (format $=$ "f") or significant digits (format = "g", = "e" or = "fg").
Default: 2 for integer, 4 for real numbers. If less than 0 , the C default of 6 digits is used.
width the total field width; if both digits and width are unspecified, width defaults to 1 , otherwise to digits +1 . width $=0$ will use width $=$ digits, width < 0 means left justify the number in this field (equivalent to flag ="-"). If necessary, the result will have more characters than width.
format equal to "d" (for integers), "f", "e", "E", "g", "G", "fg" (for reals), or "s" (for strings). Default is "d" for integers, "g" for reals.
"f" gives numbers in the usual xxx.xxx format; "e" and "E" give $\mathrm{n} . \mathrm{ddde}^{+n n}$ or $\mathrm{n} . \mathrm{dddE}_{\mathrm{n}} \mathrm{nn}$ (scientific format); " g " and " G " put x [i] into scientific format only if it saves space to do so.
" fg " uses fixed format as " f ", but digits as number of significant digits. Note that this can lead to quite long result strings, see examples below.
flag format modifier as in Kernighan and Ritchie (1988, page 243). "0" pads leading zeros; "-" does left adjustment, others are "+", " ", and "\#".
mode "double" (or "real"), "integer" or "character". Default: Determined from the storage mode of $x$.
big.mark, big.interval, small.mark, small.interval, decimal.mark used for prettying longer decimal sequences, passed to prettyNum the help page explains details.

## Details

If you set format it over-rides the setting of mode, so formatC(123.45, mode="double", format="d") gives 123.

The rendering of scientific format is platform-dependent: some systems use n . ddde+nnn or $n$. dddenn rather than $n$.ddde +nn .
formatC does not necessarily align the numbers on the decimal point, so formatC (c (6.11, 13.1), digits=2, format="fg") gives c("6.1", " 13"). If you want common formatting for several numbers, use format.

## Value

A character object of same size and attributes as x . Unlike format, each number is formatted individually. Looping over each element of $x$, $\operatorname{sprintf}(. .$.$) is called (inside the C$ function str_signif).
format. char (x) and formatC, for character x , do simple (left or right) padding with white space.

## Author(s)

Originally written by Bill Dunlap, later much improved by Martin Maechler, it was first adapted for R by Friedrich Leisch.

## References

Kernighan, B. W. and Ritchie, D. M. (1988) The C Programming Language. Second edition. Prentice Hall.

## See Also

format, sprintf for more general C like formatting.

## Examples

```
xx <- pi * 10^(-5:4)
dd <- options(digits = 4) # only for format
cbind(format(xx), formatC(xx))
cbind(formatC(xx, wid = 9, flag = "-"))
cbind(formatC(xx, dig = 5, wid = 8, format = "f", flag = "0"))
format.char(c("a", "Abc", "no way"), wid = -7) # <=> flag = "-"
formatC( c("a", "Abc", "no way"), wid = -7) # <<> flag = "-"
formatC(c((-1:1)/0,c(1,100)*pi), wid=8, dig=1)
xx <- c(1e-12,-3.98765e-10,1.45645e-69,1e-70,pi*1e37,3.44e4)
## 1 1 2 % 3 % 4
formatC(xx)
formatC(xx, format="fg") # special "fixed" format.
formatC(xx, format="f", dig=80)#>> also long strings
options(dd) # reset
```

formatDL

## Description

Format vectors of items and their descriptions as 2-column tables or LaTeX-style description lists.

## Usage

```
formatDL(x, y, style = c("table", "list"),
    width = 0.9 * getOption("width"), indent = NULL)
```


## Arguments

x
a vector giving the items to be described, or a list of length 2 or a matrix with 2 columns giving both items and descriptions.
y a vector of the same length as x with the corresponding descriptions. Only used if x does not already give the descriptions.
style a character string specifying the rendering style of the description information. If "table", a two-column table with items and descriptions as columns is produced (similar to Texinfo's @table environment. If "list", a LaTeX-style tagged description list is obtained.
width a positive integer giving the target column for wrapping lines in the output.
indent a positive integer specifying the indentation of the second column in table style, and the indentation of continuation lines in list style. Must not be greater than width/2, and defaults to width/3 for table style and width/9 for list style.

## Details

After extracting the vectors of items and corresponding descriptions from the arguments, both are coerced to character vectors.
In table style, items with more than indent - 3 characters are displayed on a line of their own.

## Value

a character vector with the formatted entries.

## Examples

```
## Use R to create the 'INDEX' for package 'eda' from its 'CONTENTS'
x <- read.dcf(file = system.file("CONTENTS", package = "eda"),
    fields = c("Entry", "Description"))
x <- as.data.frame(x)
writeLines(formatDL(x$Entry, x$Description))
## or equivalently: writeLines(formatDL(x))
## Same information in tagged description list style:
writeLines(formatDL(x$Entry, x$Description, style = "list"))
## or equivalently: writeLines(formatDL(x, style = "list"))
```


## formula Model Formulae

## Description

The generic function formula and its specific methods provide a way of extracting formulae which have been included in other objects.
as.formula is almost identical, additionally preserving attributes when object already inherits from "formula". The default value of the env argument is used only when the formula would otherwise lack an environment.

## Usage

```
y ~ model
formula(x, ...)
as.formula(object, env = parent.frame())
```


## Arguments

```
x, object an object
... further arguments passed to or from other methods.
env the environment to associate with the result.
```


## Details

The models fit by, e.g., the lm and glm functions are specified in a compact symbolic form. The ~ operator is basic in the formation of such models. An expression of the form y ~ model is interpreted as a specification that the response $y$ is modelled by a linear predictor specified symbolically by model. Such a model consists of a series of terms separated by + operators. The terms themselves consist of variable and factor names separated by : operators. Such a term is interpreted as the interaction of all the variables and factors appearing in the term.
In addition to + and :, a number of other operators are useful in model formulae. The * operator denotes factor crossing: $a * b$ interpreted $a s a+b+a: b$. The - operator indicates crossing to the specified degree. For example $(a+b+c) \wedge 2$ is identical to $(a+b+c) *(a+b+c)$ which in turn expands to a formula containing the main effects for $a$, $b$ and $c$ together with their second-order interactions. The \%in\% operator indicates that the terms on its left are nested within those on the right. For example $a+b \% i n \% a$ expands to the formula $a+a: b$. The - operator removes the specified terms, so that $(a+b+c)^{\wedge} 2-a: b$ is identical to $a+b$ $+c+b: c+a: c$. It can also used to remove the intercept term: $y^{\sim} x-1$ is a line through the origin. A model with no intercept can be also specified as $\mathrm{y}^{\sim} \mathrm{x}+0$ or $0+\mathrm{y}^{\sim} \mathrm{x}$.
While formulae usually involve just variable and factor names, they can also involve arithmetic expressions. The formula $\log (y) \sim a+\log (x)$ is quite legal. When such arithmetic expressions involve operators which are also used symbolically in model formulae, there can be confusion between arithmetic and symbolic operator use.
To avoid this confusion, the function I () can be used to bracket those portions of a model formula where the operators are used in their arithmetic sense. For example, in the formula $y \sim a+I(b+c)$, the term $b+c$ is to be interpreted as the sum of $b$ and $c$.

## Value

All the functions above produce an object of class "formula" which contains a symbolic model formula.

## Environments

A formula object has an associated environment, and this environment (rather than the parent environment) is used by model.frame to evaluate variables that are not found in the supplied data argument.

Formulas created with the $\sim$ operator use the environment in which they were created. Formulas created with as.formula will use the env argument for their environment. Preexisting formulas extracted with as.formula will only have their environment changed if env is explicitly given.

## See Also

I.

For formula manipulation: terms, and all.vars; for typical use: lm, glm, and coplot.

## Examples

```
class(fo <- y ~ x1*x2) # "formula"
fo
typeof(fo)# R internal : "language"
terms(fo)
environment(fo)
environment(as.formula("y ~ x"))
environment(as.formula("y ~ x",env=new.env()))
## Create a formula for a model with a large number of variables:
xnam <- paste("x", 1:25, sep="")
(fmla <- as.formula(paste("y ~ ", paste(xnam, collapse= "+"))))
```

fourfoldplot Fourfold Plots

## Description

Creates a fourfold display of a 2 by 2 by $k$ contingency table on the current graphics device, allowing for the visual inspection of the association between two dichotomous variables in one or several populations (strata).

## Usage

```
fourfoldplot(x, color = c("#99CCFF", "#6699CC"), conf.level = 0.95,
    std = c("margins", "ind.max", "all.max"),
    margin = c(1, 2), space = 0.2, main = NULL,
    mfrow = NULL, mfcol = NULL)
```


## Arguments

x
color a vector of length 2 specifying the colors to use for the smaller and larger diagonals of each 2 by 2 table.
conf.level confidence level used for the confidence rings on the odds ratios. Must be a single nonnegative number less than 1 ; if set to 0 , confidence rings are suppressed.
std a character string specifying how to standardize the table. Must be one of "margins", "ind.max", or "all.max", and can be abbreviated by the initial letter. If set to "margins", each 2 by 2 table is standardized to equate the margins specified by margin while preserving the odds ratio. If "ind.max" or "all.max", the tables are either individually or simultaneously standardized to a maximal cell frequency of 1 .
$\operatorname{margin} \quad$ a numeric vector with the margins to equate. Must be one of 1,2 , or $c(1$, 2) (the default), which corresponds to standardizing the row, column, or both margins in each 2 by 2 table. Only used if std equals "margins".
space the amount of space (as a fraction of the maximal radius of the quarter circles) used for the row and column lebals.
main character string for the fourfold title.
mfrow a numeric vector of the form $\mathrm{c}(\mathrm{nr}, \mathrm{nc})$, indicating that the displays for the 2 by 2 tables should be arranged in an nr by nc layout, filled by rows.
$\mathrm{mfcol} \quad$ a numeric vector of the form $\mathrm{c}(\mathrm{nr}, \mathrm{nc})$, indicating that the displays for the 2 by 2 tables should be arranged in an nr by nc layout, filled by columns.

## Details

The fourfold display is designed for the display of 2 by 2 by $k$ tables.
Following suitable standardization, the cell frequencies $f_{i j}$ of each 2 by 2 table are shown as a quarter circle whose radius is proportional to $\sqrt{f_{i j}}$ so that its area is proportional to the cell frequency. An association (odds ratio different from 1) between the binary row and column variables is indicated by the tendency of diagonally opposite cells in one direction to differ in size from those in the other direction; color is used to show this direction. Confidence rings for the odds ratio allow a visual test of the null of no association; the rings for adjacent quadrants overlap iff the observed counts are consistent with the null hypothesis.
Typically, the number $k$ corresponds to the number of levels of a stratifying variable, and it is of interest to see whether the association is homogeneous across strata. The fourfold display visualizes the pattern of association. Note that the confidence rings for the individual odds ratios are not adjusted for multiple testing.

## References

Friendly, M. (1994). A fourfold display for 2 by 2 by $k$ tables. Technical Report 217, York University, Psychology Department. http://hotspur.psych.yorku.ca/ftp/sas/ catdata/4fold.ps.gz

## See Also

## Examples

```
data(UCBAdmissions)
## Use the Berkeley admission data as in Friendly (1995).
x <- aperm(UCBAdmissions, c(2, 1, 3))
dimnames(x)[[2]] <- c("Yes", "No")
names(dimnames(x)) <- c("Sex", "Admit?", "Department")
ftable(x)
## Fourfold display of data aggregated over departments, with
## frequencies standardized to equate the margins for admission
## and sex.
## Figure 1 in Friendly (1994).
fourfoldplot(margin.table(x, c(1, 2)))
## Fourfold display of x, with frequencies in each table
## standardized to equate the margins for admission and sex.
## Figure 2 in Friendly (1994).
fourfoldplot(x)
## Fourfold display of x, with frequencies in each table
## standardized to equate the margins for admission. but not
## for sex.
## Figure 3 in Friendly (1994).
fourfoldplot(x, margin = 2)
```

frame $\quad$ Create / Start a New Plot Frame

## Description

This function (frame is an alias for plot.new) causes the completion of plotting in the current plot (if there is one) and an advance to a new graphics frame. This is used in all high-level plotting functions and also useful for skipping plots when a multi-figure region is in use.

## Usage

```
plot.new()
frame()
```


## See Also

```
plot.window, plot.default.
```

```
freeny Freeny's Revenue Data
```


## Description

Freeny's data on quarterly revenue and explanatory variables.

## Usage

data(freeny)

## Format

There are three 'freeny' data sets.
freeny.y is a time series with 39 observations on quarterly revenue from (1962,2Q) to (1971,4Q).
freeny. $x$ is a matrix of explanatory variables. The columns are freeny.y lagged 1 quarter, price index, income level, and market potential.

Finally, freeny is a data frame with variables y, lag.quarterly.revenue, price.index, income.level, and market. potential obtained from the above two data objects.

## Source

A. E. Freeny (1977) A Portable Linear Regression Package with Test Programs. Bell Laboratories memorandum.

## Examples

```
data(freeny)
summary(freeny)
pairs(freeny, main = "freeny data")
summary(fm1 <- lm(y ~ ., data = freeny))
opar <- par(mfrow = c(2, 2), oma = c(0, 0, 1.1, 0),
    mar = c(4.1, 4.1, 2.1, 1.1))
plot(fm1)
par(opar)
```

ftable Flat Contingency Tables

## Description

Create "flat" contingency tables.

## Usage

```
ftable(..., exclude = c(NA, NaN), row.vars = NULL, col.vars = NULL)
```


## Arguments

$$
\begin{array}{ll}
\ldots & \begin{array}{l}
\text { R objects which can be interpreted as factors (including character strings), } \\
\text { or a list (or data frame) whose components can be so interpreted, or a } \\
\text { contingency table object of class "table" or "ftable". }
\end{array} \\
\text { exclude } & \begin{array}{l}
\text { values to use in the exclude argument of factor when interpreting non- } \\
\text { factor objects. }
\end{array} \\
\text { row.vars } & \begin{array}{l}
\text { a vector of integers giving the numbers of the variables, or a character } \\
\text { vector giving the names of the variables to be used for the rows of the flat } \\
\text { contingency table. }
\end{array} \\
\text { col.vars } & \begin{array}{l}
\text { a vector of integers giving the numbers of the variables, or a character } \\
\text { vector giving the names of the variables to be used for the columns of the } \\
\text { flat contingency table. }
\end{array}
\end{array}
$$

## Details

ftable creates "flat" contingency tables. Similar to the usual contingency tables, these contain the counts of each combination of the levels of the variables (factors) involved. This information is then re-arranged as a matrix whose rows and columns correspond to unique combinations of the levels of the row and column variables (as specified by row. vars and col.vars, respectively). The combinations are created by looping over the variables in reverse order (so that the levels of the "left-most" variable vary the slowest). Displaying a contingency table in this flat matrix form (via print.ftable, the print method for objects of class "ftable") is often preferable to showing it as a higher-dimensional array.
ftable is a generic function. Its default method, ftable.default, first creates a contingency table in array form from all arguments except row.vars and col.vars. If the first argument is of class "table", it represents a contingency table and is used as is; if it is a flat table of class "ftable", the information it contains is converted to the usual array representation using as.ftable. Otherwise, the arguments should be R objects which can be interpreted as factors (including character strings), or a list (or data frame) whose components can be so interpreted, which are cross-tabulated using table. Then, the arguments row.vars and col.vars are used to collapse the contingency table into flat form. If neither of these two is given, the last variable is used for the columns. If both are given and their union is a proper subset of all variables involved, the other variables are summed out.

Function ftable.formula provides a formula method for creating flat contingency tables.

## Value

ftable returns an object of class "ftable", which is a matrix with counts of each combination of the levels of variables with information on the names and levels of the (row and columns) variables stored as attributes "row.vars" and "col.vars".

## See Also

ftable.formula for the formula interface (which allows a data = . argument); read.ftable for information on reading, writing and coercing flat contingency tables; table for "ordinary" cross-tabulation; xtabs for formula-based cross-tabulation.

## Examples

```
## Start with a contingency table
data(Titanic)
ftable(Titanic, row.vars = 1:3)
```

```
ftable(Titanic, row.vars = 1:2, col.vars = "Survived")
ftable(Titanic, row.vars = 2:1, col.vars = "Survived")
## Start with a data frame.
data(mtcars)
x <- ftable(mtcars[c("cyl", "vs", "am", "gear")])
x
ftable(x, row.vars = c(2, 4))
```

ftable.formula Formula Notation for Flat Contingency Tables

## Description

Produce or manipulate a flat contingency table using formula notation.

## Usage

ftable(formula, data = NULL, subset, na.action, ...)

## Arguments

| formula | a formula object with both left and right hand sides specifying the column <br> and row variables of the flat table. |
| :--- | :--- |
| data | a data frame, list or environment containing the variables to be cross- <br> tabulated, or a contingency table (see below). |
| subset | an optional vector specifying a subset of observations to be used. Ignored <br> if data is a contingency table. |
| na.action | a function which indicates what should happen when the data contain <br> NAs. Ignored if data is a contingency table. |
| $\ldots$ | further arguments to the default ftable method may also be passed as <br> arguments, see ftable.default. |

## Details

This is a method of the generic function ftable.
The left and right hand side of formula specify the column and row variables, respectively, of the flat contingency table to be created. Only the + operator is allowed for combining the variables. A . may be used once in the formula to indicate inclusion of all the "remaining" variables.

If data is an object of class "table" or an array with more than 2 dimensions, it is taken as a contingency table, and hence all entries should be nonnegative. Otherwise, if it is not a flat contingency table (i.e., an object of class "ftable"), it should be a data frame or matrix, list or environment containing the variables to be cross-tabulated. In this case, na.action is applied to the data to handle missing values, and, after possibly selecting a subset of the data as specified by the subset argument, a contingency table is computed from the variables.
The contingency table is then collapsed to a flat table, according to the row and column variables specified by formula.

## Value

A flat contingency table which contains the counts of each combination of the levels of the variables, collapsed into a matrix for suitably displaying the counts.

## See Also

```
ftable, ftable.default; table.
```


## Examples

```
data(Titanic)
Titanic
x <- ftable(Survived ~ ., data = Titanic)
x
ftable(Sex ~ Class + Age, data = x)
```

function Function Definition

## Description

These functions provide the base mechanisms for defining new functions in the R language.

## Usage

```
function( arglist ) expr
```

return(value)

## Arguments

arglist Empty or one or more name or name=expression terms.
value An expression, or a series of expressions separated by commas.

## Details

In $R$ (unlike $S$ ) the names in an argument list cannot be quoted non-standard names.
If value is a series of expressions, the value returned is a list of the evaluated expressions, with names set to the expressions where these are the names of R objects.

## See Also

args and body for accessing the arguments and body of a function. debug for debugging; invisible for return(.)ing invisibly.

## Examples

```
norm <- function(x) sqrt(x%*%x)
norm(1:4)
## An anonymous function:
(function(x,y){ z<- x^2 + y^2; x+y+z }) (0:7, 1)
```

```
GammaDist The Gamma Distribution
```


## Description

Density, distribution function, quantile function and random generation for the Gamma distribution with parameters shape and scale.

## Usage

```
dgamma(x, shape, rate = 1, scale = 1/rate, log = FALSE)
pgamma(q, shape, rate = 1, scale = 1/rate, lower.tail = TRUE, log.p = FALSE)
qgamma(p, shape, rate = 1, scale = 1/rate, lower.tail = TRUE, log.p = FALSE)
rgamma(n, shape, rate = 1, scale = 1/rate)
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
$\mathrm{n} \quad$ number of observations. If length $(\mathrm{n})>1$, the length is taken to be the number required.
rate an alternative way to specify the scale.
shape, scale shape and scale parameters.
$\log , \log . \mathrm{p} \quad$ logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

If scale is omitted, it assumes the default value of 1.
The Gamma distribution with parameters shape $=\alpha$ and scale $=\sigma$ has density

$$
f(x)=\frac{1}{\sigma^{\alpha} \Gamma(\alpha)} x^{\alpha-1} e^{-x / \sigma}
$$

for $x>0, \alpha>0$ and $\sigma>0$. The mean and variance are $E(X)=\alpha \sigma$ and $\operatorname{Var}(X)=\alpha \sigma^{2}$.

## Value

dgamma gives the density, pgamma gives the distribution function qgamma gives the quantile function, and rgamma generates random deviates.

## Note

The $S$ parametrization is via shape and rate: S has no scale parameter. Prior to 1.4.0 R only had scale.
The cumulative hazard $H(t)=-\log (1-F(t))$ is -pgamma(t, ..., lower = FALSE, log $=$ TRUE).

## See Also

gamma for the Gamma function, dbeta for the Beta distribution and dchisq for the chisquared distribution which is a special case of the Gamma distribution.

## Examples

-log(dgamma(1:4, shape=1))
p <- (1:9)/10
pgamma (qgamma (p,shape=2), shape=2)
1 - $1 / \exp (q g a m m a(p$, shape=1))
gc
Garbage Collection

## Description

A call of gc causes a garbage collection to take place. gcinfo sets a flag so that automatic collection is either silent (verbose=FALSE) or prints memory usage statistics (verbose=TRUE).

## Usage

gc(verbose = getOption("verbose"))
gcinfo(verbose)

## Arguments

verbose logical; if TRUE, the garbage collection prints statistics about cons cells and the vector heap.

## Details

A call of gc causes a garbage collection to take place. This takes place automatically without user intervention, and the primary purpose of calling gc is for the report on memory usage. However, it can be useful to call gc after a large object has been removed, as this may prompt R to return memory to the operating system.

## Value

gc returns a matrix with rows "Ncells" (cons cells, usually 28 bytes each on 32 -bit systems and 56 bytes on 64 -bit systems, and "Vcells" (vector cells, 8 bytes each), and columns "used" and "gc trigger", each also interpreted in megabytes (rounded up to the next 0.1 Mb ).

If maxima have been set for either "Ncells" or "Vcells", a fifth column is printed giving the current limits in Mb (with NA denoting no limit).
gcinfo returns the previous value of the flag.

## See Also

Memory on R's memory management and gctorture if you are an R hacker.

## Examples

```
gc() #- do it now
gcinfo(TRUE) #-- in the future, show when R does it
x <- integer(100000); for(i in 1:18) x <- c(x,i)
gcinfo(verbose = FALSE)#-- don't show it anymore
gc(TRUE)
```

gc.time Report Time Spent in Garbage Collection

## Description

This function reports the time spent in garbage collection so far in the R session.

## Usage

```
gc.time()
```


## Value

A numerical vector of length 5 giving the user CPU time, the system CPU time, the elapsed time and children's user and system CPU times (normally both zero).

## Warnings

This is experimental functionality, likely to be removed as soon as the next release.
The timings are rounded up by the sampling interval for timing processes, and so are likely to be over-estimates.

## See Also

gc, proc.time for the timings for the session.

## Examples

```
gc.time()
```

gctorture $\quad$ Torture Garbage Collector

## Description

Provokes garbage collection on (nearly) every memory allocation. Intended to ferret out memory protection bugs. Also makes R run very slowly, unfortunately.

## Usage

```
gctorture(on = TRUE)
```


## Arguments

on
logical; turning it on/off.

## Value

Previous value.

## Author(s)

Peter Dalgaard

## Geometric The Geometric Distribution

## Description

Density, distribution function, quantile function and random generation for the geometric distribution with parameter prob.

## Usage

```
dgeom (x, prob, log = FALSE)
pgeom(q, prob, lower.tail = TRUE, log.p = FALSE)
qgeom(p, prob, lower.tail = TRUE, log.p = FALSE)
rgeom(n, prob)
```


## Arguments

```
x, q vector of quantiles representing the number of failures in a sequence of Bernoulli trials before success occurs.
p vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the
    number required.
prob probability of success in each trial.
log, log.p logical; if TRUE, probabilities p are given as log(p).
lower.tail logical; if TRUE (default), probabilities are P[X\leqx], otherwise, P[X>
    x].
```


## Details

The geometric distribution with prob $=p$ has density

$$
p(x)=p(1-p)^{x}
$$

for $x=0,1,2, \ldots$
If an element of x is not integer, the result of pgeom is zero, with a warning.
The quantile is defined as the smallest value $x$ such that $F(x) \geq p$, where $F$ is the distribution function.

## Value

dgeom gives the density, pgeom gives the distribution function, qgeom gives the quantile function, and rgeom generates random deviates.

## See Also

dnbinom for the negative binomial which generalizes the geometric distribution.

## Examples

```
pp <- sort(c((1:9)/10, 1 - .2^(2:8)))
print(qg <- qgeom(pp, prob = .2))
## test that qgeom is an inverse of pgeom
print(qg1 <- qgeom(pgeom(qg, prob=.2), prob =.2))
all(qg == qg1)
Ni <- rgeom(20, prob = 1/4); table(factor(Ni, 0:max(Ni)))
```


## get Return a Variable's Value

## Description

Search for an R object with a given name and return it if found.

## Usage

```
get(x, pos=-1, envir=as.environment(pos), mode="any", inherits=TRUE)
```


## Arguments

| x | a variable name (given as a quoted character string). |
| :--- | :--- |
| pos | where to look for the object (see the details section); if omitted, the <br> function will search, as if the name of the object appeared in unquoted in <br> an expression. |
| envir | an alternative way to specify an environment to look in; see the details <br> section. |
| mode | the mode of object sought. |
| inherits | should the enclosing frames of the environment be inspected? |

## Details

The pos argument can specify the environment in which to look for the object in any of several ways: as an integer (the position in the search list); as the character string name of an element in the search list; or as an environment (including using sys.frame to access the currently active function calls). The envir argument is an alternative way to specify an environment, but is primarily there for back compatibility.

The mode includes collections such as "numeric" and "function": any member of the collection will suffice.

## Value

This function searches the specified environment for a bound variable whose name is given by the character string $x$. If the variable's value is not of the correct mode, it is ignored.

If inherits is FALSE, only the first frame of the specified environment is inspected. If inherits is TRUE, the search is continued up through the parent frames until a bound value of the right mode is found.

Using a NULL environment is equivalent to using the current environment.

## See Also

exists.

## Examples

```
get("%%%")
```

getNativeSymbolInfo Obtain a description of a native (C/Fortran) symbol

## Description

This finds and returns as comprehensive a description of a dynamically loaded or "exported" built-in native symbol. It returns information about the name of the symbol, the library in which it is located and, if available, the number of arguments it expects and by which interface it should be called (i.e .Call, .C, .Fortran, or .External). Additionally, it returns the address of the symbol and this can be passed to other C routines which can invoke. Specifically, this provides a way to explicitly share symbols between different dynamically loaded package libraries. Also, it provides a way to query where symbols were resolved, and aids diagnosing strange behavior associated with dynamic resolution.

## Usage

getNativeSymbolInfo(name, PACKAGE)

## Arguments

name the name of the native symbol as used in a call to is.loaded, etc.
PACKAGE an optional argument that specifies to which dynamically loaded library we restrict the search for this symbol. If this is "base", we search in the R executable itself.

## Details

This uses the same mechanism for resolving symbols as is used in all the native interfaces (.Call, etc.). If the symbol has been explicitly registered by the shared library in which it is contained, information about the number of arguments and the interface by which it should be called will be returned. Otherwise, a generic native symbol object is returned.

## Value

If the symbol is not found, an error is raised. Otherwise, the value is a list containing the following elements:
name the name of the symbol, as given by the name argument.
address the native memory address of the symbol which can be used to invoke the routine, and also compare with other symbol address. This is an external pointer object and of class NativeSymbol.
package a list containing 3 elements:
name the short form of the library name which can be used as the value of the PACKAGE argument in the different native interface functions.
path the fully qualified name of the shared library file.
dynamicLookup a logical value indicating whether dynamic resolution is used when looking for symbols in this library, or only registered routines can be located.
numParameters the number of arguments that should be passed in a call to this routine.
Additionally, the list will have an additional class, being CRoutine, CallRoutine, FortranRoutine or ExternalRoutine corresponding to the R interface by which it should be invoked.

## Note

One motivation for accessing this reflectance information is to be able to pass native routines to C routines as "function pointers" in C. This allows us to treat native routines and R functions in a similar manner, such as when passing an $R$ function to $C$ code that makes callbacks to that function at different points in its computation (e.g. nls). Additionally, we can resolve the symbol just once and avoid resolving it repeatedly or using the internal cache. In the future, one may be able to treat NativeSymbol objects as directly callback objects.

## Author(s)

Duncan Temple Lang

## References

For information about registering native routines, see "In Search of C/C $++\&$ FORTRAN Routines", R News, volume 1, number 3, 2001, p20-23 (http://CRAN.R-project.org/ doc/Rnews/).

## See Also

is.loaded, .C, .Fortran, .External, .Call, dyn.load.

## Examples

```
getNativeSymbolInfo("dansari")
```

library (mva)
getNativeSymbolInfo(symbol.For("hcass2"))
getNumCConverters Management of.C argument conversion list

## Description

These functions provide facilities to manage the extensible list of converters used to translate $R$ objects to $C$ pointers for use in . C calls. The number and a description of each element in the list can be retrieved. One can also query and set the activity status of individual elements, temporarily ignoring them. And one can remove individual elements.

## Usage

```
getNumCConverters()
getCConverterDescriptions()
getCConverterStatus()
setCConverterStatus(id, status)
removeCConverter(id)
```


## Arguments

id either a number or a string identifying the element of interest in the converter list. A string sis matched against the description strings for each element to identify the element. Integers are specified starting at 1 (rather than 0).
status a logical value specifying whether the element is to be considered active (TRUE) or not (FALSE).

## Details

The internal list of converters is potentially used when converting individual arguments in a . C call. If an argument has a non-trivial class attribute, we iterate over the list of converters looking for the first that "matches". If we find a matching converter, we have it create the C -level pointer corresponding to the R object. When the call to the C routine is complete, we use the same converter for that argument to reverse the conversion and create an R object from the current value in the C pointer. This is done separately for all the arguments.
The functions documented here provide R user-level capabilities for investigating and managing the list of converters. There is currently no mechanism for adding an element to the converter list within the R language. This must be done in C code using the routine R_addToCConverter ().

## Value

getNumCConverters returns an integer giving the number of elements in the list, both active and inactive.
getCConverterDescriptions returns a character vector containing the description string of each element of the converter list.
getCConverterStatus returns a logical vector with a value for each element in the converter list. Each value indicates whether that converter is active (TRUE) or inactive (FALSE). The names of the elements are the description strings returned by getCConverterDescriptions.
setCConverterStatus returns the logical value indicating the activity status of the specified element before the call to change it took effect. This is TRUE for active and FALSE for inactive. removeCConverter returns TRUE if an element in the converter list was identified and removed. In the case that no such element was found, an error occurs.

## Author(s)

Duncan Temple Lang

## References

http://developer.R-project.org/CObjectConversion.pdf

## See Also

. C

## Examples

```
getNumCConverters()
getCConverterDescriptions()
getCConverterStatus()
old <- setCConverterStatus(1,FALSE)
setCConverterStatus(1,old)
```

removeCConverter(1)
removeCConverter(getCConverterDescriptions() [1])

```
getwd Get or Set Working Directory
```


## Description

getwd returns an absolute filename representing the current working directory of the R process; setwd (dir) is used to set the working directory to dir.

## Usage

getwd()
setwd(dir)

## Arguments

dir A character string.

Value
getwd returns a character vector, or NULL if the working directory is not available on that platform.
setwd returns NULL invisibly. It will give an error if it does not succeed.

## Note

These functions are not implemented on all platforms.

## Examples

```
(WD <- getwd())
```

if (!is.null(WD)) setwd(WD)
gl Generate Factor Levels

## Description

Generate factors by specifying the pattern of their levels.

## Usage

```
gl(n, k, length = n*k, labels = 1:n, ordered = FALSE)
```


## Arguments

n an integer giving the number of levels.
$\mathrm{k} \quad$ an integer giving the number of replications.
length an integer giving the length of the result.
labels an optional vector of labels for the resulting factor levels.
ordered a logical indicating whether the result should be ordered or not.

## Value

The result has levels from 1 to n with each value replicated in groups of length k out to a total length of length.
gl is modelled on the GLIM function of the same name.

## See Also

The underlying factor().

## Examples

```
## First control, then treatment:
gl(2, 8, label = c("Control", "Treat"))
## 20 alternating 1s and 2s
gl(2, 1, 20)
## alternating pairs of 1s and 2s
gl(2, 2, 20)
```

```
glm Fitting Generalized Linear Models
```


## Description

glm is used to fit generalized linear models, specified by giving a symbolic description of the linear predictor and a description of the error distribution.

## Usage

```
glm(formula, family = gaussian, data, weights = NULL, subset = NULL,
    na.action, start = NULL, offset = NULL,
    control = glm.control(...), model = TRUE, method = "glm.fit",
    \(\mathrm{x}=\) FALSE, \(\mathrm{y}=\) TRUE, contrasts \(=\) NULL, ...)
glm.fit(x, y, weights = rep(1, nrow(x)),
    start = NULL, etastart \(=\) NULL, mustart \(=\) NULL,
    offset \(=\) rep(0, nrow(x)),
    family = gaussian(), control = glm.control(),
    intercept = TRUE)
glm.fit.null(x, y, weights \(=\) rep(1, nrow(x)),
    start = NULL, etastart \(=\) NULL, mustart \(=\) NULL,
    offset \(=\) rep(0, nrow(x)),
    family = gaussian(), control = glm.control(),
    intercept = FALSE)
weights(object, type = c("prior", "working"), ...)
```


## Arguments

formula a symbolic description of the model to be fit. The details of model specification are given below.
family a description of the error distribution and link function to be used in the model. See family for details.
data an optional data frame containing the variables in the model. By default the variables are taken from environment (formula), typically the environment from which glm is called.
weights an optional vector of weights to be used in the fitting process.
subset an optional vector specifying a subset of observations to be used in the fitting process.
na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The "factory-fresh" default is na.omit.
start starting values for the parameters in the linear predictor.
etastart starting values for the linear predictor.
mustart starting values for the vector of means.
offset this can be used to specify an a priori known component to be included in the linear predictor during fitting.

| control | a list of parameters for controlling the fitting process. See the documen- <br> tation for glm. control for details. |
| :--- | :--- |
| model | a logical value indicating whether model frame should be included as a <br> component of the returned value. |
| method | the method to be used in fitting the model. The default (and presently <br> only) method glm.fit uses iteratively reweighted least squares (IWLS). |
| $\mathrm{x}, \mathrm{y}$ | For glm: logical values indicating whether the response vector and model <br> matrix used in the fitting process should be returned as components of <br> the returned value. |
|  | For glm.fit: x is a design matrix of dimension $\mathrm{n} * \mathrm{p}$, and y is a vector <br> of observations of length n. |
| contrasts | an optional list. See the contrasts.arg of model.matrix. default. <br> object |
| an object inheriting from class "glm". |  |
| type | character, partial matching allowed. Type of weights to extract from the <br> fitted model object. |
| intercept | logical. Should an intercept be included? <br> further arguments passed to or from other methods. |

## Details

A typical predictor has the form response ~ terms where response is the (numeric) response vector and terms is a series of terms which specifies a linear predictor for response. For binomial models the response can also be specified as a factor (when the first level denotes failure and all others success) or as a two-column matrix with the columns giving the numbers of successes and failures. A terms specification of the form first + second indicates all the terms in first together with all the terms in second with duplicates removed.

A specification of the form first:second indicates the the set of terms obtained by taking the interactions of all terms in first with all terms in second. The specification first*second indicates the cross of first and second. This is the same as first + second + first:second.
glm.fit and glm.fit.null are the workhorse functions: the former calls the latter for a null model (with no intercept).
If more than one of etastart, start and mustart is specified, the first in the list will be used.

## Value

glm returns an object of class glm which inherits from the class lm. See later in this section. The function summary (i.e., summary.glm) can be used to obtain or print a summary of the results and the function anova (i.e., anova.glm) to produce an analysis of variance table.
The generic accessor functions coefficients, effects, fitted.values and residuals can be used to extract various useful features of the value returned by glm.
weights extracts a vector of weights, one for each case in the fit (after subsetting and na.action).
An object of class "glm" is a list containing at least the following components:
coefficients a named vector of coefficients

| residuals | the working residuals, that is the residuals in the final iteration of the |
| :--- | :--- |
| IWLS fit. |  |

In addition, non-null fits will have components $q r, R$ and effects relating to the final weighted linear fit.

Objects of class "glm" are normally of class c ("glm", "lm"), that is inherit from class "lm", and well-designed methods for class "lm" will be applied to the weighted linear model at the final iteration of IWLS. However, care is needed, as extractor functions for class "glm" such as residuals and weights do not just pick out the component of the fit with the same name.
If a binomial glm model is specified by giving a two-column response, the weights returned by prior.weights are the total numbers of cases (factored by the supplied case weights) and the component $y$ of the result is the proportion of successes.

## References

McCullagh P. and Nelder, J. A. (1989) Generalized Linear Models. London: Chapman and Hall.
Dobson, A. J. (1990) An Introduction to Generalized Linear Models. London: Chapman and Hall.
Venables, W. N. and Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. New York: Springer.

## See Also

anova.glm, summary.glm, etc. for glm methods, and the generic functions anova, summary, effects, fitted.values, and residuals. Further, lm for non-generalized linear models. esoph, infert and predict.glm have examples of fitting binomial glms.

## Examples

```
## Dobson (1990) Page 93: Randomized Controlled Trial :
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
print(d.AD <- data.frame(treatment, outcome, counts))
glm.D93 <- glm(counts ~ outcome + treatment, family=poisson())
anova(glm.D93)
summary(glm.D93)
## an example with offsets from Venables & Ripley (1999, pp.217-8)
## Need the anorexia data from a recent version of the package MASS:
library(MASS)
data(anorexia)
anorex.1 <- glm(Postwt ~ Prewt + Treat + offset(Prewt),
    family = gaussian, data = anorexia)
summary(anorex.1)
# A Gamma example, from McCullagh & Nelder (1989, pp. 300-2)
clotting <- data.frame(
    u = c(5,10, 15, 20, 30,40,60, 80, 100),
    lot1 = c(118,58,42,35,27,25,21,19,18),
    lot2 = c(69,35,26,21,18,16,13,12,12))
summary(glm(lot1 ~ log(u), data=clotting, family=Gamma))
summary(glm(lot2 ~ log(u), data=clotting, family=Gamma))
```

glm.control Auxiliary for Controlling GLM Fitting

## Description

Auxiliary function as user interface for glm fitting. Typically only used when calling glm or glm.fit.

## Usage

```
glm.control(epsilon=1e-04, maxit=10, trace=FALSE)
```


## Arguments

epsilon positive convergence tolerance epsilon; the iterations converge when $\mid$ dev - devold $\mid /(|\operatorname{dev}|+0.1)<$ epsilon.
maxit integer giving the maximal number of IWLS iterations.
trace logical indicating if output should be produced for each iteration.

## Details

If epsilon is small, it is also used as the tolerance for the least squares solution.
When trace is true, calls to cat produce the output for each IWLS iteration. Hence, options (digits $=*$ ) can be used to increase the precision, see the example.

## Value

A list with the arguments as components.

## See Also

 glm.fit, the fitting procedure used by glm.
## Examples

```
### A variation on example(glm) :
## Annette Dobson's example ...
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
oo <- options(digits = 12) # to see more when tracing :
glm.D93X <- glm(counts ~ outcome + treatment, family=poisson(),
    trace = TRUE, epsilon = 1e-14)
options(oo)
coef(glm.D93X) # the last two are closer to 0 than in ?glm's glm.D93
```

```
glm.summaries Accessing Generalized Linear Model Fits
```


## Description

These functions are all methods for class glm or summary.glm objects.

## Usage

```
coefficients(x) ; coef(x)
family(object, ...)
fitted.values(x) ; fitted(x)
residuals(object, type = c("deviance", "pearson", "working",
                                    "response", "partial"), ...)
```


## Arguments

```
object, x an object of class glm, typically the result of a call to glm.
test a character string, matching one of "Chisq", "F" or "Cp". See
stat.anova.
type the type of residuals which should be returned. The alternatives
    are: "deviance" (default), "pearson", "working", "response", and
    "partial".
... further arguments passed to or from other methods.
```


## See Also

glm for computing glm.obj, anova.glm; the corresponding generic functions, summary.glm, coefficients, deviance, df.residual, effects, fitted.values, residuals.

$$
\text { glm.summary } \quad \text { Summarizing Generalized Linear Model Fits }
$$

## Description

These functions are all methods for class glm or summary.glm objects.

## Usage

```
summary(object, dispersion = NULL, correlation = FALSE, ...)
print(x, digits = max(3, getOption("digits") - 3),
na.print = "", symbolic.cor = p > 4,
signif.stars = getOption("show.signif.stars"), ...)
```


## Arguments

object an object of class "glm", usually, a result of a call to glm.
$\mathrm{x} \quad$ an object of class "summary.glm", usually, a result of a call to summary.glm.
dispersion the dispersion parameter for the fitting family. By default it is obtained from object.
correlation logical; if TRUE, the correlation matrix of the estimated parameters is returned and printed.
digits the number of significant digits to use when printing.
na.print Unused.
symbolic.cor logical. If TRUE, print the correlations in a symbolic form (see symnum rather than as numbers.
signif.stars logical. If TRUE, "significance stars" are printed for each coefficient.
... further arguments passed to or from other methods.

## Details

print.summary.glm tries to be smart about formatting the coefficients, standard errors, etc. and additionally gives "significance stars" if signif.stars is TRUE.

See Also
glm, summary.

## Examples

```
## --- Continuing the Example from '(?glm'):
```

summary (glm.D93)

Gnome GNOME Desktop Graphics Device

## Description

gnome starts a GNOME compatible device driver. 'GNOME' is an acronym for GNU Network Object Model Environment.

## Usage

```
gnome(display="", width=7, height=7, pointsize=12)
```

GNOME(display="", width=7, height=7, pointsize=12)

## Arguments

display the display on which the graphics window will appear. The default is to use the value in the user's environment variable DISPLAY.
width the width of the plotting window in inches.
height the height of the plotting window in inches.
pointsize the default pointsize to be used.

## Note

This is still in development state.
The GNOME device is only available when explicitly desired at configure/compile time, see the toplevel 'INSTALL' file.

## Author(s)

Lyndon Drake 〈lyndon@stat.auckland.ac.nz〉

## References

http://www.gnome.org and http://www.gtk.org for the GTK+ (GIMP Tool Kit) libraries.

## See Also

x11, Devices.

## Examples

```
gnome(width=9)
```

gray
Gray Level Specification

## Description

Create a vector of colors from a vector of gray levels.

## Usage

gray(level)
grey(level)

## Arguments

level a vector of desired gray levels between 0 and 1 ; zero indicates "black" and one indicates "white".

## Details

The values returned by gray can be used with a col= specification in graphics functions or in par.
grey is an alias for gray.

## Value

A vector of "colors" of the same length as level.

## See Also

rainbow, hsv, rgb.

## Examples

```
gray(0:8 / 8)
```

```
grep
Pattern Matching and Replacement
```


## Description

grep searches for matches to pattern (its first argument) within the character vector x (second argument). regexpr does too, but returns more detail in a different format.
sub and gsub perform replacement of matches determined by regular expression matching.

## Usage

grep(pattern, $x$, ignore.case=FALSE, extended=TRUE, perl=FALSE, value=FALSE)
sub(pattern, replacement, x,
ignore.case=FALSE, extended=TRUE, perl=FALSE)
gsub(pattern, replacement, x,
ignore.case=FALSE, extended=TRUE, perl=FALSE)
regexpr(pattern, text, extended=TRUE, perl=FALSE)

## Arguments

| pattern | character string containing a regular expression to be matched in the given <br> character vector. |
| :--- | :--- |
| x, text | a character vector where matches are sought. |
| ignore.case | if FALSE, the pattern matching is case sensitive and if TRUE, case is ignored <br> during matching. |
| extended | if TRUE, extended regular expression matching is used, and if FALSE basic <br> regular expressions are used. |
| perl | logical. Should perl-compatible regexps be used if available? Has priority <br> over extended. |
| value | if FALSE, a vector containing the (integer) indices of the matches deter- <br> mined by grep is returned, and if TRUE, a vector containing the matching <br> elements themselves is returned. |
| replacement | a replacement for matched pattern in sub and gsub. |

## Details

The two *sub functions differ only in that sub replaces only the first occurrence of a pattern whereas gsub replaces all occurrences.

The regular expressions used are those specified by POSIX 1003.2, either extended or basic, depending on the value of the extended argument, unless perl $=$ TRUE when they are those of PCRE, ftp://ftp.csx.cam.ac.uk/pub/software/programming/pcre/.

## Value

For grep a vector giving either the indices of the elements of x that yielded a match or, if value is TRUE, the matched elements.

For sub and gsub a character vector of the same length as the original.
For regexpr an integer vector of the same length as text giving the starting position of the first match, or -1 if there is none, with attribute "match. length" giving the length of the matched text (or -1 for no match).

## Note

perl=TRUE will only be available if $R$ was compiled against PCRE: this is detected at configure time.

## See Also

agrep for approximate matching. tolower, toupper and chartr for character translations. charmatch, pmatch, match. apropos uses regexps and has nice examples.

## Examples

```
grep("[a-z]", letters)
txt <- c("arm","foot","lefroo", "bafoobar")
if(any(i <- grep("foo",txt)))
    cat("'foo' appears at least once in\n\t",txt,"\n")
i # 2 and 4
```

```
txt[i]
## Double all 'a' or 'b's; "\" must be escaped, i.e. 'doubled'
gsub("([ab])", "\\1_\\1_", "abc and ABC")
txt <- c("The", "licenses", "for", "most", "software", "are",
    "designed", "to", "take", "away", "your", "freedom",
    "to", "share", "and", "change", "it.",
        "", "By", "contrast,", "the", "GNU", "General", "Public", "License",
        "is", "intended", "to", "guarantee", "your", "freedom", "to",
        "share", "and", "change", "free", "software", "--",
        "to", "make", "sure", "the", "software", "is",
        "free", "for", "all", "its", "users")
( i <- grep("[gu]", txt) ) # indices
stopifnot( txt[i] == grep("[gu]", txt, value = TRUE) )
(ot <- sub("[b-e]",".", txt))
txt[ot != gsub("[b-e]",".", txt)]#- gsub does "global" substitution
txt[gsub("g","#", txt) !=
        gsub("g","#", txt, ignore.case = TRUE)] # the "G" words
regexpr("en", txt)
## trim trailing white space
str = 'Now is the time
sub(' +$', '', str) ## spaces only
sub('[[:space:]]+$', ',, str) ## white space, POSIX-style
if(capabilities("PCRE"))
    sub('\\s+$', ',, str, perl = TRUE) ## perl-style white space
```

grid Add Grid to a Plot

## Description

grid adds an $n x$ by ny rectangular grid to an existing plot, using lines of type lty and color col.

If more fine tuning is required, use abline ( $\mathrm{h}=., \mathrm{v}=$. ) directly.

## Usage

```
grid(nx = NULL, ny = nx, col="lightgray", lty="dotted", lwd = NULL,
    equilogs = TRUE)
```


## Arguments

$n x, n y$
col
lty
lwd non-negative numeric giving line width of the grid lines; defaults to par("lwd").
equilogs logical, only used when $\log$ coordinates and alignment with the axis tick marks are active. Settingequilogs $=$ FALSE in that case gives non equidistant tick aligned grid lines.

## See Also

## Examples

```
plot(1:3)
grid(NA,5, lwd = 2) # grid only in y-direction
data(iris)
## maybe change the desired number of tick marks: par(lab=c(mx,my,7))
op <- par(mfcol = 1:2)
with(iris,
    {
    plot(Sepal.Length, Sepal.Width, col = as.integer(Species),
        xlim = c(4, 8), ylim = c(2, 4.5), panel.first = grid(),
        main = "with(iris, plot(...., panel.first = grid(), ..) )")
    plot(Sepal.Length, Sepal.Width, col = as.integer(Species),
        panel.first = grid(3, lty=1,lwd=2),
        main = "... panel.first = grid(3, lty=1,lwd=2), ..")
    }
    )
par(op)
```

gtk GTK+ Graphics Device

## Description

A variant of the X11 graphics device for use with the GNOME GUI.

## Usage

```
gtk(display="", width=7, height=7, pointsize=12)
GTK(display="", width=7, height=7, pointsize=12)
```


## Arguments

display the display on which the graphics window will appear. The default is to use the value in the user's environment variable DISPLAY.
width the width of the plotting window in inches.
height the height of the plotting window in inches.
pointsize the default pointsize to be used.

## Details

The GTK device is only available when running under UNIX with --gui=gnome. Under that circumstance it is the default (auto-launched) device, but x11 can also be used.
The device has a toolbar in the GNOME style.

## Author(s)

Lyndon Drake 〈lyndon@stat.auckland.ac.nz〉

## References

http://www.gtk.org for the GTK+ (GIMP Tool Kit) libraries.

## See Also

x11, Devices.

HairEyeColor Hair and Eye Color of Statistics Students

## Description

Distribution of hair and eye color and sex in 592 statistics students.

## Usage

data(HairEyeColor)

## Format

A 3-dimensional array resulting from cross-tabulating 592 observations on 3 variables. The variables and their levels are as follows:

| No | Name | Levels |
| ---: | :--- | :--- |
| 1 | Hair | Black, Brown, Red, Blond |
| 2 | Eye | Brown, Blue, Hazel, Green |
| 3 | Sex | Male, Female |

## Details

This data set is useful for illustrating various techniques for the analysis of contingency tables, such as the standard chi-squared test or, more generally, log-linear modelling, and graphical methods such as mosaic plots, sieve diagrams or association plots.

## References

Snee, R. D. (1974), Graphical display of two-way contingency tables. The American Statistician, 28, 9-12.
Friendly, M. (1992), Graphical methods for categorical data. SAS User Group International Conference Proceedings, 17, 190-200. http://www.math.yorku.ca/SCS/sugi/ sugi17-paper.html
Friendly, M. (1992), Mosaic displays for loglinear models. Proceedings of the Statistical Graphics Section, American Statistical Association, pp. 61-68. http://www.math.yorku. ca/SCS/Papers/asa92.html

## See Also

chisq.test, loglin, mosaicplot

## Examples

```
data(HairEyeColor)
## Full mosaic
mosaicplot(HairEyeColor)
## Aggregate over sex:
x <- apply(HairEyeColor, c(1, 2), sum)
x
mosaicplot(x, main = "Relation between hair and eye color")
```

help Documentation

## Description

These functions provide access to documentation. Documentation on a topic with name name (typically, an R object or a data set) can be printed with either help(name) or ?name.

## Usage

```
help(topic, offline = FALSE, package = .packages(),
    lib.loc = NULL, verbose = getOption("verbose"),
    try.all.packages = getOption("help.try.all.packages"),
    htmlhelp = getOption("htmlhelp"),
    pager = getOption("pager"))
?topic
type?topic
```


## Arguments

| topic | a name or character string on which documentation is sought (but not a <br> variable containing a character string!). |
| :--- | :--- |
| offline | a logical indicating whether documentation should be displayed on-line to <br> the screen (the default) or hardcopy of it should be produced. <br> a name or character vector giving the packages to look into for documen- <br> tation. By default, all packages in the search path are used. <br> a character vector of directory names of R libraries, or NULL. The default |
| lib.loc | value of NULL corresponds to all libraries currently known. If the default <br> is used, the loaded packages are searched before the libraries. |
| verbose | logical; if TRUE, the file name is reported. |
| try.all.packages |  |
| logical; see Notes. |  |

## Details

In the case of unary and binary operators and control-flow special forms, the name may need to be quoted.
If offline is TRUE, hardcopy of the documentation is produced by running the LaTeX version of the help page through latex (note that LaTeX 2e is needed) and dvips. Depending on your dvips configuration, hardcopy will be sent to the printer or saved in a file. If the programs are in non-standard locations and hence were not found at compile time, you can either set the options latexcmd and dvipscmd, or the environment variables R_LATEXCMD and R_DVIPSCMD appropriately. The appearance of the output can be customized through a file 'Rhelp.cfg' somewhere in your LaTeX search path.

## Note

Unless lib.loc is specified explicitly, the loaded packages are searched before those in the specified libraries. This ensures that if a library is loaded from a library not in the known library trees, then the help from the loaded library is used. If lib.loc is specified explicitly, the loaded packages are not searched.
If this search fails and argument try.all.packages is TRUE and neither packages nor lib.loc is specified, then all the packages in the known library trees are searched for help on topic and a list of (any) packages where help may be found is printed (but no help is shown). N.B. searching all packages can be slow.

The help files can be many small files. On some file systems it is desirable to save space, and the text files in the 'help' directory of an installed package can be zipped up as a zip archive 'Rhelp.zip'. Ensure that file 'AnIndex' remains un-zipped. Similarly, all the files in the 'latex' directory can be zipped to 'Rhelp.zip'.

## See Also

help.search() for finding help pages on a "vague" topic. help.start() which opens the HTML version of the R help pages; library () for listing available packages and the userlevel objects they contain; data() for listing available data sets; methods().
See prompt () to get a prototype for writing help pages of private packages.

## Examples

```
help()
help(help) # the same
help(lapply)
?lapply # the same
help("for") # or ?"for", but the quotes are needed
?"+"
help(package = stepfun) # get help even when package is not loaded
data() # list all available data sets
?women # information about data set "women"
topi <- "women"
help(topi) ##--> Error: No documentation for 'topi'
try(help("bs", try.all.packages=FALSE)) # reports not found (an error)
```

```
help("bs", try.all.packages=TRUE) # reports can be found in package 'splines'
```

help.search Search the Help System

## Description

Allows for searching the help system for documentation matching a given character string in the (file) name, alias, title, or keyword entries (or any combination thereof), using either fuzzy matching or regular expression matching. Names and titles of the matched help entries are nicely displayed.

## Usage

```
help.search(pattern, fields = c("alias", "title"),
    apropos, keyword, whatis, ignore.case = TRUE,
    package = NULL, lib.loc = NULL,
    help.db = getOption("help.db"),
    verbose = getOption("verbose"),
    rebuild = FALSE, agrep = NULL)
```


## Arguments

pattern a character string to be matched in the specified fields. If this is given, the arguments apropos, keyword, and whatis are ignored.
fields a character vector specifying the fields of the help data bases to be searched. The entries must be abbreviations of "name", "alias", "title", and "keyword", corresponding to the help page's (file) name, the topics it provides documentation for, its title, and the keywords it can be classified to.
apropos a character string to be matched in the help page topics and title.
keyword a character string to be matched in the help page keywords.
whatis a character string to be matched in the help page topics.
ignore.case a logical. If TRUE, case is ignored during matching; if FALSE, pattern matching is case sensitive.
package a character vector with the names of packages to search through, or NULL in which case all available packages in the specified library trees lib.loc are searched.
lib.loc a character vector describing the location of $R$ library trees to search through, or NULL. The default value of NULL corresponds to all libraries currently known.
help.db a character string giving the file path to a previously built and saved help data base, or NULL.
verbose logical; if TRUE, the search process is traced.
rebuild a logical indicating whether the help data base should be rebuilt.
agrep if NULL (the default) and the character string to be matched consists of alphanumeric characters, whitespace or a dash only, approximate (fuzzy) matching via agrep is used; otherwise, it is taken to contain a regular expression to be matched via grep. If FALSE, approximate matching is not used. Otherwise, one can give a numeric or a list specifying the maximal distance for the approximate match, see argument max.distance in the documentation for agrep.

## Details

Upon installation of a package, the Perl script 'Rd2contents.pl' creates a 'CONTENTS' data base which contains the information on name, aliases, title and keywords (as well as the URL of the HTML version of the help file) in Debian Control Format. This is the data base searched by help.search().

The arguments apropos and whatis play a role similar to the Unix commands with the same names.
If possible, the help data base is saved to the file 'help. db ' in the '. $R$ ' subdirectory of the user's home directory or the current working directory.

Note that currently, the aliases in the matching help files are not displayed.

## Value

The results are returned in an object of class "hsearch", which has a print method for nicely displaying the results of the query. This mechanism is experimental, and may change in future versions of $R$.

## See Also

help; help.start for starting the hypertext (currently HTML) version of R's online documentation, which offers a similar search mechanism. apropos uses regexps and has nice examples.

## Examples

```
help.search("linear models") # In case you forgot how to fit linear
    # models
help.search("non-existent topic")
help.search("print") # All help pages with topics or title
    # matching 'print'
help.search(apropos = "print") # The same
help.search(keyword = "hplot") # All help pages documenting high-level
    # plots.
## Help pages with documented topics starting with 'try'.
help.search("\\btry", fields = "alias")
## Do not use '~' or '$' when matching aliases or keywords.
```

help.start Hypertext Documentation

## Description

Start the hypertext (currently HTML) version of R's online documentation.

## Usage

```
    help.start(gui = "irrelevant", browser = getOption("browser"),
    remote \(=\) NULL)
```


## Arguments

gui just for compatibility with S-PLUS.
browser the name of the program to be used as hypertext browser. It should be in the PATH, or a full path specified.
remote A character giving a valid URL for the '\$R_HOME' directory on a remote location.

## Details

All the packages in the known library trees are linked to directory '. $R$ ' in the per-session temporary directory. The links are re-made each time help.start is run, which should be done after packages are installed, updated or removed.

If the browser given by the browser argument is different from the default browser as specified by options("browser"), the default is changed to the given browser so that it gets used for all future help requests.

## See Also

help() for on- and off-line help in ASCII/Editor or PostScript format. browseURL for how the help file is displayed.

## Examples

```
help.start()
```

Hershey Hershey Vector Fonts in $R$

## Description

If the vfont argument to one of the text-drawing functions (text, mtext, title, axis, and contour) is a character vector of length 2 , hershey vector fonts are used to render the text.
These fonts have two advantages:

1. vector fonts describe each character in terms of a set of points; $R$ renders the character by joining up the points with straight lines. This intimate knowledge of the outline of each character means that R can arbitrarily transform the characters, which can mean that the vector fonts look better for rotated and 3d text.
2. this implementation was adapted from the GNU libplot library which provides support for non-ASCII and non-English fonts. This means that it is possible, for example, to produce weird plotting symbols and Japanese characters.

Drawback:
You cannot use mathematical expressions (plotmath) with Hershey fonts.

## Usage

Hershey

## Details

The Hershey characters are organised into a set of fonts, which are specified by a typeface (e.g., serif or sans serif) and a fontindex or "style" (e.g., plain or italic). The first element of vfont specifies the typeface and the second element specifies the fontindex. The first table produced by example(Hershey) shows the character a produced by each of the different fonts.
The available typeface and fontindex values are available as list components of the variable Hershey. The allowed pairs for (typeface, fontindex) are:

| serif | plain |
| :--- | :--- |
| serif | italic |
| serif | bold |
| serif | bold italic |
| serif | cyrillic |
| serif | oblique cyrillic |
| serif | EUC |
| sans serif | plain |
| sans serif | italic |
| sans serif | bold |
| sans serif | bold italic |
| script | plain |
| script | italic |
| script | bold |
| gothic english | plain |
| gothic german | plain |
| gothic italian | plain |
| serif symbol | plain |
| serif symbol | italic |
| serif symbol | bold |
| serif symbol | bold italic |

$$
\begin{array}{ll}
\text { sans serif symbol } & \text { plain } \\
\text { sans serif symbol } & \text { italic }
\end{array}
$$

and the indices of these are available as Hershey\$allowed.
Escape sequences: The string to be drawn can include escape sequences, which all begin with a $\backslash$. When R encounters a $\backslash$, rather than drawing the $\backslash$, it treats the subsequent character(s) as a coded description of what to draw.
One useful escape sequence (in the current context) is of the form: $\backslash 123$. The three digits following the $\backslash$ specify an octal code for a character. For example, the octal code for $p$ is 160 so the strings " $p$ " and " $\backslash 160$ " are equivalent. This is useful for producing characters when there is not an appropriate key on your keyboard.
The other useful escape sequences all begin with $\backslash \backslash$. These are described below.
Symbols: an entire string of Greek symbols can be produced by selecting the Serif Symbol or Sans Serif Symbol typeface. To allow Greek symbols to be embedded in a string which uses a non-symbol typeface, there are a set of symbol escape sequences of the form <br>ab. For example, the escape sequence <br>*a produces a Greek alpha. The second table in example(Hershey) shows all of the symbol escape sequences and the symbols that they produce.
ISO Latin-1: further escape sequences of the form $\backslash \backslash a b$ are provided for producing ISO Latin- 1 characters (for example, if you only have a US keyboard). Another option is to use the appropriate octal code. The (non-ASCII) ISO Latin-1 characters are in the range $241 \ldots 377$. For example, $\backslash 366$ produces the character o with an umlaut. The third table in example (Hershey) shows all of the ISO Latin-1 escape sequences.
Special Characters: a set of characters are provided which do not fall into any standard font. These can only be accessed by escape sequence. For example, $\backslash \backslash L I$ produces the zodiac sign for Libra, and $\backslash \backslash J U$ produces the astronomical sign for Jupiter. The fourth table in example (Hershey) shows all of the special character escape sequences.
Cyrillic Characters: cyrillic characters are implemented according to the K018-R encoding. On a US keyboard, these can be produced using the Serif typeface and Cyrillic (or Oblique Cyrillic) fontindex and specifying an octal code in the range 300 to 337 for lower case characters or 340 to 377 for upper case characters. The fifth table in example (Hershey) shows the octal codes for the available cyrillic characters.

Japanese Characters: 83 Hiragana, 86 Katakana, and 603 Kanji characters are implemented according to the EUC (Extended Unix Code) encoding. Each character is idenitified by a unique hexadecimal code. The Hiragana characters are in the range $0 \times 2421$ to $0 \times 2473$, Katakana are in the range $0 \times 2521$ to $0 \times 2576$, and Kanji are (scattered about) in the range $0 \times 3021$ to $0 \times 6 \mathrm{~d} 55$.
When using the Serif typeface and EUC fontindex, these characters can be produced by a pair of octal codes. Given the hexadecimal code (e.g., 0x2421), take the first two digits and add $0 \times 80$ and do the same to the second two digits (e.g., $0 \times 21$ and $0 \times 24$ become 0xa4 and 0xa1), then convert both to octal (e.g., 0xa4 and 0xa1 become 244 and 241). For example, the first Hiragana character is produced by $\backslash 244 \backslash 241$.
It is also possible to use the hexadecimal code directly. This works for all non-EUC fonts by specifying an escape sequence of the form $\backslash \backslash \# J 1234$. For example, the first Hiragana character is produced by <br>\#J2421.
The Kanji characters may be specified in a third way, using the so-called "Nelson Index", by specifying an escape sequence of the form $\backslash \backslash \# N 1234$. For example, the Kanji for "one" is produced by $\backslash \backslash \#$ N0001.

Raw Hershey Glyphs: all of the characters in the Hershey fonts are stored in a large array. Some characters are not accessible in any of the Hershey fonts. These characters can only be accessed via an escape sequence of the form $\backslash \backslash \# H 1234$. For example, the fleur-de-lys is produced by $\backslash \backslash \# H 0746$. The sixth and seventh tables of example(Hershey) shows all of the available raw glyphs.

## References

http://www.gnu.org/software/plotutils/plotutils.html

## See Also

```
text, contour, Japanese
```


## Examples

```
str(Hershey)
######
# create tables of vector font functionality
######
make.table <- function(nr, nc) {
    savepar <- par(mar=rep(0, 4), pty="s")
    plot(c(0, nc*2 + 1), c(0, -(nr + 1)),
        type="n", xlab="", ylab="", axes=FALSE)
    savepar
}
get.r <- function(i, nr) i %% nr + 1
get.c <- function(i, nr) i %/% nr + 1
draw.title <- function(title, i = 0, nr, nc) {
    r <- get.r(i, nr)
    c <- get.c(i, nr)
    text((nc*2 + 1)/2, 0, title, font=2)
}
draw.sample.cell <- function(typeface, fontindex, string, i, nr) {
    r <- get.r(i, nr)
    c <- get.c(i, nr)
    text(2*(c - 1) + 1, -r, paste(typeface, fontindex))
    text(2*c, -r, string, vfont=c(typeface, fontindex), cex=1.5)
    rect(2*(c - 1) + .5, -(r - . 5), 2*c + . 5, -(r + . 5), border="grey")
}
draw.vf.cell <- function(typeface, fontindex, string, i, nr, raw.string=NULL) {
    r <- get.r(i, nr)
    c <- get.c(i, nr)
    if (is.null(raw.string))
        raw.string <- paste("\\", string, sep="")
    text(2*(c - 1) + 1, -r, raw.string, col="grey")
    text(2*c, -r, string, vfont=c(typeface, fontindex))
    rect(2*(c - 1) + . 5, -(r - . 5), ( 2*c + . 5), -(r + . 5), border="grey")
}
nr <- 23
nc <- 1
```

```
oldpar <- make.table(nr, nc)
i <- 0
draw.title("Sample 'a' for each available font", i, nr, nc)
draw.sample.cell("serif", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "italic", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "bold", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "bold italic", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "cyrillic", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "oblique cyrillic", "a", i, nr); i <- i + 1
draw.sample.cell("serif", "EUC", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif", "italic", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif", "bold", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif", "bold italic", "a", i, nr); i <- i + 1
draw.sample.cell("script", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("script", "italic", "a", i, nr); i <- i + 1
draw.sample.cell("script", "bold", "a", i, nr); i <- i + 1
draw.sample.cell("gothic english", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("gothic german", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("gothic italian", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("serif symbol", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("serif symbol", "italic", "a", i, nr); i <- i + 1
draw.sample.cell("serif symbol", "bold", "a", i, nr); i <- i + 1
draw.sample.cell("serif symbol", "bold italic", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif symbol", "plain", "a", i, nr); i <- i + 1
draw.sample.cell("sans serif symbol", "italic", "a", i, nr); i <- i + 1
nr <- 25
nc <- 6
tf <- "serif"
fi <- "plain"
make.table(nr, nc)
i <- 0
draw.title("Symbol (incl. Greek) Escape Sequences", i, nr, nc)
## Greek alphabet in order
draw.vf.cell(tf, fi, "\\*A", i, nr); i<-i+1; { "Alpha"}
draw.vf.cell(tf, fi, "\\*B", i, nr); i<-i+1; { "Beta"}
draw.vf.cell(tf, fi, "\\*G", i, nr); i<-i+1; { "Gamma"}
draw.vf.cell(tf, fi, "\\*D", i, nr); i<-i+1; { "Delta"}
draw.vf.cell(tf, fi, "\\*E", i, nr); i<-i+1; { "Epsilon"}
draw.vf.cell(tf, fi, "\\*Z", i, nr); i<-i+1; { "Zeta"}
draw.vf.cell(tf, fi, "\\*Y", i, nr); i<-i+1; { "Eta"}
draw.vf.cell(tf, fi, "\\*H", i, nr); i<-i+1; { "Theta"}
draw.vf.cell(tf, fi, "\\*I", i, nr); i<-i+1; { "Iota"}
draw.vf.cell(tf, fi, "\\*K", i, nr); i<-i+1; { "Kappa"}
draw.vf.cell(tf, fi, "\\*L", i, nr); i<-i+1; { "Lambda"}
draw.vf.cell(tf, fi, "\\*M", i, nr); i<-i+1; { "Mu"}
draw.vf.cell(tf, fi, "\\*N", i, nr); i<-i+1; { "Nu"}
draw.vf.cell(tf, fi, "\\*C", i, nr); i<-i+1; { "Xi"}
draw.vf.cell(tf, fi, "\\*O", i, nr); i<-i+1; { "Omicron"}
draw.vf.cell(tf, fi, "\\*P", i, nr); i<-i+1; { "Pi"}
draw.vf.cell(tf, fi, "\\*R", i, nr); i<-i+1; { "Rho"}
draw.vf.cell(tf, fi, "\\*S", i, nr); i<-i+1; { "Sigma"}
draw.vf.cell(tf, fi, "\\*T", i, nr); i<-i+1; { "Tau"}
draw.vf.cell(tf, fi, "\\*U", i, nr); i<-i+1; { "Upsilon"}
draw.vf.cell(tf, fi, "\\+U", i, nr); i<-i+1; { "Upsilon1"}
draw.vf.cell(tf, fi, "\\*F", i, nr); i<-i+1; { "Phi"}
```

```
draw.vf.cell(tf, fi, "\\*X", i, nr); i<-i+1; { "Chi"}
draw.vf.cell(tf, fi, "\\*Q", i, nr); i<-i+1; { "Psi"}
draw.vf.cell(tf, fi, "\\*W", i, nr); i<-i+1; { "Omega"}
#
draw.vf.cell(tf, fi, "\\*a", i, nr); i<-i+1; { "alpha"}
draw.vf.cell(tf, fi, "\\*b", i, nr); i<-i+1; { "beta"}
draw.vf.cell(tf, fi, "\\*g", i, nr); i<-i+1; { "gamma"}
draw.vf.cell(tf, fi, "\\*d", i, nr); i<-i+1; { "delta"}
draw.vf.cell(tf, fi, "\\*e", i, nr); i<-i+1; { "epsilon"}
draw.vf.cell(tf, fi, "\\*z", i, nr); i<-i+1; { "zeta"}
draw.vf.cell(tf, fi, "\\*y", i, nr); i<-i+1; { "eta"}
draw.vf.cell(tf, fi, "\\*h", i, nr); i<-i+1; { "theta"}
draw.vf.cell(tf, fi, "\\+h", i, nr); i<-i+1; { "theta1"}
draw.vf.cell(tf, fi, "\\*i", i, nr); i<-i+1; { "iota"}
draw.vf.cell(tf, fi, "\\*k", i, nr); i<-i+1; { "kappa"}
draw.vf.cell(tf, fi, "\\**", i, nr); i<-i+1; { "lambda"}
draw.vf.cell(tf, fi, "\\*m", i, nr); i<-i+1; { "mu"}
draw.vf.cell(tf, fi, "\\*n", i, nr); i<-i+1; { "nu"}
draw.vf.cell(tf, fi, "\\*c", i, nr); i<-i+1; { "xi"}
draw.vf.cell(tf, fi, "\\*o", i, nr); i<-i+1; { "omicron"}
draw.vf.cell(tf, fi, "\\*p", i, nr); i<-i+1; { "pi"}
draw.vf.cell(tf, fi, "\\*r", i, nr); i<-i+1; { "rho"}
draw.vf.cell(tf, fi, "\\*s", i, nr); i<-i+1; { "sigma"}
draw.vf.cell(tf, fi, "\\ts", i, nr); i<-i+1; { "sigma1"}
draw.vf.cell(tf, fi, "\\*t", i, nr); i<-i+1; { "tau"}
draw.vf.cell(tf, fi, "\\*u", i, nr); i<-i+1; { "upsilon"}
draw.vf.cell(tf, fi, "\\*f", i, nr); i<-i+1; { "phi"}
draw.vf.cell(tf, fi, "\\+f", i, nr); i<-i+1; { "phi1"}
draw.vf.cell(tf, fi, "\\*x", i, nr); i<-i+1; { "chi"}
draw.vf.cell(tf, fi, "\\*q", i, nr); i<-i+1; { "psi"}
draw.vf.cell(tf, fi, "\\*w", i, nr); i<-i+1; { "omega"}
draw.vf.cell(tf, fi, "\\+p", i, nr); i<-i+1; { "omega1"}
#
draw.vf.cell(tf, fi, "\\fa", i, nr); i<-i+1; { "universal"}
draw.vf.cell(tf, fi, "\\te", i, nr); i<-i+1; { "existential"}
draw.vf.cell(tf, fi, "\\st", i, nr); i<-i+1; { "suchthat"}
draw.vf.cell(tf, fi, "\\**", i, nr); i<-i+1; { "asteriskmath"}
draw.vf.cell(tf, fi, "\\=~", i, nr); i<-i+1; { "congruent"}
draw.vf.cell(tf, fi, "\\tf", i, nr); i<-i+1; { "therefore"}
draw.vf.cell(tf, fi, "\\pp", i, nr); i<-i+1; { "perpendicular"}
draw.vf.cell(tf, fi, "\\ul", i, nr); i<-i+1; { "underline"}
draw.vf.cell(tf, fi, "\\rx", i, nr); i<-i+1; { "radicalex"}
draw.vf.cell(tf, fi, "\\ap", i, nr); i<-i+1; { "similar"}
draw.vf.cell(tf, fi, "\\fm", i, nr); i<-i+1; { "minute"}
draw.vf.cell(tf, fi, "\\<=", i, nr); i<-i+1; { "lessequal"}
draw.vf.cell(tf, fi, "\\f/", i, nr); i<-i+1; { "fraction"}
draw.vf.cell(tf, fi, "\\if", i, nr); i<-i+1; { "infinity"}
draw.vf.cell(tf, fi, "\\Fn", i, nr); i<-i+1; { "florin"}
draw.vf.cell(tf, fi, "\\CL", i, nr); i<-i+1; { "club"}
draw.vf.cell(tf, fi, "\\DI", i, nr); i<-i+1; { "diamond"}
draw.vf.cell(tf, fi, "\\HE", i, nr); i<-i+1; { "heart"}
draw.vf.cell(tf, fi, "\\SP", i, nr); i<-i+1; { "spade"}
draw.vf.cell(tf, fi, "\\<>", i, nr); i<-i+1; { "arrowboth"}
draw.vf.cell(tf, fi, "\\<-", i, nr); i<-i+1; { "arrowleft"}
draw.vf.cell(tf, fi, "\\ua", i, nr); i<-i+1; { "arrowup"}
draw.vf.cell(tf, fi, "\\->", i, nr); i<-i+1; { "arrowright"}
```

```
draw.vf.cell(tf, fi, "\\da", i, nr); i<-i+1; { "arrowdown"}
draw.vf.cell(tf, fi, "\\de", i, nr); i<-i+1; { "degree"}
draw.vf.cell(tf, fi, "\\+-", i, nr); i<-i+1; { "plusminus"}
draw.vf.cell(tf, fi, "\\sd", i, nr); i<-i+1; { "second"}
draw.vf.cell(tf, fi, "\\>=", i, nr); i<-i+1; { "greaterequal"}
draw.vf.cell(tf, fi, "\\mu", i, nr); i<-i+1; { "multiply"}
draw.vf.cell(tf, fi, "\\pt", i, nr); i<-i+1; { "proportional"}
draw.vf.cell(tf, fi, "\\pd", i, nr); i<-i+1; { "partialdiff"}
draw.vf.cell(tf, fi, "\\bu", i, nr); i<-i+1; { "bullet"}
draw.vf.cell(tf, fi, "\\di", i, nr); i<-i+1; { "divide"}
draw.vf.cell(tf, fi, "\\!=", i, nr); i<-i+1; { "notequal"}
draw.vf.cell(tf, fi, "\\==", i, nr); i<-i+1; { "equivalence"}
draw.vf.cell(tf, fi, "\\~~", i, nr); i<-i+1; { "approxequal"}
draw.vf.cell(tf, fi, "\\..", i, nr); i<-i+1; { "ellipsis"}
draw.vf.cell(tf, fi, "\\an", i, nr); i<-i+1; { "arrowhorizex"}
draw.vf.cell(tf, fi, "\\CR", i, nr); i<-i+1; { "carriagereturn"}
draw.vf.cell(tf, fi, "\\Ah", i, nr); i<-i+1; { "aleph"}
draw.vf.cell(tf, fi, "\\Im", i, nr); i<-i+1; { "Ifraktur"}
draw.vf.cell(tf, fi, "\\Re", i, nr); i<-i+1; { "Rfraktur"}
draw.vf.cell(tf, fi, "\\wp", i, nr); i<-i+1; { "weierstrass"}
draw.vf.cell(tf, fi, "\\c*", i, nr); i<-i+1; { "circlemultiply"}
draw.vf.cell(tf, fi, "\\c+", i, nr); i<-i+1; { "circleplus"}
draw.vf.cell(tf, fi, "\\es", i, nr); i<-i+1; { "emptyset"}
draw.vf.cell(tf, fi, "\\ca", i, nr); i<-i+1; { "cap"}
draw.vf.cell(tf, fi, "\\cu", i, nr); i<-i+1; { "cup"}
draw.vf.cell(tf, fi, "\\SS", i, nr); i<-i+1; { "superset"}
draw.vf.cell(tf, fi, "\\ip", i, nr); i<-i+1; { "reflexsuperset"}
draw.vf.cell(tf, fi, "\\n<", i, nr); i<-i+1; { "notsubset"}
draw.vf.cell(tf, fi, "\\SB", i, nr); i<-i+1; { "subset"}
draw.vf.cell(tf, fi, "\\ib", i, nr); i<-i+1; { "reflexsubset"}
draw.vf.cell(tf, fi, "\\mo", i, nr); i<-i+1; { "element"}
draw.vf.cell(tf, fi, "\\nm", i, nr); i<-i+1; { "notelement"}
draw.vf.cell(tf, fi, "\\/_", i, nr); i<-i+1; { "angle"}
draw.vf.cell(tf, fi, "\\gr", i, nr); i<-i+1; { "nabla"}
draw.vf.cell(tf, fi, "\\rg", i, nr); i<-i+1; { "registerserif"}
draw.vf.cell(tf, fi, "\\co", i, nr); i<-i+1; { "copyrightserif"}
draw.vf.cell(tf, fi, "\\tm", i, nr); i<-i+1; { "trademarkserif"}
draw.vf.cell(tf, fi, "\\PR", i, nr); i<-i+1; { "product"}
draw.vf.cell(tf, fi, "\\sr", i, nr); i<-i+1; { "radical"}
draw.vf.cell(tf, fi, "\\md", i, nr); i<-i+1; { "dotmath"}
draw.vf.cell(tf, fi, "\\no", i, nr); i<-i+1; { "logicalnot"}
draw.vf.cell(tf, fi, "\\\AN", i, nr); i<-i+1; { "logicaland"}
draw.vf.cell(tf, fi, "\\OR", i, nr); i<-i+1; { "logicalor"}
draw.vf.cell(tf, fi, "\\hA", i, nr); i<-i+1; { "arrowdblboth"}
draw.vf.cell(tf, fi, "\\lA", i, nr); i<-i+1; { "arrowdblleft"}
draw.vf.cell(tf, fi, "\\uA", i, nr); i<-i+1; { "arrowdblup"}
draw.vf.cell(tf, fi, "\\rA", i, nr); i<-i+1; { "arrowdblright"}
draw.vf.cell(tf, fi, "\\dA", i, nr); i<-i+1; { "arrowdbldown"}
draw.vf.cell(tf, fi, "\\lz", i, nr); i<-i+1; { "lozenge"}
draw.vf.cell(tf, fi, "\\la", i, nr); i<-i+1; { "angleleft"}
draw.vf.cell(tf, fi, "\\RG", i, nr); i<-i+1; { "registersans"}
draw.vf.cell(tf, fi, "\\CO", i, nr); i<-i+1; { "copyrightsans"}
draw.vf.cell(tf, fi, "\\TM", i, nr); i<-i+1; { "trademarksans"}
draw.vf.cell(tf, fi, "\\SU", i, nr); i<-i+1; { "summation"}
draw.vf.cell(tf, fi, "\\lc", i, nr); i<-i+1; { "bracketlefttp"}
draw.vf.cell(tf, fi, "\\lf", i, nr); i<-i+1; { "bracketleftbt"}
draw.vf.cell(tf, fi, "\\ra", i, nr); i<-i+1; { "angleright"}
```

draw.vf.cell(tf, fi, "<br>is", i, nr); i<-i+1; \{ "integral"\} draw.vf.cell(tf, fi, "<br>rc", i, nr); i<-i+1; \{ "bracketrighttp"\} draw.vf.cell(tf, fi, "<br>rf", i, nr); i<-i+1; \{ "bracketrightbt"\} draw.vf.cell(tf, fi, "<br>~=", i, nr); i<-i+1; \{ "congruent"\} draw.vf.cell(tf, fi, "<br>pr", i, nr); i<-i+1; \{ "minute"\} draw.vf.cell(tf, fi, "<br>in", i, nr); i<-i+1; \{ "infinity"\} draw.vf.cell(tf, fi, "<br>n=", i, nr); i<-i+1; \{ "notequal"\} draw.vf.cell(tf, fi, "<br>dl", i, nr); i<-i+1; \{ "nabla"\}

```
nr <- 25
nc <- 4
make.table(nr, nc)
i <- 0
draw.title("ISO Latin-1 Escape Sequences", i, nr, nc)
draw.vf.cell(tf, fi, "\\r!", i, nr); i<-i+1; { "exclamdown"}
draw.vf.cell(tf, fi, "\\ct", i, nr); i<-i+1; { "cent"}
draw.vf.cell(tf, fi, "\\Po", i, nr); i<-i+1; { "sterling"}
draw.vf.cell(tf, fi, "\\Ye", i, nr); i<-i+1; { "yen"}
draw.vf.cell(tf, fi, "\\bb", i, nr); i<-i+1; { "brokenbar"}
draw.vf.cell(tf, fi, "\\sc", i, nr); i<-i+1; { "section"}
draw.vf.cell(tf, fi, "\\ad", i, nr); i<-i+1; { "dieresis"}
draw.vf.cell(tf, fi, "\\co", i, nr); i<-i+1; { "copyright"}
draw.vf.cell(tf, fi, "\\Of", i, nr); i<-i+1; { "ordfeminine"}
draw.vf.cell(tf, fi, "\\no", i, nr); i<-i+1; { "logicalnot"}
draw.vf.cell(tf, fi, "\\hy", i, nr); i<-i+1; { "hyphen"}
draw.vf.cell(tf, fi, "\\rg", i, nr); i<-i+1; { "registered"}
draw.vf.cell(tf, fi, "\\a-", i, nr); i<-i+1; { "macron"}
draw.vf.cell(tf, fi, "\\de", i, nr); i<-i+1; { "degree"}
draw.vf.cell(tf, fi, "\\+-", i, nr); i<-i+1; { "plusminus"}
draw.vf.cell(tf, fi, "\\S2", i, nr); i<-i+1; { "twosuperior"}
draw.vf.cell(tf, fi, "\\S3", i, nr); i<-i+1; { "threesuperior"}
draw.vf.cell(tf, fi, "\\aa", i, nr); i<-i+1; { "acute"}
draw.vf.cell(tf, fi, "\\*m", i, nr); i<-i+1; { "mu"}
draw.vf.cell(tf, fi, "\\md", i, nr); i<-i+1; { "periodcentered"}
draw.vf.cell(tf, fi, "\\S1", i, nr); i<-i+1; { "onesuperior"}
draw.vf.cell(tf, fi, "\\Om", i, nr); i<-i+1; { "ordmasculine"}
draw.vf.cell(tf, fi, "\\14", i, nr); i<-i+1; { "onequarter"}
draw.vf.cell(tf, fi, "\\12", i, nr); i<-i+1; { "onehalf"}
draw.vf.cell(tf, fi, "\\34", i, nr); i<-i+1; { "threequarters"}
draw.vf.cell(tf, fi, "\\\r", i, nr); i<-i+1; { "questiondown"}
draw.vf.cell(tf, fi, "\\'A", i, nr); i<-i+1; { "Agrave"}
draw.vf.cell(tf, fi, "\\'A", i, nr); i<-i+1; { "Aacute"}
draw.vf.cell(tf, fi, "\\^A", i, nr); i<-i+1; { "Acircumflex"}
draw.vf.cell(tf, fi, "\\~~A", i, nr); i<-i+1; { "Atilde"}
draw.vf.cell(tf, fi, "\\:A", i, nr); i<-i+1; { "Adieresis"}
draw.vf.cell(tf, fi, "\\oA", i, nr); i<-i+1; { "Aring"}
draw.vf.cell(tf, fi, "\\AE", i, nr); i<-i+1; { "AE"}
draw.vf.cell(tf, fi, "\\,C", i, nr); i<-i+1; { "Ccedilla"}
draw.vf.cell(tf, fi, "\\'E", i, nr); i<-i+1; { "Egrave"}
draw.vf.cell(tf, fi, "\\'E", i, nr); i<-i+1; { "Eacute"}
draw.vf.cell(tf, fi, "\\^E", i, nr); i<-i+1; { "Ecircumflex"}
draw.vf.cell(tf, fi, "\\:E", i, nr); i<-i+1; { "Edieresis"}
draw.vf.cell(tf, fi, "\\`I", i, nr); i<-i+1; { "Igrave"}
draw.vf.cell(tf, fi, "\\'I", i, nr); i<-i+1; { "Iacute"}
draw.vf.cell(tf, fi, "\\^I", i, nr); i<-i+1; { "Icircumflex"}
draw.vf.cell(tf, fi, "\\:I", i, nr); i<-i+1; { "Idieresis"}
draw.vf.cell(tf, fi, "\\ ~N", i, nr); i<-i+1; { "Ntilde"}
```

```
draw.vf.cell(tf, fi, "\\`0", i, nr); i<-i+1; { "Ograve"}
draw.vf.cell(tf, fi, "\\'O", i, nr); i<-i+1; { "Oacute"}
draw.vf.cell(tf, fi, "\\^O", i, nr); i<-i+1; { "Ocircumflex"}
draw.vf.cell(tf, fi, "\\~O", i, nr); i<-i+1; { "Otilde"}
draw.vf.cell(tf, fi, "\\:O", i, nr); i<-i+1; { "Odieresis"}
draw.vf.cell(tf, fi, "\\mu", i, nr); i<-i+1; { "multiply"}
draw.vf.cell(tf, fi, "\\/O", i, nr); i<-i+1; { "Oslash"}
draw.vf.cell(tf, fi, "\\`U", i, nr); i<-i+1; { "Ugrave"}
draw.vf.cell(tf, fi, "\\'U", i, nr); i<-i+1; { "Uacute"}
draw.vf.cell(tf, fi, "\\^U", i, nr); i<-i+1; { "Ucircumflex"}
draw.vf.cell(tf, fi, "\\:U", i, nr); i<-i+1; { "Udieresis"}
draw.vf.cell(tf, fi, "\\'Y", i, nr); i<-i+1; { "Yacute"}
draw.vf.cell(tf, fi, "\\ss", i, nr); i<-i+1; { "germandbls"} # WRONG!
draw.vf.cell(tf, fi, "\\'a", i, nr); i<-i+1; { "agrave"}
draw.vf.cell(tf, fi, "\\'a", i, nr); i<-i+1; { "aacute"}
draw.vf.cell(tf, fi, "\\^a", i, nr); i<-i+1; { "acircumflex"}
draw.vf.cell(tf, fi, "\\~a", i, nr); i<-i+1; { "atilde"}
draw.vf.cell(tf, fi, "\\:a", i, nr); i<-i+1; { "adieresis"}
draw.vf.cell(tf, fi, "\\oa", i, nr); i<-i+1; { "aring"}
draw.vf.cell(tf, fi, "\\ae", i, nr); i<-i+1; { "ae"}
draw.vf.cell(tf, fi, "\\,c", i, nr); i<-i+1; { "ccedilla"}
draw.vf.cell(tf, fi, "\\'e", i, nr); i<-i+1; { "egrave"}
draw.vf.cell(tf, fi, "\\'e", i, nr); i<-i+1; { "eacute"}
draw.vf.cell(tf, fi, "\\^e", i, nr); i<-i+1; { "ecircumflex"}
draw.vf.cell(tf, fi, "\\:e", i, nr); i<-i+1; { "edieresis"}
draw.vf.cell(tf, fi, "\\`i", i, nr); i<-i+1; { "igrave"}
draw.vf.cell(tf, fi, "\\'i", i, nr); i<-i+1; { "iacute"}
draw.vf.cell(tf, fi, "\\^i", i, nr); i<-i+1; { "icircumflex"}
draw.vf.cell(tf, fi, "\\:i", i, nr); i<-i+1; { "idieresis"}
draw.vf.cell(tf, fi, "\\~n", i, nr); i<-i+1; { "ntilde"}
draw.vf.cell(tf, fi, "\\`o", i, nr); i<-i+1; { "ograve"}
draw.vf.cell(tf, fi, "\\'o", i, nr); i<-i+1; { "oacute"}
draw.vf.cell(tf, fi, "\\^o", i, nr); i<-i+1; { "ocircumflex"}
draw.vf.cell(tf, fi, "\\~o", i, nr); i<-i+1; { "otilde"}
draw.vf.cell(tf, fi, "\\:o", i, nr); i<-i+1; { "odieresis"}
draw.vf.cell(tf, fi, "\\di", i, nr); i<-i+1; { "divide"}
draw.vf.cell(tf, fi, "\\/o", i, nr); i<-i+1; { "oslash"}
draw.vf.cell(tf, fi, "\\'u", i, nr); i<-i+1; { "ugrave"}
draw.vf.cell(tf, fi, "\\'u", i, nr); i<-i+1; { "uacute"}
draw.vf.cell(tf, fi, "\\^u", i, nr); i<-i+1; { "ucircumflex"}
draw.vf.cell(tf, fi, "\\:u", i, nr); i<-i+1; { "udieresis"}
draw.vf.cell(tf, fi, "\\'y", i, nr); i<-i+1; { "yacute"}
draw.vf.cell(tf, fi, "\\:y", i, nr); i<-i+1; { "ydieresis"}
nr <- 25
nc <- 2
make.table(nr, nc)
i <- 0
draw.title("Special Escape Sequences", i, nr, nc)
draw.vf.cell(tf, fi, "\\AR", i, nr); i<-i+1; { "aries"}
draw.vf.cell(tf, fi, "\\TA", i, nr); i<-i+1; { "taurus"}
draw.vf.cell(tf, fi, "\\GE", i, nr); i<-i+1; { "gemini"}
draw.vf.cell(tf, fi, "\\CA", i, nr); i<-i+1; { "cancer"}
draw.vf.cell(tf, fi, "\\LE", i, nr); i<-i+1; { "leo"}
draw.vf.cell(tf, fi, "\\VI", i, nr); i<-i+1; { "virgo"}
draw.vf.cell(tf, fi, "\\LI", i, nr); i<-i+1; { "libra"}
draw.vf.cell(tf, fi, "\\SC", i, nr); i<-i+1; { "scorpio"}
```

```
draw.vf.cell(tf, fi, "\\SG", i, nr); i<-i+1; { "sagittarius"}
draw.vf.cell(tf, fi, "\\CP", i, nr); i<-i+1; { "capricornus"}
draw.vf.cell(tf, fi, "\\AQ", i, nr); i<-i+1; { "aquarius"}
draw.vf.cell(tf, fi, "\\PI", i, nr); i<-i+1; { "pisces"}
draw.vf.cell(tf, fi, "\\\~", i, nr); i<-i+1; { "modifiedcongruent"}
draw.vf.cell(tf, fi, "\\hb", i, nr); i<-i+1; { "hbar"}
draw.vf.cell(tf, fi, "\\IB", i, nr); i<-i+1; { "interbang"}
draw.vf.cell(tf, fi, "\\Lb", i, nr); i<-i+1; { "lambdabar"}
draw.vf.cell(tf, fi, "\\UD", i, nr); i<-i+1; { "undefined"}
draw.vf.cell(tf, fi, "\\SO", i, nr); i<-i+1; { "sun"}
draw.vf.cell(tf, fi, "\\ME", i, nr); i<-i+1; { "mercury"}
draw.vf.cell(tf, fi, "\\VE", i, nr); i<-i+1; { "venus"}
draw.vf.cell(tf, fi, "\\EA", i, nr); i<-i+1; { "earth"}
draw.vf.cell(tf, fi, "\\MA", i, nr); i<-i+1; { "mars"}
draw.vf.cell(tf, fi, "\\JU", i, nr); i<-i+1; { "jupiter"}
draw.vf.cell(tf, fi, "\\SA", i, nr); i<-i+1; { "saturn"}
draw.vf.cell(tf, fi, "\\UR", i, nr); i<-i+1; { "uranus"}
draw.vf.cell(tf, fi, "\\NE", i, nr); i<-i+1; { "neptune"}
draw.vf.cell(tf, fi, "\\PL", i, nr); i<-i+1; { "pluto"}
draw.vf.cell(tf, fi, "\\LU", i, nr); i<-i+1; { "moon"}
draw.vf.cell(tf, fi, "\\CT", i, nr); i<-i+1; { "comet"}
draw.vf.cell(tf, fi, "\\ST", i, nr); i<-i+1; { "star"}
draw.vf.cell(tf, fi, "\\AS", i, nr); i<-i+1; { "ascendingnode"}
draw.vf.cell(tf, fi, "\\DE", i, nr); i<-i+1; { "descendingnode"}
draw.vf.cell(tf, fi, "\\s-", i, nr); i<-i+1; { "s1"}
draw.vf.cell(tf, fi, "\\dg", i, nr); i<-i+1; { "dagger"}
draw.vf.cell(tf, fi, "\\dd", i, nr); i<-i+1; { "daggerdbl"}
draw.vf.cell(tf, fi, "\\li", i, nr); i<-i+1; { "line integral"}
draw.vf.cell(tf, fi, "\\-+", i, nr); i<-i+1; { "minusplus"}
draw.vf.cell(tf, fi, "\\||", i, nr); i<-i+1; { "parallel"}
draw.vf.cell(tf, fi, "\\rn", i, nr); i<-i+1; { "overscore"}
draw.vf.cell(tf, fi, "\\ul", i, nr); i<-i+1; { "underscore"}
```

nr <- 25
nc <- 3
make.table(nr, nc)
code <- c $(300: 307,310: 317,320: 327,330: 337,340: 347,350: 357,360: 367,370: 377$,
$243,263)$
string <- c
"\300", "\301", "\302", "\303", "\304", "\305", "\306", "\307",
"\310", "\311", "\312", "\313", "\314", "\315",
"\316", "\317", "\320", "\321", "\322", "\323",
"\} 3 2 4 " , " \backslash 3 2 5 " , " \backslash 3 2 6 " , " \backslash 3 2 7 " , " \backslash 3 3 0 " , " \backslash 3 3 1 " ,
"\332", "\333", "\334", "\335", "\336", "\337",
"\340", "\341", "\342", "\343", "\344", "\345", "\346", "\347",
"\350", "\351", "\352", "\353", "\354", "\355",
"\356", "\357", "\360", "\361", "\362", "\363",
"\364","\365", "\366", "\367", "\370", "\371",
"\372", "\373", "\374", "\375", "\376", "\377", "\243", "\263")
draw.title("Cyrillic Octal Codes", i = 0, nr ,nc)
for (i in 1:66)
draw.vf.cell(tf, "cyrillic", string[i], i-1, nr ,
raw.string=paste("<br>", as.character(code[i]), sep=""))
nr <- 25
nc <- 3
make.table(nr, nc)

```
code <- c(252,254,256,262:269,275,278:281,284,745,746,750:768,796:802,
        804:807,809,814:828,830:834,840:844)
    draw.title("Raw Hershey Escape Sequences", i=0, nr, nc)
    for (i in 1:75)
        draw.vf.cell(tf, fi, paste("\\#H",formatC(code[i],wid=4,flag=0),sep=""),
                        i-1, nr)
make.table(nr, nc)
code <- c(845:847,850:856,860:874,899:909,2296:2299,2318:2332,2367:2382,
        4014,4109)
    draw.title("More Raw Hershey Escape Sequences", i=0, nr, nc)
    for (i in 1:73)
        draw.vf.cell(tf, fi, paste("\\#H",formatC(code[i],wid=4,flag=0),sep=""),
                        i-1, nr)
    par(oldpar)
```

hist Histograms

## Description

The generic function hist computes a histogram of the given data values. If plot=TRUE, the resulting object of class "histogram" is plotted by plot.histogram, before it is returned.

## Usage

```
hist(x, ...)
hist.default(x, breaks = "Sturges", freq = NULL, probability = !freq,
        include.lowest \(=\) TRUE, right \(=\) TRUE,
        density \(=\) NULL, angle \(=45\), col \(=\) NULL, border \(=\) NULL,
        main = paste("Histogram of" , xname),
        xlim = range(breaks), ylim = NULL,
        xlab = xname, ylab,
        axes = TRUE, plot = TRUE, labels = FALSE,
        nclass = NULL, ...)
```


## Arguments

| x | a vector of values for which the histogram is desired. |
| :--- | :--- |
| breaks | one of: |
| - a vector giving the breakpoints between histogram cells, |  |
| - a single number giving the number of cells for the histogram, |  |
| - a character string naming an algorithm to compute the number of |  |
| cells (see Details), |  |
| - a function to compute the number of cells. |  |

include.lowest
logical; if TRUE, an 'x[i]' equal to the 'breaks' value will be included in the first (or last, for right $=$ FALSE) bar.
right logical; if TRUE, the histograms cells are right-closed (left open) intervals. density the density of shading lines, in lines per inch. The default value of NULL means that no shading lines are drawn. Non-positive values of 'density' also inhibit the drawing of shading lines.
angle the slope of shading lines, given as an angle in degrees (counter-clockwise).
col a colour to be used to fill the bars. The default of NULL yields unfilled bars.
border the color of the border around the bars. The default is to use the standard foreground color.
main, xlab, ylab
these arguments to title have useful defaults here.
xlim, ylim the range of x and y values with sensible defaults.
axes logical. If TRUE (default), axes are draw if the plot is drawn.
plot logical. If TRUE (default), a histogram is plotted, otherwise a list of breaks and counts is returned.
labels logical or character. Additionally draw labels on top of bars, if not FALSE; see plot.histogram.
nclass numeric (integer). For S(-PLUS) compatibility only, nclass is equivalent to breaks for a scalar or character argument.
... further graphical parameters to title and axis.

## Details

If right = TRUE (default), the histogram cells are intervals of the form (a, b], i.e. they include their right-hand endpoint, but not their left one, with the exception of the first cell when include.lowest is TRUE.
For right = FALSE, the intervals are of the form [a, b), and include.lowest really has the meaning of "include highest".
The default for breaks is "Sturges": see nclass.Sturges. Other names for which algorithms are supplied are "Scott" and "FD" / "Friedman-Diaconis". Case is ignored and partial matching is used. Alternatively, a function can be supplied which will compute the intended number of breaks as a function of x .

## Value

an object of class "histogram" which is a list with components:
breaks the $n+1$ cell boundaries ( $=$ breaks if that was a vector).
counts $\quad n$ integers; for each cell, the number of x[] inside.
density values $\hat{f}\left(x_{i}\right)$, as estimated density values. If all (diff(breaks) == 1), they are the relative frequencies counts $/ \mathrm{n}$ and in general satisfy $\sum_{i} \hat{f}\left(x_{i}\right)\left(b_{i+1}-b_{i}\right)=1$, where $b_{i}=$ breaks [i].
intensities same as density. Deprecated, but retained for compatibility.
mids the $n$ cell midpoints.
xname a character string with the actual x argument name.
equidist logical, indicating if the distances between breaks are all the same.

## Note

The resulting value does not depend on the values of the arguments freq (or probability) or plot. This is intentionally different from S.

## References

Venables, W. N. and Ripley. B. D. (1999) Modern Applied Statistics with S-PLUS. Springer.

## See Also

nclass.Sturges, stem, density.

## Examples

```
data(islands)
op <- par(mfrow=c(2, 2))
hist(islands)
str(hist(islands, col="gray", labels = TRUE))
hist(sqrt(islands), br = 12, col="lightblue", border="pink")
##-- For non-equidistant breaks, counts should NOT be graphed unscaled:
r <- hist(sqrt(islands), br = c(4*0:5, 10*3:5, 70, 100, 140), col='blue1')
text(r$mids, r$density, r$counts, adj=c(.5, -.5), col='blue3')
sapply(r[2:3], sum)
sum(r$density * diff(r$breaks)) # == 1
lines(r, lty = 3, border = "purple") # -> lines.histogram(*)
par(op)
str(hist(islands, plot= FALSE)) #-> 5 breaks
str(hist(islands, br=12, plot= FALSE)) #-> 10 (~ = 12) breaks
str(hist(islands, br=c(12,20,36,80,200,1000,17000), plot = FALSE))
    hist(islands, br=c(12,20,36,80,200,1000,17000), freq = TRUE,
                main = "WRONG histogram") # and warning
```

hist.POSIXt Histogram of a Date-Time Object

## Description

Method for hist applied to date-time objects.

## Usage

```
hist(x, breaks, ..., plot = TRUE, freq = FALSE,
    start.on.monday = TRUE, format)
```


## Arguments

$\mathrm{x} \quad$ an object inheriting from class "POSIXt".
breaks a vector of cut points or number giving the number of intervals which x is to be cut into or an interval specification, one of "secs", "mins", "hours", "days", "weeks", "months" or "years".
... graphical parameters.
plot logical. If TRUE (default), a histogram is plotted, otherwise a list of breaks and counts is returned.
freq logical; if TRUE, the histogram graphic is a representation of frequencies, i.e, the counts component of the result; if FALSE, relative frequencies ("probabilities") are plotted.
start.on.monday
logical. If breaks = "weeks", should the week start on Mondays or Sundays?
format for the x -axis labels. See strptime.

## Value

An object of class "histogram": see hist.

## See Also

seq.POSIXt, axis.POSIXct, hist

## Examples

```
hist(.leap.seconds, "years", freq = TRUE)
hist(.leap.seconds,
    seq(ISOdate(1970, 1, 10), ISOdate(2002, 1, 1), "5 years"))
## 100 random dates in a 10-week period
random.dates <- ISOdate(2001, 1, 1) + 70*86400*runif(100)
hist(random.dates, "weeks", format = "%d %b")
```

hsv HSV Color Specification

## Description

Create a vector of colors from vectors specifying hue, saturation and value.

## Usage

$\mathrm{hsv}(\mathrm{h}=1, \mathrm{~s}=1, \mathrm{v}=1$, gamma=1)

## Arguments

$h, s, v \quad$ numeric vectors of values in the range [0,1] for "hue", "saturation" and "value" to be combined to form a vector of colors. Values in shorter arguments are recycled.
gamma a "gamma correction"

## Value

This function creates a vector of "colors" corresponding to the given values in HSV space. The values returned by hsv can be used with a col= specification in graphics functions or in par.

## See Also

rainbow, rgb, gray.

## Examples

```
hsv(.5,.5,.5)
## Look at gamma effect:
n <- 20; y <- -sin(3*pi*((1:n)-1/2)/n)
op <- par(mfrow=c(3,2),mar=rep (1.5,4))
for(gamma in c(.4, .6, .8, 1, 1.2, 1.5))
    plot(y, axes = FALSE, frame.plot = TRUE,
            xlab = "", ylab = "", pch = 21, cex = 30,
            bg = rainbow(n, start=.85, end=.1, gamma = gamma),
            main = paste("Red tones; gamma=",format(gamma)))
par(op)
```

Hyperbolic Hyperbolic Functions

## Description

These functions give the obvious hyperbolic functions. They respectively compute the hyperbolic cosine, sine, tangent, arc-cosine, arc-sine, arc-tangent.

## Usage

$\cosh (\mathrm{x})$
$\sinh (x)$
$\tanh (x)$
$\operatorname{acosh}(x)$
$\operatorname{asinh}(x)$
$\operatorname{atanh}(\mathrm{x})$

## Arguments

x a numeric vector

See Also

```
cos, sin, tan, acos, asin, atan.
```


## Examples

```
Ceps <- .Machine$double.eps # ''Computer epsilon''
x <- seq(-3, 3, len=200)
stopifnot(
    abs(\operatorname{cosh}(x)-(exp(x) + exp(-x))/2) < 20*Ceps,
    abs(\operatorname{sinh}(x) - (exp(x) - exp(-x))/2) < 20*Ceps,
    Mod(cosh(x) - cos(1i*x)) < 20*Ceps,
    Mod(sinh(x) - sin(1i*x)/1i) < 20*Ceps,
    abs(tanh (x)*\operatorname{cosh}(x) - sinh (x)) < 20*Ceps
)
```

```
## Inverse:
all(abs(asinh(sinh(x)) - x) < 10*Ceps)
x[abs(acosh(cosh(x)) - abs(x)) > 100*Ceps] #- imprecise for small x
all(abs(atanh(tanh (x)) - x) < 100*Ceps)
all(abs(asinh(x) - log(x + sqrt(x^2 + 1))) < 10*Ceps)
cx <- cosh(x)
all(abs(acosh(cx) - log(cx + sqrt(cx^2 - 1))) < 1000*Ceps)
```

Hypergeometric The Hypergeometric Distribution

## Description

Density, distribution function, quantile function and random generation for the hypergeometric distribution.

## Usage

```
dhyper(x, m, n, k, log = FALSE)
phyper(q, m, n, k, lower.tail = TRUE, log.p = FALSE)
qhyper(p, m, n, k, lower.tail = TRUE, log.p = FALSE)
rhyper(nn, m, n, k)
```


## Arguments

$x, q \quad$ vector of quantiles representing the number of white balls drawn without replacement from an urn which contains both black and white balls.
m
the number of white balls in the urn.
n the number of black balls in the urn.
$\mathrm{k} \quad$ the number of balls drawn from the urn.
$\mathrm{p} \quad$ probability, it must be between 0 and 1.
$\mathrm{nn} \quad$ number of observations. If length $(\mathrm{nn})>1$, the length is taken to be the number required.
$\log , \log \cdot \mathrm{p} \quad \operatorname{logical}$; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The hypergeometric distribution is used for sampling without replacement. The density of this distribution with parameters $\mathrm{m}, \mathrm{n}$ and k ( named $N p, N-N p$, and $n$, respectively in the reference below) is given by

$$
p(x)=\binom{m}{x}\binom{n}{k-x} /\binom{m+n}{k}
$$

for $x=0, \ldots, k$.

## Value

dhyper gives the density, phyper gives the distribution function, qhyper gives the quantile function, and rhyper generates random deviates.

## References

Johnson, N. L., Kotz, S., and Kemp, A. W. (1992) Univariate Discrete Distributions, Second Edition. New York: Wiley.

## Examples

```
m <- 10; n <- 7; k <- 8
x <- 0:(k+1)
rbind(phyper(x, m, n, k), dhyper(x,m, n, k))
all(phyper(x, m, n, k) == cumsum(dhyper(x, m, n, k)))# FALSE
## but error is very small:
signif(phyper(x, m, n, k) - cumsum(dhyper(x, m, n, k)), dig=3)
```

identical Test Objects for Exact Equality

## Description

The safe and reliable way to test two objects for being exactly equal. It returns TRUE in this case, FALSE in every other case.

## Usage

identical(x, y)

## Arguments

$\mathrm{x}, \mathrm{y} \quad$ any R objects.

## Details

A call to identical is the way to test exact equality in if and while statements, as well as in logical expressions that use \&\& or ||. In all these applications you need to be assured of getting a single logical value.
Users often use the comparison operators, such as $==$ or $!=$, in these situations. It looks natural, but it is not what these operators are designed to do in $R$. They return an object like the arguments. If you expected $x$ and $y$ to be of length 1 , but it happened that one of them wasn't, you will not get a single FALSE. Similarly, if one of the arguments is NA, the result is also NA. In either case, the expression if ( $\mathrm{x}=\mathrm{y}$ y) .... won't work as expected.
The function all.equal is also sometimes used to test equality this way, but it was intended for something different. First, it tries to allow for "reasonable" differences in numeric results. Second, it returns a descriptive character vector instead of FALSE when the objects do not match. Therefore, it is not the right function to use for reliable testing either. (If you do want to allow for numeric fuzziness in comparing objects, you can combine all.equal and identical, as shown in the examples below.)

The computations in identical are also reliable and usually fast. There should never be an error. The only known way to kill identical is by having an invalid pointer at
the C level, generating a memory fault. It will usually find inequality quickly. Checking equality for two large, complicated objects can take longer if the objects are identical or nearly so, but represent completely independent copies. For most applications, however, the computational cost should be negligible.

As from R 1.6.0, identical sees NaN as different from as.double(NA), but all NaNs are equal (and all NA of the same type are equal).

## Value

A single logical value, TRUE or FALSE, never NA and never anything other than a single value.

## Author(s)

John Chambers

## See Also

all. equal for descriptions of how two objects differ; Comparison for operators that generate elementwise comparisons.

## Examples

```
identical(1, NULL) ## FALSE -- don't try this with ==
identical(1, 1.) ## TRUE in R (both are stored as doubles)
identical(1, as.integer(1)) ## FALSE, stored as different types
## how to test for object equality allowing for numeric fuzz
identical(all.equal(x, y), TRUE)
## If all.equal thinks the objects are different, it returns a
## character string, and this expression evaluates to FALSE
# even for unusual R objects :
identical(.GlobalEnv, environment())
```

identify Identify Points in a Scatter Plot

## Description

identify reads the position of the graphics pointer when the (first) mouse button is pressed. It then searches the coordinates given in x and y for the point closest to the pointer. If this point is close to the pointer, its index will be returned as part of the value of the call.

## Usage

```
identify(x, ...)
identify.default(x, y = NULL, labels = seq(along = x), pos = FALSE,
n = length(x), plot = TRUE, offset = 0.5, ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | coordinates of points in a scatter plot. Alternatively, any object which <br> defines coordinates (a plotting structure, time series etc.) can be given as <br> x and y left undefined. |
| :--- | :--- |
| labels | an optional vector, the same length as x and y, giving labels for the points. |
| pos | if pos is TRUE, a component is added to the return value which indicates <br> where text was plotted relative to each identified point ( $1=$ below, $2=$ left,, <br> $3=$ above and $4=$ right). |
| n | the maximum number of points to be identified. |
| plot | if plot is TRUE, the labels are printed at the points and if FALSE they are <br> omitted. |
| offset | the distance (in character widths) which separates the label from identified <br> points. |
| $\ldots$ | further arguments to par (.). |

## Details

If in addition, plot is TRUE, the point is labelled with the corresponding element of text.
The labels are placed either below, to the left, above or to the right of the identified point, depending on where the cursor was.

The identification process is terminated by pressing any mouse button other than the first.
If the window is resized or hidden and then exposed before the identification process has terminated, any labels drawn by identify will disappear. These will reappear once the identification process has terminated and the window is resized or hidden and exposed again. This is because the labels drawn by identify are not recorded in the device's display list until the identification process has terminated.

## Value

If pos is FALSE, an integer vector containing the indexes of the identified points.
If pos is TRUE, a list containing a component ind, indicating which points were identified and a component pos, indicating where the labels were placed relative to the identified points.

## See Also

## ifelse Conditional Element Selection

## Description

ifelse returns a value with the same shape as test which is filled with elements selected from either yes or no depending on whether the element of test is TRUE or FALSE. If yes or no are too short, their elements are recycled.

## Usage

```
ifelse(test, yes, no)
```


## Arguments

| test | a logical vector |
| :--- | :--- |
| yes | return values for true elements of test. |
| no | return values for false elements of test. |

## See Also

if.

## Examples

```
x <- c(6:-4)
sqrt(x)#- gives warning
sqrt(ifelse(x >= 0, x, NA))# no warning
## Note: the following also gives the warning !
ifelse(x >= 0, sqrt(x), NA)
```

image Display a Color Image

## Description

Creates a grid of colored or gray-scale rectangles with colors corresponding to the values in z. This can be used to display three-dimensional or spatial data aka "images". This is a generic function.
The functions heat.colors, terrain.colors and topo.colors create heat-spectrum (red to white) and topographical color schemes suitable for displaying ordered data, with n giving the number of colors desired.

## Usage

```
image(x, ...)
image.default(x, y, z, zlim, xlim, ylim, col = heat.colors(12),
        add = FALSE, xaxs = "i", yaxs = "i", xlab, ylab,
        breaks, oldstyle = FALSE, ...)
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ locations of grid lines at which the values in z are measured. These must be in (strictly) ascending order. By default, equally spaced values from 0 to 1 are used. If x is a list, its components $\mathrm{x} \$ \mathrm{x}$ and $\mathrm{x} \$ \mathrm{y}$ are used for x and y , respectively. If the list has component z this is used for z .
z
a matrix containing the values to be plotted (NAs are allowed). Note that $x$ can be used instead of $z$ for convenience.
zlim the minimum and maximum $z$ values for which colors should be plotted. Each of the given colors will be used to color an equispaced interval of this range. The midpoints of the intervals cover the range, so that values just outside the range will be plotted.
xlim, ylim
col a list of colors such as that generated by rainbow, heat.colors, topo. colors, terrain. colors or similar functions.
add logical; if TRUE, add to current plot (and disregard the following arguments). This is rarely useful because image "paints" over existing graphics.
xaxs, yaxs style of $x$ and $y$ axis. The default " $i$ " is appropriate for images. See par.
$\mathrm{xlab}, \mathrm{ylab}$ each a character string giving the labels for the x and y axis. Default to the 'call names' of x or y , or to " " if these where unspecified.
breaks a set of breakpoints for the colours: must give one more breakpoint than colour.
oldstyle logical. If true the midpoints of the colour intervals are equally spaced, and zlim[1] and zlim[2] were taken to be midpoints. (This was the default prior to R 1.1.0.) The current default is to have colour intervals of equal lengths between the limits.
... graphical parameters for plot may also be passed as arguments to this function.

## Details

The length of $x$ should be equal to the nrow $(x)+1$ or nrow $(x)$. In the first case $x$ specifies the boundaries between the cells: in the second case $x$ specifies the midpoints of the cells. Similar reasoning applies to y. It probably only makes sense to specify the midpoints of an equally-spaced grid. If you specify just one row or column and a length-one $x$ or $y$, the whole user area in the corresponding direction is filled.
If breaks is specified then zlim is unused and the algorithm used follows cut, so intervals are closed on the right and open on the left except for the lowest interval.

## Note

Based on a function by Thomas Lumley 〈tlumley@u.washington.edu〉.

## See Also

```
contour, heat.colors, topo.colors, terrain.colors, rainbow, hsv, par.
```


## Examples

```
x <- y <- seq(-4*pi, 4*pi, len=27)
r <- sqrt(outer(x^2, y^2, "+"))
image(z = z <- cos(r^2)*exp(-r/6), col=gray((0:32)/32))
image(z, axes = FALSE, main = "Math can be beautiful ...",
    xlab = expression(cos(r^2) * e^{-r/6}))
contour(z, add = TRUE, drawlabels = FALSE)
data(volcano)
```

```
x <- 10*(1:nrow(volcano))
y <- 10*(1:ncol(volcano))
image(x, y, volcano, col = terrain.colors(100), axes = FALSE)
contour(x, y, volcano, levels = seq(90, 200, by=5), add = TRUE, col = "peru")
axis(1, at = seq(100, 800, by = 100))
axis(2, at = seq(100, 600, by = 100))
box()
title(main = "Maunga Whau Volcano", font.main = 4)
```

index.search Search Indices for Help Files

## Description

Used to search the indices for help files, possibly under aliases.

## Usage

```
index.search(topic, path, file="AnIndex", type = "help")
```


## Arguments

topic The keyword to be searched for in the indices.
path The path(s) to the packages to be searched.
file The index file to be searched. Normally '"AnIndex"'.
type $\quad$ The type of file required.

## Details

For each package in path, examine the file file in directory 'type', and look up the matching file stem for topic topic, if any.

## Value

A character vector of matching files, as if they are in directory type of the corresponding package. In the special cases of type = "html", "R-ex" and "latex" the file extensions ".html", ".R" and ".tex" are added.

## See Also

```
help, example
```


## infert Infertility after Spontaneous and Induced Abortion

## Description

This is a matched case-control study dating from before the availability of conditional logistic regression.

## Usage

data(infert)

## Format

| 1. | Education | $0=0-5$ years |
| :---: | :---: | :---: |
|  |  | $1=6-11$ years |
|  |  | $2=12+$ years |
| 2. | age | age in years of case |
| 3. | parity | count |
| 4. | number of prior | $0=0$ |
|  | induced abortions | $1=1$ |
|  |  | $2=2$ or more |
| 5. | case status | 1 = case |
|  |  | $0=$ control |
| 6. | number of prior | $0=0$ |
|  | spontaneous abortions | $1=1$ |
|  |  | $2=2$ or more |
| 7. | matched set number | 1-83 |
| 8. | stratum number | 1-63 |

## Note

One case with two prior spontaneous abortions and two prior induced abortions is omitted.

## Source

Trichopoulos et al. (1976) Br. J. of Obst. and Gynaec. 83, 645-650.

## Examples

```
data(infert)
model1 <- glm(case ~ spontaneous+induced, data=infert,family=binomial())
summary(model1)
## adjusted for other potential confounders:
summary(model2 <- glm(case ~ age+parity+education+spontaneous+induced,
    data=infert,family=binomial()))
## Really should be analysed by conditional logistic regression
## which is in the survival package
if(require(survival)){
    oT <- T; oF <- F; T <- TRUE; F <- FALSE # survival fails otherwise
```

```
    model3 <- clogit(case~ spontaneous+induced+strata(stratum), data=infert)
    summary(model3)
    detach()# survival (conflicts)
    T <- oT; F <- oF
}
```

influence.measures Regression Diagnostics

## Description

This suite of functions can be used to compute some of the regression diagnostics discussed in Belsley, Kuh and Welsch (1980), and in Cook and Weisberg (1982).

## Usage

influence.measures(lm.obj)
rstandard (lm.obj,
infl = lm.influence(lm.obj),
res = weighted.residuals(lm.obj),
sd $=$ sqrt(deviance(lm.obj)/df.residual(lm.obj)))
rstudent (lm.obj, infl = ..., res = ...)
dffits (lm.obj, infl = ..., res = ...)
dfbetas (lm.obj, infl = ...)
covratio (lm.obj, infl = ..., res = ...)
cooks.distance(lm.obj, infl = ..., res = ..., sd = ...)
hat ( x , intercept $=$ TRUE)

## Arguments

| $l m . o b j$ | the resulting object returned by lm. |
| :--- | :--- |
| infl | influence structure as returned by lm.influence. |
| res | (possibly weighted) residuals, with proper default. |
| sd | standard deviation to use, see default. |
| x | the $X$ or design matrix. |
| intercept | should an intercept column be pre-prended to $\mathrm{x} ?$ |

## Details

The primary function is influence.measures which produces a class "infl" object tabular display showing the DFBETAS for each model variable, DFFITS, covariance ratios, Cook's distances and the diagonal elements of the hat matrix. Cases which are influential with respect to any of these measures are marked with an asterisk.
The functions dfbetas, dffits, covratio and cooks.distance provide direct access to the corresponding diagnostic quantities. Functions rstandard and rstudent give the standardized and Studentized residuals respectively. (These re-normalize the residuals to have unit variance, using an overall and leave-one-out measure of the error variance respectively.)

The optional infl, res and sd arguments are there to encourage the use of these direct access functions, in situations where, e.g., the underlying basic influence measures (from lm.influence) are already available.
Note that cases with weights $==0$ are dropped from all these functions, but that if a linear model has been fitted with na.action = na.exclude, suitable values are filled it for the cases excluded during fitting.
The function hat () exists mainly for $S$ (version 2) compatibility.

## References

Belsley, D. A., Kuh, E. and Welsch, R. E. (1980) Regression Diagnostics. New York: Wiley. Cook, R. D. and Weisberg, S. (1982) Residuals and Influence in Regression. London: Chapman and Hall.

See Also<br>lm.influence.

## Examples

```
## Analysis of the life-cycle savings data
## given in Belsley, Kuh and Welsch.
data(LifeCycleSavings)
lm.SR <- lm(sr ~ pop15 + pop75 + dpi + ddpi, data = LifeCycleSavings)
summary(inflm.SR <- influence.measures(lm.SR))
inflm.SR
which(apply(inflm.SR$is.inf, 1, any)) # which observations 'are' influential
dim(dfb <- dfbetas(lm.SR)) # the 1st columns of influence.measures
all(dfb == inflm.SR$infmat[, 1:5])
rstandard(lm.SR)
rstudent(lm.SR)
dffits(lm.SR)
covratio(lm.SR)
## Huber's data [Atkinson 1985]
xh <- c(-4:0, 10)
yh <- c(2.48, .73, -.04, -1.44, -1.32, 0)
summary(lmH <- lm(yh ~ xh))
influence.measures(lmH)
```

InsectSprays Effectiveness of Insect Sprays

## Description

The counts of insects in agricultural experimental units treated with different insecticides.

## Usage

data(InsectSprays)

## Format

A data frame with 72 observations on 2 variables.

| $[, 1]$ | count | numeric | Insect count |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | spray | factor | The type of spray |

## Source

Beall, G., (1942) The Transformation of data from entomological field experiments, Biometrika, 29, 243-262.

## References

McNeil, D. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(InsectSprays)
boxplot(count ~ spray, data = InsectSprays,
        xlab = "Type of spray", ylab = "Insect count",
        main = "InsectSprays data", varwidth = TRUE, col = "lightgray")
fm1 <- aov(count ~ spray, data = InsectSprays)
summary(fm1)
opar <- par(mfrow = c(2,2), oma = c(0, 0, 1.1, 0))
plot(fm1)
fm2 <- aov(sqrt(count) ~ spray, data = InsectSprays)
summary(fm2)
plot(fm2)
par(opar)
```


## INSTALL Install Add-on Packages

## Description

Utility for installing add-on packages.

## Usage

R CMD INSTALL [options] [-1 lib] pkgs

## Arguments

pkgs A list with the path names of the packages to be installed.
lib the path name of the R library tree to install to.
options a list of options through which in particular the process for building the help files can be controlled.

## Details

If used as R CMD INSTALL pkgs without explicitly specifying lib, packages are installed into the library tree rooted at the first directory given in the environment variable R_LIBS if this is set and non-null, and to the default library tree (which is rooted at '\$R_HOME/library') otherwise.

To install into the library tree lib, use R CMD INSTALL -l lib pkgs.
Both lib and the elements of pkgs may be absolute or relative path names. pkgs can also contain name of package archive files of the form 'pkg_version.tar.gz' as obtained from CRAN, these are then extracted in a temporary directory.

Some package sources contain a 'configure' script that can be passed arguments or variables via the option --configure-args and --configure-vars, respectively, if necessary. The latter is useful in particular if libraries or header files needed for the package are in nonsystem directories. In this case, one can use the configure variables LIBS and CPPFLAGS to specify these locations (and set these via --configure-vars), see section 'Configuration variables' in "R Installation and Administration" for more information. One can also bypass the configure mechanism using the option --no-configure.
If --no-docs is given, no help files are built. Options --no-text, --no-html, and --nolatex suppress creating the text, HTML, and LaTeX versions, respectively. The default is to build help files in all three versions.

If the option --save is used, the installation procedure creates a binary image of the package code, which is then loaded when the package is attached, rather than evaluating the package source at that time. Having a file 'install.R' in the package directory makes this the default behavior for the package (option --no-save overrides). You may need --save if your package requires other packages to evaluate its own source. If the file 'install.R' is non-empty, it should contain $R$ expressions to be executed when the package is attached, after loading the saved image. Options to be passed to $R$ when creating the save image can be specified via --save=ARGS.

Use R CMD INSTALL --help for more usage information.

## Packages using the methods package

Packages that require the methods package, and that use functions such as setMethod or setClass, should be installed by creating a binary image.

The package should require the methods package, both during installation and when the user attaches the package. A good solution for most cases is to include the line require("methods") twice, once at the beginning of the package's R source, and once in the file 'install. $R$ ' in the package directory (the top-level directory, not in the ' $R$ ' directory below that). The 'install.R' file causes an image to be saved, and the contents will ensure that methods are available when the package is attached.

## See Also

REMOVE, update.packages for automatic update of packages using the internet; the chapter on "Creating R packages" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).
integer Integer Vectors

## Description

Creates or tests for objects of type "integer".

## Usage

```
integer(length = 0)
as.integer(x, ...)
is.integer(x)
```


## Arguments

```
length desired length.
x object to be coerced or tested.
... further arguments passed to or from other methods.
```


## Value

integer creates a integer vector of the specified length. Each element of the vector is equal to 0 . Integer vectors exist so that data can be passed to C or Fortran code which expects them.
as.integer attempts to coerce its argument to be of integer type. The answer will be NA unless the coercion succeeds. Real values larger in modulus than the largest integer are coerced to NA (unlike $S$ which gives the most extreme integer of the same sign).
is.integer returns TRUE or FALSE depending on whether its argument is of integer type or not.
integrate Integration of One-Dimensional Functions

## Description

Adaptive quadrature of functions of one variable over a finite or infinite interval.

## Usage

integrate(f, lower, upper, subdivisions=100, rel.tol $=$.Machine\$double.eps^0.25, abs.tol = rel.tol, stop.on.error $=$ TRUE, keep. $\mathrm{xy}=$ FALSE, aux $=$ NULL,.. )

## Arguments

| $f$ | An R function taking a numeric first argument and returning a numeric <br> vector of the same length. Returning a non-finite element will generate <br> an error. |
| :--- | :--- |
| lower, upper | The limits of integration. Can be infinite. |
| subdivisions | the maximum number of subintervals. |
| rel.tol | relative accuracy requested. |
| abs.tol | absolute accuracy requested. |
| stop.on.error | logical. If true (the default) an error stops the function. If false some <br> errors will give a result with a warning in the message component. |
| keep.xy | unused. For compatibility with S. <br> aux |
| unused. For compatibility with S. |  |

## Details

If one or both limits are infinite, the infinite range is mapped onto a finite interval.
For a finite interval, globally adaptive interval subdivision is used in connection with extrapolation by the Epsilon algorithm.
rel.tol cannot be less than $\max (50 *$. Machine\$double.eps, $0.5 \mathrm{e}-28$ ) if abs.tol $<=0$.

## Value

```
A list of class "integrate" with components
value the final estimate of the integral.
abs.error estimate of the modulus of the absolute error.
subdivisions the number of subintervals produced in the subdivision process.
message "OK" or a character string giving the error message.
call the matched call.
```


## Note

Like all numerical integration routines, these evaluate the function on a finite set of points. If the function is approximately constant (in particular, zero) over nearly all its range it is possible that the result and error estimate may be seriously wrong.

When integrating over infinite intervals do so explicitly, rather than just using a large number as the endpoint. This increases the chance of a correct answer - any function whose integral over an infinite interval is finite must be near zero for most of that interval.

## References

Based on QUADPACK routines dqags and dqagi by R. Piessens and E. deDonckerKapenga, available from Netlib.

See
R. Piessens, E. deDoncker-Kapenga, C. Uberhuber, D. Kahaner (1983) Quadpack: a Subroutine Package for Automatic Integration; Springer Verlag.

## See Also

The function adapt in the adapt package on CRAN, for multivariate integration.

## Examples

```
integrate(dnorm, -1.96, 1.96)
integrate(dnorm, -Inf, Inf)
## a slowly-convergent integral
integrand <- function(x) {1/((x+1)*sqrt(x))}
integrate(integrand, lower = 0, upper = Inf)
## don't do this if you really want the integral from 0 to Inf
integrate(integrand, lower = 0, upper = 10)
integrate(integrand, lower = 0, upper = 100000)
integrate(integrand, lower = 0, upper = 1000000, stop.on.error = FALSE)
try(integrate(function(x) 2, 0, 1)) ## no vectorizable function
integrate(function(x) rep(2, length(x)), 0, 1) ## correct
## integrate can fail if misused
integrate(dnorm,0,2)
integrate(dnorm,0,20)
integrate(dnorm,0,200)
integrate(dnorm,0,2000)
integrate(dnorm,0,20000) ## fails on many systems
integrate(dnorm,0,Inf) ## works
```

interaction Compute Factor Interactions

## Description

interaction computes a factor which represents the interaction of the given factors. The result of interaction is always unordered.

## Usage

```
interaction(..., drop=FALSE)
```


## Arguments

\(\left.\begin{array}{ll}··· . \& The factors for which interaction is to be computed, or a single list giving <br>

those factors.\end{array}\right]\)| If drop is TRUE, empty factor levels are dropped from the result. The |
| :--- |
| default is to retain all factor levels. |

## Value

A factor which represents the interaction of the given factors.

## See Also

factor.

## Examples

```
a <- gl(2, 2, 8)
b <- gl(2, 4, 8)
interaction(a, b)
```

```
interaction.plot Two-way Interaction Plot
```


## Description

Plots the mean (or other summary) of the response for two-way combinations of factors, thereby illustrating possible interactions.

## Usage

```
interaction.plot(x.factor, trace.factor, response, fun = mean,
                type = c("l", "p"), legend = TRUE,
                trace.label = deparse(substitute(trace.factor)),
                fixed = FALSE, xlab, ylab, ylim, lty, col = 1,
                        pch = c(1:9, 0, letters), ...)
```


## Arguments

| x.factor | a factor whose levels will form the $x$ axis. |
| :--- | :--- |
| trace.factor | another factor whose levels will form the traces. |
| response | a numeric variable giving the response |
| fun | the function to compute the summary. Should return a single real value. |
| type | the type of plot: lines or points. |
| legend | logical. Should a legend be included? |
| trace.label | overall label for the legend. |
| fixed | Should the legend be in the order of the levels of trace.factor or in the <br> order of the traces at their right-hand ends? |
| xlab | the x label of the plot. |
| ylab | the y label of the plot. |
| ylim | numeric of length 2 giving the y limits for the plot. |
| lty | line type for the lines drawn, with sensible default. |
| col | the color to be used for plotting. |
| pch | a vector of plotting symbols or characters, with sensible default. |
| $\ldots$ | graphics parameters to be passed to the plotting routines. |

## Details

 space left at the right for the legend (if specified). If x.factor is an ordered factor and the levels are numeric, these numeric values are used for the x axis.
The response and hence its summary can contain missing values. If so, the missing values and the line segments joining them are omitted from the plot (and this can be somewhat disconcerting).
The graphics parameters xlab, ylab, ylim, lty, col and pch are given suitable defaults (and xlim and xaxs are set and cannot be overriden). The defaults are to cycle through the line types, use the foreground colour, and to use the symbols $1: 9,0$, and the capital letters to plot the traces.

## Note

Some of the argument names and the precise behaviour are chosen for S-compatibility.

## Examples

```
data(ToothGrowth)
attach(ToothGrowth)
interaction.plot(dose, supp, len, fixed=TRUE)
dose <- ordered(dose)
interaction.plot(dose, supp, len, fixed=TRUE)
detach()
data(OrchardSprays)
attach(OrchardSprays)
interaction.plot(treatment, rowpos, decrease)
interaction.plot(rowpos, treatment, decrease)
## order the rows by their mean effect
rowpos <- factor(rowpos, levels=sort.list(tapply(decrease, rowpos, mean)))
interaction.plot(rowpos, treatment, decrease)
detach()
data(esoph)
attach(esoph)
interaction.plot(agegp, alcgp, ncases/ncontrols)
interaction.plot(agegp, tobgp, ncases/ncontrols, trace.label="tobacco",
    fixed=TRUE)
detach()
```

interactive Is R Running Interactively?

## Description

Return TRUE when $R$ is being used interactively and FALSE otherwise.

## Usage

```
interactive()
```

See Also

```
source, .First
```


## Examples

```
.First <- function() if(interactive()) x11()
```

Internal Call an Internal Function

## Description

. Internal performs a call to an internal code which is built in to the R interpreter. Only true $R$ wizards should even consider using this function.

## Usage

.Internal(call)

## Arguments

```
call a call expression
```


## See Also

.Primitive, . C, .Fortran.

```
invisible Change the Print Mode to Invisible
```


## Description

Return a (temporarily) invisible copy of an object.

## Usage

invisible(x)

## Arguments

X an arbitrary R object.

## Details

This function can be useful when it is desired to have functions return values which can be assigned, but which do not print when they are not assigned.

## See Also

```
return, function.
```


## Examples

```
# These functions both return their argument
f1 <- function(x) x
f2 <- function(x) invisible(x)
f1(1)# prints
f2(1)# does not
```

IQR The Interquartile Range

## Description

computes interquartile range of the x values.

## Usage

$\operatorname{IQR}(x$, na.rm $=$ FALSE $)$

## Arguments

| x | a numeric vector. |
| :--- | :--- |
| na.rm | logical. Should missing values be removed? |

## Details

Note that this function computes the quartiles using the quantile function rather than following Tukey's recommendations, i.e., $\operatorname{IQR}(x)=$ quantile ( $x, 3 / 4$ ) - quantile ( $x, 1 / 4$ ).
For normally $N(m, 1)$ distributed $X$, the expected value of $\operatorname{IQR}(\mathrm{X})$ is $2 *$ qnorm $(3 / 4)=$ 1.3490, i.e., for a normal-consistent estimate of the standard deviation, use IQR(x) / 1.349.

## References

Tukey, J. W. (1977). Exploratory Data Analysis. Reading: Addison-Wesley.

## See Also

fivenum, mad which is more robust, range, quantile.

## Examples

```
data(rivers)
IQR(rivers)
```


## Description

This famous (Fisher's or Anderson's) iris data set gives the measurements in centimeters of the variables sepal length and width and petal length and width, respectively, for 50 flowers from each of 3 species of iris. The species are Iris setosa, versicolor, and virginica.

```
Usage
    data(iris)
    data(iris3)
```


## Format

iris is a data frame with 150 cases (rows) and 5 variables (columns) named Sepal.Length, Sepal.Width, Petal.Length, Petal.Width, and Species.
iris3 gives the same data arranged as a 3-dimensional array of size 50 by 4 by 3 , as represented by S-PLUS. The first dimension gives the case number within the species subsample, the second the measurements with names Sepal L., Sepal W., Petal L., and Petal W., and the third the species.

## Source

Fisher, R. A. (1936) The use of multiple measurements in taxonomic problems. Annals of Eugenics, 7, Part II, 179-188.

The data were collected by Anderson, Edgar (1935). The irises of the Gaspe Peninsula, Bulletin of the American Iris Society, 59, 2-5.

## See Also

matplot some examples of which use iris.

## Examples

```
data(iris3)
dni3 <- dimnames(iris3)
ii <- data.frame(matrix(aperm(iris3, c(1,3,2)), ncol=4,
            dimnames=list(NULL, sub(" L.",".Length",
                                    sub(" W.",".Width", dni3[[2]])))),
    Species = gl(3,50,lab=sub("S","s",sub("V","v",dni3[[3]]))))
data(iris)
all.equal(ii, iris) # TRUE
```

```
is.empty.model Check if a Model is Empty
```


## Description

R model notation allows models with no intercept and no predictors. These require special handling internally. is.empty.model() checks whether an object describes an empty model.

## Usage

is.empty.model(x)

## Arguments

X
A terms object or an object with a terms method.

## Value

TRUE if the model is empty

## See Also

lm,glm

## Examples

```
y <- rnorm(20)
is.empty.model(y ~ 0)
is.empty.model(y ~ -1)
is.empty.model(lm(y ~ 0))
```


## Description

is.finite and is.infinite return a vector of the same length as x , indicating which elements are finite or not.

Inf and -Inf are positive and negative 'infinity' whereas NaN means "Not a Number".

## Usage

```
is.finite(x)
is.infinite(x)
Inf
NaN
is.nan(x)
```


## Arguments

x (numerical) object to be tested.

## Details

is.finite returns a vector of the same length as $x$ the $j$ th element of which is TRUE if $x[j]$ is finite (i.e. it is not one of the values NA, NaN, Inf or -Inf).
is.infinite returns a vector of the same length as $x$ the jth element of which is TRUE if $\mathrm{x}[\mathrm{j}]$ is infinite (i.e. equal to one of $\operatorname{Inf}$ or $-\operatorname{Inf}$ ).

## Note

In R, basically all mathematical functions (including basic Arithmetic), are supposed to work properly with +/- Inf and NaN as input or output.

The basic rule should be that calls and relations with Infs really are statements with a proper mathematical limit, see the many examples below.

## References

ANSI/IEEE 754 Floating-Point Standard.
Currently (6/2002), Bill Metzenthen's 〈billm@suburbia.net〉 tutorial and examples at http://www.suburbia.net/~billm/

## See Also

NA, 'Not Available' which is not a number as well, however usually used for missing values.

## Examples

```
pi / 0 ## = Inf a non-zero number divided by zero creates infinity
0 / 0 ## = NaN
1/0 + 1/0# Inf
1/0 - 1/0# NaN
stopifnot(
    1/0 == Inf,
    1/Inf == 0
)
exp(-Inf) == 0
## (actually, the last one seems to give NA on not-very-new
## versions of Linux, which is a Linux bug and seems to be
## corrected in newer 'libc6' based Linuxen).
stopifnot(
    is.na(0/0),
    !is.na(Inf),
    is.nan(0/0),
    !is.nan(NA) && !is.infinite(NA) && !is.finite(NA),
        is.nan(NaN) && !is.infinite(NaN) && !is.finite(NaN),
    !is.nan(c(1,NA)),
    c(FALSE,TRUE,FALSE) == is.nan(c (1,NaN,NA)),
    c(FALSE,TRUE,FALSE) == is.nan(list(1,NaN,NA))#-> FALSE in older versions
)
```

```
lgamma(Inf) == Inf
Inf + Inf == Inf
Inf - Inf == NaN # NA --- should test with 'is.nan()
(1/0) * (1/0)# Inf
(1/0) / (1/0)# NaN
pm <- c(-1,1) # 'pm' = plus/minus
log(0) == - 1/0
exp(-Inf) == 0
sin(Inf)
cos(Inf)
tan(Inf)
all(atan(Inf*pm) == pm*pi/2) # TRUE
x <- c(100,-1e-13,Inf,-Inf, NaN, pi, NA)
x # 1.000000 -3.000000 Inf -Inf NA 3.141593 NA
names(x) <- formatC(x, dig=3)
is.finite(x)
##- 100 -1e-13 Inf -Inf NaN 3.14 NA
##- T T . . . T .
is.na(x)
##- 100 -1e-13 Inf -Inf NaN 3.14 NA
##- . . . . T . T
which(is.na(x) & !is.nan(x))# only 'NA': 7
is.na(x) | is.finite(x)
##- 100 -1e-13 Inf -Inf NaN 3.14 NA
##- T T . . T T T
is.infinite(x)
##- 100 -1e-13 Inf -Inf NaN 3.14 NA
##- . . T T
##-- either finite or infinite or NA:
all(is.na(x) != is.finite(x) | is.infinite(x)) # TRUE
all(is.nan(x) != is.finite(x) | is.infinite(x)) # FALSE: have 'real' NA
##--- Integer
(ix <- structure(as.integer(x), names= names(x)))
##- 100-1e-13 Inf -Inf NaN 3.14 NA
##- 100 0 NA NA NA 3 NA
all(is.na(ix) != is.finite(ix) | is.infinite(ix)) # TRUE (still)
ix[3] == (iI <- as.integer(Inf))#> warning: NAs introduced by coercion
ix[4] == (imI<- as.integer(-Inf))
iI == .Machine$integer.max # TRUE
imI == -.Machine$integer.max # TRUE
##--- Overflow in simple integer arithmetic:
as.integer(2)*iI # -2
as.integer(3)*iI # 2147483645
as.integer(3)*iI == iI-2 # TRUE
```

```
storage.mode(ii <- -3:5)
storage.mode(zm <- outer(ii,ii, FUN="*"))# integer
storage.mode(zd <- outer(ii,ii, FUN="/"))# double
range(zd, na.rm=TRUE)# -Inf Inf
zd[,ii==0]
(storage.mode(print(1:1 / 0:0)))# Inf "double"
(storage.mode(print(1:1 / 1:1)))# 1 "double"
(storage.mode(print(1:1 + 1:1)))# 2 "integer"
(storage.mode(print(2:2 * 2:2)))# 4 "integer"
```

is.function Is an Object of Type Function?

## Description

Checks whether its argument is a function.

## Usage

```
is.function(x)
```


## Arguments

X
an R object.

## Value

TRUE if $x$ is a function, and FALSE otherwise.

```
is.language Is an Object a Language Object?
```


## Description

is.language returns TRUE if x is either a variable name, a call, or an expression.

## Usage

is.language(x)

## Arguments

$x \quad$ object to be tested.

## Examples

```
ll <- list(a = expression(x^2 - 2*x + 1), b = as.name("Jim"),
    c = as.expression(exp(1)), d = call("sin", pi))
sapply(ll, typeof)
sapply(ll, mode)
stopifnot(sapply(ll, is.language))
```

```
is.object Is an Object "internally classed"?
```


## Description

A function rather for internal use. It returns TRUE if the object $x$ has the $R$ internal OBJECT attribute set, and FALSE otherwise.

## Usage

is.object( x )

## Arguments

$x \quad$ object to be tested.

## See Also

class, and methods.

## Examples

```
is.object(1) # FALSE
is.object(as.factor(1:3)) # TRUE
```

```
is.R Are we using R, rather than S?
```


## Description

Test if running under R .

## Usage

is. $R()$

## Details

The function has been written such as to correctly run in all versions of R, S and S-PLUS. In order for code to be runnable in both $R$ and $S$ dialects, either your the code must define is.R or use it as

```
if (exists("is.R") && is.function(is.R) && is.R()) {
```

\#\# R-specific code
\} else \{
\#\# S-version of code
\}

## Value

is. $R$ returns TRUE if we are using $R$ and FALSE otherwise.

See Also
R.version, system.

## Examples

```
x <- runif(20); small <- x < 0.4
# 'which()' only exists in R:
if(is.R()) which(small) else seq(along=small) [small]
```

is.recursive Is an Object Atomic or Recursive?

## Description

is.atomic returns TRUE if x does not have a list structure and FALSE otherwise.
is.recursive returns TRUE if x has a recursive (list-like) structure and FALSE otherwise.

## Usage

```
is.atomic(x)
is.recursive(x)
```


## Arguments

$x \quad$ object to be tested.

## See Also

 is.list, is.language, etc, and the demo("is.things").
## Examples

```
is.a.r <- function(x) c(is.atomic(x), is.recursive(x))
is.a.r(c(a=1,b=3))# TRUE FALSE
is.a.r(list()) # FALSE TRUE ??
is.a.r(list(2)) # FALSE TRUE
is.a.r(lm)
    # "
is.a.r(y ~ x) # "
is.a.r(expression(x+1))# should be F-T (not in 0.62.3!)
```

```
is.single Is an Object of Single Precision Type?
```


## Description

is.single reports an error. There are no single precision values in R.

## Usage

is.single( x )

## Arguments

$x \quad$ object to be tested.
islands Areas of the World's Major Landmasses

## Description

The areas in thousands of square miles of the landmasses which exceed 10,000 square miles.

## Usage

data(islands)

## Format

A named vector of length 48.

## Source

The World Almanac and Book of Facts, 1975, page 406.

## References

McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(islands)
dotchart(log(islands, 10),
    main = "islands data: log10(area) (log10(sq. miles))")
dotchart(log(islands[order(islands)], 10),
    main = "islands data: log10(area) (log10(sq. miles))")
```

```
Japanese Japanese characters in R
```


## Description

The implementation of Hershey vector fonts provides a large number of Japanese characters (Hiragana, Katakana, and Kanji).

## Details

Without keyboard support for typing Japanese characters, the only way to produce these characters is to use special escape sequences.
For example, the Hiragana character for the sound "ka" is produced by <br>\#J242b and the Katakana character for this sound is produced by $\backslash \backslash \# J 252 b$. The Kanji ideograph for "one" is produced by <br>\#J306c or <br>\#N0001.
The output from example(Japanese) shows tables of the escape sequences for the available Japanese characters.

## References

http://www.gnu.org/software/plotutils/plotutils.html

## See Also

Hershey, text, contour

## Examples

```
plot(1:9, type="n", axes=FALSE, frame=TRUE, ylab="",
    main= "example(Japanese)", xlab= "using Hershey fonts")
par(cex=3)
Vf <- c("serif", "plain")
text(4, 2, "\\#J2438\\#J2421\\#J2451\\#J2473", vfont = vf)
text(4, 4, "\\#J2538\\#J2521\\#J2551\\#J2573", vfont = Vf)
text(4, 6, "\\#J467c\\#J4b5c", vfont = Vf)
text(4, 8, "Japan", vfont = Vf)
par(cex=1)
text(8, 2, "Hiragana")
text(8, 4, "Katakana")
text(8, 6, "Kanji")
text(8, 8, "English")
######
# create tables of Japanese characters
######
make.table <- function(nr, nc) {
    opar <- par(mar=rep(0, 4), pty="s")
    plot(c(0, nc*(10%/%nc) + 1), c(0, -(nr + 1)),
        type="n", xlab="", ylab="", axes=FALSE)
    invisible(opar)
}
get.r <- function(i, nr) i %% nr + 1
```

```
get.c <- function(i, nr) i %/% nr + 1
Esc2 <- function(str) paste("\\", str, sep="")
draw.title <- function(title, nc)
    text((nc*(10%/%nc) + 1)/2, 0, title, font=2)
draw.vf.cell <- function(typeface, fontindex, string, i, nr, raw.string=NULL) {
        r <- get.r(i, nr)
        c <- get.c(i, nr)
        x0 <- 2*(c - 1)
        if (is.null(raw.string)) raw.string <- Esc2(string)
        text(x0 + 1.1, -r, raw.string, col="grey")
        text(x0 + 2, -r, string, vfont=c(typeface, fontindex))
        rect(x0 + .5, -(r - . 5), x0 + 2.5, -(r + .5), border="grey")
}
draw.vf.cell2 <- function(string, alt, i, nr) {
        r <- get.r(i, nr)
        c <- get.c(i, nr)
        x0 <- 3*(c - 1)
        text(x0 + 1.1, -r, Esc2(string <- Esc2(string)), col="grey")
        text(x0 + 2.2, -r, Esc2(Esc2(alt)), col="grey", cex=.6)
        text(x0 + 3, -r, string, vfont=c("serif", "plain"))
        rect(x0 + .5, -(r - .5), x0 + 3.5, -(r + .5), border="grey")
}
tf <- "serif"
fi <- "plain"
nr <- 25
nc <- 4
oldpar <- make.table(nr, nc)
index <- 0
digits <- c(0:9,"a","b","c","d","e","f")
draw.title("Hiragana : \\\\#J24nn", nc)
for (i in 2:7) {
        for (j in 1:16) {
            if (!((i == 2 && j == 1) || (i == 7 && j > 4))) {
                draw.vf.cell(tf, fi, paste("\\#J24", i, digits[j], sep=""),
                                    index, nr)
                index <- index + 1
            }
        }
}
nr <- 25
nc <- 4
make.table(nr, nc)
index <- 0
digits <- c(0:9,"a","b","c","d","e","f")
draw.title("Katakana : \\\\#J25nn", nc)
for (i in 2:7) {
    for (j in 1:16) {
        if (!((i == 2 && j == 1) || (i == 7 && j > 7))) {
                draw.vf.cell(tf, fi, paste("\\#J25", i, digits[j], sep=""),
                                    index, nr)
        index <- index + 1
        }
```

```
    }
```

\}
nr <- 26
nc <- 3
make.table(nr, nc)
i <- 0
draw.title("Kanji (1)", nc)
draw.vf.cell2("\#J3021", "\#N0043", i, nr); i <- i + 1
draw.vf.cell2("\#J3026", "\#N2829", i, nr); i <- i + 1
draw.vf.cell2("\#J302d", "\#N0062", i, nr); i <- i + 1
draw.vf.cell2("\#J3035", "\#N0818", i, nr); i <- i + 1
draw.vf.cell2("\#J303f", "\#N1802", i, nr); i <- i + 1
draw.vf.cell2("\#J3045", "\#N2154", i, nr); i <- i + 1
draw.vf.cell2("\#J304c", "\#N0401", i, nr); i <- i + 1
draw.vf.cell2("\#J3057", "\#N2107", i, nr); i <- i + 1
draw.vf.cell2("\#J3059", "\#N0138", i, nr) ; i <- i + 1
draw.vf.cell2("\#J305b", "\#N3008", i, nr); i <- i + 1
draw.vf.cell2("\#J305e", "\#N3579", i, nr); i <- i + 1
draw.vf.cell2("\#J3061", "\#N4214", i, nr); i <- i + 1
draw.vf.cell2("\#J306c", "\#NO001", i, nr); i <- i + 1
draw.vf.cell2("\#J3070", "\#N3294", i, nr); i <- i + 1
draw.vf.cell2("\#J3078", "\#N1026", i, nr); i <- i + 1
draw.vf.cell2("\#J307a", "\#N1562", i, nr); i <- i + 1
draw.vf.cell2("\#J3122", "\#N5006", i, nr); i <- i + 1
draw.vf.cell2("\#J3126", "\#N0878", i, nr); i <- i + 1
draw.vf.cell2("\#J3127", "\#N1280", i, nr); i <- i + 1
draw.vf.cell2("\#J3129", "\#N3673", i, nr); i <- i + 1
draw.vf.cell2("\#J312b", "\#N5042", i, nr); i <- i + 1
draw.vf.cell2("\#J3132", "\#N2629", i, nr); i <- i + 1
draw.vf.cell2("\#J313b", "\#N2973", i, nr); i <- i + 1
draw.vf.cell2("\#J313f", "\#N4725", i, nr); i <- i + 1
draw.vf.cell2("\#J3140", "\#N5046", i, nr); i <- i + 1
draw.vf.cell2("\#J314a", "\#N0130", i, nr); i <- i + 1
draw.vf.cell2("\#J3155", "\#N2599", i, nr); i <- i + 1
draw.vf.cell2("\#J315f", "\#N0617", i, nr); i <- i + 1
draw.vf.cell2("\#J3173", "\#N4733", i, nr); i <- i + 1
draw.vf.cell2("\#J3176", "\#N1125", i, nr); i <- i + 1
draw.vf.cell2("\#J3177", "\#N2083", i, nr); i <- i + 1
draw.vf.cell2("\#J317e", "\#N1504", i, nr); i <- i + 1
draw.vf.cell2("\#J3221", "\#N1885", i, nr); i <- i + 1
draw.vf.cell2("\#J3223", "\#N2361", i, nr); i <- i + 1
draw.vf.cell2("\#J3226", "\#N2922", i, nr); i <- i + 1
draw.vf.cell2("\#J322b", "\#N5399", i, nr); i <- i + 1
draw.vf.cell2("\#J322f", "\#N0551", i, nr); i <- i + 1
draw.vf.cell2("\#J3235", "\#N0260", i, nr); i <- i + 1
draw.vf.cell2("\#J3239", "\#N2634", i, nr); i <- i + 1
draw.vf.cell2("\#J323b", "\#N5110", i, nr); i <- i + 1
draw.vf.cell2("\#J323c", "\#N0009", i, nr); i <- i + 1
draw.vf.cell2("\#J323d", "\#NO350", i, nr); i <- i + 1
draw.vf.cell2("\#J323f", "\#N0409", i, nr); i <- i + 1
draw.vf.cell2("\#J3241", "\#N0422", i, nr); i <- i + 1
draw.vf.cell2("\#J3243", "\#N0716", i, nr); i <- i + 1
draw.vf.cell2("\#J3244", "\#N0024", i, nr); i <- i + 1
draw.vf.cell2("\#J3246", "\#N0058", i, nr); i <- i + 1
draw.vf.cell2("\#J3248", "\#N1311", i, nr); i <- i + 1
draw.vf.cell2("\#J324a", "\#N3272", i, nr); i <- i + 1
draw.vf.cell2("\#J324c", "\#N0107", i, nr); i <- i + 1 draw.vf.cell2("\#J324f", "\#N2530", i, nr); i <- i + 1 draw.vf.cell2("\#J3250", "\#N2743", i, nr); i <- i + 1 draw.vf.cell2("\#J3256", "\#N3909", i, nr); i <- i + 1 draw.vf.cell2("\#J3259", "\#N3956", i, nr); i <- i + 1 draw.vf.cell2("\#J3261", "\#N4723", i, nr); i <- i + 1 draw.vf.cell2("\#J3267", "\#N2848", i, nr); i <- i + 1 draw.vf.cell2("\#J3268", "\#NO050", i, nr); i <- i + 1 draw.vf.cell2("\#J3272", "\#N4306", i, nr); i <- i + 1 draw.vf.cell2("\#J3273", "\#N1028", i, nr); i <- i + 1 draw.vf.cell2("\#J3323", "\#N2264", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3324", "\#N2553", i, nr); i <- i + 1 draw.vf.cell2("\#J3326", "\#N2998", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3328", "\#N3537", i, nr); i <- i + 1 draw.vf.cell2("\#J332b", "\#N4950", i, nr); i <- i + 1 draw.vf.cell2("\#J332d", "\#N4486", i, nr); i <- i + 1 draw.vf.cell2("\#J3330", "\#N1168", i, nr); i <- i + 1 draw.vf.cell2("\#J3346", "\#N1163", i, nr); i <- i + 1 draw.vf.cell2("\#J334b", "\#N2254", i, nr); i <- i + 1 draw.vf.cell2("\#J3351", "\#N4301", i, nr); i <- i + 1 draw.vf.cell2("\#J3353", "\#N4623", i, nr); i <- i + 1 draw.vf.cell2("\#J3357", "\#N5088", i, nr); i <- i + 1 draw.vf.cell2("\#J3358", "\#N1271", i, nr); i <- i + 1 draw.vf.cell2("\#J335a", "\#N2324", i, nr); i <- i + 1 draw.vf.cell2("\#J3364", "\#N0703", i, nr); i <- i + 1 draw.vf.cell2("\#J3424", "\#N2977", i, nr); i <- i + 1 draw.vf.cell2("\#J3428", "\#N1322", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (2)", nc)
draw.vf.cell2("\#J342c", "\#N1466", i, nr); i <- i + 1 draw.vf.cell2("\#J3433", "\#N1492", i, nr); i <- i + 1 draw.vf.cell2("\#J3434", "\#N0790", i, nr); i <- i + 1 draw.vf.cell2("\#J3436", "\#N1731", i, nr); i <- i + 1 draw.vf.cell2("\#J3437", "\#N1756", i, nr); i <- i + 1 draw.vf.cell2("\#J3445", "\#N2988", i, nr); i <- i + 1 draw.vf.cell2("\#J3449", "\#N3416", i, nr); i <- i + 1 draw.vf.cell2("\#J3454", "\#N4750", i, nr); i <- i + 1 draw.vf.cell2("\#J3456", "\#N4949", i, nr); i <- i + 1 draw.vf.cell2("\#J3458", "\#N4958", i, nr); i <- i + 1 draw.vf.cell2("\#J346f", "\#N0994", i, nr); i <- i + 1 draw.vf.cell2("\#J3470", "\#N1098", i, nr); i <- i + 1 draw.vf.cell2("\#J3476", "\#N1496", i, nr); i <- i + 1 draw.vf.cell2("\#J347c", "\#N3785", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3521", "\#N2379", i, nr); i <- i + 1 draw.vf.cell2("\#J3522", "\#N1582", i, nr); i <- i + 1 draw.vf.cell2("\#J3524", "\#N2480", i, nr); i <- i + 1 draw.vf.cell2("\#J3525", "\#N2507", i, nr); i <- i + 1 draw.vf.cell2("\#J352d", "\#N4318", i, nr); i <- i + 1 draw.vf.cell2("\#J3530", "\#N4610", i, nr); i <- i + 1 draw.vf.cell2("\#J3534", "\#N5276", i, nr); i <- i + 1 draw.vf.cell2("\#J3535", "\#N5445", i, nr); i <- i + 1 draw.vf.cell2("\#J3546", "\#N3981", i, nr); i <- i + 1 draw.vf.cell2("\#J3555", "\#N4685", i, nr); i <- i + 1 draw.vf.cell2("\#J355a", "\#N0154", i, nr); i <- i + 1 draw.vf.cell2("\#J355b", "\#NO885", i, nr) ; i <- i + 1
draw.vf.cell2("\#J355d", "\#N1560", i, nr); i <- i + 1 draw.vf.cell2("\#J3565", "\#N2941", i, nr); i <- i + 1 draw.vf.cell2("\#J3566", "\#N3314", i, nr); i <- i + 1 draw.vf.cell2("\#J3569", "\#N3496", i, nr); i <- i + 1 draw.vf.cell2("\#J356d", "\#N2852", i, nr); i <- i + 1 draw.vf.cell2("\#J356e", "\#N1051", i, nr); i <- i + 1 draw.vf.cell2("\#J356f", "\#N1387", i, nr); i <- i + 1 draw.vf.cell2("\#J3575", "\#N4109", i, nr); i <- i + 1 draw.vf.cell2("\#J3577", "\#N4548", i, nr); i <- i + 1 draw.vf.cell2("\#J357b", "\#N5281", i, nr) ; i <- i + 1 draw.vf.cell2("\#J357e", "\#NO295", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3621", "\#NO431", i, nr); i <- i + 1 draw.vf.cell2("\#J3626", "\#N0581", i, nr); i <- i + 1 draw.vf.cell2("\#J362d", "\#N1135", i, nr); i <- i + 1 draw.vf.cell2("\#J362f", "\#N1571", i, nr); i <- i + 1 draw.vf.cell2("\#J3635", "\#N2052", i, nr); i <- i + 1 draw.vf.cell2("\#J3636", "\#N2378", i, nr); i <- i + 1 draw.vf.cell2("\#J364a", "\#N0103", i, nr); i <- i + 1 draw.vf.cell2("\#J364b", "\#N2305", i, nr); i <- i + 1 draw.vf.cell2("\#J364c", "\#N2923", i, nr); i <- i + 1 draw.vf.cell2("\#J3651", "\#N1065", i, nr); i <- i + 1 draw.vf.cell2("\#J3661", "\#N4671", i, nr); i <- i + 1 draw.vf.cell2("\#J3662", "\#N4815", i, nr); i <- i + 1 draw.vf.cell2("\#J3664", "\#N4855", i, nr); i <- i + 1 draw.vf.cell2("\#J3665", "\#N0146", i, nr); i <- i + 1 draw.vf.cell2("\#J3671", "\#N3128", i, nr); i <- i + 1 draw.vf.cell2("\#J3675", "\#N3317", i, nr); i <- i + 1 draw.vf.cell2("\#J367e", "\#N1386", i, nr); i <- i + 1 draw.vf.cell2("\#J3738", "\#N0449", i, nr); i <- i + 1 draw.vf.cell2("\#J3739", "\#N0534", i, nr); i <- i + 1 draw.vf.cell2("\#J373e", "\#N2937", i, nr); i <- i + 1 draw.vf.cell2("\#J373f", "\#N1077", i, nr); i <- i + 1 draw.vf.cell2("\#J3741", "\#N1589", i, nr); i <- i + 1 draw.vf.cell2("\#J3742", "\#N1602", i, nr); i <- i + 1 draw.vf.cell2("\#J374f", "\#N0195", i, nr); i <- i + 1 draw.vf.cell2("\#J3750", "\#N3523", i, nr); i <- i + 1 draw.vf.cell2("\#J3757", "\#N4312", i, nr); i <- i + 1 draw.vf.cell2("\#J375a", "\#N4620", i, nr); i <- i + 1 draw.vf.cell2("\#J3767", "\#N2412", i, nr); i <- i + 1 draw.vf.cell2("\#J3768", "\#N2509", i, nr); i <- i + 1 draw.vf.cell2("\#J376a", "\#N3313", i, nr); i <- i + 1 draw.vf.cell2("\#J376b", "\#N3540", i, nr); i <- i + 1 draw.vf.cell2("\#J376c", "\#N4205", i, nr); i <- i + 1 draw.vf.cell2("\#J376e", "\#N2169", i, nr); i <- i + 1 draw.vf.cell2("\#J3777", "\#N1045", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3824", "\#N2868", i, nr); i <- i + 1 draw.vf.cell2("\#J3826", "\#N3180", i, nr); i <- i + 1 draw.vf.cell2("\#J3828", "\#N3543", i, nr); i <- i + 1 draw.vf.cell2("\#J382b", "\#N4284", i, nr); i <- i + 1 draw.vf.cell2("\#J3833", "\#N5220", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (3)", nc)
draw.vf.cell2("\#J3835", "\#N0275", i, nr); i <- i + 1
draw.vf.cell2("\#J3836", "\#NO825", i, nr); i <- i + 1
draw.vf.cell2("\#J3839", "\#N1568", i, nr); i <- i + 1
draw.vf.cell2("\#J383a", "\#N2637", i, nr); i <- i + 1 draw.vf.cell2("\#J383b", "\#N2656", i, nr); i <- i + 1 draw.vf.cell2("\#J383d", "\#N2943", i, nr); i <- i + 1 draw.vf.cell2("\#J3840", "\#N4309", i, nr); i <- i + 1 draw.vf.cell2("\#J3842", "\#N4987", i, nr); i <- i + 1 draw.vf.cell2("\#J3845", "\#NO770", i, nr); i <- i + 1 draw.vf.cell2("\#J3847", "\#N1036", i, nr); i <- i + 1 draw.vf.cell2("\#J384c", "\#N1567", i, nr); i <- i + 1 draw.vf.cell2("\#J384d", "\#N1817", i, nr); i <- i + 1 draw.vf.cell2("\#J384e", "\#N2044", i, nr); i <- i + 1 draw.vf.cell2("\#J385d", "\#N5415", i, nr) ; i <- i + 1 draw.vf.cell2("\#J385e", "\#N0015", i, nr); i <- i + 1 draw.vf.cell2("\#J3861", "\#N0162", i, nr); i <- i + 1 draw.vf.cell2("\#J3865", "\#N1610", i, nr); i <- i + 1 draw.vf.cell2("\#J3866", "\#N1628", i, nr); i <- i + 1 draw.vf.cell2("\#J386c", "\#N4374", i, nr); i <- i + 1 draw.vf.cell2("\#J3872", "\#NO290", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3877", "\#N1358", i, nr); i <- i + 1 draw.vf.cell2("\#J3878", "\#N0579", i, nr); i <- i + 1 draw.vf.cell2("\#J387d", "\#N0868", i, nr); i <- i + 1 draw.vf.cell2("\#J387e", "\#N0101", i, nr); i <- i + 1 draw.vf.cell2("\#J3929", "\#N1451", i, nr); i <- i + 1 draw.vf.cell2("\#J3931", "\#N1683", i, nr); i <- i + 1 draw.vf.cell2("\#J393d", "\#N2343", i, nr); i <- i + 1 draw.vf.cell2("\#J3943", "\#N0092", i, nr); i <- i + 1 draw.vf.cell2("\#J394d", "\#N3684", i, nr); i <- i + 1 draw.vf.cell2("\#J3954", "\#N4213", i, nr); i <- i + 1 draw.vf.cell2("\#J3955", "\#N1641", i, nr); i <- i + 1 draw.vf.cell2("\#J395b", "\#N4843", i, nr) ; i <- i + 1 draw.vf.cell2("\#J395d", "\#N4883", i, nr); i <- i + 1 draw.vf.cell2("\#J395f", "\#N4994", i, nr); i <- i + 1 draw.vf.cell2("\#J3960", "\#N1459", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3961", "\#N5188", i, nr); i <- i + 1 draw.vf.cell2("\#J3962", "\#N5248", i, nr); i <- i + 1 draw.vf.cell2("\#J3966", "\#NO882", i, nr); i <- i + 1 draw.vf.cell2("\#J3967", "\#N0383", i, nr); i <- i + 1 draw.vf.cell2("\#J3971", "\#N1037", i, nr); i <- i + 1 draw.vf.cell2("\#J3975", "\#N5403", i, nr); i <- i + 1 draw.vf.cell2("\#J397c", "\#N5236", i, nr) ; i <- i + 1 draw.vf.cell2("\#J397e", "\#N4660", i, nr); i <- i + 1 draw.vf.cell2("\#J3a21", "\#N2430", i, nr); i <- i + 1 draw.vf.cell2("\#J3a23", "\#NO352", i, nr); i <- i + 1 draw.vf.cell2("\#J3a2c", "\#N2261", i, nr); i <- i + 1 draw.vf.cell2("\#J3a38", "\#N1455", i, nr); i <- i + 1 draw.vf.cell2("\#J3a39", "\#N3662", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3a42", "\#N1515", i, nr); i <- i + 1 draw.vf.cell2("\#J3a46", "\#NO035", i, nr); i <- i + 1 draw.vf.cell2("\#J3a47", "\#N2146", i, nr); i <- i + 1 draw.vf.cell2("\#J3a59", "\#N3522", i, nr); i <- i + 1 draw.vf.cell2("\#J3a5f", "\#N1055", i, nr); i <- i + 1 draw.vf.cell2("\#J3a6e", "\#N0407", i, nr); i <- i + 1 draw.vf.cell2("\#J3a72", "\#N2119", i, nr); i <- i + 1 draw.vf.cell2("\#J3a79", "\#N2256", i, nr); i <- i + 1 draw.vf.cell2("\#J3b2e", "\#N3113", i, nr); i <- i + 1 draw.vf.cell2("\#J3b30", "\#N0008", i, nr); i <- i + 1 draw.vf.cell2("\#J3b33", "\#N1407", i, nr); i <- i + 1 draw.vf.cell2("\#J3b36", "\#N2056", i, nr) ; i <- i + 1
draw.vf.cell2("\#J3b3b", "\#N3415", i, nr); i <- i + 1 draw.vf.cell2("\#J3b40", "\#N4789", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3b45", "\#N0362", i, nr); i <- i + 1 draw.vf.cell2("\#J3b4d", "\#N1025", i, nr); i <- i + 1 draw.vf.cell2("\#J3b4e", "\#N1160", i, nr); i <- i + 1 draw.vf.cell2("\#J3b4f", "\#N1208", i, nr); i <- i + 1 draw.vf.cell2("\#J3b52", "\#N1264", i, nr); i <- i + 1 draw.vf.cell2("\#J3b54", "\#NO284", i, nr); i <- i + 1 draw.vf.cell2("\#J3b57", "\#N3001", i, nr); i <- i + 1 draw.vf.cell2("\#J3b58", "\#N1904", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3b59", "\#N2039", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3b5e", "\#N2211", i, nr); i <- i + 1 draw.vf.cell2("\#J3b5f", "\#N2429", i, nr); i <- i + 1 draw.vf.cell2("\#J3b60", "\#N2439", i, nr); i <- i + 1 draw.vf.cell2("\#J3b61", "\#N2478", i, nr); i <- i + 1 draw.vf.cell2("\#J3b64", "\#N3265", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (4)", nc)
draw.vf.cell2("\#J3b65", "\#N3492", i, nr); i <- i + 1
draw.vf.cell2("\#J3b66", "\#N3510", i, nr); i <- i + 1
draw.vf.cell2("\#J3b6a", "\#N3845", i, nr); i <- i + 1
draw.vf.cell2("\#J3b73", "\#N2435", i, nr); i <- i + 1
draw.vf.cell2("\#J3b75", "\#N5428", i, nr); i <- i + 1
draw.vf.cell2("\#J3b76", "\#NO272", i, nr); i <- i + 1
draw.vf.cell2("\#J3b7a", "\#N1281", i, nr); i <- i + 1
draw.vf.cell2("\#J3b7d", "\#N1903", i, nr); i <- i + 1
draw.vf.cell2("\#J3b7e", "\#N2126", i, nr); i <- i + 1
draw.vf.cell2("\#J3c21", "\#N0638", i, nr); i <- i + 1
draw.vf.cell2("\#J3c27", "\#N3209", i, nr); i <- i + 1
draw.vf.cell2("\#J3c28", "\#N3228", i, nr); i <- i + 1
draw.vf.cell2("\#J3c2a", "\#N3697", i, nr); i <- i + 1
draw.vf.cell2("\#J3c2b", "\#N3841", i, nr); i <- i + 1
draw.vf.cell2("\#J3c2d", "\#N3860", i, nr); i <- i + 1
draw.vf.cell2("\#J3c2f", "\#N5375", i, nr); i <- i + 1
draw.vf.cell2("\#J3c30", "\#N1556", i, nr); i <- i + 1
draw.vf.cell2("\#J3c34", "\#N4619", i, nr); i <- i + 1
draw.vf.cell2("\#J3c37", "\#NO261", i, nr) ; i <- i + 1
draw.vf.cell2("\#J3c3c", "\#N1300", i, nr); i <- i + 1
draw.vf.cell2("\#J3c3e", "\#N2631", i, nr); i <- i + 1
draw.vf.cell2("\#J3c41", "\#N4518", i, nr); i <- i + 1
draw.vf.cell2("\#J3c42", "\#N1297", i, nr); i <- i + 1
draw.vf.cell2("\#J3c4d", "\#N4603", i, nr); i <- i + 1
draw.vf.cell2("\#J3c50", "\#N2074", i, nr) ; i <- i + 1
draw.vf.cell2("\#J3c54", "\#N3685", i, nr); i <- i + 1
draw.vf.cell2("\#J3c56", "\#N4608", i, nr); i <- i + 1
draw.vf.cell2("\#J3c5c", "\#N1377", i, nr); i <- i + 1
draw.vf.cell2("\#J3c61", "\#N4809", i, nr); i <- i + 1
draw.vf.cell2("\#J3c63", "\#N3926", i, nr); i <- i + 1
draw.vf.cell2("\#J3c67", "\#NO285", i, nr); i <- i + 1
draw.vf.cell2("\#J3c68", "\#N3699", i, nr); i <- i + 1
draw.vf.cell2("\#J3c6a", "\#N1827", i, nr); i <- i + 1
draw.vf.cell2("\#J3c6f", "\#N3295", i, nr); i <- i + 1
draw.vf.cell2("\#J3c72", "\#N2573", i, nr); i <- i + 1
draw.vf.cell2("\#J3c73", "\#N5186", i, nr); i <- i + 1
draw.vf.cell2("\#J3c7e", "\#N0622", i, nr) ; i <- i + 1
draw.vf.cell2("\#J3d29", "\#N3273", i, nr); i <- i + 1 draw.vf.cell2("\#J3d2a", "\#N3521", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3d2e", "\#N3863", i, nr); i <- i + 1 draw.vf.cell2("\#J3d39", "\#N4798", i, nr); i <- i + 1 draw.vf.cell2("\#J3d3d", "\#NO768", i, nr); i <- i + 1 draw.vf.cell2("\#J3d3e", "\#N1613", i, nr); i <- i + 1 draw.vf.cell2("\#J3d44", "\#N3597", i, nr); i <- i + 1 draw.vf.cell2("\#J3d45", "\#NO224", i, nr); i <- i + 1 draw.vf.cell2("\#J3d50", "\#N0097", i, nr); i <- i + 1 draw.vf.cell2("\#J3d51", "\#N1621", i, nr); i <- i + 1 draw.vf.cell2("\#J3d55", "\#N2122", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3d60", "\#N0791", i, nr); i <- i + 1 draw.vf.cell2("\#J3d63", "\#N3509", i, nr); i <- i + 1 draw.vf.cell2("\#J3d68", "\#N1162", i, nr); i <- i + 1 draw.vf.cell2("\#J3d6b", "\#N2138", i, nr); i <- i + 1 draw.vf.cell2("\#J3d71", "\#N3719", i, nr); i <- i + 1 draw.vf.cell2("\#J3d77", "\#N1185", i, nr); i <- i + 1 draw.vf.cell2("\#J3d7c", "\#N4993", i, nr); i <- i + 1 draw.vf.cell2("\#J3e26", "\#N0321", i, nr); i <- i + 1 draw.vf.cell2("\#J3e2e", "\#N1355", i, nr); i <- i + 1 draw.vf.cell2("\#J3e2f", "\#NO166", i, nr); i <- i + 1 draw.vf.cell2("\#J3e3d", "\#N2137", i, nr); i <- i + 1 draw.vf.cell2("\#J3e3e", "\#N2212", i, nr); i <- i + 1 draw.vf.cell2("\#J3e46", "\#N2772", i, nr); i <- i + 1 draw.vf.cell2("\#J3e4b", "\#N3192", i, nr); i <- i + 1 draw.vf.cell2("\#J3e4e", "\#N3280", i, nr); i <- i + 1 draw.vf.cell2("\#J3e57", "\#N1638", i, nr); i <- i + 1 draw.vf.cell2("\#J3e5a", "\#N4341", i, nr); i <- i + 1 draw.vf.cell2("\#J3e5d", "\#N4472", i, nr); i <- i + 1 draw.vf.cell2("\#J3e65", "\#N0798", i, nr); i <- i + 1 draw.vf.cell2("\#J3e68", "\#N0223", i, nr); i <- i + 1 draw.vf.cell2("\#J3e6c", "\#N1113", i, nr); i <- i + 1 draw.vf.cell2("\#J3e6f", "\#N1364", i, nr); i <- i + 1 draw.vf.cell2("\#J3e75", "\#N2839", i, nr); i <- i + 1 draw.vf.cell2("\#J3e78", "\#N4002", i, nr); i <- i + 1 draw.vf.cell2("\#J3f22", "\#N2303", i, nr); i <- i + 1 draw.vf.cell2("\#J3f27", "\#N3889", i, nr); i <- i + 1 draw.vf.cell2("\#J3f29", "\#N5154", i, nr); i <- i + 1 draw.vf.cell2("\#J3f2d", "\#NO403", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (5)", nc)
draw.vf.cell2("\#J3f34", "\#N1645", i, nr); i <- i + 1 draw.vf.cell2("\#J3f36", "\#N1920", i, nr) ; i <- i + 1 draw.vf.cell2("\#J3f37", "\#N2080", i, nr); i <- i + 1 draw.vf.cell2("\#J3f39", "\#N2301", i, nr); i <- i + 1 draw.vf.cell2("\#J3f3f", "\#N0783", i, nr); i <- i + 1 draw.vf.cell2("\#J3f43", "\#N3837", i, nr); i <- i + 1 draw.vf.cell2("\#J3f48", "\#N4601", i, nr); i <- i + 1 draw.vf.cell2("\#J3f49", "\#N4646", i, nr); i <- i + 1 draw.vf.cell2("\#J3f4a", "\#N4709", i, nr); i <- i + 1 draw.vf.cell2("\#J3f4c", "\#N5055", i, nr); i <- i + 1 draw.vf.cell2("\#J3f4d", "\#N0339", i, nr); i <- i + 1 draw.vf.cell2("\#J3f5e", "\#N1034", i, nr); i <- i + 1 draw.vf.cell2("\#J3f62", "\#NO211", i, nr); i <- i + 1 draw.vf.cell2("\#J3f65", "\#N2482", i, nr) ; i <- i + 1
draw.vf.cell2("\#J3f69", "\#N3676", i, nr); i <- i + 1 draw.vf.cell2("\#J3f74", "\#N2057", i, nr) ; i <- i + 1 draw.vf.cell2("\#J402d", "\#N1666", i, nr); i <- i + 1 draw.vf.cell2("\#J402e", "\#N1799", i, nr); i <- i + 1 draw.vf.cell2("\#J4030", "\#N2436", i, nr); i <- i + 1 draw.vf.cell2("\#J4031", "\#N2121", i, nr); i <- i + 1 draw.vf.cell2("\#J4032", "\#N2143", i, nr); i <- i + 1 draw.vf.cell2("\#J4035", "\#N0027", i, nr); i <- i + 1 draw.vf.cell2("\#J4038", "\#N2991", i, nr); i <- i + 1 draw.vf.cell2("\#J403e", "\#N4273", i, nr); i <- i + 1 draw.vf.cell2("\#J4044", "\#N5076", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4045", "\#N5077", i, nr); i <- i + 1 draw.vf.cell2("\#J404e", "\#N2108", i, nr); i <- i + 1 draw.vf.cell2("\#J404f", "\#N2194", i, nr); i <- i + 1 draw.vf.cell2("\#J4050", "\#N3176", i, nr); i <- i + 1 draw.vf.cell2("\#J4051", "\#N3306", i, nr); i <- i + 1 draw.vf.cell2("\#J4056", "\#N4534", i, nr); i <- i + 1 draw.vf.cell2("\#J405a", "\#N0667", i, nr); i <- i + 1 draw.vf.cell2("\#J405c", "\#N1951", i, nr); i <- i + 1 draw.vf.cell2("\#J405e", "\#N1855", i, nr); i <- i + 1 draw.vf.cell2("\#J4063", "\#N5044", i, nr); i <- i + 1 draw.vf.cell2("\#J4064", "\#N3539", i, nr); i <- i + 1 draw.vf.cell2("\#J4065", "\#N3855", i, nr); i <- i + 1 draw.vf.cell2("\#J4068", "\#N0571", i, nr); i <- i + 1 draw.vf.cell2("\#J4069", "\#N0156", i, nr); i <- i + 1 draw.vf.cell2("\#J406e", "\#N1447", i, nr); i <- i + 1 draw.vf.cell2("\#J4070", "\#N1823", i, nr); i <- i + 1 draw.vf.cell2("\#J407e", "\#N3580", i, nr); i <- i + 1 draw.vf.cell2("\#J4125", "\#N3873", i, nr); i <- i + 1 draw.vf.cell2("\#J4130", "\#N0595", i, nr); i <- i + 1 draw.vf.cell2("\#J4133", "\#N2770", i, nr); i <- i + 1 draw.vf.cell2("\#J4134", "\#N0384", i, nr); i <- i + 1 draw.vf.cell2("\#J4147", "\#N3511", i, nr); i <- i + 1 draw.vf.cell2("\#J4148", "\#N3520", i, nr); i <- i + 1 draw.vf.cell2("\#J4150", "\#N0859", i, nr); i <- i + 1 draw.vf.cell2("\#J4158", "\#N1402", i, nr); i <- i + 1 draw.vf.cell2("\#J415b", "\#N1728", i, nr); i <- i + 1 draw.vf.cell2("\#J4161", "\#N2100", i, nr); i <- i + 1 draw.vf.cell2("\#J416a", "\#N2241", i, nr); i <- i + 1 draw.vf.cell2("\#J416d", "\#N3567", i, nr); i <- i + 1 draw.vf.cell2("\#J4170", "\#N3939", i, nr); i <- i + 1 draw.vf.cell2("\#J4175", "\#N4234", i, nr); i <- i + 1 draw.vf.cell2("\#J4176", "\#N4539", i, nr); i <- i + 1 draw.vf.cell2("\#J417c", "\#NO540", i, nr); i <- i + 1 draw.vf.cell2("\#J417d", "\#N1137", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4224", "\#N4701", i, nr); i <- i + 1 draw.vf.cell2("\#J4226", "\#N0509", i, nr); i <- i + 1 draw.vf.cell2("\#J422b", "\#N0196", i, nr); i <- i + 1 draw.vf.cell2("\#J422c", "\#N2632", i, nr); i <- i + 1 draw.vf.cell2("\#J422d", "\#N4546", i, nr); i <- i + 1 draw.vf.cell2("\#J422e", "\#N4700", i, nr); i <- i + 1 draw.vf.cell2("\#J4233", "\#N3544", i, nr); i <- i + 1 draw.vf.cell2("\#J4236", "\#N0590", i, nr); i <- i + 1 draw.vf.cell2("\#J4238", "\#N1267", i, nr); i <- i + 1 draw.vf.cell2("\#J423e", "\#N0361", i, nr); i <- i + 1 draw.vf.cell2("\#J423f", "\#N1169", i, nr); i <- i + 1 draw.vf.cell2("\#J4240", "\#N1172", i, nr); i <- i + 1
draw.vf.cell2("\#J424a", "\#N2313", i, nr); i <- i + 1 draw.vf.cell2("\#J424e", "\#NO405", i, nr); i <- i + 1 draw.vf.cell2("\#J4250", "\#N2067", i, nr); i <- i + 1 draw.vf.cell2("\#J4256", "\#N1743", i, nr); i <- i + 1 draw.vf.cell2("\#J4265", "\#NO364", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (6)", nc)
draw.vf.cell2("\#J4267", "\#N1171", i, nr); i <- i + 1 draw.vf.cell2("\#J4268", "\#N3385", i, nr) ; i <- i + 1 draw.vf.cell2("\#J426a", "\#N2164", i, nr); i <- i + 1 draw.vf.cell2("\#J426c", "\#N2655", i, nr); i <- i + 1 draw.vf.cell2("\#J4274", "\#N2503", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4323", "\#N4721", i, nr); i <- i + 1 draw.vf.cell2("\#J432b", "\#N4458", i, nr); i <- i + 1 draw.vf.cell2("\#J432f", "\#N4384", i, nr); i <- i + 1 draw.vf.cell2("\#J4331", "\#N0139", i, nr); i <- i + 1 draw.vf.cell2("\#J433a", "\#N1418", i, nr); i <- i + 1 draw.vf.cell2("\#J433b", "\#N3172", i, nr); i <- i + 1 draw.vf.cell2("\#J4346", "\#N1575", i, nr); i <- i + 1 draw.vf.cell2("\#J434b", "\#N2996", i, nr); i <- i + 1 draw.vf.cell2("\#J434d", "\#NO488", i, nr); i <- i + 1 draw.vf.cell2("\#J434e", "\#N3169", i, nr); i <- i + 1 draw.vf.cell2("\#J434f", "\#N1056", i, nr); i <- i + 1 draw.vf.cell2("\#J4356", "\#N3644", i, nr); i <- i + 1 draw.vf.cell2("\#J4359", "\#N4722", i, nr); i <- i + 1 draw.vf.cell2("\#J435d", "\#N3366", i, nr); i <- i + 1 draw.vf.cell2("\#J4362", "\#N3325", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4363", "\#N3940", i, nr); i <- i + 1 draw.vf.cell2("\#J4365", "\#N3665", i, nr); i <- i + 1 draw.vf.cell2("\#J4366", "\#N0081", i, nr); i <- i + 1 draw.vf.cell2("\#J4368", "\#N1291", i, nr); i <- i + 1 draw.vf.cell2("\#J436b", "\#N0053", i, nr); i <- i + 1 draw.vf.cell2("\#J436c", "\#N2236", i, nr); i <- i + 1 draw.vf.cell2("\#J436e", "\#N4115", i, nr); i <- i + 1 draw.vf.cell2("\#J442b", "\#N3788", i, nr); i <- i + 1 draw.vf.cell2("\#J442c", "\#N2702", i, nr); i <- i + 1 draw.vf.cell2("\#J4436", "\#N4543", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4439", "\#N4938", i, nr); i <- i + 1 draw.vf.cell2("\#J443b", "\#N5340", i, nr); i <- i + 1 draw.vf.cell2("\#J443e", "\#NO775", i, nr); i <- i + 1 draw.vf.cell2("\#J444c", "\#N4703", i, nr); i <- i + 1 draw.vf.cell2("\#J4463", "\#NO406", i, nr); i <- i + 1 draw.vf.cell2("\#J446a", "\#N1296", i, nr) ; i <- i + 1 draw.vf.cell2("\#J446c", "\#N1508", i, nr); i <- i + 1 draw.vf.cell2("\#J446d", "\#N1514", i, nr); i <- i + 1 draw.vf.cell2("\#J4472", "\#N1914", i, nr); i <- i + 1 draw.vf.cell2("\#J4478", "\#N3285", i, nr); i <- i + 1 draw.vf.cell2("\#J4479", "\#N3581", i, nr); i <- i + 1 draw.vf.cell2("\#J4526", "\#N1987", i, nr); i <- i + 1 draw.vf.cell2("\#J452a", "\#N3097", i, nr); i <- i + 1 draw.vf.cell2("\#J452f", "\#N0931", i, nr); i <- i + 1 draw.vf.cell2("\#J4534", "\#N4844", i, nr); i <- i + 1 draw.vf.cell2("\#J4535", "\#N0588", i, nr); i <- i + 1 draw.vf.cell2("\#J4537", "\#N0016", i, nr); i <- i + 1 draw.vf.cell2("\#J453e", "\#N4615", i, nr); i <- i + 1

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draw.vf.cell2("#J4540", "#N0804", i, nr); i <- i + 1
draw.vf.cell2("#J4544", "#N2994", i, nr); i <- i + 1
draw.vf.cell2("#J4545", "#N5050", i, nr); i <- i + 1
draw.vf.cell2("#J454c", "#N1614", i, nr); i <- i + 1
draw.vf.cell2("#J4559", "#N1511", i, nr); i <- i + 1
draw.vf.cell2("#J455a", "#N1050", i, nr); i <- i + 1
draw.vf.cell2("#J455f", "#N1161", i, nr); i <- i + 1
draw.vf.cell2("#J4561", "#N0665", i, nr); i <- i + 1
draw.vf.cell2("#J4563", "#N1109", i, nr); i <- i + 1
draw.vf.cell2("#J4567", "#NO230", i, nr); i <- i + 1
draw.vf.cell2("#J456c", "#NO213", i, nr); i <- i + 1
draw.vf.cell2("#J4574", "#N2745", i, nr); i <- i + 1
draw.vf.cell2("#J4576", "#N1359", i, nr); i <- i + 1
draw.vf.cell2("#J4579", "#N3396", i, nr); i <- i + 1
draw.vf.cell2("#J4626", "#N4465", i, nr); i <- i + 1
draw.vf.cell2("#J4630", "#N0730", i, nr); i <- i + 1
draw.vf.cell2("#J4631", "#N0619", i, nr); i <- i + 1
draw.vf.cell2("#J4633", "#N1354", i, nr); i <- i + 1
draw.vf.cell2("#J463b", "#N4724", i, nr); i <- i + 1
draw.vf.cell2("#J463c", "#N4853", i, nr); i <- i + 1
draw.vf.cell2("#J4643", "#N2860", i, nr); i <- i + 1
draw.vf.cell2("#J4649", "#N4375", i, nr); i <- i + 1
draw.vf.cell2("#J465e", "#N2160", i, nr); i <- i + 1
draw.vf.cell2("#J4662", "#N0082", i, nr); i <- i + 1
draw.vf.cell2("#J466e", "#N0778", i, nr); i <- i + 1
draw.vf.cell2("#J4671", "#N5038", i, nr); i <- i + 1
draw.vf.cell2("#J4673", "#NO273", i, nr); i <- i + 1
draw.vf.cell2("#J4679", "#N3724", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (7)", nc)
draw.vf.cell2("#J467c", "#N2097", i, nr); i <- i + 1
draw.vf.cell2("#J467e", "#N0574", i, nr); i <- i + 1
draw.vf.cell2("#J4721", "#N1189", i, nr); i <- i + 1
draw.vf.cell2("#J472e", "#N2797", i, nr); i <- i + 1
draw.vf.cell2("#J472f", "#N0188", i, nr); i <- i + 1
draw.vf.cell2("#J4733", "#N2808", i, nr); i <- i + 1
draw.vf.cell2("#J4734", "#N3472", i, nr); i <- i + 1
draw.vf.cell2("#J4748", "#N2529", i, nr); i <- i + 1
draw.vf.cell2("#J474f", "#N5191", i, nr); i <- i + 1
draw.vf.cell2("#J4769", "#N3275", i, nr); i <- i + 1
draw.vf.cell2("#J4772", "#N3095", i, nr); i <- i + 1
draw.vf.cell2("#J477e", "#N5385", i, nr); i <- i + 1
draw.vf.cell2("#J4821", "#NO049", i, nr); i <- i + 1
draw.vf.cell2("#J482c", "#N0577", i, nr); i <- i + 1
draw.vf.cell2("#J482f", "#N3092", i, nr); i <- i + 1
draw.vf.cell2("#J483e", "#N0132", i, nr); i <- i + 1
draw.vf.cell2("#J483f", "#N0817", i, nr); i <- i + 1
draw.vf.cell2("#J4841", "#N1469", i, nr); i <- i + 1
draw.vf.cell2("#J484c", "#N3865", i, nr); i <- i + 1
draw.vf.cell2("#J4856", "#N4811", i, nr); i <- i + 1
draw.vf.cell2("#J4860", "#N1604", i, nr); i <- i + 1
draw.vf.cell2("#J4866", "#N2470", i, nr); i <- i + 1
draw.vf.cell2("#J4869", "#N3109", i, nr); i <- i + 1
draw.vf.cell2("#J4873", "#N5080", i, nr); i <- i + 1
draw.vf.cell2("#J4874", "#N5152", i, nr); i <- i + 1
```

draw.vf.cell2("\#J4878", "\#N1383", i, nr); i <- i + 1 draw.vf.cell2("\#J4879", "\#N1631", i, nr) ; i <- i + 1 draw.vf.cell2("\#J487e", "\#N3658", i, nr); i <- i + 1 draw.vf.cell2("\#J4921", "\#N5421", i, nr); i <- i + 1 draw.vf.cell2("\#J492e", "\#N3397", i, nr); i <- i + 1 draw.vf.cell2("\#J4934", "\#N0033", i, nr); i <- i + 1 draw.vf.cell2("\#J4938", "\#N2359", i, nr); i <- i + 1 draw.vf.cell2("\#J4939", "\#N0131", i, nr); i <- i + 1 draw.vf.cell2("\#J493d", "\#N0108", i, nr); i <- i + 1 draw.vf.cell2("\#J4942", "\#N3042", i, nr); i <- i + 1 draw.vf.cell2("\#J4943", "\#N3271", i, nr) ; i <- i + 1 draw.vf.cell2("\#J494a", "\#N0923", i, nr); i <- i + 1 draw.vf.cell2("\#J4954", "\#N0017", i, nr); i <- i + 1 draw.vf.cell2("\#J495b", "\#N1468", i, nr); i <- i + 1 draw.vf.cell2("\#J4963", "\#N2832", i, nr); i <- i + 1 draw.vf.cell2("\#J4969", "\#N4488", i, nr); i <- i + 1 draw.vf.cell2("\#J4977", "\#N5148", i, nr); i <- i + 1 draw.vf.cell2("\#J497d", "\#N1484", i, nr); i <- i + 1 draw.vf.cell2("\#J4a23", "\#N4255", i, nr); i <- i + 1 draw.vf.cell2("\#J4a26", "\#N0173", i, nr); i <- i + 1 draw.vf.cell2("\#J4a2a", "\#N2857", i, nr); i <- i + 1 draw.vf.cell2("\#J4a2c", "\#N0578", i, nr); i <- i + 1 draw.vf.cell2("\#J4a38", "\#N2064", i, nr); i <- i + 1 draw.vf.cell2("\#J4a39", "\#N4959", i, nr); i <- i + 1 draw.vf.cell2("\#J4a3f", "\#N0026", i, nr); i <- i + 1 draw.vf.cell2("\#J4a42", "\#N0589", i, nr); i <- i + 1 draw.vf.cell2("\#J4a44", "\#N4945", i, nr); i <- i + 1 draw.vf.cell2("\#J4a46", "\#N3461", i, nr); i <- i + 1 draw.vf.cell2("\#J4a50", "\#N0511", i, nr); i <- i + 1 draw.vf.cell2("\#J4a51", "\#N0306", i, nr); i <- i + 1 draw.vf.cell2("\#J4a52", "\#N2842", i, nr); i <- i + 1 draw.vf.cell2("\#J4a55", "\#N4661", i, nr); i <- i + 1 draw.vf.cell2("\#J4a6c", "\#N2466", i, nr); i <- i + 1 draw.vf.cell2("\#J4a7c", "\#N2084", i, nr); i <- i + 1 draw.vf.cell2("\#J4a7d", "\#N2082", i, nr); i <- i + 1 draw.vf.cell2("\#J4b21", "\#N2535", i, nr); i <- i + 1 draw.vf.cell2("\#J4b26", "\#N3749", i, nr); i <- i + 1 draw.vf.cell2("\#J4b4c", "\#N0751", i, nr); i <- i + 1 draw.vf.cell2("\#J4b4f", "\#N5404", i, nr); i <- i + 1 draw.vf.cell2("\#J4b5c", "\#N0096", i, nr); i <- i + 1 draw.vf.cell2("\#J4b63", "\#N5390", i, nr); i <- i + 1 draw.vf.cell2("\#J4b68", "\#N2467", i, nr); i <- i + 1 draw.vf.cell2("\#J4b74", "\#N0855", i, nr); i <- i + 1 draw.vf.cell2("\#J4b7c", "\#N0007", i, nr); i <- i + 1 draw.vf.cell2("\#J4c23", "\#N0913", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4c24", "\#N0179", i, nr); i <- i + 1 draw.vf.cell2("\#J4c29", "\#N1316", i, nr); i <- i + 1 draw.vf.cell2("\#J4c35", "\#N2773", i, nr); i <- i + 1 draw.vf.cell2("\#J4c37", "\#N3164", i, nr); i <- i + 1 draw.vf.cell2("\#J4c3e", "\#N1170", i, nr); i <- i + 1 draw.vf.cell2("\#J4c40", "\#N2110", i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title("Kanji (8)", nc)
draw.vf.cell2("\#J4c4c", "\#N5087", i, nr); i <- i + 1 draw.vf.cell2("\#J4c53", "\#N2473", i, nr) ; i <- i + 1
draw.vf.cell2("\#J4c5a", "\#N2170", i, nr); i <- i + 1 draw.vf.cell2("\#J4c5c", "\#N3127", i, nr); i <- i + 1 draw.vf.cell2("\#J4c64", "\#N4944", i, nr); i <- i + 1 draw.vf.cell2("\#J4c67", "\#N4940", i, nr); i <- i + 1 draw.vf.cell2("\#J4c6b", "\#NO298", i, nr); i <- i + 1 draw.vf.cell2("\#J4c70", "\#N3168", i, nr); i <- i + 1 draw.vf.cell2("\#J4c72", "\#N1598", i, nr); i <- i + 1 draw.vf.cell2("\#J4c74", "\#N4074", i, nr); i <- i + 1 draw.vf.cell2("\#J4c78", "\#N2233", i, nr); i <- i + 1 draw.vf.cell2("\#J4c7d", "\#N2534", i, nr); i <- i + 1 draw.vf.cell2("\#J4d2d", "\#N3727", i, nr); i <- i + 1 draw.vf.cell2("\#J4d30", "\#N2565", i, nr); i <- i + 1 draw.vf.cell2("\#J4d3a", "\#N5030", i, nr); i <- i + 1 draw.vf.cell2("\#J4d3c", "\#N1167", i, nr); i <- i + 1 draw.vf.cell2("\#J4d3e", "\#N0408", i, nr); i <- i + 1 draw.vf.cell2("\#J4d4f", "\#N2659", i, nr); i <- i + 1 draw.vf.cell2("\#J4d51", "\#N2993", i, nr); i <- i + 1 draw.vf.cell2("\#J4d53", "\#N3656", i, nr); i <- i + 1 draw.vf.cell2("\#J4d55", "\#N4001", i, nr); i <- i + 1 draw.vf.cell2("\#J4d57", "\#N4274", i, nr); i <- i + 1 draw.vf.cell2("\#J4d5b", "\#N5012", i, nr); i <- i + 1 draw.vf.cell2("\#J4d63", "\#N3680", i, nr); i <- i + 1 draw.vf.cell2("\#J4d68", "\#NO202", i, nr); i <- i + 1 draw.vf.cell2("\#J4d6b", "\#N5049", i, nr); i <- i + 1 draw.vf.cell2("\#J4d70", "\#N3856", i, nr); i <- i + 1 draw.vf.cell2("\#J4d71", "\#N0199", i, nr); i <- i + 1 draw.vf.cell2("\#J4d72", "\#N1431", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4d78", "\#N3264", i, nr); i <- i + 1 draw.vf.cell2("\#J4d7d", "\#N2942", i, nr); i <- i + 1 draw.vf.cell2("\#J4e24", "\#N4813", i, nr); i <- i + 1 draw.vf.cell2("\#J4e25", "\#N5040", i, nr); i <- i + 1 draw.vf.cell2("\#J4e26", "\#N5005", i, nr); i <- i + 1 draw.vf.cell2("\#J4e28", "\#N0319", i, nr); i <- i + 1 draw.vf.cell2("\#J4e29", "\#N3343", i, nr); i <- i + 1 draw.vf.cell2("\#J4e2e", "\#N2576", i, nr); i <- i + 1 draw.vf.cell2("\#J4e32", "\#N3191", i, nr); i <- i + 1 draw.vf.cell2("\#J4e33", "\#N3471", i, nr); i <- i + 1 draw.vf.cell2("\#J4e35", "\#N5440", i, nr); i <- i + 1 draw.vf.cell2("\#J4e3e", "\#N0034", i, nr); i <- i + 1 draw.vf.cell2("\#J4e41", "\#N3468", i, nr); i <- i + 1 draw.vf.cell2("\#J4e49", "\#N3885", i, nr); i <- i + 1 draw.vf.cell2("\#J4e4c", "\#N2141", i, nr); i <- i + 1 draw.vf.cell2("\#J4e4f", "\#N0715", i, nr); i <- i + 1 draw.vf.cell2("\#J4e53", "\#N2210", i, nr); i <- i + 1 draw.vf.cell2("\#J4e55", "\#N2807", i, nr) ; i <- i + 1 draw.vf.cell2("\#J4e58", "\#N4630", i, nr); i <- i + 1 draw.vf.cell2("\#J4e60", "\#N5138", i, nr); i <- i + 1 draw.vf.cell2("\#J4e63", "\#N0428", i, nr); i <- i + 1 draw.vf.cell2("\#J4e64", "\#N0642", i, nr); i <- i + 1 draw.vf.cell2("\#J4e6d", "\#N5048", i, nr); i <- i + 1 draw.vf.cell2("\#J4e6e", "\#N5056", i, nr); i <- i + 1 draw.vf.cell2("\#J4e73", "\#N2438", i, nr); i <- i + 1 draw.vf.cell2("\#J4f22", "\#N4702", i, nr); i <- i + 1 draw.vf.cell2("\#J4f27", "\#N2750", i, nr); i <- i + 1 draw.vf.cell2("\#J4f29", "\#N4561", i, nr); i <- i + 1 draw.vf.cell2("\#J4f37", "\#N3683", i, nr); i <- i + 1 draw.vf.cell2("\#J4f3b", "\#NO283", i, nr); i <- i + 1

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draw.vf.cell2("#J4f40", "#N4391", i, nr); i <- i + 1
draw.vf.cell2("#J4f42", "#N3268", i, nr); i <- i + 1
draw.vf.cell2("#J4f43", "#N4358", i, nr); i <- i + 1
draw.vf.cell2("#J4f44", "#N0054", i, nr); i <- i + 1
draw.vf.cell2("#J4f47", "#N1710", i, nr); i <- i + 1
draw.vf.cell2("#J534c", "#N0973", i, nr); i <- i + 1
draw.vf.cell2("#J5879", "#N1794", i, nr); i <- i + 1
draw.vf.cell2("#J5960", "#N1942", i, nr); i <- i + 1
draw.vf.cell2("#J626f", "#N3200", i, nr); i <- i + 1
draw.vf.cell2("#J6446", "#N3458", i, nr); i <- i + 1
draw.vf.cell2("#J6647", "#N5083", i, nr); i <- i + 1
draw.vf.cell2("#J6d55", "#N4633", i, nr); i <- i + 1
par(oldpar)
```

jitter Add 'Jitter' (Noise) to Numbers

## Description

Add a small amount of noise to a numeric vector.

## Usage

```
jitter(x, factor=1, amount = NULL)
```


## Arguments

x numeric to which jitter should be added.
factor numeric
amount numeric; if positive, used as amount (see below), otherwise, if $=0$ the default is factor $* z / 50$.
Default (NULL): factor $* \mathrm{~d} / 5$ where d is about the smallest difference between x values.

## Details

The result, say $r$, is $r<-x+\operatorname{runif}(n,-a$, $a)$ where $n<-l e n g t h(x)$ and $a$ is the amount argument (if specified).
Let $z<-\max (x)-\min (x)$ (assuming the usual case). The amount a to be added is either provided as positive argument amount or otherwise computed from $\mathbf{z}$, as follows:
If amount $==0$, we set $a<-$ factor $* z / 50$ (same as S).
If amount is NULL (default), we set a <- factor $* \mathrm{~d} / 5$ where $d$ is the smallest difference between adjacent unique (apart from fuzz) x values.

## Value

jitter ( $\mathrm{x}, \ldots$ ) returns a numeric of the same length as x , but with an amount of noise added in order to break ties.

## Author(s)

Werner Stahel and Martin Maechler, ETH Zurich

## References

Chambers, J. M., Cleveland, W. S., Kleiner, B. and Tukey, P.A. (1983) Graphical Methods for Data Analysis. Wadsworth; figures 2.8, 4.22, 5.4.

## See Also

rug which you may want to combine with jitter.

## Examples

```
round(jitter(c(rep(1,3), rep(1.2, 4), rep(3,3))), 3)
## These two 'fail' with S-plus 3.x:
jitter(rep(0, 7))
jitter(rep(10000,5))
```

kappa Estimate the Condition Number

## Description

An estimate of the condition number of a matrix or of the $R$ matrix of a $Q R$ decomposition, perhaps of a linear fit. The condition number is defined as the ratio of the largest to the smallest non-zero singular value of the matrix.

## Usage

```
kappa(z, ...)
kappa.lm (z, ...)
kappa.default(z, exact = FALSE, ...)
kappa.qr (z, ...)
kappa.tri (z, exact = FALSE, ...)
```


## Arguments

z
A matrix or a the result of qr or a fit from a class inheriting from "lm".
exact logical. Should the result be exact?
... further arguments passed to or from other methods.

## Details

If exact $=$ FALSE (the default) the condition number is estimated by a cheap approximation. Following S, this uses the LINPACK routine 'dtrco.f'. However, in R (or S) the exact calculation is also likely to be quick enough.

## Value

The condition number, kappa, or an approximation if exact=FALSE.

## Author(s)

B.D. Ripley

## See Also

svd for the singular value decomposition and qr for the $Q R$ one.

## Examples

```
kappa(x1 <- cbind(1,1:10))# 15.71
kappa(x1, exact = TRUE) # 13.68
kappa(x2 <- cbind(x1,2:11))# high! [x2 is singular!]
hilbert <- function(n) { i <- 1:n; 1 / outer(i - 1, i, "+") }
sv9 <- svd(h9 <- hilbert(9))$ d
kappa(h9)# pretty high!
kappa(h9, exact = TRUE) == max(sv9) / min(sv9)
kappa(h9, exact = TRUE) / kappa(h9) # . 677 (i.e. rel.error = 32%)
```

kronecker
Kronecker products on arrays

## Description

Computes the generalised kronecker product of two arrays, X and Y. kronecker (X, Y) returns an array A with dimensions $\operatorname{dim}(X) * \operatorname{dim}(Y)$.

## Usage

kronecker (X, Y, FUN = "*", make.dimnames = FALSE, ...) X \% x\% Y

## Arguments

| X | A vector or array. |
| :--- | :--- |
| Y | A vector or array. |
| FUN | a function; it may be a quoted string. |
| make.dimnames | Provide dimnames that are the product of the dimnames of X and Y. |
| $\ldots$ | optional arguments to be passed to FUN. |

## Details

If $X$ and $Y$ do not have the same number of dimensions, the smaller array is padded with dimensions of size one. The returned array comprises submatrices constructed by taking $X$ one term at a time and expanding that term as FUN (x, Y, ...).
$\% \mathrm{x} \%$ is an alias for kronecker (where FUN is hardwired to "*").

Author(s)
Jonathan Rougier, 〈J.C.Rougier@durham.ac.uk〉

## References

Matrix Algebra Useful for Statistics, Shayle R. Searle, John Wiley and Sons, 1982.

## See Also

outer, on which kronecker is built and $\% * \%$ for usual matrix multiplication.

## Examples

```
# simple scalar multiplication
( M <- matrix(1:6, ncol=2) )
stopifnot(kronecker(4, M)==4 * M)
# Block diagonal matrix:
stopifnot(kronecker(diag(1, 3), M) == diag(1, 3) %x% M)
# ask for dimnames
fred <- matrix(1:12, 3, 4, dimnames=list(LETTERS[1:3], LETTERS[4:7]))
bill <- c("happy" = 100, "sad" = 1000)
kronecker(fred, bill, make.dimnames = TRUE)
bill <- outer(bill, c("cat"=3, "dog"=4))
kronecker(fred, bill, make.dimnames = TRUE)
```


## labels <br> Find Labels from Object

## Description

Find a suitable set of labels from an object for use in printing or plotting, for example.

## Usage

```
labels(object, ...)
labels.default(object, ...)
labels.terms(object, ...)
labels.lm(object, ...)
```


## Arguments

$\begin{array}{ll}\text { object } & \text { Any } \mathrm{R} \text { object: the function is generic. } \\ \ldots & \text { further arguments passed to or from other methods. }\end{array}$

Value
A character vector or list of such vectors. For a vector the results is the names or seq(along=x), for a data frame or array it is the dimnames (with NULL expanded to $\operatorname{seq}(l e n=d[i])$ ), for a terms object it is the term labels and for an lm object it is the term labels for estimable terms.

Author(s)
B.D. Ripley

```
lapply Apply a Function over a List or Vector
```


## Description

lapply returns a list of the same length as X. Each element of which is the result of applying FUN to the corresponding element of X .
sapply is a "user-friendly" version of lapply also accepting vectors as X, and returning a vector or array with dimnames if appropriate.

## Usage

lapply(X, FUN, ...)
sapply(X, FUN, ..., simplify = TRUE, USE. NAMES = TRUE)

## Arguments

X list or vector to be used.
FUN the function to be applied. In the case of functions like,$+ \% * \%$, etc., the function name must be quoted.
... optional arguments to FUN.
simplify logical; should the result be simplified to a vector if possible?
USE.NAMES logical; if TRUE and if $X$ is character, use $X$ as names for the result unless it had names already.

## See Also

apply, tapply.

## Examples

```
x <- list(a = 1:10, beta = exp(-3:3), logic = c(TRUE,FALSE,FALSE,TRUE))
# compute the list mean for each list element
lapply(x,mean)
# median and quartiles for each list element
lapply(x, quantile, probs = 1:3/4)
sapply(x, quantile)
str(i39 <- sapply(3:9, seq))# list of vectors
sapply(i39, fivenum)
```


## Last.value Value of Last Evaluated Expression

## Description

The value of the internal evaluation of a top-level R expression is always assigned to .Last.value (in package:base) before further processing (e.g. printing).

## Usage

.Last.value

## Details

The value of a top-level assignment is put in .Last.value, unlike S.
Do not assign to .Last.value in the workspace, because this will always mask the object of the same name in package:base.

## See Also

eval

## Examples

```
## These will not work correctly from example(),
## but they will in make check or if pasted in,
## as example() does not run them at the top level
gamma(1:15) # think of some intensive calculation...
fac14 <- .Last.value # keep them
library("eda") # returns invisibly
.Last.value # shows what library(.) above returned
```

layout Specifying Complex Plot Arrangements

## Description

layout divides the device up into as many rows and columns as there are in matrix mat, with the column-widths and the row-heights specified in the respective arguments.

## Usage

```
layout(mat,
    widths = rep(1, dim(mat)[2]),
    heights= rep(1, dim(mat)[1]),
    respect= FALSE)
layout.show(n = 1)
lcm(x)
```


## Arguments

```
mat a matrix object specifying the location of the next N figures on the output device. Each value in the matrix must be 0 or a positive integer. If \(N\) is the largest positive integer in the matrix, then the integers \(\{1, \ldots, N-1\}\) must also appear at least once in the matrix.
widths a vector of values for the widths of columns on the device. Relative widths are specified with numeric values. Absolute widths (in centimetres) are specified with the 1 cm () function (see examples).
heights a vector of values for the heights of rows on the device. Relative and absolute heights can be specified, see widths above.
respect either a logical value or a matrix object. If the latter, then it must have the same dimensions as mat and each value in the matrix must be either 0 or 1.
n
number of figures to plot.
x
```


## Details

Figure $i$ is allocated a region composed from a subset of these rows and columns, based on the rows and columns in which $i$ occurs in mat.

The respect argument controls whether a unit column-width is the same physical measurement on the device as a unit row-height.
layout. show ( n ) plots (part of) the current layout, namely the outlines of the next n figures. $l \mathrm{~cm}$ is a trivial function, to be used as the interface for specifying absolute dimensions for the widths and heights arguments of layout().

## Value

layout returns the number of figures, $N$, see above.

## Author(s)

Paul R. Murrell

## References

Murrell, P. R. (1999) Layouts: A mechanism for arranging plots on a page. Journal of Computational and Graphical Statistics, 8, 121-134. Chapter 5 of Paul Murrell's Ph.D. thesis.

## See Also

par with arguments mfrow, mfcol, or mfg.

## Examples

```
def.par <- par(no.readonly = TRUE)# save default, for resetting...
## divide the device into two rows and two columns
## allocate figure 1 all of row 1
## allocate figure 2 the intersection of column 2 and row 2
layout(matrix(c(1,1,0,2), 2, 2, byrow = TRUE))
```

```
## show the regions that have been allocated to each plot
layout.show(2)
## divide device into two rows and two columns
## allocate figure 1 and figure 2 as above
## respect relations between widths and heights
nf <- layout(matrix(c(1,1,0,2), 2, 2, byrow=TRUE), respect=TRUE)
layout.show(nf)
## create single figure which is 5cm square
nf <- layout(matrix(1), widths=lcm(5), heights=lcm(5))
layout.show(nf)
##-- Create a scatterplot with marginal histograms -----
x <- pmin(3, pmax(-3, rnorm(50)))
y <- pmin(3, pmax(-3, rnorm(50)))
xhist <- hist(x, breaks=seq(-3,3,0.5), plot=FALSE)
yhist <- hist(y, breaks=seq(-3,3,0.5), plot=FALSE)
top <- max(c(xhist$counts, yhist$counts))
xrange <- c(-3,3)
yrange <- c(-3,3)
nf <- layout(matrix(c(2,0,1,3),2,2,byrow=TRUE), c(3,1), c(1,3), TRUE)
layout.show(nf)
par(mar=c (3,3,1,1))
plot(x, y, xlim=xrange, ylim=yrange, xlab="", ylab="")
par(mar=c (0,3,1,1))
barplot(xhist$counts, axes=FALSE, ylim=c(0, top), space=0)
par(mar=c(3,0,1,1))
barplot(yhist$counts, axes=FALSE, xlim=c(0, top), space=0, horiz=TRUE)
par(def.par)#- reset to default
```

legend Add Legends to Plots

## Description

This function can be used to add legends to plots. Note that a call to the function locator can be used in place of the x and y arguments.

## Usage

```
legend(x, y, legend, fill, col = "black", lty, lwd, pch,
    angle = NULL, density = NULL,
    bty = "o", bg = par("bg"), pt.bg = NA, cex = 1,
    xjust = 0, yjust = 1, x.intersp = 1, y.intersp = 1, adj = 0,
    text.width = NULL, merge = do.lines && has.pch, trace = FALSE,
    ncol = 1, horiz = FALSE)
```


## Arguments

## $\mathrm{x}, \mathrm{y}$

fill if specified, this argument will cause boxes filled with the specified colors to appear beside the legend text.
col the color of points or lines appearing in the legend.
lty, lwd the line types and widths for lines appearing in the legend. One of these two must be specified for line drawing.
pch the plotting symbols appearing in the legend, either as vector of 1character strings, or one (multi character) string. Must be specified for symbol drawing.
angle
density
bty
bg
pt.bg
cex
xjust
yjust
x.intersp
y.intersp
adj
text.width the width of the legend text in $x$ ("user") coordinates. Defaults to the proper value computed by strwidth(legend).
merge logical; if TRUE, "merge" points and lines but not filled boxes. Defaults to TRUE if there are points and lines.
trace logical; if TRUE, shows how legend does all its magical computations.
ncol the number of columns in which to set the legend items (default is 1 , a vertical legend).
horiz logical; if TRUE, set the legend horizontally rather than vertically (specifying horiz overrides the ncol specification).

## Details

"Attribute" arguments such as col, pch, lty, etc, are recycled if necessary. merge is not.
Points are drawn after lines in order that they can cover the line with their background color pt.bg, if applicable.

## Value

A (invisible) list with list components

```
rect a list with components
    w,h positive numbers giving width and height of the legend's box.
    left,top x and y coordinates of upper left corner of the box.
text a list with components
    x,y numeric vectors of length length(legend), giving the x and y coor-
    dinates of the legend's text(s).
```


## See Also

plot, barplot which uses legend(), and text for more examples of math expressions.

## Examples

```
## Run the example in '?matplot' or the following:
leg.txt <- c("Setosa Petals", "Setosa Sepals",
            "Versicolor Petals", "Versicolor Sepals")
y.leg <- c(4.5, 3, 2.1, 1.4, .7)
cexv <- c(1.2, 1, 4/5, 2/3, 1/2)
matplot(c(1,8), c(0,4.5), type = "n", xlab = "Length", ylab = "Width",
    main = "Petal and Sepal Dimensions in Iris Blossoms")
for (i in seq(cexv)) {
    text (1, y.leg[i]-.1, paste("cex=",formatC(cexv[i])), cex=.8, adj = 0)
    legend(3, y.leg[i], leg.txt, pch = "sSvV", col = c(1, 3), cex = cexv[i])
}
## 'merge = TRUE' for merging lines & points:
x <- seq(-pi, pi, len = 65)
plot(x, sin(x), type = "l", ylim = c(-1.2, 1.8), col = 3, lty = 2)
points(x, cos(x), pch = 3, col = 4)
lines(x, tan(x), type = "b", lty = 1, pch = 4, col = 6)
title("legend(..., lty = c(2, -1, 1), pch = c(-1,3,4), merge = TRUE)",
    cex.main = 1.1)
legend(-1, 1.9, c("sin", "cos", "tan"), col = c(3,4,6),
                lty = c(2, -1, 1), pch = c(-1, 3, 4), merge = TRUE, bg='gray90')
##--- log scaled Examples
leg.txt <- c("a one", "a two")
par(mfrow = c(2,2))
for(ll in c("","x","y","xy")) {
    plot(2:10, log=ll, main=paste("log = '",ll,")", sep=""))
    abline(1,1)
    lines(2:3,3:4, col=2) #
    points(2,2, col=3) #
    rect(2,3,3,2, col=4)
    text(c(3,3),2:3, c("rect(2,3,3,2, col=4)",
                                    "text(c(3,3),2:3,\"c(rect(...)\")"), adj = c(0,.3))
    legend(list(x=2,y=8), legend = leg.txt, col=2:3, pch=1:2,
        lty=1, merge=TRUE)#, trace=TRUE)
}
par(mfrow=c(1,1))
##-- Math expressions:
```

```
plot(x, sin(x), type="l", col = 2,xlab=expression(phi),ylab=expression(f(phi)))
abline(h=-1:1, v=pi/2*(-6:6), col="gray90")
lines(x, cos(x), col = 3, lty = 2)
ex.cs1 <- expression(plain(sin) * phi, paste("cos", phi))# 2 ways
str(legend(-3, .9, ex.cs1, lty=1:2, col=2:3, adj = c(0, .6)))# adj y !
x <- rexp(100, rate = .5)
hist(x, main = "Mean and Median of a Skewed Distribution")
abline(v = mean(x), col=2, lty=2, lwd=2)
abline(v = median(x), col=3, lty=3, lwd=2)
ex12 <- expression(bar(x) == sum(over(x[i], n), i==1, n),
    hat(x) == median(x[i], i==1,n))
str(legend(4.1, 30, ex12, col = 2:3, lty=2:3, lwd=2))
## 'Filled' boxes -- for more, see example(plotfactor)
data(PlantGrowth)
plot(cut(weight, 3) ~ group, data = PlantGrowth,
        col = NULL, density = 16*(1:3))
## Using 'ncol' :
x <- 0:64/64
matplot(x, outer(x, 1:7, function(x, k) sin(k * pi * x)),
        type = "o", col = 1:7, ylim = c(-1, 1.5), pch = "*")
op <- par(bg="antiquewhite1")
legend(0, 1.5, paste("sin(",1:7,"pi * x)"), col=1:7, lty=1:7, pch = "*",
        ncol = 4, cex=.8)
legend(.8,1.2, paste("sin(",1:7,"pi * x)"), col=1:7, lty=1:7, pch = "*",cex=.8)
legend(0, -.1, paste("sin(",1:4,"pi * x)"), col=1:4, lty=1:4, ncol=2, cex=.8)
legend(0, -.4, paste("sin(",5:7,"pi * x)"), col=5:7, pch=24, ncol=2, cex=1.5,
        pt.bg="pink")
par(op)
## point covering line :
y <- sin(3*pi*x)
plot(x,y,type="l",col="blue", main = "points with bg & legend(*, pt.bg)")
points(x,y,pch=21,bg="white")
legend(.4,1,"sin(c x)",pch=21,pt.bg="white",lty=1, col = "blue")
```

length Length of a Vector or List

## Description

Get or set the length of vectors (including lists).

## Usage

length ( x )
length (x) <- n

## Arguments

x a vector or list.
n
an integer.

## Details

The replacement form can be used to reset the length of a vector. If a vector is shortened, extra values are discarded and when a vector is lengthened, it is padded out to its new length with NAs.

## Value

The length of x as an integer of length 1 , if x is (or can be coerced to) a vector or list. Otherwise, length returns NA.

## Examples

```
length(diag(4))# = 16 (4 x 4)
length(options())# 12 or more
length(y ~ x1 + x2 + x3)# 3
length(expression(x, {y <- x^2; y+2}, x^y)) # 3
```

levels Levels Attributes

## Description

levels provides access to the levels attribute of a variable. The first form returns the value of the levels of its argument and the second sets the attribute.

The assignment form ("levels<-") of levels is a generic function and new methods can be written for it. The most important method is that for factors:

## Usage

```
levels(x)
levels(x) <- value
```


## Arguments

x
an object, for example a factor.

See Also
levels<-.factor, nlevels.
levels.factor Factor Levels Assignment

## Description

levels<- provides a way to alter the levels attribute of factor. value can be a vector of character strings with length at least the the number of levels of $x$, or a named list specifying how to rename the levels.

## Usage <br> levels(x) <- value

See Also
factor, levels, levels<-, nlevels.

## Examples

```
# assign individual levels
x <- gl(2, 4, 8)
levels(x)[1] <- "low"
levels(x)[2] <- "high"
x
# or as a group
y <- gl(2, 4, 8)
levels(y) <- c("low", "high")
y
# combine some levels
z <- gl(3, 2, 12)
levels(z) <- c("A", "B", "A")
z
# same, using a named list
z <- gl(3, 2, 12)
levels(z) <- list(A=c(1,3), B=2)
z
# we can add levels this way:
f <- factor(c("a","b"))
levels(f) <- c("c", "a", "b")
f
f <- factor(c("a","b"))
levels(f) <- list(C="C", A="a", B="b")
f
```


## Description

library and require load add-on packages. .First.lib is called when a package is loaded; .Last.lib is called when a package is detached. .packages returns information about package availability. .path.package returns information about where a package was loaded from. .find.package returns the directory paths of installed packages.

## Usage

```
library(package, help, lib.loc = NULL, character.only = FALSE,
                logical.return = FALSE, warn.conflicts = TRUE,
            keep.source = getOption("keep.source.pkgs"),
            verbose = getOption("verbose"))
require(package, quietly = FALSE, warn.conflicts = TRUE,
            keep.source = getOption("keep.source.pkgs"))
    .First.lib(libname, pkgname)
    .Last.lib(libpath)
    .packages(all.available = FALSE, lib.loc = NULL)
    .path.package(package = .packages(), quiet = FALSE)
    .find.package(package, lib.loc = NULL, quiet = FALSE,
                            verbose = getOption("verbose"))
    .libPaths(new)
    .Library
    .Autoloaded
```


## Arguments

package, help name or character string giving the name of a package.
lib.loc a character vector describing the location of R library trees to search through, or NULL. The default value of NULL corresponds to all libraries currently known.
character.only
a logical indicating whether package or help can be assumed to be character strings.
logical.return
logical. If it is TRUE, FALSE or TRUE is returned to indicate success.
warn.conflicts
logical. If TRUE, warnings are printed about conflicts from attaching the new package, unless that package contains an object .conflicts.OK.
keep.source logical. If TRUE, functions "keep their source" including comments, see argument keep.source to options.
verbose a logical. If TRUE, additional diagnostics are printed.
quietly a logical. If TRUE, no message confirming package loading is printed.

| libname | a character string giving the library directory where the package was <br> found. |
| :--- | :--- |
| pkgname | a character string giving the name of the package. |
| libpath | a character string giving the complete path to the package. |
| all.available | logical; if TRUE return a character vector of all available packages in <br> lib.loc. |
| quiet | logical. For .path.package, should this not give warnings or an error if <br> the package(s) are not loaded? For .find.package, should this not give <br> warnings or an error if the package(s) are not found? |
| new | a character vector with the locations of R library trees. |

## Details

library (package) and require(package) both load the package with name package. require is designed for use inside other functions; it returns FALSE and gives a warning (rather than an error as library ()) if the package does not exist. Both functions check and update the list of currently loaded packages and do not reload code that is already loaded.
For large packages, setting keep.source = FALSE may save quite a bit of memory.
If library is called with no package or help argument, it lists all available packages in the libraries specified by lib.loc, and returns the corresponding information in an object of class "libraryIQR". The structure of this class may change in future versions. In earlier versions of $R$, only the names of all available packages were returned; use . packages (all $=$ TRUE) for obtaining these.
library (help $=$ somename) computes basic information about the package somename, and returns this in an object of class "packageInfo". The structure of this class may change in future versions.
.First.lib is called when a package is loaded by library. It is called with two arguments, the name of the library directory where the package was found (i.e., the corresponding element of lib.loc), and the name of the package (in that order). It is a good place to put calls to library.dynam which are needed when loading a package into this function (don't call library.dynam directly, as this will not work if the package is not installed in a "standard" location). .First.lib is invoked after the search path interrogated by search() has been updated, so as.environment(match("package:name"), search()) will return the environment in which the package is stored. If calling .First.lib gives an error the loading of the package is abandoned, and the package will be unavailable. Similarly, if the option ".First.lib" has a list element with the package's name, this element is called in the same manner as .First.lib when the package is loaded. This mechanism allows the user to set package "load hooks" in addition to startup code as provided by the package maintainers.
.Last.lib is called when a package is detached. Beware that it might be called if .First.lib has failed, so it should be written defensively. (It is called within try, so errors will not stop the package being detached.)
.packages() returns the "base names" of the currently attached packages invisibly whereas . packages (all.available $=$ TRUE) gives (visibly) all packages available in the library location path lib.loc.
.path.package returns the paths from which the named packages were loaded, or if none were named, for all currently loaded packages. Unless quiet $=$ TRUE it will warn if some of the packages named are not loaded, and given an error if none are. This function is not meant to be called by users, and its interface might change in future versions.
.find.package returns the paths to the locations where the given packages can be found. If lib.loc is NULL, then then attached packages are searched before the libraries. If a package is found more than once, the first match is used. Unless quiet = TRUE a warning will be given about the named packages which are not found, and an error if none are. If verbose is true, warnings about packages found more than once are given. This function is not meant to be called by users, and its interface might change in future versions.
.Autoloaded contains the "base names" of the packages for which autoloading has been promised.
.Library is a character string giving the location of the default library, the 'library' subdirectory of R_HOME. .libPaths is used for getting or setting the library trees that R knows about (and hence uses when looking for packages). If called with argument new, the library search path is set to unique(new, .Library) and this is returned. If given no argument, a character vector with the currently known library trees is returned. The library search path is initialized at startup from the environment variable R_LIBS (which should be a colon-separated list of directories at which R library trees are rooted) by calling .libPaths with the directories specified in R_LIBS.

## Value

library returns the list of loaded (or available) packages (or TRUE if logical.return is TRUE). require returns a logical indicating whether the required package is available.

## Note

library and require can only load an installed package, and this is detected by having a 'DESCRIPTION' file containing a Built: field. Packages installed prior to 1.2.0 (released in December 2000) will need to be re-installed.
Under Unix-alikes, the code checks that the package was installed under a similar operating system as given by .Platform\$canonical. name.

## Author(s)

R core; Guido Masarotto for the all.available=TRUE part of .packages.

## See Also

attach, detach, search, objects, autoload, library.dynam, data, install.packages, INSTALL, REMOVE.

## Examples

```
(.packages()) # maybe just "base"
.packages(all = TRUE) # return all available as character vector
library() # list all available packages
library(lib = .Library) # list all packages in the default library
library(help = eda) # documentation on package 'eda'
library(eda) # load package 'eda'
require(eda) # the same
(.packages()) # "eda", too
detach("package:eda")
# if the package name is in a character vector, use
pkg <- "eda"
library(pkg, character.only = TRUE)
```

```
detach(pos = match(paste("package", pkg, sep=":"), search()))
    .path.package()
    .Autoloaded # maybe "ctest"
.libPaths() # all library trees R knows about
require(nonexistent) # FALSE
## Suppose a package needs to call a shared library named 'fooEXT',
## where 'EXT' is the system-specific extension. Then you should use
    .First.lib <- function(lib, pkg) {
    library.dynam("foo", pkg, lib)
}
```

library.dynam Loading Shared Libraries

## Description

Load the specified file of compiled code if it has not been loaded already.

## Usage

```
library.dynam(chname, package = .packages(), lib.loc = NULL,
    verbose = getOption("verbose"), file.ext, ...)
.dynLibs(new)
```


## Arguments

| chname | a character string naming a shared library to load. |
| :--- | :--- |
| package | a character vector with the names of packages to search through. <br> a character vector describing the location of R library trees to search <br> through, or NULL. The default value of NULL corresponds to all libraries <br> currently known. |
| verbose | a logical value indicating whether an announcement is printed on the <br> console before loading the shared library. The default value is taken from <br> the verbose entry in the system options. |
| file.ext | the extension to append to the file name to specify the library to be <br> loaded. This defaults to the appropriate value for the operating system. <br> additional arguments needed by some libraries that are passed to the call <br> to dyn.load to control how the library is loaded. |
| new | a character vector of packages which have loaded shared libraries. |

## Details

This is designed to be used inside a package rather than at the command line, and should really only be used inside .First.lib(). The system-specific extension for shared libraries (e.g., '.so' or '.sl' on Unix systems) should not be added.
.dynLibs is used for getting or setting the packages that have loaded shared libraries (using library.dynam). Versions of R prior to 1.6 .0 used an internal global variable .Dyn.libs for storing this information: this variable is now defunct.

## Value

library.dynam returns a character vector with the names of packages which have used it in the current R session to load shared libraries. This vector is returned as invisible, unless the chname argument is missing.

## See Also

.First.lib, library, dyn.load, .packages, .libPaths
SHLIB for how to create suitable shared libraries.

## Examples

```
library.dynam()# which packages have been '(dynamically loaded''
```

license The $R$ License Terms

## Description

The license terms under which R is distributed.

## Usage

license()
licence()

## Details

R is distributed under the terms of the GNU GENERAL PUBLIC LICENSE Version 2, June 1991. A copy of this license is in '\$R_HOME/COPYING'.
A small number of files (the API header files and import library) are distributed under the LESSER GNU GENERAL PUBLIC LICENSE version 2.1. A copy of this license is in '\$R_HOME/COPYING.LIB'.

```
LifeCycleSavings Intercountry Life-Cycle Savings Data
```


## Description

Data on the savings ratio 1960-1970.

## Usage

data(LifeCycleSavings)

## Format

A data frame with 50 observations on 5 variables.

| $[, 1]$ | sr | numeric | aggregate personal savings |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | pop15 | numeric | \% of population under 15 |
| $[, 3]$ | pop75 | numeric | \% of population over 75 |
| $[, 4]$ | dpi | numeric | real per-capita disposable income |
| $[, 5]$ | ddpi | numeric | \% growth rate of dpi |

## Details

Under the life-cycle savings hypothesis as developed by Franco Modigliani, the savings ratio (aggregate personal saving divided by disposable income) is explained by per-capita disposable income, the percentage rate of change in per-capita disposable income, and two demographic variables: the percentage of population less than 15 years old and the percentage of the population over 75 years old. The data are averaged over the decade 1960-1970 to remove the business cycle or other short-term fluctuations.

## Source

The data were obtained from Belsley, Kuh and Welsch (1980). They in turn obtained the data from Sterling (1977).

## References

Sterling, Arnie (1977) Unpublished BS Thesis. Massachusetts Institute of Technology.
Belsley, D. A., E. Kuh. E. and Welsch, R. E. (1980) Regression Diagnostics. New York: Wiley.

## Examples

```
data(LifeCycleSavings)
pairs(LifeCycleSavings, panel = panel.smooth,
    main = "LifeCycleSavings data")
fm1 <- lm(sr ~ pop15 + pop75 + dpi + ddpi, data = LifeCycleSavings)
summary(fm1)
```

lines Add Connected Line Segments to a Plot

## Description

A generic function taking coordinates given in various ways and joining the corresponding points with line segments.

## Usage

lines( $x, \ldots$ )
lines.default(x, y=NULL, type="l", col=par("col"), lty=par("lty"), ...)

## Arguments

$\mathrm{x}, \mathrm{y} \quad$ coordinate vectors of points to join.
type character indicating the type of plotting; actually any of the types as in plot.
col color to use.
lty line type to use.
... Further graphical parameters (see par) may also be supplied as arguments, particularly, line type, lty and line width, 1wd.

## Details

The coordinates can be passed to lines in a plotting structure (a list with x and y components), a time series, etc. See xy.coords.

The coordinates can contain NA values. If a point contains NA it either its x or y value, it is omitted from the plot, and lines are not drawn to or from such points. Thus missing values can be used to achieve breaks in lines.

## See Also

points, plot, and the underlying "primitive" plot.xy.
par for how to specify colors.

## Examples

data(cars)
\# draw a smooth line through a scatter plot
plot(cars, main="Stopping Distance versus Speed")
lines(lowess(cars))

## LINK

Create Executable Programs

## Description

Front-end for creating executable programs.

## Usage

R CMD LINK [options] linkcmd

## Arguments

linkcmd a list of commands to link together suitable object files (include library objects) to create the executable program.
options further options to control the linking, or for obtaining information about usage and version.

## Details

The linker front-end is useful in particular when linking against the R shared library, in which case linkcmd must contain -lR but need not specify its library path.

Currently only works if the C compiler is used for linking, and no $\mathrm{C}++$ code is used.
Use R CMD LINK --help for more usage information.
list Lists - Generic and Dotted Pairs

## Description

Functions to construct, coerce and check for all kinds of R lists.

## Usage

```
list(...)
pairlist(...)
as.list(x, ...)
as.list.default(x, ...)
as.pairlist(x)
is.list(x)
is.pairlist(x)
alist(...)
```


## Arguments

```
... objects.
x
object to be coerced or tested.
```


## Details

Most lists in R internally are Generic Vectors, whereas traditional dotted pair lists (as in LISP) are still available.

The arguments to list or pairlist are of the form value or tag=value. The functions return a list composed of its arguments with each value either tagged or untagged, depending on how the argument was specified.
alist is like list, except in the handling of tagged arguments with no value. These are handled as if they described function arguments with no default (cf. formals), whereas list simply ignores them.
as.list attempts to coerce its argument to list type. For functions, this returns the concatenation of the list of formals arguments and the function body. For expressions, the list of constituent calls is returned.
is.list returns TRUE iff its argument is a list or a pairlist of length> 0, whereas is. pairlist only returns TRUE in the latter case.

An empty pairlist, pairlist() is the same as NULL. This is different from list().

## See Also

vector(., mode="list"), c, for concatenation; formals.

## Examples

```
data(cars)
# create a plotting structure
pts <- list(x=cars[,1], y=cars[,2])
plot(pts)
# Argument lists
f <- function()x
# Note the specification of a "..." argument:
formals(f) <- al <- alist(x=, y=2, ...=)
f
str(al)
str(pl <- as.pairlist(ps.options()))
## These are all TRUE:
is.list(pl) && is.pairlist(pl)
!is.null(list())
is.null(pairlist())
!is.list(NULL)
is.pairlist(pairlist())
is.null(as.pairlist(list()))
is.null(as.pairlist(NULL))
```

list.files List the Files in a Directory/Folder

## Description

This function produces a list containing the names of files in the named directory. dir is an alias.

## Usage

```
list.files(path = ".", pattern=NULL, all.files=FALSE, full.names=FALSE)
    dir(path = ".", pattern=NULL, all.files=FALSE, full.names=FALSE)
```


## Arguments

| path | a character vector of full path names. |
| :--- | :--- |
| pattern | an optional regular expression. Only file names which match the regular <br> expression will be returned. |
| all.files | a logical value. If FALSE, only the names of visible files are returned. If <br> TRUE, all file names will be returned. |
| full.names | a logical value. If TRUE, the directory path is prepended to the file names. <br> If FALSE, only the file names are returned. |

## Value

A character vector containing the names of the files in the specified directories, or " " if there were no files. If a path does not exist or is not a directory or is unreadable it is skipped, with a warning.
The files are sorted in alphabetical order, on the full path if full.names = TRUE.

## Note

File naming conventions are very platform dependent.

## Author(s)

Ross Ihaka

## See Also

file.info, file.access and files for many more file handling functions.

## Examples

```
list.files(R.home())
## Only files starting with a-l or r (*including* uppercase):
dir("../..", pattern = "^[a-lr]",full.names=TRUE)
```


## lm Fitting Linear Models

## Description

lm is used to fit linear models. It can be used to carry out regression, single stratum analysis of variance and analysis of covariance (although aov may provide a more convenient interface for these).

## Usage

```
lm(formula, data, subset, weights, na.action,
    method = "qr", model = TRUE, x = FALSE, y = FALSE, qr = TRUE,
    singular.ok = TRUE, contrasts = NULL, offset = NULL, ...)
```


## Arguments

formula a symbolic description of the model to be fit. The details of model specification are given below.
data an optional data frame containing the variables in the model. By default the variables are taken from environment (formula), typically the environment from which lm is called.
subset an optional vector specifying a subset of observations to be used in the fitting process.
weights an optional vector of weights to be used in the fitting process. If specified, weighted least squares is used with weights weights (that is, minimizing $\operatorname{sum}\left(w^{*} e^{\wedge} 2\right)$ ); otherwise ordinary least squares is used.
na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The "factory-fresh" default is na.omit.
method the method to be used; for fitting, currently only method="qr" is supported; method="model.frame" returns the model frame (the same as with model = TRUE, see below).
model, $x, y, q r$
logicals. If TRUE the corresponding components of the fit (the model frame, the model matrix, the response, the QR decomposition) are returned.
singular.ok logical, defaulting to TRUE. FALSE is not yet implemented.
contrasts an optional list. See the contrasts.arg of model.matrix.default.
offset this can be used to specify an a priori known component to be included in the linear predictor during fitting. An offset term can be included in the formula instead or as well, and if both are specified their sum is used.
... additional arguments to be passed to the low level regression fitting functions (see below).

## Details

Models for 1 m are specified symbolically. A typical model has the form response ~ terms where response is the (numeric) response vector and terms is a series of terms which specifies a linear predictor for response. A terms specification of the form first + second indicates all the terms in first together with all the terms in second with duplicates removed. A specification of the form first: second indicates the set of terms obtained by taking the interactions of all terms in first with all terms in second. The specification first*second indicates the cross of first and second. This is the same as first + second + first:second.
lm calls the lower level functions lm.fit, etc, see below, for the actual numerical computations. For programming only, you may consider doing likewise.

## Value

lm returns an object of class "lm" or for multiple responses of class c("mlm", "lm").
The functions summary and anova are used to obtain and print a summary and analysis of variance table of the results. The generic accessor functions coefficients, effects, fitted.values and residuals extract various useful features of the value returned by lm.
An object of class "lm" is a list containing at least the following components:

```
coefficients a named vector of coefficients
residuals the residuals, that is response minus fitted values.
fitted.values the fitted mean values.
rank the numeric rank of the fitted linear model.
weights (only for weighted fits) the specified weights.
df.residual the residual degrees of freedom.
call the matched call.
terms the terms object used.
contrasts (only where relevant) the contrasts used.
xlevels (only where relevant) a record of the levels of the factors used in fitting.
y if requested, the response used.
x if requested, the model matrix used.
model if requested (the default), the model frame used.
```

In addition, non-null fits will have components assign, effects and (unless not requested) qr relating to the linear fit, for use by extractor functions such as summary and effects.

## Note

Offsets specified by offset will not be included in predictions by predict.lm, whereas those specified by an offset term in the formula will be.

## See Also

summary.lm for summaries and anova.lm for the ANOVA table; aov for a different interface. The generic functions coefficients, effects, residuals, fitted.values. predict.lm (via predict) for prediction, including confidence and prediction intervals. lm.influence for regression diagnostics, and glm for generalized linear models.
The underlying low level functions, lm.fit for plain, and lm.wfit for weighted regression fitting.

## Examples

```
## Annette Dobson (1990) "An Introduction to Generalized Linear Models".
## Page 9: Plant Weight Data.
ctl <- c(4.17,5.58,5.18,6.11,4.50,4.61,5.17,4.53,5.33,5.14)
trt <- c(4.81,4.17,4.41,3.59,5.87,3.83,6.03,4.89,4.32,4.69)
group <- gl(2,10,20, labels=c("Ctl","Trt"))
weight <- c(ctl, trt)
anova(lm.D9 <- lm(weight ~ group))
summary(lm.D90 <- lm(weight ~ group - 1))# omitting intercept
summary(resid(lm.D9) - resid(lm.D90)) #- residuals almost identical
opar <- par(mfrow = c(2,2), oma = c(0, 0, 1.1, 0))
plot(lm.D9, las = 1) # Residuals, Fitted, ...
par(opar)
## model frame :
stopifnot(identical(lm(weight ~ group, method = "model.frame"),
model.frame(lm.D9)))
```

lm.fit Fitter Functions for Linear Models

## Description

These are the basic computing engines called by lm used to fit linear models. These should usually not be used directly unless by experienced users.

## Usage

```
lm.fit (x, y, offset = NULL, method = "qr", tol = 1e-7, ...)
lm.wfit(x, y, w, offset = NULL, method = "qr", tol = 1e-7, ...)
lm.fit.null (x, y, method = "qr", tol = 1e-7, ...)
lm.wfit.null(x, y, w, method = "qr", tol = 1e-7, ...)
```


## Arguments

$\mathrm{x} \quad$ design matrix of dimension $\mathrm{n} * \mathrm{p}$.
$y \quad$ vector of observations of length $n$.
$\mathrm{w} \quad$ vector of weights (length n ) to be used in the fitting process for the wfit functions. Weighted least squares is used with weights w, i.e., sum(w * $e^{\wedge} 2$ ) is minimized.
offset numeric of length n). This can be used to specify an a priori known component to be included in the linear predictor during fitting.
method currently, only method="qr" is supported.
tol tolerance for the qr decomposition. Default is $1 \mathrm{e}-7$.
... currently disregarded.

## Details

The functions lm. $\{\mathrm{w}\}$ fit.null are called by lm.fit or lm.wfit respectively, when x has zero columns.

## Value

a list with components

| coefficients | $p$ vector |
| :--- | :--- |
| residuals | $n$ vector |
| fitted.values | $n$ vector |
| effects | $n$ vector; ...... |
| weights | $n$ vector - only for the *wfit* functions. |
| rank | integer, giving the rank |
| df.residual | degrees of freedom of residuals |
| qr | the QR decomposition, see qr. |

See Also
lm which you should use for linear least squares regression, unless you know better.

## Examples

```
set.seed(129)
n <- 7 ; p <- 2
X <- matrix(rnorm(n * p), n,p) # no intercept!
y <- rnorm(n)
w <- rnorm(n)^2
str(lmw <- lm.wfit(x=X, y=y, w=w))
str(lm. <- lm.fit (x=X, y=y))
str(lm0 <- lm.fit.null (x=X, y=y))
str(lmw0 <- lm.wfit.null(x=X, y=y,w=w))
```

lm.influence Regression Diagnostics

## Description

This function provides the basic quantities which are used in forming a wide variety of diagnostics for checking the quality of regression fits.

## Usage

lm.influence(lm.obj)

## Arguments

lm.obj an object as returned by lm.

## Details

The influence.measures() and other functions listed in See Also provide a more user oriented way of computing a variety of regression diagnostics.

## Value

A list containing the following components:
hat a vector containing the diagonal of the "hat" matrix.
coefficients the change in the estimated coefficients which results when the i-th case is dropped from the regression is contained in the i-th row of this matrix.
sigma a vector whose i-th element contains the estimate of the residual standard deviation obtained when the i-th case is dropped from the regression.

## Note

The coefficients returned by the $R$ version of $1 m$.influence differ from those computed by S. Rather than returning the coefficients which result from dropping each case, we return the changes in the coefficients. This is more directly useful in many diagnostic measures.
Note that cases with weights $==0$ are dropped (contrary to the situation in S).
if a model has been fitted with na.action=na.exclude (see na.exclude), cases excluded in the fit are considered here.

## References

Belsley, D. A., Kuh, E. and Welsch, R. E. (1980) Regression Diagnostics. New York: Wiley.

```
See Also
    summary.lm for summary and related methods;
    influence.measures,
    hat for the hat matrix diagonals,
    dfbetas, dffits, covratio, cooks.distance, lm.
```


## Examples

```
## Analysis of the life-cycle savings data
## given in Belsley, Kuh and Welsch.
data(LifeCycleSavings)
summary(lm.SR <- lm(sr ~ pop15 + pop75 + dpi + ddpi,
                                    data = LifeCycleSavings),
    corr = TRUE)
str(lmI <- lm.influence(lm.SR))
## For more 'user level' examples, use example(influence.measures)
```

lm.summaries Accessing Linear Model Fits

## Description

All these functions are methods for class "lm" objects.

## Usage

```
coefficients(object, ...) ; \method
```


## Arguments

object, $x \quad$ an object of class lm, usually, a result of a call to lm.
... further arguments passed to or from other methods.

## Details

The generic accessor functions coefficients, effects, fitted.values and residuals can be used to extract various useful features of the value returned by lm.

## See Also

The model fitting function lm, anova. 1m.
coefficients, deviance, df.residual, effects, fitted.values, glm for generalized linear models, lm.influence for regression diagnostics, weighted.residuals, residuals, residuals.glm, summary.lm.

## Examples

```
##-- Continuing the lm(.) example:
coef(lm.D90)# the bare coefficients
## The 2 basic regression diagnostic plots [plot.lm(.) is preferred]
plot(resid(lm.D90), fitted(lm.D90))# Tukey-Anscombe's
abline(h=0, lty=2, col = 'gray')
qqnorm(residuals(lm.D90))
```

lm.summary Summarizing Linear Model Fits

## Description

summary method for class "lm".

## Usage

```
summary(object, correlation = FALSE, ...)
print(x, digits = max(3, getOption("digits") - 3),
    symbolic.cor = p > 4,
    signif.stars = getOption("show.signif.stars"), ...)
```


## Arguments

| object | an object of class "lm", usually, a result of a call to lm. |
| :--- | :--- |
| x | an object of class "summary.lm", usually, a result of a call to summary.lm. <br> correlation |
| logical; if TRUE, the correlation matrix of the estimated parameters is <br> returned and printed. |  |
| digits | the number of significant digits to use when printing. |
| symbolic.cor | logical. If TRUE, print the correlations in a symbolic form (see symnum <br> rather than as numbers. |
| signif.stars | logical. If TRUE, "significance stars" are printed for each coefficient. <br> further arguments passed to or from other methods. |

## Details

print.summary.lm tries to be smart about formatting the coefficients, standard errors, etc. and additionally gives "significance stars" if signif.stars is TRUE.

## Value

The function summary.lm computes and returns a list of summary statistics of the fitted linear model given in object, using the components (list elements) "call" and "terms" from its argument, plus

| residuals | the weighted residuals, the usual residuals rescaled by the square root of <br> the weights specified in the call to lm. |
| :--- | :--- |
| coefficients | a $p \times 4$ matrix with columns for the estimated coefficient, its standard <br> error, t-statistic and corresponding (two-sided) p-value. |
| sigma | the square root of the estimated variance of the random error |

$$
\hat{\sigma}^{2}=\frac{1}{n-p} \sum_{i} R_{i}^{2}
$$

where $R_{i}$ is the $i$-th residual, residuals [i].
df
degrees of freedom, a 3 -vector $(p, n-p, p *)$.
fstatistic (for models including non-intercept terms) a 3-vector with the value of the F-statistic with its numerator and denominator degrees of freedom.
r.squared
$R^{2}$, the "fraction of variance explained by the model",

$$
R^{2}=1-\frac{\sum_{i} R_{i}^{2}}{\sum_{i}\left(y_{i}-y^{*}\right)^{2}},
$$

where $y^{*}$ is the mean of $y_{i}$ if there is an intercept and zero otherwise.
adj.r.squared the above $R^{2}$ statistic "adjusted", penalizing for higher $p$.
cov.unscaled a $p \times p$ matrix of (unscaled) covariances of the $\hat{\beta}_{j}, j=1, \ldots, p$.
correlation the correlation matrix corresponding to the above cov.unscaled, if correlation $=$ TRUE is specified.

## See Also

The model fitting function lm, summary.

## Examples

```
##-- Continuing the lm(.) example:
coef(lm.D90)# the bare coefficients
sld90 <- summary(lm.D90 <- lm(weight ~ group -1))# omitting intercept
sld90
coef(sld90)# much more
```

load Reload Saved Datasets

## Description

Reload the datasets written to a file with the function save.

## Usage

load(file, envir = parent.frame())

## Arguments

$$
\begin{array}{ll}
\text { file } & \text { a connection or a character string giving the name of the file to load. } \\
\text { envir } & \text { the environment where the data should be loaded. }
\end{array}
$$

## See Also

save.

## Examples

```
## save all data
save(list = ls(), file= "all.Rdata")
## restore the saved values to the current environment
load("all.Rdata")
## restore the saved values to the workspace
load("all.Rdata", .GlobalEnv)
```

localeconv Find Details of the Numerical Representations in the Current Locale

## Description

Get details of the numerical representations in the current locale.

## Usage

Sys.localeconv()

## Value

A character vector with 18 named components. See your ISO C documentation for details of the meaning.
It is possible to compile R without support for locales, in which case the value will be NULL.

## See Also

locales for ways to set locales: by default R uses the C clocal for "LC_NUMERIC" and "LC_MONETARY".

## Examples

```
Sys.localeconv()
## The results in the default C locale are
## decimal_point thousands_sep grouping int_curr_symbol
## "." "" "" ""
## currency_symbol mon_decimal_point mon_thousands_sep mon_grouping
##
## positive_sign regative_sign 
## p_cs_precedes p_sep_by_space n_cs_precedes n_sep_by_space
\#\# "127" "127" "127" "127"
## p_sign_posn n_sign_posn
## "127" "127"
## Now try your default locale (which might be "C").
old <- Sys.getlocale()
Sys.setlocale(locale = "")
Sys.localeconv()
Sys.setlocale(locale = old)
read.table("foo", dec=Sys.localeconv()["decimal_point"])
```

locales Query or Set Aspects of the Locale

## Description

Get details of or set aspects of the locale for the R process.

## Usage

```
Sys.getlocale(category = "LC_ALL")
Sys.setlocale(category = "LC_ALL", locale = "")
```


## Arguments

```
category character string. Must be one of "LC_ALL", "LC_COLLATE", "LC_CTYPE",
    "LC_MONETARY", "LC_NUMERIC" or "LC_TIME".
locale character string. A valid locale name on the system in use. Normally ""
    (the default) will pick up the default locale for the system.
```


## Details

The locale describes aspects of the internationalization of a program. Initially most aspects of the locale of R are set to "C" (which is the default for the C language and reflects NorthAmerican usage). R does set "LC_CTYPE" and "LC_COLLATE", which allow the use of a different character set (typically ISO Latin 1) and alphabetic comparisons in that character set (including the use of sort) and "LC_TIME" may affect the behaviour of as.POSIXlt and strptime and functions which use them (but not date).
R can be built with no support for locales, but it is normally available on Unix and is available on Windows.
Some systems will have other locale categories, but the six described here are those specified by POSIX.

## Value

A character string of length one describing the locale in use (after setting for Sys.setlocale), or an empty character string if the locale is invalid (with a warning) or NULL if locale information is unavailable.
For category = "LC_ALL" the details of the string are system-specific: it might be a single locale or a set of locales separated by "/" (Solaris) or ";" (Windows). For portability, it is best to query categories individually. It is guaranteed that the result of foo <Sys.getlocale() can used in Sys.setlocale("LC_ALL", locale $=$ foo) on the same machine.

## Warning

Setting "LC_NUMERIC" can produce output that R cannot then read by scan or read.table with their default arguments, which are not locale-specific.

## See Also

strptime for uses of category = "LC_TIME". localeconv for details of numerical representations.

## Examples

```
Sys.getlocale()
Sys.getlocale("LC_TIME")
Sys.setlocale("LC_TIME", "de") # Solaris: details are OS-dependent
Sys.setlocale("LC_TIME", "German") # Windows
Sys.setlocale("LC_COLLATE", "C") # turn off locale-specific sorting
```

locator Graphical Input

## Description

Reads the position of the graphics cursor when the (first) mouse button is pressed.

## Usage

```
locator(n = 512, type = "n", ...)
```


## Arguments

$\mathrm{n} \quad$ the maximum number of points to locate.
type One of "n", "p", "l" or "o". If "p" or "o" the points are plotted; if "l" or "o" they are joined by lines.
... additional graphics parameters used if type $!=$ " $n$ " for plotting the locations.

## Details

Unless the process is terminated prematurely by the user (see below) at most n positions are determined.

The identification process can be terminated by pressing any mouse button other than the first.

The current graphics parameters apply just as if plot.default has been called with the same value of type. The plotting of the points and lines is subject to clipping, but locations outside the current clipping rectangle will be returned.
If the window is resized or hidden and then exposed before the input process has terminated, any lines or points drawn by locator will disappear. These will reappear once the input process has terminated and the window is resized or hidden and exposed again. This is because the points and lines drawn by locator are not recorded in the device's display list until the input process has terminated.

## Value

A list containing x and y components which are the coordinates of the identified points.

```
See Also
identify
```


## Description

$\log$ computes natural logarithms, $\log 10$ computes common (i.e., base 10) logarithms, and $\log 2$ computes binary (i.e., base 2) logarithms. The general form $\operatorname{logb}(x$, base) computes logarithms with base base (log10 and log2 are only special cases).
$\log 1 \mathrm{p}(\mathrm{x})$ computes $\log (1+x)$ accurately also for $|x| \ll 1$ (and less accurately when $x \approx-1$ ). exp computes the exponential function.
expm1 (x) computes $\exp (x)-1$ accurately also for $|x| \ll 1$.

## Usage

```
log(x, base = exp(1))
logb(x, base = exp(1))
log10(x)
log2(x)
exp(x)
expm1(x)
log1p(x)
```


## Arguments

| x | a numeric or complex vector. |
| :--- | :--- |
| base | positive number. The base with respect to which logarithms are com- <br> puted. Defaults to $e=\exp (1)$. |

## Value

A vector of the same length as x containing the transformed values. $\log (0)$ gives $-\operatorname{Inf}$ (when available).

## Note

$\log$ and $\operatorname{logb}$ are the same thing in R, but logb is preferred if base is specified, for S-PLUS compatibility.

## See Also

Trig, sqrt, Arithmetic.

## Examples

```
log(exp(3))
all.equal(log(1:10), log(1:10, exp(1)))
log10(30) == log(30, 10)
log10(1e7)# = 7
log2(2^pi) == 2^log2(pi)
Mod(pi - log(exp(pi*1i)) / 1i) < .Machine$double.eps
Mod(1+exp(pi*1i)) < .Machine$double.eps
```

```
    x <- 10^-(1+2*1:9)
    cbind(x, log(1+x), log1p(x), exp(x)-1, expm1(x))
```

Logic Logical Operators

## Description

These operators act on logical vectors.

## Usage

```
! x
x & y
x && y
x | y
x || y
xor(x, y)
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ logical vectors

## Details

! indicates logical negation (NOT).
\& and \&\& indicate logical AND and \| and \| I indicate logical OR. The shorter form performs elementwise comparisons in much the same way as arithmetic operators. The longer form evaluates left to right examining only the first element of each vector. Evaluation proceeds only until the result is determined. The longer form is appropriate for programming controlflow and typically preferred in if clauses.
xor indicates elementwise exclusive OR.
NA is a valid logical object. Where a component of x or y is NA, the result will be NA if the outcome is ambiguous. In other words NA \& TRUE evaluates to NA, but NA \& FALSE evaluates to FALSE. See the examples below.

## See Also

TRUE or logical.
Syntax for operator precedence.

## Examples

```
y <- 1 + (x <- rpois(50, lambda=1.5) / 4 - 1)
x[(x > 0) & (x < 1)] # all x values between 0 and 1
if (any(x == 0) || any(y == 0)) "zero encountered"
## construct truth tables :
x <- c(NA, FALSE, TRUE)
names(x) <- as.character(x)
outer(x, x, "&")## AND table
outer(x, x, "|")## OR table
```

```
logical Logical Vectors
```


## Description

Create or test for objects of type "logical", and the basic logical "constants".

## Usage

TRUE
FALSE
T; F
logical (length $=0$ )
as.logical(x, ...)
is.logical(x)

## Arguments

| length | desired length. |
| :--- | :--- |
| x | object to be coerced or tested. |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

TRUE and FALSE are part of the R language, where T and F are global variables set to these. All four are logical(1) vectors.

## Value

logical creates a logical vector of the specified length. Each element of the vector is equal to FALSE.
as.logical attempts to coerce its argument to be of logical type. For factors, this uses the levels (labels) and not the codes.
is.logical returns TRUE or FALSE depending on whether its argument is of logical type or not.
Logistic The Logistic Distribution

## Description

Density, distribution function, quantile function and random generation for the logistic distribution with parameters location and scale.

## Usage

```
dlogis(x, location = 0, scale = 1, log = FALSE)
plogis(q, location = 0, scale = 1, lower.tail = TRUE, log.p = FALSE)
qlogis(p, location = 0, scale = 1, lower.tail = TRUE, log.p = FALSE)
rlogis(n, location = 0, scale = 1)
```


## Arguments

$\mathrm{x}, \mathrm{q} \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n
number of observations. If length $(\mathrm{n})>1$, the length is taken to be the number required.
location, scale
location and scale parameters.
$\log$, log.p logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

If location or scale are omitted, they assume the default values of 0 and 1 respectively.
The Logistic distribution with location $=\mu$ and scale $=\sigma$ has distribution function

$$
F(x)=\frac{1}{1+e^{-(x-\mu) / \sigma}}
$$

and density

$$
f(x)=\frac{1}{\sigma} \frac{e^{(x-\mu) / \sigma}}{\left(1+e^{(x-\mu) / \sigma}\right)^{2}}
$$

It is a long-tailed distribution with mean $\mu$ and variance $\pi^{2} / 3 \sigma^{2}$.

## Value

dlogis gives the density, plogis gives the distribution function, qlogis gives the quantile function, and rlogis generates random deviates.

## Examples

```
eps <- 100 * .Machine$double.eps
x <- c(0:4, rlogis(100))
all.equal(plogis(x), 1/(1 + exp(-x)), tol = eps)
all.equal(plogis(x, lower=FALSE), exp(-x)/ (1 + exp(-x)), tol = eps)
all.equal(plogis(x, lower=FALSE, log=TRUE), -log(1 + exp(x)), tol = eps)
all.equal(dlogis(x), exp(x) * (1 + exp(x))^-2, tol = eps)
var(rlogis(4000, 0, s = 5))# approximately (+/- 3)
pi^2/3 * 5^2
```


## logLik Extract Log-Likelihood

## Description

This function is generic; method functions can be written to handle specific classes of objects. Classes which already have methods for this function include: lm, nls in package nls, and gls, lme and others in package nlme.

## Usage

logLik(object, ...)
as.data.frame(x, row.names $=$ NULL, optional $=$ FALSE $)$

## Arguments

object any object from which a log-likelihood value, or a contribution to a loglikelihood value, can be extracted.
... some methods for this generic function require additional arguments.
$\mathrm{x} \quad$ an object of class logLik.
row.names, optional
arguments to the as.data.frame method; see its documentation.

## Value

Returns an object, say $r$, of class logLik which is a number with attributes, attr (r, "df") (degrees of freedom) giving the number of parameters in the model. There's a simple print method for logLik objects.

The details depend on the method function used; see the appropriate documentation.

## Author(s)

Jose Pinheiro and Douglas Bates

## See Also

logLik.lm, logLik.nls, logLik.gls, logLik.lme, etc.

## Examples

```
## see the method function documentation
x <- 1:5
lmx <- lm(x ~ 1)
logLik(lmx) # using print.logLik() method
str(logLik(lmx))
```

logLik.glm Extract Log-Likelihood from an glm Object

## Description

Returns the log-likelihood value of the generalized linear model represented by object evaluated at the estimated coefficients.

## Usage

```
logLik(object, ...)
```


## Arguments

object an object inheriting from class "glm".
... further arguments to be passed to or from methods.

## Details

As a family does not have to specify how to calculate the log-likelihood, this is based on the family's function to compute the AIC. For gaussian, Gamma and inverse.gaussian families it assumed that the dispersion of the GLM is estimated and has been included in the AIC, and for all other families it is assumed that the dispersion is known.

Not that this procedure is not completely accurate for the gamma and inverse gaussian families, as the estimate of dispersion used is not the MLE.

## Value

the log-likelihood of the linear model represented by object evaluated at the estimated coefficients.

## See Also

```
glm, logLik.lm
```


## Examples

```
##-- Continuing the glm(.) example:
Eq <- function(x,y) all.equal(x,y, tol = 1e-12)
stopifnot(Eq(AIC(anorex.1), anorex.1$aic),
    Eq(AIC(g1), g1$aic),
    Eq(AIC(g2), g2$aic))
## next was wrong in 1.4.1
x <- 1:10
lmx <- logLik(lm(x ~ 1)); glmx <- logLik(glm(x ~ 1))
stopifnot(all.equal(as.vector(lmx), as.vector(glmx)),
    all.equal(attr(lmx, 'df'), attr(glmx, 'df')))
```

logLik.lm

## Description

If REML=FALSE, returns the log-likelihood value of the linear model represented by object evaluated at the estimated coefficients; else, the restricted log-likelihood evaluated at the estimated coefficients is returned.

## Usage

logLik(object, REML = FALSE, ...)

## Arguments

object an object inheriting from class "lm".
REML an optional logical value. If TRUE the restricted log-likelihood is returned, else, if FALSE, the log-likelihood is returned. Defaults to FALSE.
... further arguments to be passed to or from methods.

## Value

an object of class logLik, the (restricted) log-likelihood of the linear model represented by object evaluated at the estimated coefficients. Note that error variance $\sigma^{2}$ is estimated in $\operatorname{lm}$ () and hence counted as well.

## Author(s)

Jose Pinheiro and Douglas Bates

## References

Harville, D.A. (1974). Bayesian Inference for Variance Components Using Only Error Contrasts. Biometrika, 61, 383-385.

## See Also

lm

## Examples

```
data(attitude)
(fm1 <- lm(rating ~ ., data = attitude))
logLik(fm1)
logLik(fm1, REML = TRUE)
Nnlme <- is.na(match("package:nlme", search()))
if(require(nlme)) {
    data(Orthodont)
    fm1 <- lm(distance ~ Sex * age, Orthodont)
    print(logLik(fm1))
    print(logLik(fm1, REML = TRUE))
    if(Nnlme) detach( "package:nlme")
}
```


## Description

loglin is used to fit log-linear models to multidimensional contingency tables by Iterative Proportional Fitting.

## Usage

```
loglin(table, margin, start = rep(1, length(table)), fit = FALSE,
            eps = 0.1, iter = 20, param = FALSE, print = TRUE)
```


## Arguments

table a contingency table to be fit, typically the output from table.
margin a list of vectors with the marginal totals to be fit.
(Hierarchical) log-linear models can be specified in term of these marginal totals which give the "maximal" factor subsets contained in the model. For example, in a three-factor model, list $(c(1,2), c(1,3))$ specifies a model which contains parameters for the grand mean, each factor, and the 1-2 and 1-3 interactions, respectively (but no 2-3 or 1-2-3 interaction), i.e., a model where factors 2 and 3 are independent conditional on factor 1 (sometimes represented as '[12][13]').
The names of factors (i.e., names (dimnames (table))) may be used rather than numeric indices.
start a starting estimate for the fitted table. This optional argument is important for incomplete tables with structural zeros in table which should be preserved in the fit. In this case, the corresponding entries in start should be zero and the others can be taken as one.
fit a logical indicating whether the fitted values should be returned.
eps maximum deviation allowed between observed and fitted margins.
iter maximum number of iterations.
param a logical indicating whether the parameter values should be returned.
print a logical. If TRUE, the number of iterations and the final deviation are printed.

## Details

The Iterative Proportional Fitting algorithm as presented in Haberman (1972) is used for fitting the model. At most iter iterations are performed, convergence is taken to occur when the maximum deviation between observed and fitted margins is less than eps. All internal computations are done in double precision; there is no limit on the number of factors (the dimension of the table) in the model.
Assuming that there are no structural zeros, both the Likelihood Ratio Test and Pearson test statistics have an asymptotic chi-squared distribution with df degrees of freedom.
Package 'MASS' contains loglm, a front-end to loglin which allows the log-linear model to be specified and fitted in a formula-based manner similar to that of other fitting functions such as lm or glm.

## Value

A list with the following components.
lrt the Likelihood Ratio Test statistic.
pearson the Pearson test statistic (X-squared).
df the degrees of freedom for the fitted model. There is no adjustment for structural zeros.
margin list of the margins that were fit. Basically the same as the input margin, but with numbers replaced by names where possible.
fit An array like table containing the fitted values. Only returned if fit is TRUE.
param A list containing the estimated parameters of the model. The "standard" constraints of zero marginal sums (e.g., zero row and column sums for a two factor parameter) are employed. Only returned if param is TRUE.

## Author(s)

Kurt Hornik

## References

Haberman, S. J. (1972) Log-linear fit for contingency tables-Algorithm AS51. Applied Statistics, 21, 218-225.
Agresti, A. (1990) Categorical data analysis. New York: Wiley.

## See Also

table

## Examples

```
data(HairEyeColor)
## Model of joint independence of sex from hair and eye color.
fm <- loglin(HairEyeColor, list(c(1, 2), c(1, 3), c(2, 3)))
fm
1 - pchisq(fm$lrt, fm$df)
## Model with no three-factor interactions fits well.
```

Lognormal The Log Normal Distribution

## Description

Density, distribution function, quantile function and random generation for the log normal distribution whose logarithm has mean equal to meanlog and standard deviation equal to sdlog.

## Usage

```
dlnorm(x, meanlog \(=0\), sdlog \(=1, \log =\) FALSE)
plnorm(q, meanlog \(=0\), sdlog \(=1\), lower.tail = TRUE, log.p = FALSE)
qlnorm(p, meanlog \(=0\), sdlog \(=1\), lower.tail \(=\) TRUE, log. \(p=\) FALSE)
rlnorm(n, meanlog \(=0\), sdlog \(=1\) )
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the number required.
meanlog, sdlog
mean and standard deviation of the distribution on the log scale with default values of 0 and 1 respectively.
$\log , \log . \mathrm{p} \quad$ logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The log normal distribution has density

$$
f(x)=\frac{1}{\sqrt{2 \pi} \sigma x} e^{-(\log (x)-\mu)^{2} / 2 \sigma^{2}}
$$

where $\mu$ and $\sigma$ are the mean and standard deviation of the logarithm. The mean is $E(X)=$ $\exp \left(\mu+1 / 2 \sigma^{2}\right)$, and the variance $\operatorname{Var}(X)=\exp \left(2 \mu+\sigma^{2}\right)\left(\exp \left(\sigma^{2}\right)-1\right)$ and hence the coefficient of variation is $\sqrt{\exp \left(\sigma^{2}\right)-1}$ which is approximately $\sigma$ when that is small (e.g. $\sigma<1 / 2)$.

## Value

dlnorm gives the density, plnorm gives the distribution function, qlnorm gives the quantile function, and rlnorm generates random deviates.

## Note

The cumulative hazard $H(t)=-\log (1-F(t))$ is - plnorm(t, r, lower $=$ FALSE, $\log =$ TRUE).

## See Also

dnorm for the normal distribution.

## Examples

```
dlnorm(1) == dnorm(0)
x <- rlnorm(1000) # not yet always :
all(abs(x - qlnorm(plnorm(x))) < 1e4 * .Machine$double.eps * x)
```

longley Longley's Regression Data

## Description

A macroeconomic data set which provides a well-known example for a highly collinear regression.

## Usage

data(longley)

## Format

A data frame with 7 economical variables, observed yearly from 1947 to $1962(n=16)$.
GNP.deflator: GNP implicit price deflator (1954 = 100)
GNP: Gross National Product.
Unemployed: number of unemployed.
Armed.Forces: number of people in the armed forces.
Population: 'noninstitutionalized' population $\geq 14$ years of age.
Year: the year (time).
Employed: number of people employed.
The regression $\operatorname{lm}$ (Employed ~ .) is known to be highly collinear.

## Source

J. W. Longley (1967) An appraisal of least-squares programs from the point of view of the user. Journal of the American Statistical Association, 62, 819-841.

## Examples

```
## give the data set in the form it is used in S-PLUS:
data(longley)
longley.x <- data.matrix(longley[, 1:6])
longley.y <- longley[, "Employed"]
pairs(longley, main = "longley data")
summary(fm1 <- lm(Employed ~ ., data = longley))
opar <- par(mfrow = c(2, 2), oma = c(0, 0, 1.1, 0),
    mar = c(4.1, 4.1, 2.1, 1.1))
plot(fm1)
par(opar)
```

lower.tri Lower and Upper Triangular Part of a Matrix

## Description

Returns a matrix of logicals the same size of a given matrix with entries TRUE in the lower or upper triangle.

## Usage

```
lower.tri(x, diag = FALSE)
upper.tri(x, diag = FALSE)
```


## Arguments

| x | a matrix. |
| :--- | :--- |
| diag | logical. Should the diagonal be included? |

## See Also

diag, matrix.

## Examples

```
m2 <- ma <- matrix(1:20, 4, 5)
m2[lower.tri(m2)] <- NA
m2
stopifnot(lower.tri(ma) == !upper.tri(ma, diag=TRUE))
```

| lowess $\quad$ Scatter Plot Smoothing |
| :--- | :--- |

## Description

This function performs the computations for the $L O W E S S$ smoother (see the reference below). lowess returns a list containing components x and y which give the coordinates of the smooth. The smooth should be added to a plot of the original points with the function lines.

## Usage

```
lowess(x, y, f=2/3, iter=3, delta=.01*diff(range(x)))
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ vectors giving the coordinates of the points in the scatter plot. Alternatively a single plotting structure can be specified.
f
iter the number of robustifying iterations which should be performed. Using smaller values of iter will make lowess run faster.
delta values of $x$ which lie within delta of each other replaced by a single value in the output from lowess.

## References

Cleveland, W. S. (1979) Robust locally weighted regression and smoothing scatterplots. J. Amer. Statist. Assoc. 74, 829-836.

Cleveland, W. S. (1981) LOWESS: A program for smoothing scatterplots by robust locally weighted regression. The American Statistician, 35, 54.

## See Also

loess (in package modreg), a newer formula based version of lowess (with different defaults!).

## Examples

```
data(cars)
plot(cars, main = "lowess(cars)")
lines(lowess(cars), col = 2)
lines(lowess(cars, f=.2), col = 3)
legend(5, 120, c(paste("f = ", c("2/3", ".2"))), lty = 1, col = 2:3)
```

1s $\quad$ List Objects

## Description

ls and objects return a vector of character strings giving the names of the objects in the specified environment. When invoked with no argument at the top level prompt, ls shows what data sets and functions a user has defined. When invoked with no argument inside a function, ls returns the names of the functions local variables. This is useful in conjunction with browser.

## Usage

ls(name, pos $=-1$, envir = as.environment(pos),
all.names $=$ FALSE, pattern)
objects(name, pos= -1 , envir $=$ as.environment (pos), all.names $=$ FALSE, pattern)

## Arguments

| name | which environment to use in listing the available objects. Defaults to the <br> current environment. Although called name for back compatibility, in fact <br> this argument can specify the environment in any form; see the details <br> section. |
| :--- | :--- |
| pos | An alternative argument to name for specifying the environment as a po- <br> sition in the search list. Mostly there for back compatibility. |
| envir | an alternative argument to name for specifying the environment evaluation <br> environment. Mostly there for back compatibility. |
| all.names | a logical value. If TRUE, all object names are returned. If FALSE, names <br> which begin with a "." are omitted. |
| an optional regular expression, see grep. Only names matching pattern |  |
| are returned. |  |

## Details

The name argument can specify the environment from which object names are taken in one of several forms: as an integer (the position in the search list); as the character string name of an element in the search list; or as an explicit environment (including using sys.frame to access the currently active function calls). By default, the environment of the call to ls or objects is used. The pos and envir arguments are an alternative way to specify an environment, but are primarily there for back compatibility.

## See Also

apropos (or find) for finding objects in the whole search path; grep for more details on "regular expressions"; class, methods, etc. for object-oriented programming.

## Examples

```
.Ob <- 1
ls(pat="0")
ls(pat="O", all = TRUE) # also shows ".[foo]"
# shows an empty list because inside myfunc no variables are defined
myfunc <- function() {ls()}
myfunc()
# define a local variable inside myfunc
myfunc <- function() {y <- 1; ls()}
myfunc() # shows "y"
```

ls.diag Compute Diagnostics for 'Isfit' Regression Results

## Description

Computes basic statistics, including standard errors, t- and p-values for the regression coefficients.

## Usage

ls.diag(ls.out)

## Arguments

ls.out Typically the result of lsfit()

## Value

A list with the following numeric components.
std.dev The standard deviation of the errors, an estimate of $\sigma$.
hat diagonal entries $h_{i i}$ of the hat matrix $H$
std.res standardized residuals
stud.res studentized residuals
cooks Cook's distances
dfits DFITS statistics
correlation correlation matrix
std.err standard errors of the regression coefficients
cov.scaled Scaled covariance matrix of the coefficients
cov.unscaled Unscaled covariance matrix of the coefficients

## References

Belsley, D. A., Kuh, E. and Welsch, R. E. (1980) Regression Diagnostics. New York: Wiley.

## See Also

hat for the hat matrix diagonals, ls.print, lm.influence, summary.lm, anova.

## Examples

```
##-- Using the same data as the lm(.) example:
lsD9 <- lsfit(x = as.numeric(gl(2, 10, 20)), y = weight)
dlsD9 <- ls.diag(lsD9)
str(dlsD9, give.attr=FALSE)
abs(1 - sum(dlsD9$hat) / 2) < 10*.Machine$double.eps # sum(h.ii) = p
plot(dlsD9$hat, dlsD9$stud.res, xlim=c(0,0.11))
abline(h = 0, lty = 2, col = "lightgray")
```

ls.print Print 'lsfit' Regression Results

## Description

Computes basic statistics, including standard errors, t- and p-values for the regression coefficients and prints them if print.it is TRUE.

## Usage

ls.print(ls.out, digits $=4$, print.it $=$ TRUE)

## Arguments

ls.out Typically the result of lsfit()
digits The number of significant digits used for printing
print.it a logical indicating whether the result should also be printed

## Value

A list with the components
summary The ANOVA table of the regression
coef.table matrix with regression coefficients, standard errors, $t$ - and p-values

## Note

Usually, you'd rather use summary $(\operatorname{lm}(\ldots))$ and anova( $\operatorname{lm}(\ldots)$ ) for obtaining similar output.

## See Also

ls.diag, lsfit, also for examples; lm, lm.influence which usually are preferable.

## 1sfit Find the Least Squares Fit

## Description

The least squares estimate of $\beta$ in the model

$$
\boldsymbol{Y}=\boldsymbol{X} \boldsymbol{\beta}+\boldsymbol{\epsilon}
$$

is found.

## Usage

```
lsfit(x, y, wt=NULL, intercept=TRUE, tolerance=1e-07, yname=NULL)
```


## Arguments

x
y
intercept
tolerance
yname
wt an optional vector of weights for performing weighted least squares.
a matrix whose rows correspond to cases and whose columns correspond to variables.
the responses, possibly matrix valued if you want to fit multiple left hand sides. whether or not an intercept term should be used.
the tolerance to be used in the matrix decomposition.
an unused parameter for compatibility.

## Details

If weights are specified then a weighted least squares is performed with the weight given to the $j$ th case specified by the $j$ th entry in wt.
If any observation has a missing value in any field, that observation is removed before the analysis is carried out. This can be quite inefficient if there is a lot of missing data.
The implementation is via a modification of the LINPACK subroutines which allow for multiple left-hand sides.

## Value

A list with the following named components:
coef the least squares estimates of the coefficients in the model (stated below).
residuals residuals from the fit.
intercept indicates whether an intercept was fitted.
qr the QR decomposition of the design matrix.

## See Also

lm which usually is preferable; ls.print, ls.diag.

## Examples

```
##-- Using the same data as the lm(.) example:
lsD9 <- lsfit(x = codes(gl(2,10)), y = weight)
ls.print(lsD9)
```

mad Median Absolute Deviation

## Description

Compute the median absolute deviation, i.e., the (lo-/hi-) median of the absolute deviations from the median, and (by default) adjust by a factor for asymptotically normal consistency.

## Usage

```
mad(x, center = median(x), constant = 1.4826, na.rm = FALSE,
    low = FALSE, high = FALSE)
```


## Arguments

$\mathrm{x} \quad$ a numeric vector.
center Optionally, the centre: defauls to the median.
constant scale factor.
na.rm if TRUE then NA values are stripped from $x$ before computation takes place.
low if TRUE, compute the "lo-median", i.e., for even sample size, do not average the two middle values, but take the smaller one.
high if TRUE, compute the "hi-median", i.e. take the larger of the two middle values for even sample size.

## Details

The actual value calculated is constant $*$ cMedian(abs( $\mathrm{x}-\mathrm{center)}$ ) with the default value of center being median( x ), and cMedian being the usual, the "low" or "high" median, see the arguments description for low and high above.
The default constant $=1.4826$ (approximately $1 / \Phi^{-1}\left(\frac{3}{4}\right)=1 /$ qnorm(3/4) ) ensures consistency, i.e.,

$$
E\left[\operatorname{mad}\left(X_{1}, \ldots, X_{n}\right)\right]=\sigma
$$

for $X_{i}$ distributed as $N\left(\mu, \sigma^{2}\right)$ and large $n$.
If na.rm is TRUE then NA values are stripped from $x$ before computation takes place. If this is not done then an NA value in x will cause mad to return NA.

## See Also

IQR which is simpler but less robust, median, var.

## Examples

```
mad(c(1:9))
print(mad(c(1:9), constant=1)) ==
    mad(c(1:8,100), constant=1) # = 2 ; TRUE
x <- c(1,2,3, 5,7,8)
sort(abs(x - median(x)))
c(mad(x, co=1), mad(x, co=1, lo = TRUE), mad(x, co=1, hi = TRUE))
```


## mahalanobis Mahalanobis Distance

## Description

Returns the Mahalanobis distance of all rows in x and the vector $\mu=$ center with respect to $\Sigma=$ cov. This is (for vector x ) defined as

$$
D^{2}=(x-\mu)^{\prime} \Sigma^{-1}(x-\mu)
$$

## Usage

```
mahalanobis(x, center, cov, inverted=FALSE, tol.inv = 1e-7)
```


## Arguments

| x | vector or matrix of data with, say, $p$ columns. |
| :--- | :--- |
| center | mean vector of the distribution or second data vector of length $p$. |
| cov | covariance matrix $(p \times p)$ of the distribution. |
| inverted | logical. If TRUE, cov is supposed to contain the inverse of the covariance <br> matrix. |
| tol.inv | tolerance to be used for computing the inverse (if inverted is false), see <br> solve. |

## Author(s)

Friedrich Leisch

## See Also

```
cov, var
```


## Examples

```
ma <- cbind(1:6, 1:3)
(S <- var(ma))
mahalanobis(c(0,0), 1:2, S)
x <- matrix(rnorm(100*3), ncol = 3)
stopifnot(mahalanobis(x, 0, diag(ncol(x))) == rowSums(x*x))
    ##- Here, D^2 = usual Euclidean distances
Sx <- cov(x)
D2 <- mahalanobis(x, rowMeans(x), Sx)
plot(density(D2, bw=.5), main="Mahalanobis distances, n=100, p=3"); rug(D2)
qqplot(qchisq(ppoints(100), df=3), D2,
    main = expression("Q-Q plot of Mahalanobis" * ~D`2 *
        " vs. quantiles of" * ~ chi [3]^2))
abline(0, 1, col = 'gray')
```

make.link

## Description

This function is used with the family functions in $g l m()$. Given a link, it returns a link function, an inverse link function, the derivative $d \mu / d \eta$ and a function for domain checking.

## Usage

```
make.link(link)
```


## Arguments

link character or numeric; one of "logit", "probit", "cloglog", "identity", "log", "sqrt", "1/mu^2", "inverse", or number, say $\lambda$ resulting in power link $=\mu^{\lambda}$.

## Value

A list with components

| linkfun | Link function function(mu) |
| :--- | :--- |
| linkinv | Inverse link function function(eta) |
| mu.eta | Derivative function(eta) $d \mu / d \eta$ |
| valideta | function(eta) \{ TRUE if all of eta is in the domain of linkinv $\}.$ |

## See Also

glm, family.

## Examples

```
str(make.link("logit"))
12 <- make.link(2)
l2$linkfun(0:3)# 0 1 4 9
12$mu.eta(eta= 1:2)#= 1/(2*sqrt(eta))
```

```
make.names
Make Syntactically Valid Names
```


## Description

Make syntactically valid names out of character vectors.

## Usage

make.names(names, unique $=$ FALSE)

## Arguments

names character (vector) to be coerced to syntactically valid names.
unique logical; if TRUE, the resulting elements are unique. This may be desired for, e.g., column names.

## Details

A syntactically valid name consists of letters, numbers, and the dot character and starts with a letter or the dot.

All invalid characters are translated to ".". A missing value is translated to "NA".
If unique $=$ TRUE a sequence number is appended to each duplicate (after coercion).

## Value

A character vector of same length as names with each changed to a syntactically valid name.

## See Also

names, character, data.frame.

## Examples

```
make.names(c("a and b", "a_and_b"), unique=TRUE)#-> "a.and.b" "a.and.b1"
all(make.names(letters) == letters)# TRUE
data(state)
state.name[make.names(state.name) != state.name]# those 10 with a space
```

```
make.socket
Create a Socket Connection
```


## Description

With server = FALSE attempts to open a client socket to the specified port and host. With server = TRUE listens on the specified port for a connection and then returns a server socket. It is a good idea to use on.exit to ensure that a socket is closed, as you only get 64 of them.

## Usage

```
make.socket(host = "localhost", port, fail = TRUE, server = FALSE)
```


## Arguments

| host | name of remote host |
| :--- | :--- |
| port | port to connect to/listen on |
| fail | failure to connect is an error? |
| server | a server socket? |

## Value

An object of class "socket".
socket socket number. This is for internal use
port port number of the connection
host name of remote computer

## Warning

I don't know if the connecting host name returned when server $=$ TRUE can be trusted. I suspect not.

Author(s)
Thomas Lumley

## References

Adapted from Luke Tierney's code for XLISP-Stat, in turn based on code from Robbins and Robbins "Practical UNIX Programming"

## See Also

close.socket, read.socket

## Examples

```
daytime <- function(host = "localhost"){
    a <- make.socket(host, 13)
    on.exit(close.socket(a))
    read.socket(a)
}
## Offical time (UTC) from US Naval Observatory
daytime("tick.usno.navy.mil")
```

make.tables Create model.tables

## Description

These are support functions for (the methods of) model.tables and probably not much of use otherwise.

## Usage

```
make.tables.aovproj (proj.cols, mf.cols, prjs, mf,
    fun = "mean", prt = FALSE, ...)
    make.tables.aovprojlist(proj.cols, strata.cols, model.cols, projections,
    model, eff, fun = "mean", prt = FALSE, ...)
```

See Also
model.tables

## Description

A utility to help model.frame.default create the right matrices when predicting from models with terms like poly or ns.

## Usage

makepredictcall(var, call)

## Arguments

var A variable.
call The term in the formula, as a call.

## Details

This is a generic function with methods for poly, bs and ns: the default method handles scale. If model.frame. default encounters such a term when creating a model frame, it modifies the predvars attribute of the terms supplied to replace the term with one that will work for predicting new data. For example makepredictcall.ns adds arguments for the knots and intercept.
To make use of this, have your model-fitting function return the terms attribute of the model frame, or copy the predvars attribute of the terms attribute of the model frame to your terms object.
To extend this, make sure the term creates variables with a class, and write a suitable method for that class.

## Value

A replacement for call for the predvars attribute of the terms.

## See Also

```
model.frame, poly, scale, bs, ns, cars
```


## Examples

```
## using poly: this did not work in R < 1.5.0
data(women)
fm <- lm(weight ~ poly(height, 2), data = women)
plot(women, xlab = "Height (in)", ylab = "Weight (lb)")
ht <- seq(57, 73, len = 200)
lines(ht, predict(fm, data.frame(height=ht)))
## see also example(cars)
## see bs and ns for spline examples.
```

manova Multivariate Analysis of Variance

## Description

A class of multivariate analysis of variance.

## Usage

manova(...)

## Arguments

## ... Arguments to be passed to aov.

## Details

Class "manova" differs from class "aov" in selecting a different summary method. Function manova calls aov and then add class "manova" to the result object for each stratum.

## Value

See aov and the comments in Details here.

## Note

manova does not support multistratum analysis of variance, so the formula should not include an Error term.

## Author(s)

B.D. Ripley

## References

Krzanowski, W. J. (1988) Principles of Multivariate Analysis. A User's Perspective. Oxford.

Hand, D. J. and Taylor, C. C. (1987) Multivariate Analysis of Variance and Repeated Measures. Chapman and Hall.

## See Also

aov, summary.manova, the latter containing examples.

```
margin.table
Compute table margin
```


## Description

For a contingency table in array form, compute the sum of table entries for a given index.

## Usage

margin.table(x, margin=NULL)

## Arguments

| x | an array |
| :--- | :--- |
| $\operatorname{margin}$ | index number (1 for rows, etc.) |

## Details

This is really just apply ( $x$, margin, sum) packaged up for newbies, except that if margin has length zero you get sum ( x ).

## Value

The relevant marginal table. The class of x is copied to the output table, except in the summation case.

## Author(s)

Peter Dalgaard

## Examples

```
m<-matrix(1:4,2)
margin.table(m,1)
margin.table(m,2)
```

```
mat.or.vec Create a Matrix or a Vector
```


## Description

mat.or.vec creates an $n r$ by nc zero matrix if nc is greater than 1 , and a zero vector of length nr if nc equals 1 .

## Usage

```
mat.or.vec(nr, nc)
```


## Arguments

$\mathrm{nr}, \mathrm{nc} \quad$ numbers of rows and columns.

## Examples

```
mat.or.vec(3, 1)
mat.or.vec(3, 2)
```


## match Value Matching

## Description

match returns a vector of the positions of (first) matches of its first argument in its second.
$\%$ in\% is a more intuitive interface as a binary operator, which returns a logical vector indicating if there is a match or not for its left operand.

## Usage

```
match(x, table, nomatch = NA, incomparables = FALSE)
x %in% table
```


## Arguments

$\mathrm{x} \quad$ the values to be matched.
table the values to be matched against.
nomatch the value to be returned in the case when no match is found. Note that it is coerced to integer.
incomparables a vector of values that cannot be matched. Any value in x matching a value in this vector is assigned the nomatch value. Currently, FALSE is the only possible value, meaning that all values can be matched.

## Details

\%in\% is currently defined as

```
"%in%" <- function(x, table) match(x, table, nomatch = 0) > 0
```

Factors are converted to character vectors, and then $x$ and table are coerced to a common type (the later of the two types in R's ordering, logical $<$ integer $<$ numeric $<$ complex $<$ character) before matching.

## Value

In both cases, a vector of the same length as x .
match: An integer vector giving the position in table of the first match if there is a match, otherwise nomatch.
If $x[i]$ is found to equal table[j] then the value returned in the i-th position of the return value is $j$, for the smallest possible $j$. If no match is found, the value is nomatch.
$\% \mathrm{in} \%$ : A logical vector, indicating if a match was located for each element of x .

## See Also

pmatch and charmatch for (partial) string matching, match.arg, etc for function argument matching.
is.element for an S-compatible equivalent of \%in\%.

## Examples

```
## The intersection of two sets :
intersect <- function(x, y) y[match(x, y, nomatch = 0)]
intersect(1:10,7:20)
1:10 %in% c(1,3,5,9)
sstr <- c("c","ab","B","bba","c","@","bla","a","Ba", "%")
sstr[sstr %in% c(letters,LETTERS)]
"%w/o%" <- function(x,y) x[!x %in% y] #-- x without y
(1:10) %w/o% c(3,7,12)
```

```
match.arg Argument Verification Using Partial Matching
```


## Description

match.arg matches arg against a table of candidate values as specified by choices.

## Usage

match.arg(arg, choices)

## Arguments

| $\arg$ | a character string |
| :--- | :--- |
| choices | a character vector of candidate values |

## Details

In the one-argument form match. $\arg (\arg )$, the choices are obtained from a default setting for the formal argument arg of the function from which match. arg was called.
Matching is done using pmatch, so arg may be abbreviated.

## Value

The unabbreviated version of the unique partial match if there is one; otherwise, an error is signalled.

## See Also

 pmatch, match.fun, match.call.
## Examples

```
## Extends the example for 'switch'
center <- function(x, type = c("mean", "median", "trimmed")) {
    type <- match.arg(type)
    switch(type,
            mean = mean(x),
            median = median(x),
            trimmed = mean(x, trim = .1))
}
```

```
x <- rcauchy(10)
center(x, "t") # Works
center(x, "med") # Works
center(x, "m") # Error
```

```
match.call Argument Matching
```


## Description

match.call returns a call in which all of the arguments are specified by their names. The most common use is to get the call of the current function, with all arguments named.

## Usage

```
match.call(definition = NULL, call = sys.call(sys.parent()),
    expand.dots = TRUE)
```


## Arguments

definition a function, by default the function from which match.call is called.
call an unevaluated call to the function specified by definition, as generated by call.
expand.dots logical. Should arguments matching . . . in the call be included or left as a . . . argument?

Value
An object of class call.

## See Also

call, pmatch, match.arg, match.fun.

## Examples

```
match.call(get, call("get", "abc", i = FALSE, p = 3))
## -> get(x = "abc", pos = 3, inherits = FALSE)
fun <- function(x, lower = 0, upper = 1) {
    structure((x - lower) / (upper - lower), CALL = match.call())
}
fun(4 * atan(1), u = pi)
```

```
match.fun Function Verification for "Function Variables"
```


## Description

When called inside functions that take a function as argument, extract the desired function object while avoiding undesired matching to objects of other types.

## Usage

match.fun(FUN, descend $=$ TRUE)

## Arguments

FUN item to match as function.
descend logical; control whether to search past non-function objects.

## Details

match.fun is not intended to be used at the top level since it will perform matching in the parent of the caller.

If FUN is a function, it is returned. If it is a symbol or a character vector of length one, it will be looked up using get in the environment of the parent of the caller. If it is of any other mode, it is attempted first to get the argument to the caller as a symbol (using substitute twice), and if that fails, an error is declared.

If descend = TRUE, match.fun will look past non-function objects with the given name; otherwise if FUN points to a non-function object then an error is generated.

This is now used in base functions such as apply, lapply, outer, and sweep.

## Value

A function matching FUN or an error is generated.

## Bugs

The descend argument is a bit of misnomer and probably not actually needed by anything. It may go away in the future.
It is impossible to fully foolproof this. If one attaches a list or data frame containing a character object with the same name of a system function, it will be used.

## Author(s)

Peter Dalgaard and Robert Gentleman, based on an earlier version by Jonathan Rougier.

## See Also

```
match.arg, get
```


## Examples

```
# Same as get("*"):
match.fun("*")
# Overwrite outer with a vector
outer <- 1:5
match.fun(outer, descend = FALSE) #-> Error: not a function
match.fun(outer) # finds it anyway
is.function(match.fun("outer")) # as well
```

matmult Matrix Multiplication

## Description

Multiplies two matrices, if they are conformable. If one argument is a vector, it will be coerced to a either a row or column matrix to make the two arguments conformable. If both are vectors it will return the inner product.

## Usage

a $\%$ \% b

## Value

The matrix product. Use drop to get rid of dimensions which have only one level.

## See Also

matrix, Arithmetic, diag.

## Examples

```
x <- 1:4
(z <- x %*% x) # scalar ("inner") product (1 x 1 matrix)
drop(z) # as scalar
y <- diag(x)
z <- matrix(1:12, ncol = 3, nrow = 4)
y %*% z
y %*% x
x %*% z
```

```
matplot Plot Columns of Matrices
```


## Description

Plot the columns of one matrix against the columns of another.

```
Usage
    matplot(x, y, type = "p", lty = 1:5, lwd = 1, pch = NULL, col = 1:6,
        cex = NULL, xlab = NULL, ylab = NULL, xlim = NULL, ylim = NULL,
        ..., add = FALSE, verbose = getOption("verbose"))
    matpoints(x, y, type = "p", lty = 1:5, lwd = 1, pch = NULL, col = 1:6, ...)
    matlines (x, y, type = "l", lty = 1:5, lwd = 1, pch = NULL, col = 1:6, ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | vectors or matrices of data for plotting. The number of rows should match. If one of them are missing, the other is taken as $y$ and an $x$ vector of $1: n$ is used. Missing values (NAs) are allowed. |
| :---: | :---: |
| type | character string (length 1 vector) or vector of 1-character strings indicating the type of plot for each column of y , see plot for all possible types. The first character of type defines the first plot, the second character the second, etc. Characters in type are cycled through; e.g., "pl" alternately plots points and lines. |
| lty, lwd | vector of line types and widths. The first element is for the first column, the second element for the second column, etc., even if lines are not plotted for all columns. Line types will be used cyclically until all plots are drawn. |
| pch | character string or vector of 1-characters or integers for plotting characters, see points. The first character is the plotting-character for the first plot, the second for the second, etc. The default is the digits (1 through $9,0)$ then the letters. |
| col | vector of colors. Colors are used cyclically. |
| cex | vector of character expansion sizes, used cyclically. |
| xlab, ylab | titles for x and y axes, as in plot. |
| xlim, ylim | ranges of x and y axes, as in plot. |
|  | Graphical parameters (see par) and any further arguments of plot, typically plot.default, may also be supplied as arguments to this function. Hence, the high-level graphics control arguments described under par and the arguments to title may be supplied to this function. |
| add | logical. If TRUE, plots are added to current one, using points and lines. |
| verbose | logical. If TRUE, write one line of what is done. |

## Details

Points involving missing values are not plotted.
The first column of x is plotted against the first column of y , the second column of x against the second column of y , etc. If one matrix has fewer columns, plotting will cycle
back through the columns again. (In particular, either x or y may be a vector, against which all columns of the other argument will be plotted.)

The first element of col, cex, lty, lwd is used to plot the axes as well as the first line.
Because plotting symbols are drawn with lines and because these functions may be changing the line style, you should probably specify lty=1 when using plotting symbols.

## Side Effects

Function matplot generates a new plot; matpoints and matlines add to the current one.

## See Also

plot, points, lines, matrix, par.

## Examples

```
matplot((-4:5)^2, main = "Quadratic") # almost identical to plot(*)
sines <- outer(1:20, 1:4, function(x, y) sin(x / 20 * pi * y))
matplot(sines, pch = 1:4, type = "o", col = rainbow(ncol(sines)))
x <- 0:50/50
matplot(x, outer(x, 1:8, function(x, k) sin(k*pi * x)),
    ylim = c(-2,2), type = "plobcsSh",
    main= "matplot(,type = \"plobcsSh\" )")
## pch & type = vector of 1-chars :
matplot(x, outer(x, 1:4, function(x, k) sin(k*pi * x)),
    pch = letters[1:4], type = c("b","p","o"))
data(iris) # is data.frame with 'Species' factor
table(iris$Species)
iS <- iris$Species == "setosa"
iV <- iris$Species == "versicolor"
op <- par(bg = "bisque")
matplot(c(1, 8), c(0, 4.5), type= "n", xlab = "Length", ylab = "Width",
    main = "Petal and Sepal Dimensions in Iris Blossoms")
matpoints(iris[iS,c(1,3)], iris[iS,c(2,4)], pch = "sS", col = c(2,4))
matpoints(iris[iV,c(1,3)], iris[iV,c(2,4)], pch = "vV", col = c(2,4))
legend(1, 4, c(" Setosa Petals", " Setosa Sepals",
            "Versicolor Petals", "Versicolor Sepals"),
    pch = "sSvV", col = rep(c(2,4), 2))
nam.var <- colnames(iris) [-5]
nam.spec <- as.character(iris[1+50*0:2, "Species"])
iris.S <- array(NA, dim = c(50,4,3), dimnames = list(NULL, nam.var, nam.spec))
for(i in 1:3) iris.S[,,i] <- data.matrix(iris[1:50+50*(i-1), -5])
matplot(iris.S[,"Petal.Length",], iris.S[,"Petal.Width",], pch="SCV",
    col = rainbow(3, start = .8, end = .1),
    sub = paste(c("S", "C", "V"), dimnames(iris.S)[[3]],
                sep = "=", collapse= ", "),
    main = "Fisher's Iris Data")
```


## matrix Matrices

## Description

matrix creates a matrix from the given set of values.
as.matrix attempts to turn its argument into a matrix.
is.matrix tests if its argument is a (strict) matrix.

## Usage

```
matrix(data = NA, nrow = 1, ncol = 1, byrow = FALSE, dimnames = NULL)
as.matrix(x)
is.matrix(x)
```


## Arguments

| data | an optional data vector. |
| :--- | :--- |
| nrow | the desired number of rows |
| ncol | the desired number of columns |
| byrow | logical. If FALSE (the default) the matrix is filled by columns, otherwise <br> the matrix is filled by rows. |
| dimnames | A dimnames attribute for the matrix: a list of length 2. |
| x | an R object. |

## Details

If either of nrow or ncol is not given, an attempt is made to infer it from the length of data and the other parameter.
is.matrix returns TRUE if $x$ is a matrix (i.e., it is not a data.frame and has a dim attribute of length 2) and FALSE otherwise.
as.matrix is a generic function. The method for data frames will convert any non-numeric column into a character vector using format and so return a character matrix.

## See Also

data.matrix, which attempts to convert to a numeric matrix.

## Examples

```
is.matrix(as.matrix(1:10))
data(warpbreaks)
!is.matrix(warpbreaks)# data.frame, NOT matrix!
str(warpbreaks)
str(as.matrix(warpbreaks))#using as.matrix.data.frame(.) method
```


## Description

Find the maximum position for each row of a matrix, breaking ties at random.

## Usage

$\max . \operatorname{col}(m)$

## Arguments

m numerical matrix

## Details

Ties are broken at random. The determination of "tie" assumes that the entries are probabilities.

## Value

index of a maximal value for each row, an integer vector of length nrow(m).

## Author(s)

W. N. Venables and B. D. Ripley

## References

Venables, W. N. and Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. New York: Springer (3nd ed).

## See Also

 which.max for vectors.
## Examples

```
data(swiss)
table(mc <- max.col(swiss))# mostly "1" and "5", 5 x "2" and once "4"
swiss[unique(print(mr <- max.col(t(swiss)))) , ] # 3 33 45 45 33 6
```

```
mean
Arithmetic Mean
```


## Description

Generic function for the (trimmed) arithmetic mean.

## Usage

```
mean(x, ...)
mean.default(x, trim \(=0\), na.rm = FALSE, ...)
```


## Arguments

X
trim
na.rm a logical value indicating whether NA values should be stripped before the computation proceeds.
... further arguments passed to or from other methods.

## Value

For a data frame, a named vector with the appropriate method being applied column by column.

If trim is zero (the default), the arithmetic mean of the values in x is computed.
If trim is non-zero, a symmetrically trimmed mean is computed with a fraction of trim observations deleted from each end before the mean is computed.

See Also

```
weighted.mean, mean.POSIXct
```


## Examples

```
x <- c(0:10, 50)
xm <- mean(x)
c(xm, mean(x, trim = 0.10))
all.equal(mean(x, trim = 0.5), median(x))
data(USArrests)
mean(USArrests, trim = 0.2)
```

```
median Median Value
```


## Description

Compute the sample median of the vector of values given as its argument.

## Usage

```
median(x, na.rm=FALSE)
```


## Arguments

| x | a numeric vector containing the values whose median is to be computed. |
| :--- | :--- |
| na.rm | a logical value indicating whether NA values should be stripped before the <br> computation proceeds. |

## See Also

quantile for general quantiles.

## Examples

```
median(1:4)# = 2.5 [even number]
median(c(1:3,100,1000))# = 3 [odd, robust]
```

Memory Memory Available for Data Storage

## Description

Use command line options to control the memory available for R .

## Usage

```
R --min-vsize=vl --max-vsize=vu --min-nsize=nl --max-nsize=nu
mem.limits(nsize = NA, vsize = NA)
```


## Arguments

vl, vu, vsize Heap memory in bytes.
nl , nu , nsize Number of cons cells.

## Details

R has a variable-sized workspace (from version 1.2.0). There is now much less need to set memory options than previously, and most users will never need to set these. They are provided both as a way to control the overall memory usage (which can also be done by operating-system facilities such as limit on Unix), and since setting larger values of the minima will make R slightly more efficient on large tasks.

To understand the options, one needs to know that R maintains separate areas for fixed and variable sized objects. The first of these is allocated as an array of "cons cells" (Lisp programmers will know what they are, others may think of them as the building blocks of the language itself, parse trees, etc.), and the second are thrown on a "heap" of "Vcells" of 8 bytes each. Effectively, the input v is rounded up to the nearest multiple of 8 .
Each cons cell occupies 28 bytes on a 32 -bit machine, (usually) 56 bytes on a 64 -bit machine.
The --*-nsize options can be used to specify the number of cons cells and the --*-vsize options specify the size of the vector heap in bytes. Both options must be integers or integers followed by $\mathrm{G}, \mathrm{M}, \mathrm{K}$, or k meaning Giga $\left(2^{30}=1073741824\right)$ Mega $\left(2^{20}=1048576\right)$, (computer) Kilo $\left(2^{10}=1024\right)$, or regular kilo (1000).
The --min-* options set the minimal sizes for the number of cons cells and for the vector heap. These values are also the initial values, but thereafter $R$ will grow or shrink the areas depending on usage, but never exceeding the limits set by the --max-* options nor decreasing below the initial values.

The default values are currently minima of 350 k cons cells, 6 Mb of vector heap and no maxima (other than machine resources). The maxima can be changed during an R session by calling mem.limits. (If this is called with the default values, it reports the current settings.)

You can find out the current memory consumption (the heap and cons cells used as numbers and megabytes) by typing gc() at the R prompt. Note that following gcinfo(TRUE), automatic garbage collection always prints memory use statistics. Maxima will never be reduced below the current values for triggering garbage collection, and attempts to do so will be silently ignored.
When using read.table, the memory requirements are in fact higher than anticipated, because the file is first read in as one long string which is then split again. Use scan if possible in case you run out of memory when reading in a large table.

## Value

(mem.limits) an integer vector giving the current settings of the maxima, possibly NA.

## Note

For backwards compatibility, options --nsize and --vsize are equivalent to --min-nsize and --min-vsize.

## See Also

gc for information on the garbage collector, memory.profile for profiling the usage of cons cells.

## Examples

```
# Start R with 10MB of heap memory and 500k cons cells, limit to
# 100Mb and 1M cells
```

```
## Unix
R --min-vsize=10M --max-vsize=100M --min-nsize=500k --max-nsize=1M
```

memory.profile Profile the Usage of Cons Cells

## Description

Lists the usage of the cons cells by SEXPREC type.

## Usage

memory.profile()

## Details

The current types and their uses are listed in the include file 'Rinternals.h'. There will be blanks in the list corresponding to types that are no longer in use (types 11 and 12 at the time of writing). Also FUNSXP is not included.

## Value

A vector of counts, named by the types.

## See Also

gc for the overall usage of cons cells.

## Examples

```
memory.profile()
```

menu Menu Interaction Function

## Description

menu presents the user with a menu of choices labelled from 1 to the number of choices. To exit without choosing an item one can select ' 0 '.

## Usage

menu(choices, graphics = FALSE, title = "")

## Arguments

| choices | a character vector of choices |
| :--- | :--- |
| graphics | a logical indicating whether a graphics menu should be used. Currently <br> unused. |
| title | a character string to be used as the title of the menu |

## Value

The number corresponding to the selected item, or 0 if no choice was made.

## Examples

```
switch(menu(c("List letters", "List LETTERS")) + 1,
```

    cat("Nothing done\n"), letters, LETTERS)
    ```
merge Merge Two Data Frames
```


## Description

Merge two data frames by common columns or row names, or do other versions of database "join" operations.

## Usage

```
merge(x, y, by, by.x, by.y, all = FALSE, all. \(x=a l l, ~ a l l . y ~=~ a l l, ~\)
    sort \(=\) TRUE, suffixes \(\left.=c\left(" . x^{\prime \prime}, " \cdot y^{\prime \prime}\right)\right)\)
```


## Arguments

$\mathrm{x}, \mathrm{y}$ data frames, or objects to be coerced to one
by, by.x, by.y
specifications of the common columns. See Details.
all logical; all=L is shorthand for all. $\mathrm{x}=\mathrm{L}$ and all. $\mathrm{y}=\mathrm{L}$.
all.x logical; if TRUE, then extra rows will be added to the output, one for each row in $x$ that has no matching row in $y$. These rows will have NAs in those columns that are usually filled with values from y. The default is FALSE, so that only rows with data from both x and y are included in the output.
all.y logical; analogous to all.x above.
sort logical. Should the results be sorted on the by columns?
suffixes character(2) specifying the suffixes to be used for making non-by names() unique.

## Details

By default the data frames are merged on the columns with names they both have, but separate specifications of the columns can be given by by.x and by.y. Columns can be specified by name, number or by a logical vector: the name "row.names" or the number 0 specifies the row names. The rows in the two data frames that match on the specified columns are extracted, and joined together. If there is more than one match, all possible matches contribute one row each.
If the by.* vector are of length 0 , the result, $r$, is the "Cartesian product" of x and y , i.e., $\operatorname{dim}(r)=c(n r o w(x) *$ nrow, $n c o l(x)+n c o l(y))$.
If all. $x$ is true, all the non matching cases of $x$ are appended to the result as well, with NA filled in the corresponding columns of y ; analogously for all.y.
If the remaining columns in the data frames have any common names, these have suffixes (".x" and ".y" by default) appended to make the names of the result unique.

## Value

A data frame. The rows are by default lexicographically sorted on the common columns, but are otherwise in the order in which they occurred in y. The columns are the common columns followed by the remaining columns in x and then those in y . If the matching involved row names, an extra column Row.names is added at the left, and in all cases the result has no special row names.

## See Also

```
data.frame, by, cbind
```


## Examples

```
authors <- data.frame(
    surname = c("Tukey", "Venables", "Tierney", "Ripley", "McNeil"),
    nationality = c("US", "Australia", "US", "UK", "Australia"),
    deceased = c("yes", rep("no", 4)))
books <- data.frame(
    name = c("Tukey", "Venables", "Tierney",
        "Ripley", "Ripley", "McNeil", "R Core"),
    title = c("Exploratory Data Analysis",
                "Modern Applied Statistics ...",
                "LISP-STAT",
                "Spatial Statistics", "Stochastic Simulation",
                "Interactive Data Analysis",
                "An Introduction to R"),
    other.author = c(NA, "Ripley", NA, NA, NA, NA,
                "Venables & Smith"))
(m1 <- merge(authors, books, by.x = "surname", by.y = "name"))
(m2 <- merge(books, authors, by.x = "name", by.y = "surname"))
stopifnot(as.character(m1[,1]) == as.character(m2[,1]),
    all.equal(m1[, -1], m2[, -1][ names(m1)[-1] ]),
    dim(merge(m1, m2, by = integer (0))) == c(36, 10))
## "R core" is missing from authors and appears only here :
merge(authors, books, by.x = "surname", by.y = "name", all = TRUE)
```

Methods Internal and Group Methods and Generic Functions

## Description

Many R-internal functions are generic and allow methods to be written for. Group methods in particular are available for the "Math", "Ops", and "Summary" group.

## Usage

Math.data.frame(x, ...)
Math.factor (x, ...)

```
Ops.data.frame(e1, e2 = NULL)
Ops.factor(e1, e2)
Ops.ordered(e1, e2)
Summary.data.frame(x, ...)
Summary.factor(x, ...)
.Method
.Generic
.Group
.Class
```


## Arguments

```
x, e1, e2 objects.
... further arguments passed to methods.
```


## Group Dispatching

There are three groups for which methods can be written, namely the "Math", "Ops" and "Summary" groups.

A function $f$ belonging to one of these groups must be .Internal or .Primitive and will automatically be using <grp>.<class> (ob) when $f(\langle o b\rangle)$ is called, $f$ belongs to group <grp> and <ob> is of class <class>.

1. Group "Math":

- abs, sign, sqrt, floor, ceiling, trunc, round, signif
- exp, log, cos, sin, tan, acos, asin, atan cosh, sinh, tanh, acosh, asinh, atanh
- lgamma, gamma, gammaCody, digamma, trigamma, tetragamma, pentagamma
- cumsum, cumprod, cummax, cummin

2. Group "Ops":

- "+", "-", "*", "/", "~", "\%\%", "\%/\%"
- "\&", "|", "!"
- "==", "!=", "<", "<=", ">=", ">"

3. Group "Summary":

- all, any
- sum, prod
- min, max
- range


## Simple Dispatching

The following builtin functions are generic as well, i.e., you can write methods for them:
[, [ [
dimnames<-, dimnames, dim<-, dim
c, unlist, as.vector, is.na, is.nan

## References

Appendix A, Classes and Methods of Chambers, J. M. and Hastie, T. J. eds (1992) Statistical Models in S. Wadsworth \& Brooks/Cole.

## See Also

 methods for methods of non-Internal generic functions.
## Examples

```
methods("Math")
methods("Ops")
methods("Summary")
d.fr <- data.frame(x=1:9, y=rnorm(9))
data.class(1 + d.fr) == "data.frame" ##-- add to d.f. ...
```

methods
Class Methods

## Description

$R$ possesses a simple generic function mechanism which can be used for an object-oriented style of programming. Method despatch takes place based on the class of the first argument to the generic function or on the object supplied as an argument to UseMethod or NextMethod.

## Usage

```
UseMethod(generic, object)
NextMethod(generic = NULL, object = NULL, ...)
methods(generic.function, class)
```


## Arguments

```
generic a character string naming a function.
object an object whose class will determine the method to be dispatched. De- faults to the first argument of the enclosing function.
... further arguments to be passed to the method.
generic.function
a generic function, or a character string naming a generic function.
class a symbol or character string naming a class: only used if generic.function is not supplied.
```


## Details

An R "object" is a data object which has a class attribute. A class attribute is a character vector giving the names of the classes which the object "inherits" from. When a generic function fun is applied to an object with class attribute c("first", "second"), the system searches for a function called fun.first and, if it finds it, applied it to the object. If no such function is found a function called fun. second is tried. If no class name produces a suitable function, the function fun. default is used.
methods can be used to find out about the methods for a particular generic function or class. See the examples below for details.

Now for some obscure details that need to appear somewhere. These comments will be slightly different than those in Appendix A of the White S Book. UseMethod creates a "new" function call with arguments matched as they came in to the generic. Any local variables defined before the call to UseMethod are retained (!?). Any statements after the call to UseMethod will not be evaluated as UseMethod does not return.

NextMethod invokes the next method (determined by the class). It does this by creating a special call frame for that method. The arguments will be the same in number, order and name as those to the current method but their values will be promises to evaluate their name in the current method and environment. Any arguments matched to ... are handled specially. They are passed on as the promise that was supplied as an argument to the current environment. (S does this differently!) If they have been evaluated in the current (or a previous environment) they remain evaluated.

NextMethod should not be called except in methods called by UseMethod. In particular it will not work inside anonymous calling functions (eg get("print.ts")(AirPassengers)).

## Note

This scheme is called $S 3$ ( S version 3). For new projects, it is recommended to use the more flexible and robust $S 4$ scheme provided in the 'methods' package after library (methods). The function .isMethodsDispatchOn() returns TRUE if the new S4 methods are available but is meant for R internal use only.

The methods function was written by Martin Maechler.

## See Also

class

## Examples

```
methods(summary)
methods(print)
methods(class = data.frame)
methods("[") ##- does not list the C-internal ones...
```

```
missing Does a Formal Argument have a Value?
```


## Description

missing can be used to test whether a value was specified as an argument to a function.

## Usage

missing( x )

## Arguments

$\mathrm{x} \quad$ a formal argument.

## Details

missing ( x ) is only reliable if x has not been altered since entering the function: in particular it will always be false after $\mathrm{x}<-\operatorname{match} . \arg (\mathrm{x})$.

The example shows how a plotting function can be written to work with either a pair of vectors giving x and y coordinates of points to be plotted or a single vector giving y values to be plotted against their indexes.

Currently missing can only be used in the immediate body of the function that defines the argument, not in the body of a nested function or a local call. This may change in the future.

## See Also

substitute for argument expression; NA for "missing values" in data.

## Examples

```
myplot <- function(x,y) {
    if(missing(y)) {
                y <- x
                x <- 1:length(y)
        }
        plot(x,y)
    }
```

    mode The (Storage) Mode of an Object
    
## Description

Get or set the type or storage mode of an object.

## Usage

```
mode(x)
mode(x) <- "<mode>"
storage.mode(x)
storage.mode(x) <- "<mode>"
```


## Arguments

x
any R object.

## Details

Both mode and storage.mode return a character string giving the (storage) mode of the object - often the same - both relying on the output of typeof ( $x$ ), see the example below.
The two assignment versions are currently identical. Both mode(x) <- newmode and storage.mode( $x$ ) <- newmode change the mode or storage.mode of object $x$ to newmode.
As storage mode "single" is only a pseudo-mode in $R$, it will not be reported by mode or storage.mode: use attr (object, "Csingle") to examine this. However, the assignment versions can be used to set the mode to "single", which sets the real mode to "double" and the "Csingle" attribute to TRUE. Setting any other mode will remove this attribute.
Note (in the examples below) that some calls have mode " (" which is S compatible.

## See Also

typeof for the R-internal "mode", attributes.

## Examples

```
sapply(options(),mode)
cex3 <- c("NULL","1","1:1","1i","list(1)","data.frame(x=1)", "pairlist(pi)",
    "c", "lm", "formals(lm)[[1]]", "formals(lm)[[2]]",
    "y~x","expression((1))[[1]]", "(y~x)[[1]]", "expression(x <- pi)[[1]][[1]]")
lex3 <- sapply(cex3, function(x) eval(parse(text=x)))
mex3 <- t(sapply(lex3, function(x) c(typeof(x), storage.mode(x), mode(x))))
dimnames(mex3) <- list(cex3, c("typeof(.)","storage.mode(.)","mode(.)"))
mex3
## This also makes a local copy of 'pi':
storage.mode(pi) <- "complex"
storage.mode(pi)
rm(pi)
```

model.extract Extract Components from a Model Frame

## Description

Returns the response, offset, subset, weights or other special components of a model frame passed as optional arguments to model.frame.

## Usage

```
model.extract(frame, component)
model.offset(x)
model.response(data, type = "any")
model.weights(x)
```


## Arguments

```
frame, x, data
```

A model frame.
component The name of a components to extract, such as "weights", "subset".
type One of "any", "numeric", "double". Using the either of latter two co- erces the result to have storage mode "double".

## Details

model.offset and model.response are equivalent to model.frame(, "offset") and model.frame(, "response") respectively.
model.weights is slightly different from model.frame(, "weights") in not naming the vector it returns.

## Value

The specified component of the model frame, usually a vector.

## See Also

model.frame, offset

## Examples

```
data(esoph)
a <- model.frame(cbind(ncases,ncontrols) ~ agegp+tobgp+alcgp, data=esoph)
model.extract(a, "response")
stopifnot(model.extract(a, "response") == model.response(a))
a <- model.frame(ncases/(ncases+ncontrols) ~ agegp+tobgp+alcgp,
    data = esoph, weights = ncases+ncontrols)
model.response(a)
model.extract(a, "weights")
a <- model.frame(cbind(ncases,ncontrols) ~ agegp,
    something = tobgp, data = esoph)
names(a)
stopifnot(model.extract(a, "something") == esoph$tobgp)
```

model.frame Extracting the "Environment" of a Model Formula

## Description

model.frame (a generic function) and its methods return a data.frame with the variables needed to use formula and any ... arguments.

## Usage

model.frame(formula, ...)
model.frame.default(formula, data = NULL, subset $=$ NULL, na.action $=$ na.fail, drop.unused.levels = FALSE, xlev = NULL, ...)

Methods for
lm glm aovlist

## Arguments

| formula <br> data | a model formula <br> data.frame, list, environment or object coercible to data.frame con- <br> taining the variables in formula. |
| :--- | :--- |
| subset | a specification of the rows to be used. Defaults to all rows. <br> how NAs are treated. The default is first, any na.action attribute of <br> data, second a na.action setting of options, and third na.fail if that <br> is unset. The "factory-fresh" default is na.omit. |
| drop.unused.levels |  |
| should factors have unused levels dropped? Defaults to FALSE. |  |

## Details

Variables in the formula, subset and in ... are looked for first in data and then in the environment of formula: see the help for formula() for further details.
First all the variables needed are collected into a data frame. Then subset expression is evaluated, and it is is used as a row index to the data frame. Then the na.action function is applied to the data frame (and may well add attributes). The levels of any factors in the data frame are adjusted according to the drop.unused.levels and xlev arguments.

## Value

A data.frame containing the variables used in formula plus those specified ....

## See Also

model.matrix for the "design matrix", formula for formulas and expand.model.frame for model.frame manipulation.

## Examples

```
data(cars)
data.class(model.frame(dist ~ speed, data = cars))
```

model.matrix Construct Design Matrices

## Description

model.matrix creates a design matrix.

## Usage

```
model.matrix (object, ...)
model.matrix(object, data = environment(object),
contrasts.arg = NULL, xlev = NULL, ...)
```


## Arguments

object an object of an appropriate class. For the default method, a model formula or terms object.
data a data frame created with model.frame.
contrasts.arg A list, whose entries are contrasts suitable for input to the contrasts function and whose names are the names of columns of data containing factors.
xlev to be used as argument of model.frame if data has no "terms" attribute.
... further arguments passed to or from other methods.

## Details

model.matrix creates a design matrix from the description given in terms (formula), using the data in data which must contain columns with the same names as would be created by a call to model.frame(formula) or, more precisely, by evaluating attr(terms(formula), "variables"). There may be other columns and the order is not important. If contrasts is specified it overrides the default factor coding for that variable.

## Value

The design matrix for a regression model with the specified formula and data.

## References

Chambers, J. M. and Hastie, T. J. eds (1992) Statistical Models in S. Chapman \& Hall, London.

## See Also

model.frame, model.extract, terms

## Examples

```
data(trees)
ff <- log(Volume) ~ log(Height) + log(Girth)
str(m <- model.frame(ff, trees))
mat <- model.matrix(ff, m)
dd <- data.frame(a = gl(3,4), b = gl(4,1,12))# balanced 2-way
options("contrasts")
model.matrix(~ a + b, dd)
model.matrix(~ a + b, dd, contrasts = list(a="contr.sum"))
model.matrix(~ a + b, dd, contrasts = list(a="contr.sum", b="contr.poly"))
m.orth <- model.matrix(~a+b, dd, contrasts = list(a="contr.helmert"))
crossprod(m.orth)# m.orth is ALMOST orthogonal
```

model.tables

Compute Tables of Results from an Aov Model Fit.

## Description

Computes summary tables for model fits, especially complex aov fits.

## Usage

```
model.tables(x, ...)
model.tables.aov(x, type = "effects", se = FALSE, cterms, ...)
model.tables.aovlist(x, type = "effects", se = FALSE, ...)
```


## Arguments

x a model object, usually produced by aov
type type of table: currently only "effects" and "means" are implemented.
se should standard errors be computed?
cterms A character vector giving the names of the terms for which tables should be computed. The default is all tables.
... further arguments passed to or from other methods.

## Details

For type = "effects" give tables of the coefficients for each term, optionally with standard errors.

For type = "means" give tables of the mean response for each combinations of levels of the factors in a term.

## Value

An object of class "tables.aov", as list which may contain components
tables A list of tables for each requested term.
n
The replication information for each term.
se
Standard error information.

## Warning

The implementation is incomplete, and only the simpler cases have been tested thoroughly. Weighted aov fits are not supported.

## Author(s)

B.D. Ripley

## See Also

aov, proj, replications, TukeyHSD, se.contrast

## Examples

```
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
    K=factor(K), yield=yield)
npk.aov <- aov(yield ~ block + N*P*K, npk)
model.tables(npk.aov, "means", se=TRUE)
## as a test, not particularly sensible statistically
options(contrasts=c("contr.helmert", "contr.treatment"))
npk.aovE <- aov(yield ~ N*P*K + Error(block), npk)
model.tables(npk.aovE, se=TRUE)
model.tables(npk.aovE, "means")
```

morley Michaelson-Morley Speed of Light Data

## Description

The classical data of Michaelson and Morley on the speed of light. The data consists of five experiments, each consisting of 20 consecutive 'runs'. The response is the speed of light measurement, suitably coded.

## Usage

```
data(morley)
```


## Format

A data frame contains the following components:
Expt The experiment number, from 1 to 5 .
Run The run number within each experiment.
Speed Speed-of-light measurement.

## Details

The data is here viewed as a randomized block experiment with 'experiment' and 'run' as the factors. 'run' may also be considered a quantitative variate to account for linear (or polynomial) changes in the measurement over the course of a single experiment.

## Source

A. J. Weekes (1986) A Genstat Primer. London: Edward Arnold.

## Examples

```
data(morley)
morley$Expt <- factor(morley$Expt)
morley$Run <- factor(morley$Run)
attach(morley)
plot(Expt, Speed, main = "Speed of Light Data", xlab = "Experiment No.")
fm <- aov(Speed ~ Run + Expt, data = morley)
summary(fm)
fm0 <- update(fm, . ~ . - Run)
anova(fm0, fm)
detach(morley)
```

```
mosaicplot Mosaic Plots
```


## Description

Plots a mosaic on the current graphics device.

## Usage

```
mosaicplot(x, main = NULL, xlab = NULL, ylab = NULL,
    sort = NULL, off = NULL, dir = NULL,
    color = FALSE, shade = FALSE, margin = NULL,
    type = c("pearson", "deviance", "FT"), ...)
mosaicplot(formula, data = NULL, ..., subset)
```


## Arguments

| x | a contingency table in array form, with optional category labels specified <br> in the dimnames (x) attribute. The table is best created by the table() <br> command. |
| :--- | :--- |
| main | character string for the mosaic title. |
| xlab, ylab | x-and y-axis labels used for the plot; by default, the first and second <br> element of names (dimnames (X)) (i.e., the name of the first and second <br> variable in $X)$. |
| sort | vector ordering of the variables, containing a permutation of the integers <br> $1: l e n g t h(d i m(x)) ~(t h e ~ d e f a u l t) . ~$ |
| off | vector of offsets to determine percentage spacing at each level of the mo- <br> saic (appropriate values are between 0 and 20 and the default is 10 at each <br> level). There should be one offset for each dimension of the contingency <br> table. |

$\left.\begin{array}{ll}\text { dir } & \begin{array}{l}\text { vector of split directions ("v" for vertical and "h" for horizontal) for each } \\ \text { level of the mosaic, one direction for each dimension of the contingency } \\ \text { table. The default consists of alternating directions, beginning with a } \\ \text { vertical split. }\end{array} \\ \text { logical or (recycling) vector of colors for color shading, used only when } \\ \text { shade is FALSE. The default color=FALSE gives empty boxes with no } \\ \text { shading. } \\ \text { a logical indicating whether to produce extended mosaic plots, or a nu- } \\ \text { meric vector of at most } 5 \text { distinct positive numbers giving the absolute } \\ \text { values of the cut points for the residuals. By default, shade is FALSE, and } \\ \text { simple mosaics are created. Using shade = TRUE cuts absolute values at } \\ \text { 2 and 4. } \\ \text { a list of vectors with the marginal totals to be fit in the log-linear model. }\end{array}\right\}$

## Details

This is a generic function. It currently has a default method (mosaicplot.default) and a formula interface (mosaicplot.formula).
Extended mosaic displays show the standardized residuals of a loglinear model of the counts from by the color and outline of the mosaic's tiles. (Standardized residuals are often referred to a standard normal distribution.) Negative residuals are drawn in shaded of red and with broken outlines; positive ones are drawn in blue with solid outlines.
For the formula method, if data is an object inheriting from classes "table" or "ftable", or an array with more than 2 dimensions, it is taken as a contingency table, and hence all entries should be nonnegative. In this case, the left-hand side of formula should be empty, and the variables on the right-hand side should be taken from the names of the dimnames attribute of the contingency table. A marginal table of these variables is computed, and a mosaic of this table is produced.
Otherwise, data should be a data frame or matrix, list or environment containing the variables to be cross-tabulated. In this case, after possibly selecting a subset of the data as specified by the subset argument, a contingency table is computed from the variables given in formula, and a mosaic is produced from this.
See Emerson (1998) for more information and a case study with television viewer data from Nielsen Media Research.

## Author(s)

S-PLUS original by John Emerson 〈emerson@stat.yale.edu〉. Modified and enhanced for R by KH.

## References

Hartigan, J.A., and Kleiner, B. (1984) A mosaic of television ratings. The American Statistician, 38, 32-35.
Emerson, J. W. (1998) Mosaic displays in S-PLUS: a general implementation and a case study. Statistical Computing and Graphics Newsletter (ASA), 9, 1, 17-23.

Friendly, M. (1994) Mosaic displays for multi-way contingency tables. Journal of the American Statistical Association, 89, 190-200.

The home page of Michael Friendly (http://www.math.yorku.ca/SCS/friendly.html) provides information on various aspects of graphical methods for analyzing categorical data, including mosaic plots.

## See Also

assocplot, loglin.

## Examples

```
data(Titanic)
mosaicplot(Titanic, main = "Survival on the Titanic", color = TRUE)
## Formula interface for tabulated data:
mosaicplot(~ Sex + Age + Survived, data = Titanic, color = TRUE)
data(HairEyeColor)
mosaicplot(HairEyeColor, shade = TRUE)
## Independence model of hair and eye color and sex. Indicates that
## there are significantly more blue eyed blond females than expected
## in the case of independence (and too few brown eyed blond females).
mosaicplot(HairEyeColor, shade = TRUE, margin = list(c(1,2), 3))
## Model of joint independence of sex from hair and eye color. Males
## are underrepresented among people with brown hair and eyes, and are
## overrepresented among people with brown hair and blue eyes, but not
## ''significantly''.
## Formula interface for raw data: visualize crosstabulation of numbers
## of gears and carburettors in Motor Trend car data.
data(mtcars)
mosaicplot(~ gear + carb, data = mtcars, color = TRUE)
mosaicplot(~ gear + carb, data = mtcars, color = 2:3)# color recycling
```

```
mtcars Motor Trend Road Tests
```


## Description

The data was extracted from the 1974 Motor Trend US magazine, and comprises fuel consumption and 10 aspects of automobile design and performance for 32 automobiles (1973-74 models).

## Usage

data(mtcars)

## Format

A data frame with 32 observations on 11 variables.

| $[, 1]$ | mpg | Miles/(US) gallon |
| :--- | :--- | :--- |
| $[, 2]$ | cyl | Number of cylinders |
| $[, 3]$ | disp | Displacement (cu.in.) |
| $[, 4]$ | hp | Gross horsepower |
| $[, 5]$ | drat | Rear axle ratio |
| $[, 6]$ | wt | Weight (lb/1000) |
| $[, 7]$ | qsec | $1 / 4$ mile time |
| $[, 8]$ | vs | V/S |
| $[, 9]$ | am | Transmission $(0=$ automatic, $1=$ manual $)$ |
| $[, 10]$ | gear | Number of forward gears |
| $[, 11]$ | carb | Number of carburettors |

## Source

Henderson and Velleman (1981), Building multiple regression models interactively. Biometrics, 37, 391-411.

## Examples

```
data(mtcars)
pairs(mtcars, main = "mtcars data")
coplot(mpg ~ disp | as.factor(cyl), data = mtcars,
    panel = panel.smooth, rows = 1)
```

mtext
Write Text into the Margins of a Plot

## Description

Text is written in one of the four margins of the current figure region or one of the outer margins of the device region.

## Usage

```
mtext(text, side = 3, line = 0, outer = FALSE, at = NA,
    adj = NA, cex = NA, col = NA, font = NA, vfont = NULL, ...)
```


## Arguments

text one or more character strings or expressions.
side $\quad$ on which side of the plot $(1=$ bottom, $2=$ left, $3=$ top, $4=$ right $)$.
line on which MARgin line, starting at 0 counting outwards.
outer use outer margins if available.
at give location in user-coordinates. If length(at)==0 (the default), the location will be determined by adj.
adj adjustment for each string. For strings parallel to the axes, adj=0 means left or bottom alignment, and adj $=1$ means right or top aligment. If adj is not a finite value (the default), the value par("las") determines the adjustment. For strings plotted parallel to the axis the default is to centre the string.
... Further graphical parameters (see text and par) ; currently supported are:
cex $\quad$ character expansion factor $($ default $=1)$.
col color to use.
font font for text.
vfont vector font for text.

## Details

The "user coordinates" in the outer margins always range from zero to one, and are not affected by the user coordinates in the figure region(s) - R is differing here from other implementations of S.
The arguments side, line, at, at, adj, the further graphical parameters and even outer can be vectors, and recycling will take place to plot as many strings as the longest of the vector arguments. Note that a vector adj has a different meaning from text.
adj $=0.5$ will centre the string, but for outer=TRUE on the device region rather than the plot region.

Parameter las will determine the orientation of the string(s). For strings plotted perpendicular to the axis the default justifcation is to place the end of the string nearest the axis on the specified line.

Note that if the text is to be plotted perpendicular to the axis, adj determines the justification of the string and the position along the axis unless at is specified.

## Side Effects

The given text is written onto the current plot.

## See Also

title, text, plot, par; plotmath for details on mathematical annotation.

## Examples

```
plot(1:10, (-4:5)^2, main="Parabola Points", xlab="xlab")
mtext("10 of them")
for(s in 1:4)
    mtext(paste("mtext(..., line= -1, {side, col, font} = ",s,
        ", cex = ", (1+s)/2, ")"), line = -1,
        side=s, col=s, font=s, cex= (1+s)/2)
mtext("mtext(..., line= -2)", line = -2)
mtext("mtext(..., line= -2, adj = 0)", line = -2, adj =0)
##--- log axis :
plot(1:10, exp(1:10), log='y', main="log='y'", xlab="xlab")
for(s in 1:4) mtext(paste("mtext(...,side=",s,")"), side=s)
```

```
n2mfrow
```


## Description

Easy Setup for plotting multiple figures (in a rectangular layout) on one page. It allows to specify a main title and uses smart defaults for several par calls.

## Usage

n2mfrow(nr.plots)

## Arguments

nr.plots integer; the number of plot figures you'll want to draw.

## Value

A length two integer vector $\mathrm{nr}, \mathrm{nc}$ giving the number of rows and columns, fulfilling nr $>=\mathrm{nc}>=1$ and $\mathrm{nr} * \mathrm{nc}>=\mathrm{nr}$.plots.

## Author(s)

Martin Maechler

## See Also

par, layout.

## Examples

```
n2mfrow(8) # 3 x 3
n <- 5 ; x <- seq(-2,2, len=51)
## suppose now that 'n' is not known {inside function}
op <- par(mfrow = n2mfrow(n))
for (j in 1:n)
    plot(x, x^j, main = substitute(x^ exp, list(exp = j)), type='l', col="blue")
sapply(1:10, n2mfrow)
```

NA Not Available / "Missing" Values

## Description

NA is a logical constant of length 1 which contains a missing value indicator. NA can be freely coerced to any other vector type.
The generic function is.na returns a logical vector of the same "form" as its argument x , containing TRUE for those elements marked NA or NaN (!) and FALSE otherwise. dim, dimnames and names attributes are preserved.
The generic function is.na<- sets elements to NA.

## Usage

NA
is.na(x)
is.na.data.frame(x)
is.na(x) <- value

## Arguments

| x | an R object to be tested. |
| :--- | :--- |
| value | a suitable index vector for use with x. |

## Details

For character vectors the value "NA" represents missingness.
is.na( x ) works elementwise when x is a list. The method dispatching is C-internal, rather than via UseMethod.

Function is.na<- may provide a safer way to set missingness. It behaves differently for factors, for example.

## See Also

NaN, is.nan, etc. and the utility function complete.cases.
na.action, na.omit, na.fail on how methods can be tuned to deal with missing values.

## Examples

```
is.na(c(1,NA)) #> F TRUE
is.na(paste(c(1,NA)))#> F FALSE
```

na.action $\quad N A$ Action

## Description

na.action is a generic function, and na.action.default its default method.

## Usage

na.action(object, ...)
na.action.default(object, ...)

## Arguments

object any object whose NA action is given.
... further arguments special methods could require.

## Value

The "NA action" which should be applied to object whenever NAs are not desired.

## See Also

```
options("na.action"), na.omit, na.fail
```


## Examples

```
na.action(c(1, NA))
```

```
na.fail Handle Missing Values in Objects
```


## Description

These generic functions are useful for dealing with NAs in e.g., data frames. na.fail returns the object if it does not contain any missing values, and signals an error otherwise. na.omit returns the object with incomplete cases removed. na.pass returns the object unchanged.

## Usage

```
na.fail(object, ...)
na.omit(object, ...)
na.exclude(object, ...)
na.pass(object, ...)
```


## Arguments

object an R object, typically a data frame
... further arguments special methods could require.

## Details

At present these will handle vectors, matrices and data frames comprising vectors and matrices (only).

If na.omit removes cases, the row numbers of the cases form the "na.action" attribute of the result, of class "omit".
na.exclude differs from na.omit only in the class of the "na.action" attribute of the result, which is "exclude". This gives different behaviour in functions making use of naresid and napredict: when na.exclude is used the residuals and predictions are padded to the correct length by inserting NAs for cases omitted by na. exclude.

## See Also

na.action; options with argument na.action for setting "NA actions"; and lm and glm for functions using these.

## Examples

```
DF <- data.frame(x = c(1, 2, 3), y = c(0, 10, NA))
na.omit(DF)
m <- as.matrix(DF)
na.omit(m)
stopifnot(all(na.omit(1:3) == 1:3)) # does not affect objects with no NA's
try(na.fail(DF))#> Error: missing values in ...
options("na.action")
```

name Variable Names or Symbols, respectively

## Description

as.symbol coerces its argument to be a symbol, or equivalently, a name. The argument must be of mode "character". as.name is an alias for as.symbol.
is.symbol (and is.name equivalently) returns TRUE or FALSE depending on whether its argument is a symbol (i.e. name) or not.

## Usage

```
as.symbol(x)
is.symbol(y)
as.name ( x )
is. name ( y )
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ objects to be coerced or tested.

## Note

The term "symbol" is from the lisp background of R, whereas "name" has been the standard S term for this.

See Also
call, is.language. For the internal object mode, typeof.

## Examples

```
an <- as.name("arrg")
is.name(an) # TRUE
str(an)# symbol
```


## Description

Functions to get or set the names of an object.

## Usage

```
names(x)
names(x) <- value
```


## Arguments

| $x$ | an $R$ object. |
| :--- | :--- |
| value | a character vector of the same length as $x$, or NULL. |

## Details

names is a generic accessor function, and names<- is a generic replacement function. The default methods get and set the "names" attribute of a vector or list.

It is possible to update just part of the names attribute via the general rules: see the examples. This works because the expression there is evaluated as $z<-$ "names<-" $(z$, " [<-"(names(z), 3, "c2")).

## Value

For names, NULL or a character vector of the same length as x .
For names<-, the updated object. (Note that the value of names $(x)$ <- value is that of the assignment, value, not the return value from the left-hand side.)

## Examples

```
# print the names attribute of the islands data set
names(islands)
# remove the names attribute
names(islands) <- NULL
z <- list(a=1, b="c", c=1:3)
names(z)
# change just the name of the third element.
names(z)[3] <- "c2"
z
```


## namespace Name Spaces

## Description

Name Space Support Functions.

## Usage

pkg: :name

## Arguments

| pgk | package name symbol or string literal. |
| :--- | :--- |
| name | variable name symbol or string literal. |

## Details

The expression pkg: :name returns the value of the exported variable name in package pkg. Currently only the base name space is supported.
Assignment into name spaces is not supported.

## Examples

```
base::log
base: :"+"
```

naprint Adjust for Missing Values

## Description

Use missing value information to report the effects of an na.action.

## Usage

naprint(x, ...)

## Arguments

x
An object produced by an na.action function.
...
further arguments passed to or from other methods.

## Details

This is a generic function, and the exact information differs by method. naprint.omit reports the number of rows omitted: naprint. default reports an empty string.

## Value

A character string providing information on missing values, for example the number.

## naresid Adjust for Missing Values

## Description

Use missing value information to adjust residuals and predictions.

## Usage

naresid(omit, x, ...)
napredict(omit, x, ...)

## Arguments

omit An object produced by an na.action function.
x
A vector, data frame, or matrix to be adjusted based upon the missing value information.
... further arguments passed to or from other methods.

## Details

These are utility functions used to allow predict and resid methods for modelling functions to compensate for the removal of NAs in the fitting process. There are used by the default, "lm" and "glm" methods, and by further methods in packages MASS, rpart and survival. The default methods do nothing. The method for the na.exclude action to pad the object with NAs in the correct positions to have the same number of rows as the original data frame.
Currently naresid and napredict are identical, but future methods need not be. naresid is used for residuals, and napredict for fitted values and predictions.

## Value

These return a similar object to x .

## Note

Packages rpart and survival5 used to contain versions of these functions that had an na. omit action equivalent to that now used for na.exclude.
nargs $\quad$ The Number of Arguments to a Function

## Description

When used inside a function body, nargs returns the number of arguments supplied to that function, including positional arguments left blank.

## Usage

nargs()

See Also
args, formals and sys.call.

## Examples

```
tst <- function(a, b = 3, ...) {nargs()}
tst() # 0
tst(clicketyclack) # 1 (even non-existing)
tst(c1, a2, rr3) # 3
foo <- function(x, y, z, w) {
    cat("call was", deparse(match.call()), "\n")
    nargs()
}
foo() # 0
foo(,,3) # 3
foo(z=3) # 1, even though this is the same call
nargs()# not really meaningful
```

```
nchar Count the Number of Characters
```


## Description

nchar takes a character vector as an argument and returns a vector whose elements contain the number of characters in the corresponding element of x .

## Usage

nchar (x)

## Arguments

x
character vector, or a vector to be coerced to a character vector.

## Details

The internal equivalent of as.character is performed on $x$. If you want to operate on non-vector objects passing them through deparse first will be required.

## Value

The number of characters as the string will be printed (integer 2 for a missing string).

## See Also

strwidth giving width of strings for plotting; paste, substr, strsplit

## Examples

```
x <- c("asfef","qwerty","yuiop[","b","stuff.blah.yech")
nchar(x)
# 5 6 6 1 15
nchar(deparse(mean))
# 23 1 1645 111 64 2 17 50 43 2 17 1
```

```
nclass
Compute the Number of Classes for a Histogram
```


## Description

Compute the number of classes for a histogram, for use internally in hist.

## Usage

nclass.Sturges(x)
nclass.scott( x )
nclass.FD(x)

## Arguments

x
A data vector.

## Details

nclass. Sturges uses Sturges' formula, implicitly basing bin sizes on the range of the data. nclass.scott uses Scott's choice for a normal distribution based on the estimate of the standard error.
nclass.FD uses the Freedman-Diaconis choice based on the inter-quartile range.

## Value

The suggested number of classes.

## Note

For consistency with earlier versions of R, nclass.Sturges rounds down. This is incompatible with S-PLUS, and probably wrong: however the other algorithms are to be preferred.

## References

Venables, W. N. and Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. Springer, pages 118-9.

Freedman, D. and Diaconis, P. (1981) On the histogram as a density estimator: $L_{2}$ theory. Zeitschrift für Wahrscheinlichkeitstheorie und verwandte Gebiete 57, 453-476.

Scott, D. W. (1979) On optimal and data-based histograms. Biometrika 66, 605-610.
Scott, D. W. (1992) Multivariate Density Estimation. Theory, Practice, and Visualization. Wiley.

See Also

hist

## NegBinomial The Negative Binomial Distribution

## Description

Density, distribution function, quantile function and random generation for the negative binomial distribution with parameters size and prob.

## Usage

dnbinom(x, size, prob, mu, log = FALSE)
pnbinom(q, size, prob, mu, lower.tail = TRUE, log.p = FALSE)
qnbinom(p, size, prob, mu, lower.tail = TRUE, log.p = FALSE)
rnbinom(n, size, prob, mu)

## Arguments

x
q vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
n number of observations. If length $(\mathrm{n})>1$, the length is taken to be the number required.
size target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution).
prob probability of success in each trial.
$\mathrm{mu} \quad$ alternative parametrization via mean: see Details
$\log , \log . \mathrm{p} \quad$ logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The negative binomial distribution with size $=n$ and prob $=p$ has density

$$
p(x)=\frac{\Gamma(x+n)}{\Gamma(n) x!} p^{n}(1-p)^{x}
$$

for $x=0,1,2, \ldots$
This represents the number of failures which occur in a sequence of Bernoulli trials before a target number of successes is reached.
A negative binomial distribution can arise as a mixture of Poisson distributions with mean distributed as a gamma (pgamma) distribution with scale parameter (1 - prob)/prob and shape parameter size. (This definition allows non-integer values of size.) In this model prob $=$ scale $(1+$ scale $)$, and the mean is size * (1 - prob)/prob)

The alternative parametrization (often used in ecology) is by the mean mu, and size, the dispersion parameter, where prob $=\mathrm{size} /(\mathrm{size}+\mathrm{mu})$. In this parametrization the variance is $m u+m u \wedge 2 / s i z e$.

If an element of x is not integer, the result of dnbinom is zero, with a warning.
The quantile is defined as the smallest value $x$ such that $F(x) \geq p$, where $F$ is the distribution function.

## Value

dnbinom gives the density, pnbinom gives the distribution function, qnbinom gives the quantile function, and rnbinom generates random deviates.

## See Also

dbinom for the binomial, dpois for the Poisson and dgeom for the geometric distribution, which is a special case of the negative binomial.

## Examples

```
x <- 0:11
dnbinom(x, size = 1, prob = 1/2) * 2^(1 + x) # == 1
126 / dnbinom(0:8, size = 2, prob = 1/2) #- theoretically integer
## Cumulative ('p') = Sum of discrete prob.s ('d'); Relative error :
summary(1 - cumsum(dnbinom(x, size = 2, prob = 1/2)) /
    pnbinom(x, size = 2, prob = 1/2))
x <- 0:15
size <- (1:20)/4
persp(x,size, dnb <- outer(x,size,function(x,s)dnbinom(x,s, pr= 0.4)),
    xlab = "x", ylab = "s", zlab="density", theta = 150)
title(tit <- "negative binomial density(x,s, pr = 0.4) vs. x & s")
image (x,size, log10(dnb), main= paste("log [",tit,"]"))
contour(x,size, log10(dnb),add=TRUE)
## Alternative parametrization
x1 <- rnbinom(500, mu = 4, size = 1)
x2 <- rnbinom(500, mu = 4, size = 10)
x3 <- rnbinom(500, mu = 4, size = 100)
h1 <- hist(x1, breaks = 20, plot = FALSE)
h2 <- hist(x2, breaks = h1$breaks, plot = FALSE)
h3 <- hist(x3, breaks = h1$breaks, plot = FALSE)
barplot(rbind(h1$counts, h2$counts, h3$counts),
    beside = TRUE, col = c("red","blue","cyan"),
    names.arg = round(h1$breaks[-length(h1$breaks)]))
```

    nextn Highly Composite Numbers
    
## Description

nextn returns the smallest integer, greater than or equal to $n$, which can be obtained as a product of powers of the values contained in factors. nextn is intended to be used to find a suitable length to zero-pad the argument of fft to so that the transform is computed quickly. The default value for factors ensures this.

## Usage

```
nextn(n, factors=c(2,3,5))
```


## Arguments

n
an integer.
factors a vector of positive integer factors.

## See Also

convolve, fft.

## Examples

```
nextn(1001) # 1024
table(sapply(599:630, nextn))
```

nhtemp Average Yearly Temperatures in New Haven

## Description

The mean annual temperature in degrees Fahrenheit in New Haven, Connecticut, from 1912 to 1971.

## Usage

data(nhtemp)

## Format

A time series of 60 observations.

## Source

Vaux, J. E. and Brinker, N. B. (1972) Cycles, 1972, 117-121.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(nhtemp)
plot(nhtemp, main = "nhtemp data",
    ylab = "Mean annual temperature in New Haven, CT (deg. F)")
```

```
nlevels The Number of Levels of a Factor
```


## Description

Return the number of levels which its argument has.

## Usage

nlevels(x)

## Arguments

x
an object, usually a factor.

## Details

If the argument is not a factor, NA is returned.
The actual factor levels (if they exist) can be obtained with the levels function.

## Examples

nlevels $(\mathrm{gl}(3,7))$ \# = 3
nlm Non-Linear Minimization

## Description

This function carries out a minimization of the function $f$ using a Newton-type algorithm. See the references for details.

## Usage

```
nlm(f, p, hessian = FALSE, typsize=rep(1, length(p)), fscale=1,
    print.level = 0, ndigit=12, gradtol = 1e-6,
    stepmax = max(1000 * sqrt(sum((p/typsize)^2)), 1000),
    steptol = 1e-6, iterlim = 100, check.analyticals = TRUE, ...)
```


## Arguments

$\mathrm{f} \quad$ the function to be minimized. If the function value has an attribute called gradient or both gradient and hessian attributes, these will be used in the calculation of updated parameter values. Otherwise, numerical derivatives are used. deriv returns a function with suitable gradient attribute. This should be a function a vector of the length of $p$ followed by any other arguments specified in dots.
p starting parameter values for the minimization.
hessian if TRUE, the hessian of $f$ at the minimum is returned.
typsize an estimate of the size of each parameter at the minimum.

| fscale | an estimate of the size of $f$ at the minimum. |
| :---: | :---: |
| print.level | this argument determines the level of printing which is done during the minimization process. The default value of 0 means that no printing occurs, a value of 1 means that initial and final details are printed and a value of 2 means that full tracing information is printed. |
| ndigit | the number of significant did |
| gradtol | a positive scalar giving the tolerance at which the scaled gradient is considered close enough to zero to terminate the algorithm. The scaled gradient is a measure of the relative change in $f$ in each direction $p$ [i] divided by the relative change in p [i]. |
| stepmax | a positive scalar which gives the maximum allowable scaled step length stepmax is used to prevent steps which would cause the optimization function to overflow, to prevent the algorithm from leaving the area of interest in parameter space, or to detect divergence in the algorithm. stepmax would be chosen small enough to prevent the first two of these occurrences, but should be larger than any anticipated reasonable step. |
| steptol | A positive scalar providing the minimum allowable relative step length. |
| iterlim | a positive integer specifying the maximum number of iterations to be performed before the program is terminated. |
| check.analyticals |  |
|  | a logical scalar specifying whether the analytic gradients and Hessians if they are supplied, should be checked against numerical derivatives at the initial parameter values. This can help detect incorrectly formulated gradients or Hessians. <br> additional arguments to $f$. |

## Details

If a gradient or hessian is supplied but evaluates to the wrong mode or length, it will be ignored if check. analyticals = TRUE (the default) with a warning. The hessian is not even checked unless the gradient is present and passes the sanity checks.
From the three methods available in the original source, we always use method " 1 " which is line search.

## Value

A list containing the following components:

| minimum | the value of the estimated minimum of $f$. |
| :--- | :--- |
| estimate | the point at which the mininum value of $f$ is obtained. |
| gradient | the gradient at the estimated minimum of $f$. |
| hessian | the hessian at the estimated minimum of $f$ (if requested). |
| code | an integer indicating why the optimization process terminated. |
| 1: relative gradient is close to zero, current iterate is probably solution. |  |
| 2: successive iterates within tolerance, current iterate is probably solu- |  |
| $\quad$ tion. |  |
| 3: last global step failed to locate a point lower than estimate. Ei- |  |
| $\quad$ ther estimate is an approximate local minimum of the function or |  |
| steptol is too small. |  |

4: iteration limit exceeded.
5: maximum step size stepmax exceeded five consecutive times. Either the function is unbounded below, becomes asymptotic to a finite value from above in some direction or stepmax is too small.
iterations the number of iterations performed.

## References

Dennis, J. E. and Schnabel, R. B. (1983) Numerical Methods for Unconstrained Optimization and Nonlinear Equations. Prentice-Hall, Englewood Cliffs, NJ.
Schnabel, R. B., Koontz, J. E. and Weiss, B. E. (1985) A modular system of algorithms for unconstrained minimization. ACM Trans. Math. Software, 11, 419-440.

## See Also

optim. optimize for one-dimensional minimization and uniroot for root finding. deriv to calculate analytical derivatives.
For nonlinear regression, nls (in package nls), may be of better use.

## Examples

```
f <- function(x) sum((x-1:length(x))^2)
nlm(f, c(10,10))
nlm(f, c(10,10), print.level = 2)
str(nlm(f, c(5), hessian = TRUE))
f <- function(x, a) sum((x-a) ^2)
nlm(f, c(10,10), a=c(3,5))
f <- function(x, a)
{
    res <- sum((x-a)^2)
    attr(res, "gradient") <- 2*(x-a)
    res
}
nlm(f, c(10,10), a=c(3,5))
## more examples, including the use of derivatives.
demo(nlm)
```

```
noquote Class for "no quote" Printing of Strings
```


## Description

These functions exist both as utilities and as an example of using class and object orientation.

## Usage

```
noquote(obj)
print.noquote(x, ...)
obj[j]
```


## Arguments

obj, x any R object; typically a vector of character strings.

Value
noquote returns its argument as an object of class "noquote". The function "[.noquote" ensures that the class is not lost by subsetting.

For (default) printing, print.noquote will be used which prints characters without quotes ("...").

## Author(s)

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## See Also

methods, class, print.

## Examples

```
letters
nql <- noquote(letters)
nql
nql[1:4] <- "oh"
nq1[1:12]
cmp.logical <- function(log.v)
{
    ## Purpose: compact printing of logicals
    log.v <- as.logical(log.v)
    noquote(if(length(log.v)==0)"()" else c(".","|")[1+log.v])
}
cmp.logical(runif(20) > 0.8)
```

Normal The Normal Distribution

## Description

Density, distribution function, quantile function and random generation for the normal distribution with mean equal to mean and standard deviation equal to sd.

## Usage

```
dnorm(x, mean=0, sd=1, log = FALSE)
pnorm(q, mean=0, sd=1, lower.tail = TRUE, log.p = FALSE)
qnorm(p, mean=0, sd=1, lower.tail = TRUE, log.p = FALSE)
rnorm(n, mean=0, sd=1)
```


## Arguments

| $\mathrm{x}, \mathrm{q}$ | vector of quantiles. |
| :--- | :--- |
| p | vector of probabilities. |
| n | number of observations. If length $(\mathrm{n})>1$, the length is taken to be the <br> number required. |
| mean | vector of means. <br> sd <br> log, log.p |
| vector of standard deviations. <br> lowical; if TRUE, probabilities p are given as $\log (\mathrm{p})$. |  |
|  | logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ |

## Details

If mean or sd are not specified they assume the default values of 0 and 1 , respectively.
The normal distribution has density

$$
f(x)=\frac{1}{\sqrt{2 \pi} \sigma} e^{-(x-\mu)^{2} / 2 \sigma^{2}}
$$

where $\mu$ is the mean of the distribution and $\sigma$ the standard deviation.
qnorm is based on Wichura's algorithm AS 241 which provides precise results up to about 16 digits.

## Value

dnorm gives the density, pnorm gives the distribution function, qnorm gives the quantile function, and rnorm generates random deviates.

## References

Wichura, M. J. (1988) Algorithm AS 241: The Percentage Points of the Normal Distribution. Applied Statistics, 37, 477-484.

## See Also

runif and .Random.seed about random number generation, and dlnorm for the Lognormal distribution.

## Examples

```
dnorm(0) == 1/ sqrt(2*pi)
dnorm(1) == exp(-1/2)/ sqrt(2*pi)
dnorm(1) == 1/ sqrt(2*pi*exp(1))
## Using "log = TRUE" for an extended range :
par(mfrow=c (2,1))
plot(function(x)dnorm(x, log=TRUE), -60, 50, main = "log { Normal density }")
curve(log(dnorm(x)), add=TRUE, col="red",lwd=2)
mtext("dnorm(x, log=TRUE)", adj=0); mtext("log(dnorm(x))", col="red", adj=1)
plot(function(x)pnorm(x, log=TRUE), -50, 10, main = "log { Normal Cumulative }")
curve(log(pnorm(x)), add=TRUE, col="red",lwd=2)
mtext("pnorm(x, log=TRUE)", adj=0); mtext("log(pnorm(x))", col="red", adj=1)
```

NotYet Not Yet Implemented Functions and Unused Arguments

## Description

In order to pinpoint missing functionality, the R core team uses these functions for missing $R$ functions and not yet used arguments of existing $R$ functions (which are typically there for compatibility purposes).
You are very welcome to contribute your code ...

## Usage

.NotYetImplemented()
.NotYetUsed(arg, error = TRUE)

## Arguments

$$
\begin{array}{ll}
\text { arg } & \text { an argument of a function that is not yet used. } \\
\text { error } & \text { a logical. If TRUE, an error is signalled; if FALSE; only a warning is given. }
\end{array}
$$

## See Also

the contrary, Deprecated and Defunct for outdated code.

## Examples

```
plot.mlm # to see how the ''NotYetImplemented''
    # reference is made automagically
try(plot.mlm())
barplot(1:5, inside = TRUE) # 'inside' is not yet used
```

```
nrow The Number of Rows/Columns of an Array
```


## Description

nrow and ncol return the number of rows or columns present in $x$. NCOL and NROW do the same treating a vector as 1 -column matrix.

## Usage

nrow (x)
ncol(x)
NCOL (x)
NROW ( x )

## Arguments

x
a vector, array or data frame

## Value

an integer of length 1 or NULL.

## See Also

dim which returns all dimensions; array, matrix.

## Examples

```
ma <- matrix(1:12, 3, 4)
nrow(ma) # 3
ncol(ma) # 4
ncol(array(1:24, dim = 2:4)) # 3, the second dimension
NCOL(1:12) # 1
NROW(1:12) # 12
```

nsl

## Description

Interface to gethostbyname.

## Usage

nsl(hostname)

## Arguments

hostname the name of the host.

## Value

The IP address, as a character string, or NULL if the call fails.

## Note

This was included as a test of internet connectivity, to fail if the node running R is not connected. It will also return NULL if BSD networking is not supported, including the header file 'arpa/inet.h'.

## Examples

```
nsl("www.r-project.org")
```

NULL
The Null Object

## Description

NULL represents the null object in R. NULL is used mainly to represent the lists with zero length, and is often returned by expressions and functions whose value is undefined.
as.null ignores its argument and returns the value NULL.
is.null returns TRUE if its argument is NULL and FALSE otherwise.

## Usage

NULL
as.null (x, ...)
is.null(x)

## Arguments

X an object to be tested or coerced. ... ignored.

## Examples

```
is.null(list()) # FALSE (on purpose!)
is.null(integer(0))# F
is.null(logical(0))# F
as.null(list(a=1,b='c'))
```

```
numeric Numeric Vectors
```


## Description

numeric creates a real vector of the specified length. The elements of the vector are all equal to 0 .
as.numeric attempts to coerce its argument to numeric type (either integer or real).
is.numeric returns TRUE if its argument is of type real or type integer and FALSE otherwise.

## Usage

numeric (length $=0$ )
as.numeric(x, ...)
is.numeric(x)

## Arguments

| length | desired length. |
| :--- | :--- |
| x | object to be coerced or tested. |
| $\ldots$ | further arguments passed to or from other methods. |

## Note

R has no single precision data type. All real numbers are stored in double precision format. While as.numeric is a generic function, user methods must be written for as.double, which it calls
as.numeric for factors yields the codes underlying the factor levels, not the numeric representation of the labels.

## Examples

```
as.numeric(c("-.1"," 2.7 ","B")) # (-0.1, 2.7, NA) + warning
as.numeric(factor(5:10))
```

object.size
Report the Space Allocated for an Object

## Description

Provides an estimate of the memory that is being used to store an R object.

## Usage

object.size(x)

## Arguments

## x <br> An R object.

## Details

Exactly which parts of the memory allocation should be attributed to which object is not clear-cut. This function merely provides a rough indication. For example, it will not detect if character storage for character strings are shared between identical elements (which it will be if rep was used, for example).
The calculation is of the size of the object, and excludes the space needed to store its name in the symbol table.

## Value

An estimate of the memory allocation attributable to the object, in bytes.

## Examples

```
object.size(letters)
object.size(ls)
## find the 10 largest objects in base
z <- sapply(ls("package:base"), function(x) object.size(get(x)))
as.matrix(rev(sort(z)) [1:10])
```


## octmode Display Numbers in Octal

## Description

Convert or print integers in octal format, with as many digits as are needed to display the largest, using leading zeroes as necessary.

## Usage

as.character (x, ...)
format(x, ...)
print(x, ...)

## Arguments

$x \quad$ An object inheriting from class "octmode".
... further arguments passed to or from other methods.

## Details

Class "octmode" consists of integer vectors with that class atttribute, used merely to ensure that they are printed in octal notation, specifically for Unix-like file permissions such as 755.

## See Also

These are auxiliary functions for file.info

```
offset Include an Offset in a Model Formula
```


## Description

An offset is a term to be added to a linear predictor, such as in a generalised linear model, with known coefficient 1 rather than an estimated coefficient.

## Usage

```
offset(object)
```


## Arguments

object An offset to be included in a model frame

## Value

The input value.

## See Also

model.offset, model.frame, glm

```
on.exit
Function Exit Code
```


## Description

on.exit records the expression given as its argument as needing to be executed when the current function exits (either naturally or as the result of an error). This is useful for resetting graphical parameters or performing other cleanup actions.

If no expression is provided, i.e. the call is on.exit() then the current on.exit code is removed.

## Usage

on.exit(expr, add = FALSE)

## Arguments

expr an expression to be executed.
add if TRUE, add expr to be executed after any previously set expressions.

## See Also

sys.on.exit to see the current expression.

## Examples

```
opar <- par(mai = c(1,1,1,1))
on.exit(par(opar))
```

optim General-purpose Optimization

## Description

General-purpose optimization based on Nelder-Mead, quasi-Newton and conjugate-gradient algorithms. It includes an option for box-constrained optimization.

## Usage

```
optim(par, fn, gr = NULL,
    method = c("Nelder-Mead", "BFGS", "CG", "L-BFGS-B", "SANN"),
    lower = -Inf, upper = Inf,
    control = list(), hessian = FALSE, ...)
```


## Arguments

| par | Initial values for the parameters to be optimized over. <br> A function to be minimized (or maximized), with first argument the vector <br> of parameters over which minimization is to take place. It should return <br> a scalar result. |
| :--- | :--- |
| gr | A function to return the gradient. Not needed for the "Nelder-Mead" <br> and "SANN" method. If it is NULL and it is needed, a finite-difference <br> approximation will be used. |
| method | The method to be used. See Details. |
| lower, upper | Bounds on the variables for the "L-BFGS-B" method. |
| control | A list of control parameters. See Details. |
| hessian | Logical. Should a numerically differentiated Hessian matrix be returned? |
| .. | Further arguments to be passed to fn and gr. |

## Details

By default this function performs minimization, but it will maximize if control\$fnscale is negative.
The default method is an implementation of that of Nelder and Mead (1965), that uses only function values and is robust but relatively slow. It will work reasonably well for non-differentiable functions.
Method "BFGS" is a quasi-Newton method (also known as a variable metric algorithm), specifically that published simultaneously in 1970 by Broyden, Fletcher, Goldfarb and Shanno. This uses function values and gradients to build up a picture of the surface to be optimized.
Method "CG" is a conjugate gradients method based on that by Fletcher and Reeves (1964) (but with the option of Polak-Ribiere or Beale-Sorenson updates). Conjugate gradient methods will generally be more fragile that the BFGS method, but as they do not store a matrix they may be successful in much larger optimization problems.
Method "L-BFGS-B" is that of Byrd et. al. (1994) which allows box constraints, that is each variable can be given a lower and/or upper bound. The initial value must satisfy the constraints. This uses a limited-memory modification of the BFGS quasi-Newton method. If non-trivial bounds are supplied, this method will be selected, with a warning.

Nocedal and Wright (1999) is a comprehensive reference for the previous three methods.
Method "SANN" is a variant of simulated annealing given in Belisle (1992). Simulatedannealing belongs to the class of stochastic global optimization methods. It uses only function values but is relatively slow. It will also work for non-differentiable functions. This implementation uses the Metropolis function for the acceptance probability. The next candidate point is generated from a Gaussian Markov kernel with scale proportional to the actual temperature. Temperatures are decreased according to the logarithmic cooling schedule as given in Belisle (1992, p. 890). Note that the "SANN" method depends critically on the settings of the control parameters. It is not a general-purpose method but can be very useful in getting to a good value on a very rough surface.
Function fn can return NA or Inf if the function cannot be evaluated at the supplied value, but the initial value must have a computable finite value of fn . (Except for method "L-BFGS-B" where the values should always be finite.)
optim can be used recursively, and for a single parameter as well as many.
The control argument is a list that can supply any of the following components:
trace Integer. If positive, tracing information on the progress of the optimization is produced. Higher values may produce more tracing information: for method "L-BFGS-B" there are six levels of tracing. (To understand exactly what these do see the source code: higher levels give more detail.)
fnscale An overall scaling to be applied to the value of fn and gr during optimization. If negative, turns the problem into a maximization problem. Optimization is performed on $f n(p a r) / f n s c a l e$.
parscale A vector of scaling values for the parameters. Optimization is performed on par/parscale and these should be comparable in the sense that a unit change in any element produces about a unit change in the scaled value.
ndeps A vector of step sizes for the finite-difference approximation to the gradient, on par/parscale scale. Defaults to 1e-3.
maxit The maximum number of iterations. Defaults to 100 for the derivative-based methods, and 500 for "Nelder-Mead". For "SANN" maxit gives the total number of function evaluations. There is no other stopping criterion. Defaults to 10000.
abstol The absolute convergence tolerance. Only useful for non-negative functions, as a tolerance for reaching zero.
reltol Relative convergence tolerance. The algorithm stops if it is unable to reduce the value by a factor of reltol * (abs (val) + reltol) at a step. Defaults to sqrt (. Machine\$double.eps), typically about $1 \mathrm{e}-8$.
alpha, beta, gamma Scaling parameters for the "Nelder-Mead" method. alpha is the reflection factor (default 1.0), beta the contraction factor (0.5) and gamma the expansion factor (2.0).
REPORT The frequency of reports for the "BFGS" and "L-BFGS-B" methods if control\$trace is positive. Defaults to every 10 iterations.
type for the conjugate-gradients method. Takes value 1 for the Fletcher-Reeves update, 2 for Polak-Ribiere and 3 for Beale-Sorenson.
1 mm is an integer giving the number of BFGS updates retained in the "L-BFGS-B" method, It defaults to 5 .
factr controls the convergence of the "L-BFGS-B" method. Convergence occurs when the reduction in the objective is within this factor of the machine tolerance. Default is 1 e 7 , that is a tolerance of about $1 \mathrm{e}-8$.
pgtol helps controls the convergence of the "L-BFGS-B" method. It is a tolerance on the projected gradient in the current search direction. This defaults to zero, when the check is suppressed.
temp controls the "SANN" method. It is the starting temperature for the cooling schedule. Defaults to 10 .
tmax is the number of function evaluations at each temperature for the "SANN" method. Defaults to 10 .

## Value

A list with components:
par $\quad$ The best set of parameters found.
value The value of fn corresponding to par.
counts A two-element integer vector giving the number of calls to fn and gr respectively. This excludes those calls needed to compute the Hessian, if requested, and any calls to fn to compute a finite-difference approximation to the gradient.

```
convergence An integer code. 0 indicates successful convergence. Error codes are
    1 indicates that the iteration limit maxit had been reached.
    10 indicates degeneracy of the Nelder-Mead simplex.
    51 indicates a warning from the "L-BFGS-B" method; see component
    message for further details.
52 indicates an error from the "L-BFGS-B" method; see component
    message for further details.
message A character string giving any additional information returned by the op-
timizer, or NULL.
hessian Only if argument hessian is true. A symmetric matrix giving an estimate of the Hessian at the solution found. Note that this is the Hessian of the unconstrained problem even if the box constraints are active.
```


## Note

optim will work with one-dimensional pars, but the default method does not work well (and will warn). Use optimize instead.
The code for methods "Nelder-Mead", "BFGS" and "CG" was based originally on Pascal code in Nash (1990) that was translated by p2c and then hand-optimized. Dr Nash has agreed that the code can be made freely available.
The code for method "L-BFGS-B" is based on Fortran code by Zhu, Byrd, Lu-Chen and Nocedal obtained from Netlib (file opt/lbfgs_bcm.shar: another version is in toms/778).
The code for method "SANN" was contributed by A. Trapletti.

## References

Belisle, C. J. P. (1992) Convergence theorems for a class of simulated annealing algorithms on $R^{d}$. J Applied Probability, 29, 885-895.

Byrd, R. H., Lu, P., Nocedal, J. and Zhu, C. (1995) A limited memory algorithm for bound constrained optimization. SIAM J. Scientific Computing, 16, 1190-1208.

Fletcher, R. and Reeves, C. M. (1964) Function minimization by conjugate gradients. Computer Journal 7, 148-154.
Nash, J. C. (1990) Compact Numerical Methods for Computers. Linear Algebra and Function Minimisation. Adam Hilger.
Nelder, J. A. and Mead, R. (1965) A simplex algorithm for function minimization. Computer Journal 7, 308-313.

Nocedal, J. and Wright, S. J. (1999) Numerical Optimization. Springer.

## See Also

nlm, optimize

## Examples

```
fr <- function(x) { ## Rosenbrock Banana function
    x1 <- x[1]
    x2 <- x[2]
    100 * (x2 - x1 * x1)^2 + (1 - x1)^2
}
grr <- function(x) { ## Gradient of 'fr'
```

```
    x1 <- x[1]
    x2 <- x[2]
    c(-400 * x1 * (x2 - x1 * x1) - 2 * (1 - x1),
    200 * (x2 - x1 * x1))
}
optim(c(-1.2,1), fr)
optim(c(-1.2,1), fr, grr, method = "BFGS")
optim(c(-1.2,1), fr, NULL, method = "BFGS", hessian = TRUE)
optim(c(-1.2,1), fr, grr, method = "CG")
optim(c(-1.2,1), fr, grr, method = "CG", control=list(type=2))
optim(c(-1.2,1), fr, grr, method = "L-BFGS-B")
flb <- function(x)
    { p <- length(x); sum(c(1, rep(4, p-1)) * (x - c(1, x[-p]) ^2)^2) }
## 25-dimensional box constrained
optim(rep(3, 25), flb, NULL, "L-BFGS-B",
    lower=rep(2, 25), upper=rep(4, 25)) # par[24] is *not* at boundary
## "wild" function , global minimum at about -15.81515
fw <- function (x)
    10*sin}(0.3*x)*\operatorname{sin}(1.3*x^2) + 0.00001*x^4 + 0.2*x+80
plot(fw, -50, 50, n=1000, main = "optim() minimising 'wild function'")
res <- optim(50, fw, method="SANN",
    control=list(maxit=20000, temp=20, parscale=20))
res
## Now improve locally
(r2 <- optim(res$par, fw, method="BFGS"))
points(r2$par, r2$val, pch = 8, col = "red", cex = 2)
```


## optimize One Dimensional Optimization

## Description

The function optimize searches the interval from lower to upper for a minimum or maximum of the function $f$ with respect to its first argument.
It uses Fortran code (from Netlib) based on algorithms given in the reference.
optimise is an alias for optimize.

## Usage

```
optimize(f = , interval = , lower = min(interval),
    upper = max(interval), maximum = FALSE,
    tol = .Machine$double.eps^0.25, ...)
optimise(f = , interval = , lower = min(interval),
    upper = max(interval), maximum = FALSE,
    tol = .Machine$double.eps^0.25, ...)
```


## Arguments

the function to be optimized. The function is either minimized or maximized over its first argument depending on the value of maximum.

| interval | a vector containing the end-points of the interval to be searched for the <br> minimum. |
| :--- | :--- |
| lower | the lower end point of the interval to be searched. |
| upper | the upper end point of the interval to be searched. |
| maximum | logical. Should we maximize or minimize (the default)? |
| tol | the desired accuracy. |
| $\ldots$ | additional arguments to f. |

## Value

A list with components minimum (or maximum) and objective which give the location of the minimum (or maximum) and the value of the function at that point.

## References

Brent, R. (1973) Algorithms for Minimization without Derivatives. Englewood Cliffs N.J.: Prentice-Hall.

## See Also

nlm, uniroot.

## Examples

```
f <- function (x,a) (x-a)^2
xmin <- optimize(f, c(0, 1), tol = 0.0001, a = 1/3)
xmin
```

```
options
Options Settings
```


## Description

options allows the user to set and examine a variety of global "options" which affect the way in which R computes and displays its results.

## Usage

```
options(...)
getOption(x)
.Options
```


## Arguments

| $\ldots$. | any options can be defined, using name = value. |
| :--- | :--- |
|  | However, only the ones below are used in "base R". |
|  | Further, options ('name') == options () ['name'], see the example. |
| prompt | a string, used for R's prompt; should usually end in a blank (" "). <br> continue |

width controls the number of characters on a line. You may want to change this if you re-size the window that $R$ is running in. Valid values are $10 \ldots 10000$ with default normally 80 . (The valid values are in file 'Print.h' and can be changed by re-compiling R.)
digits controls the number of digits to print when printing numeric values. It is a suggestion only. Valid values are $1 . . .22$ with default 7. See print.default.
editor sets the default text editor, e.g., for edit. Set from the environment variable VISUAL on UNIX.
pager the (stand-alone) program used for displaying ASCII files on R's console. Defaults to '\$R_HOME/bin/pager'.
browser default HTML browser used by help.start () on UNIX, or a non-default browser on Windows.
mailer default mailer used by bug.report(). can be "none".
contrasts the default contrasts used in model fitting such as with aov or lm. A character vector of length two, the first giving the function to be used with unordered factors and the second the function to be used with ordered factors.
expressions sets a limit on the number of nested expressions that will be evaluated. This is especially important on the Macintosh since stack overflow is likely if this is set too high. Valid values are $25 . . .100000$ with default 500 .
keep.source When TRUE, the source code for functions (newly defined or loaded) is stored in their "source" attribute (see attr) allowing comments to be kept in the right places.
The default is interactive(), i.e., TRUE for interactive use.
keep.source.pkgs
As for keep.source, for functions in packages loaded by library or require. Defaults to FALSE unless the environment variable R_KEEP_PKG_SOURCE is set to yes.
na.action the name of a function for treating missing values (NA's) for certain situations.
papersize the default paper format used by postscript; set by environment variable R_PAPERSIZE when R is started and defaulting to "a4" if that is unset or invalid.
printcmd the command used by postscript for printing; set by environment variable R_PRINTCMD when $R$ is started. This should be a command that expects either input to be piped to 'stdin' or to be given a single filename argument.
latexcmd, dvipscmd
character strings giving commands to be used in off-line printing of help pages.
show.signif.stars, show.coef.Pvalues
logical, affecting P value printing, see print.coefmat.
ts.eps the relative tolerance for certain time series (ts) computations.
error either a function or an expression governing the handling of noncatastrophic errors such as those generated by stop as well as by signals and internally detected errors. If the option is a function, a call to that
function, with no arguments, is generated as the expression. The default value is NULL: see stop for the behaviour in that case. The function dump.frames provides one alternative that allows post-mortem debugging.
a logical. Should error messages be printed? Intended for use with try or a user-installed error handler.
warn sets the handling of warning messages. If warn is negative all warnings are ignored. If warn is zero (the default) warnings are stored until the toplevel function returns. If fewer than 10 warnings were signalled they will be printed otherwise a message saying how many (max 50 ) were signalled. A top-level variable called last.warning is created and can be viewed through the function warnings. If warn is one, warnings are printed as they occur. If warn is two or larger all warnings are turned into errors.
warning.length
sets the truncation limit for error and warning messages. A non-negative integer, with allowed values 100-8192, default 1000 .
warning.expression
an R code expression to be called if a warning is generated, replacing the standard message. If non-null is called irrespective of the value of option warn.
check.bounds logical, defaulting to FALSE. If true, a warning is produced whenever a "generalized vector" (atomic or list) is extended, by something like x <$1: 3$; $x[5]<-6$.
echo logical. Only used in non-interactive mode, when it controls whether input is echoed. Command-line option --slave sets this initially to FALSE.
verbose logical. Should R report extra information on progress? Set to TRUE by the command-line option --verbose.
device a character string giving the default device for that session. This defaults to the normal screen device (e.g. x11, windows or gtk) for an interactive session, and postscript in batch use or if a screen is not available.

X11colortype The default colour type for X11 devices.
CRAN The URL of the preferred CRAN node for use by update.packages. Defaults to http://cran.r-project.org.
download.file.method
Method to be used for download.file. Currently download methods "internal", "wget" and "lynx" are available. There is no default for this option, when method = "auto" is chosen: see download.file.
unzip the command used for unzipping help files. Defaults to the value of R_UNZIPCMD, which is set in 'etc/Renviron' if an unzip command was found during configuration.
de.cellwidth integer: the cell widths (number of characters) to be used in the data editor dataentry. If this is unset, 0 , negative or NA, variable cell widths are used.
encoding An integer vector of length 256 holding an input encoding. Defaults to native.enc ( $=0: 255$ ). See connections.
timeout integer. The timeout for Internet operations, in seconds. Default 60 seconds.
internet.info The minimum level of information to be printed on url downloads etc. Default is 2 , for failure causes. Set to 1 or 0 to get more information.

X a character string holding one of the above option names.

## Details

Invoking options () with no arguments returns a list with the current values of the options. Note that not all options listed above are set initially. To access the value of a single option, one should use getOption("width"), e.g., rather than options("width") which is a list of length one.
.Options also always contains the options() list, for S compatibility. You must use it "read only" however.
The default settings of some of these options are

| prompt | "> " | continue | "+ " |
| :--- | :--- | :--- | :--- |
| width | 80 | digits | 7 |
| expressions | 500 | keep.source | TRUE |
| show.signif.stars | TRUE | show.coef.Pvalues | TRUE |
| na.action | na.omit | ts.eps | 1 e-5 |
| error | NULL | show.error.messags | TRUE |
| warn | 0 | warning.length | 1000 |
| echo | TRUE | verbose | FALSE |

Others are set from environment variables or are platform-dependent.

## Value

A list (in any case) with the previous values of the options changed, or all options when no arguments were given.

## Examples

```
options() # printing all current options
op <- options(); str(op) # nicer printing
# .Options is the same:
all(sapply(1:length(op), function(i) all(.Options[[i]] == op[[i]])))
options('width')[[1]] == options()$width # the latter needs more memory
options(digits=20)
pi
# set the editor, and save previous value
old.o <- options(editor="nedit")
old.o
options(check.bounds = TRUE)
x <- NULL; x[4] <- "yes" # gives a warning
options(op) # reset (all) initial options
options('digits')
## set contrast handling to be like S
options(contrasts=c("contr.helmert", "contr.poly"))
```

```
## on error, terminate the R session with error status 66
options(error=quote(q("no", status=66, runLast=FALSE)))
stop("test it")
## set an error action for debugging: see ?debugger.
options(error=dump.frames)
## A possible setting for non-interactive sessions
options(error=quote({dump.frames(to.file=TRUE); q()}))
```

OrchardSprays Potency of Orchard Sprays

## Description

An experiment was conducted to assess the potency of various constituents of orchard sprays in repelling honeybees, using a Latin square design.

## Usage

data(OrchardSprays)

## Format

A data frame with 64 observations on 4 variables.

| $[, 1]$ | rowpos | numeric | Row of the design |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | colpos | numeric | Column of the design |
| $[, 3]$ | treatment | factor | Treatment level |
| $[, 4]$ | decrease | numeric | Response |

## Details

Individual cells of dry comb were filled with measured amounts of lime sulphur emulsion in sucrose solution. Seven different concentrations of lime sulphur ranging from a concentration of $1 / 100$ to $1 / 1,562,500$ in successive factors of $1 / 5$ were used as well as a solution containing no lime sulphur.
The responses for the different solutions were obtained by releasing 100 bees into the chamber for two hours, and then measuring the decrease in volume of the solutions in the various cells.

An $* \times 8$ Latin square design was used and the treatments were coded as follows:

$$
\begin{array}{cl}
\text { A } & \text { highest level of lime sulphur } \\
\text { B } & \text { next highest level of lime sulphur } \\
\cdot & \\
\cdot & \\
\cdot & \\
\text { G } & \text { lowest level of lime sulphur } \\
\text { H } & \text { no lime sulphur }
\end{array}
$$

## Source

Finney, D. J. (1947) Probit Analysis. Cambridge.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(OrchardSprays)
pairs(OrchardSprays, main = "OrchardSprays data")
```


## order

Ordering Permutation

## Description

order returns a permutation which rearranges its first argument into ascending or descending order, breaking ties by further arguments. sort.list is the same, using only one argument.

## Usage

order(..., na.last = TRUE, decreasing = FALSE)
sort.list(x, partial = NULL, na.last = TRUE, decreasing = FALSE)

## Arguments

... a sequence of vectors, all of the same length.
$x$ a vector.
partial vector of indices for partial sorting.
decreasing logical. Should the sort order be increasing or decreasing?
na.last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed.

## Details

In the case of ties in the first vector, values in the second are used to break the ties. If the values are still tied, values in the later arguments are used to break the tie (see the first example).
partial is supplied for compatibility with other implementations of $S$, but no other values are accepted and ordering is always complete.

## See Also

sort and rank.

## Examples

```
(ii <- order(x <- c(1,1,3:1,1:4,3), y <- c(9,9:1), z <-c(2,1:9)))
## 6 5 5 2 1 1 7 4 10 8 3 9
rbind(x,y,z)[,ii] # shows the reordering (ties via 2nd & 3rd arg)
## Suppose we wanted descending order on y. A simple solution is
rbind(x,y,z)[, order(x, -y, z)]
## For character vectors we can make use of rank:
cy <- as.character(y)
rbind(x,y,z)[, order(x, -rank(y), z)]
## rearrange matched vectors so that the first is in ascending order
x <- c(5:1, 6:8, 12:9)
y <- (x - 5) ^2
o <- order(x)
rbind(x[o], y[o])
## tests of na.last
a <- c(4, 3, 2, NA, 1)
b <- c(4, NA, 2, 7, 1)
z <- cbind(a, b)
(o <- order(a, b)); z[o, ]
(o <- order(a, b, na.last = FALSE)); z[o, ]
(o <- order(a, b, na.last = NA)); z[o, ]
```

```
outer Outer Product of Arrays
```


## Description

The outer product of the arrays X and Y is the array A with dimension $\mathrm{c}(\operatorname{dim}(\mathrm{X})$, $\operatorname{dim}(Y))$ where element A[c(arrayindex.x, arrayindex.y)] = FUN(X[arrayindex.x], Y[arrayindex.y], ...).

## Usage

outer (X, Y, FUN="*", ...)
x \% $\%$ y

## Arguments

| X | A vector or array. |
| :--- | :--- |
| Y | A vector or array. |
| FUN | a function to use on the outer products, it may be a quoted string. |
| $\ldots$ | optional arguments to be passed to FUN. |

## Details

FUN must be a function (or the name of it) which expects at least two arguments and which operates elementwise on arrays.

Where they exist, the [dim]names of X and Y will be preserved.
$\% \%$ is an alias for outer (where FUN cannot be changed from "*").

## Author(s)

Jonathan Rougier

## See Also

$\% * \%$ for usual (inner) matrix vector multiplication; kronecker which is based on outer.

## Examples

```
x <- 1:9; names(x) <- x
# Multiplication & Power Tables
x %o% x
y <- 2:8; names(y) <- paste(y,":",sep="")
outer(y, x, "^")
outer(month.abb, 1999:2003, FUN = "paste")
## three way multiplication table:
x %%% x %%% y [1:3]
```

p.adjust Adjust p-values for multiple comparisons

## Description

Given a set of p-values, returns p-values adjusted using one of several methods.

## Usage

p.adjust(p, method=p.adjust.methods, n=length(p))
p.adjust.methods

## Arguments

| p | vector of p-values |
| :--- | :--- |
| method | correction method |
| n | number of comparisons |

## Details

The adjustment methods include the Bonferroni correction ("bonferroni") in which the p-values are multiplied by the number of comparisons. Four less conservative corrections are also included by Holm (1979) ("holm"), Hochberg (1988) ("hochberg"), Hommel (1988) ("hommel") and Benjamini \& Hochberg (1995) ("fdr"), respectively. A pass-through option ("none") is also included. The set of methods are contained in the p.adjust.methods vector for the benefit of methods that need to have the method as an option and pass it on to p.adjust.

The first four methods are designed to give strong control of the family wise error rate. There seems no reason to use the unmodified Bonferroni correction because it is dominated by Holm's method, which is also valid under arbitrary assumptions.
Hochberg's and Hommel's methods are valid when the hypothesis tests are independent or when they are non-negatively associated (Sarkar, 1998; Sarkar and Chang, 1997). Hommel's
method is more powerful than Hochberg's, but the difference is usually small and the Hochberg p-values are faster to compute.

The "fdr" method of Benjamini and Hochberg (1995) controls the false discovery rate, the expected proportion of false discoveries amongst the rejected hypotheses. The false discovery rate is a less stringent condition than the family wise error rate, so Benjamini and Hochberg's method is more powerful than the other methods.

## Value

A vector of corrected $p$-values.

## References

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. Journal of the Royal Statistical Society Series B, 57, 289-300.

Holm, S. (1979). A simple sequentially rejective multiple test procedure. Scandinavian Journal of Statistics, 6, 65-70.

Hommel, G. (1988). A stagewise rejective multiple test procedure based on a modified Bonferroni test. Biometrika, 75, 383-386.

Hochberg, Y. (1988). A sharper Bonferroni procedure for multiple tests of significance. Biometrika, 75, 800-803.

Shaffer, J. P. (1995). Multiple hypothesis testing. Annual Review of Psychology, 46, 561576. (An excellent review of the area.)

Sarkar, S. (1998). Some probability inequalities for ordered MTP2 random variables: a proof of Simes conjecture. Annals of Statistics, 26, 494-504.

Sarkar, S., and Chang, C. K. (1997). Simes' method for multiple hypothesis testing with positively dependent test statistics. Journal of the American Statistical Association, 92, 1601-1608.

Wright, S. P. (1992). Adjusted P-values for simultaneous inference. Biometrics, 48, 10051013. (Explains the adjusted P-value approach.)

## See Also

```
pairwise.* functions in the ctest package, such as pairwise.t.test.
```


## Examples

```
x <- rnorm(50, m=c(rep (0,25),rep (3,25)))
p <- 2*pnorm( -abs(x))
round(p, 3)
round(p.adjust(p), 3)
round(p.adjust(p,"bonferroni"), 3)
round(p.adjust(p,"fdr"), 3)
```

package.contents Package Contents and Description

## Description

Parses and returns the 'CONTENTS' and 'DESCRIPTION' file of a package.

## Usage

```
package.contents(pkg, lib.loc = NULL)
package.description(pkg, lib.loc = NULL, fields = NULL)
```


## Arguments

| pkg | a character string with the package name. |
| :--- | :--- |
| lib.loc | a character vector describing the location of R library trees to search <br> through, or NULL. The default value of NULL corresponds to all libraries <br> currently known. |
| fields | a character vector giving the tags of fields to return (if other fields occur <br> in the file they are ignored). |

## Value

package.contents returns NA if there is no 'CONTENTS' file for the given package; otherwise, a character matrix with column names c("Entry", "Keywords", "Description") and rows giving the corresponding entries in the CONTENTS data base for each Rd file in the package.
If a 'DESCRIPTION' for the given package is found and can successfully be read, package.description returns a named character vector with the values of the (given) fields as elements and the tags as names. If not, it returns a named vector of NAs with the field tags as names if fields is not null, and NA otherwise.

## See Also

read.dcf

## Examples

```
package.contents("mva")
package.contents("mva")[, c("Entry", "Description")]
package.description("ts")
package.description("ts")[c("Package", "Version")]
## NOTE: No subscripting using '$' or abbreviated field tags!
```

package.dependencies Check Package Dependencies

## Description

Parses and checks the dependencies of a package against the currently installed version of R [and other packages].

## Usage

package.dependencies( x , check=FALSE)

## Arguments

$$
\begin{array}{ll}
\mathrm{x} & \text { A matrix of package descriptions as returned by CRAN. packages. } \\
\text { check } & \text { If TRUE, return logical vector of check results. If FALSE, return parsed list } \\
\text { of dependencies. }
\end{array}
$$

## Details

Currently we only check if the package conforms with the currently running version of R . IN the future we might add checks for inter-package dependencies.

```
See Also
    update.packages
```

package.skeleton Create a skeleton for a new package

## Description

package.skeleton automates some of the setup for a new package. It creates directories, saves functions and data to appropriate places, and creates skeleton help files and README files describing further steps in packaging.

## Usage

```
package.skeleton(name="anRpackage", list, environment=.GlobalEnv,
    path=".", force=FALSE)
```


## Arguments

| name | directory name for your package |
| :--- | :--- |
| list | vector of names of R objects to put in the package |
| environment | if list is omitted, the contents of this environment are packaged |
| path | path to put the package directories in |
| force | If FALSE will not overwrite an existing directory |

## Value

used for its side-effects.

## References

Read the "Writing R Extensions" manual for more details

## See Also

install.packages

## Examples

```
f<-function(x,y) x+y
g<-function(x,y) x-y
d<-data.frame(a=1,b=2)
e<-rnorm(1000)
package.skeleton(list=c("f","g","d","e"),name="AnExample")
```

```
packageStatus Package Management Tools
```


## Description

Summarize information about installed packages and packages available at various repositories, and automatically upgrade outdated packages. These tools will replace update. packages and friends in the future and are currently work in progress.

## Usage

```
packageStatus(lib.loc = NULL, repositories = getOption("repositories"))
summary(object, ...)
update(object, lib.loc=levels(object$inst$LibPath),
    repositories=levels(object$avail$Repository), ...)
upgrade(object, ask=TRUE, ...)
```


## Arguments

| lib.loc | a character vector describing the location of R library trees to search <br> through, or NULL. The default value of NULL corresponds to all libraries <br> currently known. |
| :--- | :--- |
| repositories | a character vector of URLs describing the location of R package reposito- <br> ries on the Internet or on the local machine. |
| object | return value of packageStatus. |
| ask | if TRUE, the user is prompted which packages should be upgraded and <br> which not. |
| $\ldots$ | currently not used. |

page

## Examples

```
x <- packageStatus()
print(x)
summary(x)
upgrade(x)
x <- update(x)
print(x)
```

page Invoke a Pager on an $R$ Object

## Description

Displays a representation of the object named by x in a pager.

## Usage

page(x, method = c("dput", "print"), ...)

## Arguments

x
the name of an R object.
method The default method is to dump the object via dput. An alternative is to print to a file.
... additional arguments for file.show. Intended for setting pager as title and delete.file are already used.

## Author(s)

B. D. Ripley

## See Also

file.show, edit, fix.
pairs Scatterplot Matrices

## Description

A matrix of scatterplots is produced.

## Usage

```
pairs(x, ...)
pairs.default(x, labels = colnames(x), panel = points, ...,
lower.panel = panel, upper.panel = panel,
diag.panel = NULL, text.panel = textPanel,
label.pos = 0.5 + has.diag/3,
cex.labels = NULL, font.labels = 1,
row1attop = TRUE, gap=1)
```


## Arguments

```
x the coordinates of points given as columns of a matrix.
labels the names of the variables.
panel function(x,y,\ldots) which is used to plot the contents of each panel of
    the display.
... graphical parameters can be given as arguments to plot.
lower.panel, upper.panel
    separate panel functions to be used below and above the diagonal respec-
    tively.
diag.panel optional function(x, ...) to be applied on the diagonals.
text.panel optional function(x, y, labels, cex, font, ...) to be applied on
    the diagonals.
label.pos y position of labels in the text panel.
cex.labels, font.labels
    graphics parameters for the text panel.
row1attop logical. Should the layout be matrix-like with row 1 at the top, or graph-
    like with row 1 at the bottom?
gap Distance between subplots, in margin lines.
```


## Details

The $i j$ th scatterplot contains $\mathrm{x}[, \mathrm{i}]$ plotted against $\mathrm{x}[, \mathrm{j}]$. The "scatterplot' can be customised by setting panel functions to appear as something completely different. The offdiagonal panel functions are passed the appropriate columns of $x$ as $x$ and $y$ : the diagonal panel function (if any) is passed a single column, and the text. panel function is passed a single ( $x, y$ ) location and the column name.

The graphical parameters pch and col can be used to specify a vector of plotting symbols and colors to be used in the plots.
The graphical parameter oma will be set by pairs. default unless supplied as an argument.

## Author(s)

Enhancements for R 1.0.0 contributed by Dr. Jens Oehlschlaegel-Akiyoshi and R-core members.

## Examples

```
data(iris)
pairs(iris[1:4], main = "Anderson's Iris Data -- 3 species",
    pch = 21, bg = c("red", "green3", "blue")[codes(iris$Species)])
data(USJudgeRatings)
pairs(USJudgeRatings)
## put histograms on the diagonal
panel.hist <- function(x, ...)
{
    usr <- par("usr"); on.exit(par(usr))
    par(usr = c(usr[1:2], 0, 1.5) )
    h <- hist(x, plot = FALSE)
```

```
    breaks <- h$breaks; nB <- length(breaks)
    y <- h$counts; y <- y/max(y)
    rect(breaks[-nB], 0, breaks[-1], y, col="cyan", ...)
}
pairs(USJudgeRatings[1:5], panel=panel.smooth,
    cex = 1.5, pch = 24, bg="light blue",
    diag.panel=panel.hist, cex.labels = 2, font.labels=2)
## put (absolute) correlations on the upper panels,
## with size proportional to the correlations.
panel.cor <- function(x, y, digits=2, prefix="", cex.cor)
{
    usr <- par("usr"); on.exit(par(usr))
    par(usr = c(0, 1, 0, 1))
    r <- abs(cor(x, y))
    txt <- format(c(r, 0.123456789), digits=digits)[1]
    txt <- paste(prefix, txt, sep="")
    if(missing(cex.cor)) cex <- 0.8/strwidth(txt)
    text(0.5, 0.5, txt, cex = cex * r)
}
pairs(USJudgeRatings, lower.panel=panel.smooth, upper.panel=panel.cor)
```

```
pairs.formula Formula Notation for Scatterplot Matrices
```


## Description

Produce a matrix of scatterplots using formula notation.

## Usage

```
pairs(formula, data = NULL, ..., subset)
```


## Arguments

| formula | a formula, such as $y \sim x$. |
| :--- | :--- |
| data | a data.frame (or list) from which the variables in formula should be taken. |
| $\ldots$ | arguments to the default pairs method and graphical parameters may also <br> be passed as arguments, see par. |
| subset | an optional vector specifying a subset of observations to be used for plot- <br> ting. |

## Details

This is a method of the generic function pairs. It operates by setting up the data from the formula specification, and then calling pairs.default.

## See Also

pairs.default

## Examples

```
data(swiss)
pairs(~ Fertility + Education + Catholic, data = swiss,
    subset = Education < 20, main = "Swiss data, Education < 20")
```

palette Set or View the Graphics Palette

## Description

View or manipulate the color palette which is used when a col= has a numeric index.

## Usage

```
palette(value)
```


## Arguments

value an optional character vector.

## Details

If value has length 1 , it is taken to be the name of a built in color palette. If value has length greater than 1 it is assumed to contain a description of the colors which are to make up the new palette (either by name or by RGB levels).
If value is omitted or has length 0 , no change is made the current palette.
Currently, the only built-in palette is "default".

## Value

The palette which was in effect. This is invisible unless the argument is omitted.

## See Also

colors for the vector of built-in "named" colors; hsv, gray, rainbow, terrain.colors,... to construct colors;
col2rgb for translating colors to RGB 3-vectors.

## Examples

```
palette() # obtain the current palette
palette(rainbow(6)) # six color rainbow
(palette(gray(seq(0,.9,len=25)))) # gray scales; print old palette
matplot(outer(1:100,1:30), type='l', lty=1,lwd=2, col=1:30,
        main = "Gray Scales Palette",
        sub = "palette(gray(seq(0,.9,len=25)))")
palette("default") # reset back to the default
```


## Palettes Color Palettes

## Description

Create a vector of n "contiguous" colors.

## Usage

rainbow(n, $s=1, v=1$, start $=0$, end $=\max (1, \mathrm{n}-1) / \mathrm{n}$, gamma = 1)
heat.colors(n)
terrain.colors(n)
topo.colors(n)
cm.colors(n)

## Arguments

n
s, v
start
end
gamma
the number of colors $(\geq 1)$ to be in the palette.
the "saturation" and "value" to be used to complete the HSV color descriptions. scriptions.
the (corrected) hue in $[0,1]$ at which the rainbow begins.
the (corrected) hue in $[0,1]$ at which the rainbow ends.
the gamma correction, see argument gamma in hsv.

## Details

Conceptually, all of these functions actually use (parts of) a line cut out of the 3-dimensional color space, parametrized by $\mathrm{hsv}(\mathrm{h}, \mathrm{s}, \mathrm{v}$, gamma), where gamma $=1$ for the foo.colors function, and hence, equispaced hues in RGB space tend to cluster at the red, green and blue primaries.

Some applications such as contouring require a palette of colors which do not "wrap around" to give a final color close to the starting one.

With rainbow, the parameters start and end can be used to specify particular subranges of hues. The following values can be used when generating such a subrange: red $=0$, yellow $=\frac{1}{6}$, green $=\frac{2}{6}$, cyan $=\frac{3}{6}$, blue $=\frac{4}{6}$ and magenta $=\frac{5}{6}$.

## Value

A character vector, cv, of color names. This can be used either to create a user-defined color palette for subsequent graphics by palette (cv), a col= specification in graphics functions or in par.

## See Also

colors, palette, hsv, rgb, gray and col2rgb for translating to RGB numbers.

## Examples

```
# A Color Wheel
pie(rep(1,12), col=rainbow(12))
##------ Some palettes -------------
ch.col <- c("rainbow(n, start=.7, end=.1)", "heat.colors(n)",
    "terrain.colors(n)", "topo.colors(n)", "cm.colors(n)")
n <- if(.Device == "postscript") 64 else 16
    # Since for screen, larger n may give color allocation problem
nt <- length(ch.col)
i <- 1:n; j <- n / nt; d <- j/6; dy <- 2*d
plot(i,i+d, type="n", yaxt="n", ylab="", main=paste("color palettes; n=",n))
for (k in 1:nt) {
    rect(i-.5,(k-1)*j+ dy, i+.4, k*j, col=eval(parse(text=ch.col[k])))
    text(2*j, k * j +dy/4, ch.col[k])
}
```

panel.smooth Simple Panel Plot

## Description

An example of a simple useful panel function to be used as argument in e.g., coplot or pairs.

## Usage

```
panel.smooth(x, y, col, bg=NA, pch, cex = 1, col.smooth = "red",
    span = 2/3, iter=3, ...)
```


## Arguments

```
\(\mathrm{x}, \mathrm{y} \quad\) numeric vectors of the same length
col,bg,pch, cex
                    numeric or character codes for the color(s), point type and size of points;
                    see also par.
col.smooth color to be used by lines for drawing the smooths.
span smoothing parameter \(f\) for lowess, see there.
iter number of robustness iterations for lowess.
... further arguments to lines.
```


## See Also

coplot and pairs where panel.smooth is typically used; lowess.

## Examples

```
data(swiss)
pairs(swiss, panel = panel.smooth, pch = ".")# emphasize the smooths
pairs(swiss, panel = panel.smooth, lwd = 2, cex= 1.5, col="blue")# hmm...
```


## Description

par can be used to set or query graphical parameters. Parameters can be set by specifying them as arguments to par in tag = value form, or by passing them as a list of tagged values.

## Usage

```
par(..., no.readonly = FALSE)
<highlevel plot> (..., <tag> = <value>)
```


## Arguments

... arguments in tag = value form, or a list of tagged values. The tags must come from the graphical parameters described below.
no.readonly logical; if TRUE and there are no other arguments, only parameters are returned which can be set by a subsequent par() call.

## Details

Parameters are queried by giving one or more character vectors to par.
par() (no arguments) or par(no.readonly=TRUE) is used to get all the graphical parameters (as a named list). Their names are currently taken from the variable .Pars. .Pars.readonly contains the names of the par arguments which are readonly.
$\boldsymbol{R} . \boldsymbol{O}$. indicates read-only arguments: These may only be used in queries, i.e., they do not set anything.

All but these $\boldsymbol{R} . \boldsymbol{O}$. and the following low-level arguments can be set as well in high-level and mid-level plot functions, such as plot, points, lines, axis, title, text, mtext:

- "ask"
- "fig", "fin"
- "mai", "mar", "mex"
- "mfrow", "mfcol", "mfg"
- "new"
- "oma", "omd", "omi"
- "pin", "plt", "ps", "pty"
- "usr"
- "xlog", "ylog"


## Value

When parameters are set, their former values are returned in an invisible named list. Such a list can be passed as an argument to par to restore the parameter values. Use $\operatorname{par}$ (no.readonly $=$ TRUE) for the full list of parameters that can be restored.
When just one parameter is queried, the value is a character string. When two or more parameters are queried, the result is a list of character strings, with the list names giving the parameters.
Note the inconsistency: setting one parameter returns a list, but querying one parameter returns a vector.

## Graphical Parameters

adj The value of adj determines the way in which text strings are justified. A value of 0 produces left-justified text, 0.5 centered text and 1 right-justified text. (Any value in $[0,1]$ is allowed, and on most devices values outside that interval will also work.) Note that the adj argument of text also allows adj $=c(x, y)$ for different adjustment in x - and y - direction.
ann If set to FALSE, high-level plotting functions do not annotate the plots they produce with axis and overall titles. The default is to do annotation.
ask logical. If TRUE, the user is asked for input, before a new figure is drawn.
bg The color to be used for the background of plots. A description of how colors are specified is given below.
bty A character string which determined the type of box which is drawn about plots. If bty is one of "०", "l", "7", "c", "u", or "]" the resulting box resembles the corresponding upper case letter. A value of " n " suppresses the box.
cex A numerical value giving the amount by which plotting text and symbols should be scaled relative to the default.
cex.axis The magnification to be used for axis annotation relative to the current.
cex.lab The magnification to be used for x and y labels relative to the current.
cex.main The magnification to be used for main titles relative to the current.
cex.sub The magnification to be used for sub-titles relative to the current.
cin $\boldsymbol{R} . \boldsymbol{O}$.; character size (width, height) in inches.
col A specification for the default plotting color. A description of how colors are specified is given below.
col.axis The color to be used for axis annotation.
col.lab The color to be used for x and y labels.
col.main The color to be used for plot main titles.
col.sub The color to be used for plot sub-titles.
cra R.O.; size of default character (width, height) in "rasters" (pixels).
crt A numerical value specifying (in degrees) how single characters should be rotated. It is unwise to expect values other than multiples of 90 to work. Compare with srt which does string rotation.
csi $\boldsymbol{R} . \boldsymbol{O}$.; height of (default sized) characters in inches.
cxy R.O.; size of default character (width, height) in user coordinate units. par("cxy") is $\operatorname{par}($ "cin") $/ \operatorname{par}($ "pin") scaled to user coordinates. Note that c(strwidth(ch), strwidth(ch)) for a given string ch is usually much more precise.
din $\boldsymbol{R} . \boldsymbol{O} . ;$ the device dimensions in inches.
err (Unimplemented; R is silent when points outside the plot region are not plotted.) The degree of error reporting desired.
fg The color to be used for the foreground of plots. This is the default color is used for things like axes and boxes around plots. A description of how colors are specified is given below.
fig A numerical vector of the form $c(x 1, x 2, y 1, y 2)$ which gives the (NDC) coordinates of the figure region in the display region of the device.
fin A numerical vector of the form $c(x, y)$ which gives the size of the figure region in inches.
font An integer which specifies which font to use for text. If possible, device drivers arrange so that 1 corresponds to plain text, 2 to bold face, 3 to italic and 4 to bold italic.
font.axis The font to be used for axis annotation.
font.lab The font to be used for x and y labels.
font.main The font to be used for plot main titles.
font.sub The font to be used for plot sub-titles.
gamma the gamma correction, see argument gamma to hsv.
lab A numerical vector of the form $\mathrm{c}(\mathrm{x}, \mathrm{y}, \mathrm{len})$ which modifies the way that axes are annotated. The values of $x$ and $y$ give the (approximate) number of tickmarks on the x and y axes and len specifies the label size. The default is $\mathrm{c}(5,5,7)$. Currently, len is unimplemented.
las numeric in $\{0,1,2,3\}$; the style of axis labels.
0: always parallel to the axis [default],
1: always horizontal,
2: always perpendicular to the axis,
3: always vertical.
Note that other string/character rotation (via argument srt to par) does not affect the axis labels.
lty The line type. Line types can either be specified as an integer ( $0=$ blank, $1=$ solid, $2=$ dashed, $3=$ dotted, $4=$ dotdash, $5=$ longdash, $6=$ twodash) or as one of the character strings "blank", "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash", where "blank" uses ‘invisible lines’ (i.e., doesn’t draw them).
Alternatively, a string of up to 8 characters (from c(0:9, "A":"F")) may be given, giving the length of line segments which are alternatively drawn and skipped. See section 'Line Type Specification' below.
lwd The line width, a positive number, defaulting to 1.
mai A numerical vector of the form c (bottom, left, top, right) which gives the margin size specified in inches.
mar A numerical vector of the form c(bottom, left, top, right) which gives the lines of margin to be specified on the four sides of the plot. The default is $c(5,4,4,2)$ +0.1 .
mex mex is a character size expansion factor which is used to describe coordinates in the margins of plots.
mfcol, mfrow $A$ vector of the form $\mathrm{c}(\mathrm{nr}, \mathrm{nc})$. Subsequent figures will be drawn in an nr -by-nc array on the device by columns ( mfcol ), or rows (mfrow), respectively. Consider the alternatives, layout and split.screen.
mfg A numerical vector of the form $\mathrm{c}(\mathrm{i}, \mathrm{j})$ where i and j indicate which figure in an array of figures is to be drawn next (if setting) or is being drawn (if enquiring). The array must already have been set by mfcol or mfrow.
For compatibility with S , the form $\mathrm{c}(\mathrm{i}, \mathrm{j}, \mathrm{nr}, \mathrm{nc}$ ) is also accepted, when nr and nc should be the current number of rows and number of columns. Mismatches will be ignored, with a warning.
mgp The margin line (in mex units) for the axis title, axis labels and axis line. The default is $c(3,1,0)$.
mkh The height in inches of symbols to be drawn when the value of pch is an integer. Completely ignored currently.
new logical, defaulting to FALSE. If set to TRUE, the next high-level plotting command (actually plot.new) should not clean the frame before drawing "as if it was on a new device".
oma A vector of the form c(bottom, left, top, right) giving the size of the outer margins in lines of text.
omd A vector of the form $\mathrm{c}(\mathrm{x} 1, \mathrm{x} 2, \mathrm{y} 1, \mathrm{y} 2)$ giving the outer margin region in NDC $(=$ normalized device coordinates), i.e., as fraction (in $[0,1]$ ) of the device region.
omi A vector of the form c(bottom, left, top, right) giving the size of the outer margins in inches.
pch Either an integer specifying a symbol or a single character to be used as the default in plotting points.
pin The width and height of the current plot in inches.
plt A vector of the form $\mathrm{c}(\mathrm{x} 1, \mathrm{x} 2, \mathrm{y} 1, \mathrm{y} 2)$ giving the coordinates of the plot region as fractions of the current figure region.
ps integer; the pointsize of text and symbols.
pty A character specifying the type of plot region to be used; "s" generates a square plotting region and " m " generates the maximal plotting region.
smo (Unimplemented) a value which indicates how smooth circles and circular arcs should be.
srt The string rotation in degrees.
tck The length of tick marks as a fraction of the smaller of the width or height of the plotting region. If $t c k=1$, grid lines are drawn. The default setting ( $t c k=N A$ ) is to use tcl $=-0.5$ (see below).
tcl The length of tick marks as a fraction of the height of a line of text. The default value is -0.5 ; setting $\mathrm{tcl}=$ NA sets $\mathrm{tck}=-0.01$ which is $\mathrm{S}^{\prime}$ default.
tmag A number specifying the enlargement of text of the main title relative to the other annotating text of the plot.
type character; the default plot type desired, see plot.default(type=...), defaulting to " p ".
usr A vector of the form $\mathrm{c}(\mathrm{x} 1, \mathrm{x} 2, \mathrm{y} 1, \mathrm{y} 2)$ giving the extremes of the user coordinates of the plotting region. When a logarithmic scale is in use (i.e., par ("xlog") is true, see below), then the x-limits will be 10 ~ $\operatorname{par}($ "usr") [1:2]. Similarly for the y-axis.
$\operatorname{xaxp} \mathrm{A}$ vector of the form $\mathrm{c}(\mathrm{x} 1, \mathrm{x} 2, \mathrm{n})$ giving the coordinates of the extreme tick marks and the number of intervals between tick-marks when par ("xlog") is false. Otherwise, when $\log$ coordinates are active, the three values have a different meaning: For a small range, n is negative, and the ticks are as in the linear case, otherwise, n is in 1:3, specifying a case number, and x1 and x2 are the lowest and highest power of 10 inside the user coordinates, par("usr") [1:2]. See axTicks() for more details.
xaxs The style of axis interval calculation to be used for the x-axis. Possible values are "r", "i", "e", "s", "d". The styles are generally controlled by the range of data or xlim, if given. Style "r" (regular) first extends the data range by 4 percent and then finds an axis with pretty labels that fits within the range. Style "i" (internal) just finds an axis with pretty labels that fits within the original data range. Style "s" (standard) finds an axis with pretty labels within which the original data range fits. Style "e" (extended) is like style "s", except that it is also ensured that there is room for plotting symbols within the bounding box. Style "d" (direct) specifies that the current axis should be used on subsequent plots. (Only " $r$ " and " $i$ " styles are currently implemented)
xaxt A character which specifies the axis type. Specifying " $n$ " causes an axis to be set up, but not plotted. The standard value is "s": for compatibility with $S$ values "l" and "e" are accepted but are equivalent to "s".
xlog R.O.; logical value (see log in plot.default). If TRUE, a logarithmic scale is in use (e.g., after plot (*, $\log =" \mathrm{x} ")$ ). For a new device, it defaults to FALSE, i.e., linear scale.
xpd A logical value or NA. If FALSE, all plotting is clipped to the plot region, if TRUE, all plotting is clipped to the figure region, and if NA, all plotting is clipped to the device region.
yaxp A vector of the form $c(y 1, y 2, n)$ giving the coordinates of the extreme tick marks and the number of intervals between tick-marks unless for log coordinates, see xaxp above.
yaxs The style of axis interval calculation to be used for the $y$-axis. See xaxs above.
yaxt A character which specifies the axis type. Specifying " n " causes an axis to be set up, but not plotted.
ylog R.O.; a logical value; see xlog above.

## Color Specification

Colors can be specified in several different ways. The simplest way is with a character string giving the color name (e.g., "red"). A list of the possible colors can be obtained with the function colors. Alternatively, colors can be specified directly in terms of there RGB components with a string of the form "\#RRGGBB" where each of the pairs RR, GG, BB consist of two hexadecimal digits giving a value in the range 00 to FF . Colors can also be specified by giving an index into a small table of colors, the palette. This provides compatibility with S. Index 0 corresponds to the background color.

Additionally, "transparent" or (integer) NA is transparent, useful for filled areas (such as the background!), and just invisible for things like lines or text.
The functions rgb, hsv, gray and rainbow provide additional ways of generating colors.

## Line Type Specification

Line types can either be specified by giving an index into a small built in table of line types ( $1=$ solid, $2=$ dashed, etc, see lty above) or directly as the lengths of on/off stretches of line. This is done with a string of up to eight characters, namely (hexadesimal) digits which give the lengths in consecutive positions in the string. For example, the string "33" specifies three units on followed by three off and "3313" specifies three units on followed by three off followed by one on and finally three off. The 'units' here are (on most devices) proportional to 1 wd , and with $l_{\mathrm{wd}}=1$ are in pixels or points.

## Note

The effect of restoring all the (settable) graphics parameters as in the examples is hard to predict if the device has been resized. Several of them are attempting to set the same things in different ways, and those last in the alphabet will win. In particular, the settings of mai, mar, pin, plt and pty interact, as do the outer margin settings, the figure layout and figure region size.

## See Also

plot.default for some high-level plotting parameters; colors, gray, rainbow, rgb; options for other setup parameters; graphic devices x11, postscript and setting up device regions by layout and split.screen.

## Examples

```
op <- par(mfrow = c(2, 2), # 2 x 2 pictures on one plot
        pty = "s") # square plotting region,
    # independent of device size
## At end of plotting, reset to previous settings:
par(op)
## Alternatively,
op <- par(no.readonly = TRUE) # the whole list of settable par's.
## do lots of plotting and par(.) calls, then reset:
par(op)
par("ylog") # FALSE
plot(1 : 12, log = "y")
par("ylog") # TRUE
plot(1:2, xaxs = "i") # 'inner axis' w/o extra space
stopifnot(par("xaxp")[1:2] == 1:2 &&
        par("usr") [1:2] == 1:2)
( nr.prof <-
    c(prof.pilots=16,lawyers=11,farmers=10, salesmen=9,physicians=9,
        mechanics=6,policemen=6,managers=6, engineers=5,teachers=4,
        housewives=3,students=3,armed.forces=1))
par(las = 3)
barplot(rbind(nr.prof)) # R 0.63.2: shows alignment problem
par(las = 0)# reset to default
ex <- function() {
    old.par <- par(no.readonly = TRUE) # all par settings which
                                    # could be changed.
    on.exit(par(old.par))
    ## ...
    ## ... do lots of par() settings and plots
    ## ...
    invisible() #-- now, par(old.par) will be executed
}
ex()
```

Paren Parentheses and Braces

## Description

Open parenthesis, (, and open brace, \{, are .Primitive functions in R.
Effectively, (is semantically equivalent to the identity function(x) $x$, whereas $\{$ is slightly more interesting, see examples.

## Usage

( ... )
\{ ... \}

## See Also

if, return, etc for other objects used in the R language itself.
Syntax for operator precedence.

## Examples

```
f <- get("(")
e <- expression(3 + 2 * 4)
f(e) == e # TRUE
do <- get("{")
do(x <- 3, y <- 2*x-3, 6-x-y); x; y
```

parse Parse Expressions

## Description

parse returns the parsed but unevaluated expressions in a list. Each element of the list is of mode expression.

## Usage

parse(file = "", n = NULL, text = NULL, prompt = "?", white = FALSE)

## Arguments

file a connection, or a character string giving the name of a file or a URL to read the expressions from. If file is "" and text is missing or NULL then input is taken from the console.
n
the number of statements to parse. If n is negative the file is parsed in its entirety.
text character vector. The text to parse. Elements are treated as if they were lines of a file.

```
prompt the prompt to print when parsing from the keyboard. NULL means to use
    R's prompt, options("prompt")[[1]].
white if TRUE then any white space separates expressions otherwise only newlines or semicolons do.
```


## Details

All versions of R accept input from a connection with end of line marked by LF (as used on Unix), CRLF (as used on DOS/Windows) or CR (as used on Mac). The final line can be incomplete, that is missing the final EOL marker.
See source for the limits on the size of functions that can be parsed (by default).

## See Also

scan, source, eval, deparse.

## Examples

```
cat("x <- c(1,4)\n x - 3 -10 ; outer(1:7,5:9)\n", file="xyz.Rdmped")
# parse 3 statements from the file "xyz.Rdmped"
parse(file = "xyz.Rdmped", n = 3)
unlink("xyz.Rdmped")
```

```
paste Concatenate Strings
```


## Description

Concatenate vectors after converting to character.

## Usage

```
paste(..., sep = " ", collapse = NULL)
```


## Arguments

| $\ldots$. | one or more R objects, to be coerced to character vectors. |
| :--- | :--- |
| sep | a character string to separate the terms. |
| collapse | an optional character string to separate the results. |

## Details

paste converts its arguments to character strings, and concatenates them (separating them by the string given by sep). If the arguments are vectors, they are concatenated term-byterm to give a character vector result.

If a value is specified for collapse, the values in the result are then concatenated into a single string, with the elements being separated by the value of collapse.

## Value

A character vector of the concatenated values.

## See Also

String manipulation with as.character, substr, nchar, strsplit; further, cat which concatenates and writes to a file, and sprintf for C like string construction.

## Examples

```
paste(1:12) # same as as.character(1:12)
paste("A", 1:6, sep = "")
paste("Today is", date())
```

pdf PDF Graphics Device

## Description

pdf starts the graphics device driver for producing PDF graphics.

## Usage

```
pdf(file = ifelse(onefile, "Rplots.pdf", "Rplot%03d.pdf"),
    width = 6, height = 6, onefile = TRUE, family = "Helvetica",
    encoding, bg, fg, pointsize)
```


## Arguments

| file | a character string giving the name of the file. |
| :--- | :--- |
| width, height |  |
| onefile | the width and height of the graphics region in inches. <br> logical: if true (the default) allow multiple figures in one file. If false, <br> generate a file name containing the page number. |
| family | the font family to be used, one of "AvantGarde", "Bookman", "Courier", <br> "Helvetica", "Helvetica-Narrow", "NewCenturySchoolbook", <br> "Palatino" or "Times". |
| encoding | the name of an encoding file. Defaults to "ISOLatin1.enc" in the <br> 'R_HOME/afm' directory, which is used if the path does not contain a |
| pointsize | path separator. An extension ".enc" can be omitted. <br> the default point size to be used. |
| fg | the default background color to be used. |
|  | the default foreground color to be used. |

## Details

pdf() opens the file file and the PDF commands needed to plot any graphics requested are sent to that file.
See postscript for details of encodings, as the internal code is shared between the drivers. The native PDF encoding is given in file 'PDFDoc.enc'.
pdf writes uncompressed PDF. It is primarily intended for producing PDF graphics for inclusion in other documents, and PDF-includers such as pdftex are usually able to handle compression.
At present the PDF is fairly simple, with each page being represented as a single stream. The $R$ graphics model does not distinguish graphics objects at the level of the driver interface.

## Note

Acrobat Reader does not use the fonts specified but rather emulates them from multiplemaster fonts. This can be seen in imprecise centring of characters, for example the multiply and divide signs in Helvetica.

## See Also

Devices, postscript

## Examples

```
## Test function for encodings
TestChars <- function(encoding="ISOLatin1")
{
    pdf(encoding=encoding)
    par(pty="s")
    plot(c(0,15), c(0,15), type="n", xlab="", ylab="")
    title(paste("Centred chars in encoding", encoding))
    grid(15, 15, lty=1)
    for(i in c(32:255)) {
        x <- i
        y <- i
        points(x, y, pch=i)
    }
    dev.off()
}
## there will be many warnings.
TestChars("ISOLatin2")
## doesn't view properly in US-spec Acrobat 5.05, but gs7.04 works.
## Lots of characters are not centred.
```

    persp Perspective Plots
    
## Description

This function draws perspective plots of surfaces over the $\mathrm{x}-\mathrm{y}$ plane. persp is a generic function.

## Usage

```
persp(x, ...)
persp.default(x = seq(0, 1, len = nrow(z)), y = seq(0, 1, len = ncol(z)), z,
    xlim = range(x), ylim = range(y), zlim = range(z, na.rm = TRUE),
    xlab = NULL, ylab = NULL, zlab = NULL, main = NULL, sub = NULL,
    theta = 0, phi = 15, r = sqrt(3), d = 1, scale = TRUE, expand = 1,
    col = "white", border = NULL, ltheta = -135, lphi = 0, shade = NA,
    box = TRUE, axes = TRUE, nticks = 5, ticktype = "simple",
    ...)
```


## Arguments



## Details

The plots are produced by first transforming the coordinates to the interval $[0,1]$. The surface is then viewed by looking at the origin from a direction defined by theta and phi. If theta and phi are both zero the viewing direction is directly down the negative y axis. Changing theta will vary the azimuth and changing phi the colatitude.

## Value

The viewing transformation matrix, say VT, a $4 \times 4$ matrix suitable for projecting 3D coordinates $(x, y, z)$ into the 2 D plane using homogenous 4 D coordinates $(x, y, z, t)$. It can be used to superimpose additional graphical elements on the 3D plot, by lines () or points (), e.g. using the function trans3d given in the last examples section below.

## See Also

contour and image.

## Examples

```
## More examples in demo(persp) !!
##
# (1) The Obligatory Mathematical surface.
# Rotated sinc function.
x <- seq(-10, 10, length= 30)
y <- x
f<- function(x,y) { r <- sqrt(x^2+y^2); 10* sin(r)/r }
z <- outer(x, y, f)
z[is.na(z)] <- 1
op <- par(bg = "white")
persp(x, y, z, theta = 30, phi = 30, expand = 0.5, col = "lightblue")
persp(x, y, z, theta = 30, phi = 30, expand = 0.5, col = "lightblue",
    ltheta = 120, shade = 0.75, ticktype = "detailed",
    xlab = "X", ylab = "Y", zlab = "Sinc( r )"
) -> res
round(res, 3)
# (2) Add to existing persp plot :
trans3d <- function(x,y,z, pmat) {
    tr <- cbind(x,y,z,1) %*% pmat
    list(x = tr[,1]/tr[,4], y= tr[,2]/tr[,4])
}
xE <- c(-10,10); xy <- expand.grid(xE, xE)
points(trans3d(xy[,1], xy[,2], 6, pm = res), col = 2, pch =16)
lines (trans3d(x, y=10, z= 6 + sin(x), pm = res), col = 3)
phi <- seq(0, 2*pi, len = 201)
r1 <- 7.725 # radius of 2nd maximum
xr <- r1 * cos(phi)
yr <- r1 * sin(phi)
lines(trans3d(xr,yr, f(xr,yr), res), col = "pink", lwd=2)## (no hidden lines)
# (3) Visualizing a simple DEM model
```

```
data(volcano)
z <- 2 * volcano # Exaggerate the relief
x <- 10 * (1:nrow(z)) # 10 meter spacing (S to N)
y <- 10 * (1:ncol(z)) # 10 meter spacing (E to W)
## Don't draw the grid lines : border = NA
par(bg = "slategray")
persp(x, y, z, theta = 135, phi = 30, col = "green3", scale = FALSE,
    ltheta = -120, shade = 0.75, border = NA, box = FALSE)
par(op)
```

phones

The World's Telephones

## Description

The number of telephones in various regions of the world (in thousands).

## Usage

data(phones)

## Format

A matrix with 7 rows and 8 columns. The columns of the matrix give the figures for a given region, and the rows the figures for a year.

The regions are: North America, Europe, Asia, South America, Oceania, Africa, Central America.

The years are: 1951, 1956, 1957, 1958, 1959, 1960, 1961.

## Source

AT\&T (1961) The World's Telephones.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(phones)
matplot(rownames(phones), phones, type = "b", log = "y",
    xlab = "Year", ylab = "Number of telephones (1000's)")
legend(1951.5, 80000, colnames(phones), col = 1:7, lty = 1:7, pch = rep(21, 7))
title(main = "phones data: log scale for response")
```

```
pictex A PicTeX Graphics Driver
```


## Description

This function produces graphics suitable for inclusion in TeX and LaTeX documents.

## Usage

```
pictex(file = "Rplots.tex", width = 5, height = 4, debug = FALSE,
    bg = "white", fg = "black")
```


## Arguments

| file | the file where output will appear. |
| :--- | :--- |
| width | The width of the plot in inches. |
| height | the height of the plot in inches. |
| debug | should debugging information be printed. |
| bg | the background color for the plot. |
| fg | the foreground color for the plot. |

## Details

This driver does not have any font metric information, so the use of plotmath is not supported.
Multiple plots will be placed as separate environments in the output file.

## Author(s)

This driver was provided by Valerio Aimale 〈valerio@svpop.com.dist.unige.it〉 of the Department of Internal Medicine, University of Genoa, Italy.

## References

Knuth, D. E. (1984) The TeXbook. Reading, MA: Addison-Wesley.
Lamport, L. (1994) LATEX: A Document Preparation System. Reading, MA: AddisonWesley.

Goossens, M., Mittelbach, F. and Samarin, A. (1994) The LATEX Companion. Reading, MA: Addison-Wesley.

## See Also

```
postscript, Devices.
```


## Examples

```
pictex()
plot(1:11,(-5:5)^2, type='b', main="Simple Example Plot")
dev.off()
##---------------------
%% LaTeX Example
\documentclass{article}
\usepackage{pictex}
\begin{document}
% . . .
\begin{figure}[h]
    \centerline{\input{Rplots.tex}}
    \caption{}
\end{figure}
% . . .
\end{document}
%%-- TeX Example --
\input pictex
$$ \input Rplots.tex $$
##--------------------
unlink("Rplots.tex")
```

pie Pie Charts

## Description

Draw a pie chart.

## Usage

```
pie(x, labels \(=\) names(x), edges \(=200\), radius \(=0.8\),
        density \(=\) NULL, angle \(=45\), col \(=\) NULL, border \(=\) NULL, lty \(=\) NULL,
        main = NULL, ...)
```


## Arguments

| x | a vector of positive quantities. The values in x are displayed as the areas <br> of pie slices. <br> a vector of character strings giving names for the slices. For empty or NA <br> labels, no pointing line is drawn either. |
| :--- | :--- |
| edges | the circular outline of the pie is approximated by a polygon with this <br> many edges. |
| radius | the pie is drawn centered in a square box whose sides range from -1 to <br> 1. If the character strings labeling the slices are long it may be necessary <br> to use a smaller radius. |
| density | the density of shading lines, in lines per inch. The default value of NULL <br> means that no shading lines are drawn. Non-positive values of 'density' <br> also inhibit the drawing of shading lines. |

angle the slope of shading lines, given as an angle in degrees (counter-clockwise).
col a vector of colors to be used in filling or shading the slices. If missing a set of 6 pastel colours is used, unless density is specified when $\operatorname{par}(" f g ")$ is used.
border, lty (possibly vectors) arguments passed to polygon which draws each slice.
main an overall title for the plot.
graphical parameters can be given as arguments to pie.

## Note

Pie charts are a very bad way of displaying information. The eye is good at judging linear measures and bad at judging relative areas. A bar chart or dot chart is a preferable way of displaying this type of data.

Cleveland (1985), p. 264: "Data that can be shown by pie charts always can be shown by a dot chart. This means that judgements of position along a common scale can be made instead of the less accurate angle judgements." This statement is based on the empirical investigations of Cleveland and McGill as well as investigations by perceptual psychologists.

Prior to R 1.5.0 this was known as piechart, which is the name of a Trellis function, so the name was changed to be compatible with S .

## References

Cleveland, W. S. (1985) The elements of graphing data. Wadsworth: Monterey, CA, USA.

## See Also

```
dotchart.
```


## Examples

```
pie(rep(1, 24), col = rainbow(24), radius = 0.9)
pie.sales <- c(0.12, 0.3, 0.26, 0.16, 0.04, 0.12)
names(pie.sales) <- c("Blueberry", "Cherry",
    "Apple", "Boston Cream", "Other", "Vanilla Cream")
pie(pie.sales) # default colours
pie(pie.sales,
    col = c("purple", "violetred1", "green3", "cornsilk", "cyan", "white"))
pie(pie.sales, col = gray(seq(0.4,1.0,length=6)))
pie(pie.sales, density = 10, angle = 15 + 10 * 1:6)
n <- 200
pie(rep(1,n), labels="", col=rainbow(n), border=NA,
    main = "pie(*, labels=\"\", col=rainbow(n), border=NA,..")
```

PkgUtils Utilities for Building and Checking Add-on Packages

## Description

Utilities for checking whether the sources of an R add-on package work correctly, and for building a source or binary package from them.

## Usage

R CMD build [options] pkgdirs
R CMD check [options] pkgdirs

## Arguments

pkgdirs a list of names of directories with sources of R add-on packages.
options further options to control the processing, or for obtaining information about usage and version of the utility.

## Details

R CMD check checks R add-on packages from their sources, performing a wide variety of diagnostic checks.
R CMD build builds $R$ source or binary packages from their sources.
Use R CMD foo --help to obtain usage information on utility foo.

## See Also

The chapter "Processing Rd format" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).
PlantGrowth Results from an Experiment on Plant Growth

## Description

Results from an experiment to compare yields (as measured by dried weight of plants) obtained under a control and two different treatment conditions.

## Usage

data(PlantGrowth)

## Format

A data frame of 30 cases on 2 variables.

| $[, 1]$ | weight | numeric |
| :--- | :--- | :--- |
| $[, 2]$ | group | factor |

The levels of group are 'ctrl', 'trt1', and 'trt2'.

## Source

Dobson, A. J. (1983) An Introduction to Statistical Modelling. London: Chapman and Hall.

## Examples

```
## One factor ANOVA example from Dobson's book, cf. Table 7.4:
data(PlantGrowth)
boxplot(weight ~ group, data = PlantGrowth, main = "PlantGrowth data",
    ylab = "Dried weight of plants", col = "lightgray",
    notch = TRUE, varwidth = TRUE)
anova(lm(weight ~ group, data = PlantGrowth))
```

```
plot Generic X-Y Plotting
```


## Description

Generic function for plotting of R objects. For more details about the graphical parameter arguments, see par.

## Usage

plot(x, y, xlim=range(x), ylim=range(y), type="p", main, xlab, ylab, ...)

## Arguments

x
y
xlim, ylim
type
the coordinates of points in the plot. Alternatively, a single plotting structure, function or any R object with a plot method can be provided. the y coordinates of points in the plot, optional if x is an appropriate structure.
the ranges to be encompassed by the x and y axes.
what type of plot should be drawn. Possible types are

- "p" for points,
- "l" for lines,
- "b" for both,
- "c" for the lines part alone of "b",
- "o" for both "overplotted",
- "h" for "histogram" like (or "high-density") vertical lines,
- "s" for stair steps,
- "S" for other steps, see Details below,
- " n " for no plotting.

All other types give a warning or an error; using, e.g., type = "punkte" being equivalent to type $=$ " $p$ " for $S$ compatibility.
main an overall title for the plot.
$\mathrm{xlab} \quad$ a title for the x axis.
$\mathrm{ylab} \quad$ a title for the y axis.
... graphical parameters can be given as arguments to plot.

## Details

For simple scatter plots, plot.default will be used. However, there are plot methods for many R objects, including functions, data.frames, density objects, etc. Use methods (plot) and the documentation for these.
The two step types differ in their $\mathrm{x}-\mathrm{y}$ preference: Going from $(x 1, y 1)$ to $(x 2, y 2)$ with $x 1<x 2$, type $=$ "s" moves first horizontal, then vertical, whereas type $=$ "S" moves the other way around.

## See Also

plot.default, plot.formula and other methods; points, lines, par.

## Examples

```
data(cars)
plot(cars)
lines(lowess(cars))
plot(sin, -pi, 2*pi)
## Discrete Distribution Plot:
plot(table(rpois(100,5)), type = "h", col = "red", lwd=10,
    main="rpois(100,lambda=5)")
## Simple quantiles/ECDF, see ecdf() {library(stepfun)} for a better one:
plot(x <- sort(rnorm(47)), type = "s", main = "plot(x, type = \"s\")")
points(x, cex = .5, col = "dark red")
```

```
plot.default The Default Scatterplot Function
```


## Description

Draw a scatter plot with "decorations" such as axes and titles in the active graphics window.

## Usage

```
plot.default(x, y = NULL, type = "p", xlim = NULL, ylim = NULL,
    log = "", main = NULL, sub = NULL, xlab = NULL, ylab = NULL,
    ann = par("ann"), axes = TRUE, frame.plot = axes,
    panel.first = NULL, panel.last = NULL,
    col = par("fg"), bg = NA, pch = par("pch"),
    cex = par("cex"), lty = par("lty"), lab = par("lab"),
    lwd = par("lwd"), asp = NA, ...)
```


## Arguments

$\mathrm{x}, \mathrm{y}$
the x and y arguments provide the x and y coordinates for the plot. Any reasonable way of defining the coordinates is acceptable. See the function xy. coords for details.

| type | 1-character string giving the type of plot desired. The following values <br> are possible, for details, see plot: "p" for points, "l" for lines, "o" for <br> overplotted points and lines, "b", "c") for (empty if "c") points joined <br> by lines, "s" and "S" for stair steps and "h" for histogram-like vertical <br> lines. Finally, "n" does not produce any points or lines. |
| :--- | :--- |
| xlim | the x limits (min,max) of the plot. |
| ylim | the y limits of the plot. |
| a character string which contains "x" if the x axis is to be logarithmic, |  |
| log if the y axis is to be logarithmic and "xy" or "yx" if both axes are to |  |

## References

Cleveland, W. S. (1985) The Elements of Graphing Data. Monterey, CA: Wadsworth.

## See Also

```
plot, plot.window, xy.coords.
```


## Examples

```
data(cars)
Speed <- cars$speed
Distance <- cars$dist
plot(Speed, Distance, panel.first = grid(8,8),
    pch = 0, cex = 1.2, col = "blue")
plot(Speed, Distance,
    panel.first = lines(lowess(Speed, Distance), lty = "dashed"),
    pch = 0, cex = 1.2, col = "blue")
## Show the different plot types
x <- 0:12
y <- sin(pi/5 * x)
op <- par(mfrow = c(3,3), mar = . 1+ c(2,2,3,1))
for (tp in c("p","l","b", "c","o","h", "s","S","n")) {
    plot(y ~ x, type = tp,
        main = paste("plot(*, type = \"",tp,"\")",sep=""))
        if(tp == "S") {
            lines(x,y, type = "s", col = "red", lty = 2)
            mtext("lines(*, type = \"s\", ...)", col = "red", cex=.8)
        }
}
par(op)
##--- Log-Log Plot with custom axes
lx <- seq(1,5, length=41)
yl <- expression(e^{-frac(1,2) * {log[10](x) }^2})
y <- exp(-.5*lx^2)
op <- par(mfrow=c(2,1), mar=par("mar")+c(0,1,0,0))
plot(10^lx, y, log="xy", type="l", col="purple",
    main="Log-Log plot", ylab=yl, xlab="x")
plot(10^lx, y, log="xy", type="o", pch='.', col="forestgreen",
    main="Log-Log plot with custom axes", ylab=yl, xlab="x",
    axes = FALSE, frame.plot = TRUE)
axis(1, at = my.at <- 10^(1:5), labels = formatC(my.at, format="fg"))
at.y <- 10^(-5:-1)
axis(2, at = at.y, labels = formatC(at.y, format="fg"), col.axis="red")
par(op)
```

plot.density Plot Method for Kernel Density Estimation

## Description

The plot method for density objects.

## Usage

```
plot(x, main = NULL, xlab = NULL, ylab = "Density", type = "l",
    zero.line = TRUE, ...)
```


## Arguments

```
\(x \quad a\) "density" object.
main, xlab, ylab, type
    plotting parameters with useful defaults.
    ... further plotting parameters.
    zero.line logical; if TRUE, add a base line at \(y=0\)
```


## Value

None.

## References

## See Also

density.

```
plot.factor Plotting Factor Variables
```


## Description

This functions implements a "scatterplot" method for factor arguments of the generic plot function. Actually, boxplot or barplot are used when appropriate.

## Usage

```
plot.factor(x, y, legend.text = levels(y), ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | numeric or factor. y may be missing. |
| :--- | :--- |
| legend.text | a vector of text used to construct a legend for the plot. Only used if y is <br> present and a factor. |
| $\ldots$ | Further arguments to plot, see also par. |

## See Also

```
plot.default, plot.formula, barplot, boxplot.
```


## Examples

```
data(PlantGrowth)
plot(PlantGrowth) # -> plot.data.frame
plot(weight ~ group, data = PlantGrowth) # numeric vector ~ factor
plot(cut(weight, 2) ~ group, data = PlantGrowth) # factor ~ factor
## passing "..." to barplot() eventually:
plot(cut(weight, 3) ~ group, data = PlantGrowth, density = 16*(1:3),col=NULL)
plot(PlantGrowth$group, axes=FALSE, main="no axes")# extremly silly
```

```
plot.formula Formula Notation for Scatterplots
```


## Description

Specify a scatterplot or add points or lines via a formula.

## Usage

```
plot(formula, data \(=\) parent.frame(), ..., subset,
            ylab = varnames[response], ask = TRUE)
points(formula, data = parent.frame(), ..., subset)
lines(formula, data = parent.frame(), ..., subset)
```


## Arguments

| formula | a formula, such as $\mathrm{y} \sim \mathrm{x}$. |
| :--- | :--- |
| data | a data.frame (or list) from which the variables in formula should be taken. |
| $\ldots$ | Further graphical parameters may also be passed as arguments, see par. |
| subset | an optional vector specifying a subset of observations to be used in the <br> fitting process. |
| ylab | the y label of the plot(s). <br> ask |
|  | logical, see par. |

## Details

Both the terms in the formula and the . . . arguments are evaluated in data enclosed in parent.frame() if data is a list or a data frame. The terms of the formula and those arguments in ... that are of the same length as data are subjected to the subsetting specified in subset. If the formula in plot.formula contains more than one non-response term, a series of plots of y against each term is given. A plot against the running index can be specified as plot ( $\mathrm{y}^{\sim} 1$ ).
If y is an object (ie. has a class attribute) then plot.formula looks for a plot method for that class first.

## Value

These functions are invoked for their side effect of drawing in the active graphics device.

## See Also

```
plot.default, plot.factor.
```


## Examples

```
data(airquality)
op <- par(mfrow=c(2,1))
plot(Ozone ~ Wind, data = airquality, pch=as.character(Month))
plot(Ozone ~ Wind, data = airquality, pch=as.character(Month),
    subset = Month != 7)
par(op)
```

```
plot.histogram Plot Histograms
```


## Description

These are methods for objects of class "histogram", typically produced by hist.

## Usage

```
plot(x, freq = equidist, density = NULL, angle = 45,
    col = NULL, border = par("fg"),
    lty = NULL, main = paste("Histogram of", x$xname),
    xlim = range(x$breaks), ylim = range(y, 0),
    xlab = x$xname, ylab, axes = TRUE, labels = FALSE,
    add = FALSE, ...)
lines(x, ...)
```


## Arguments

x
a histogram object, or a list with components intensities, mid, etc, see hist for information about the components of x .
freq logical; if TRUE, the histogram graphic is to present a representation of frequencies, i.e, $\mathrm{x} \$$ counts; if FALSE, relative frequencies ("probabilities"), i.e., $x \$$ intensities, are plotted. The default is true for equidistant breaks and false otherwise.
col a colour to be used to fill the bars. The default of NULL yields unfilled bars.
border the color of the border around the bars.
angle, density
select shading of bars by lines: see rect.
lty the line type used for the bars, see also lines.
xlim, ylim the range of x and y values with sensible defaults.
main, xlab, ylab
these arguments to title have useful defaults here.
axes logical, indicating if axes should be drawn.
labels logical or character. Additionally draw labels on top of bars, if not FALSE; if TRUE, draw the counts or rounded intensities; if labels is a character, draw itself.
add logical. If TRUE, only the bars are added to the current plot. This is what lines.histogram(*) does.
... further graphical parameters to title and axis.

## Details

lines.histogram(*) is the same as plot.histogram(*, add = TRUE).

## See Also

hist, stem, density.

## Examples

```
data(women)
str(wwt <- hist(women$weight, nc= 7, plot = FALSE))
plot(wwt, labels = TRUE) # default main & xlab using wwt$xname
plot(wwt, border = "dark blue", col = "light blue",
    main = "Histogram of 15 women's weights", xlab = "weight [pounds]")
## Fake "lines" example, using non-default labels:
w2 <- wwt; w2$counts <- w2$counts - 1
lines(w2, col = "Midnight Blue", labels = ifelse(w2$counts, "> 1", "1"))
```

```
plot.lm
```

Plot Diagnostics for an lm Object

## Description

Four plots (selectable by which) are currently provided: a plot of residuals against fitted values, a Scale-Location plot of $\sqrt{\mid \text { residuals } \mid}$ against fitted values, a Normal Q-Q plot, and a plot of Cook's distances versus row labels.

## Usage

```
plot(x, which = 1:4,
    caption = c("Residuals vs Fitted", "Normal Q-Q plot",
                "Scale-Location plot", "Cook's distance plot"),
    panel = points,
    sub.caption = deparse(x$call), main = "",
    ask = prod(par("mfcol")) < length(which) && dev.interactive(),
    id.n = 3, labels.id = names(residuals(x)), cex.id = 0.25)
```


## Arguments

| x | lm object, typically result of lm or glm. |
| :--- | :--- |
| which | If a subset of the plots is required, specify a subset of the numbers 1:4. |
| caption | Captions to appear above the plots |
| panel | Panel function. A useful alternative to points is panel.smooth. <br> sub.caption <br> common title -above figures if there are multiple; used as sub (s.title) <br> otherwise. <br> title to each plot—in addition to the above caption. |
| main | logical; if TRUE, the user is asked before each plot, see par (ask=.). <br> ask |
| other parameters to be passed through to plotting functions. |  |
| id.n | number of points to be labelled in each plot, starting with the most ex- <br> treme. |
| labels.id | vector of labels, from which the labels for extreme points will be chosen. <br> nULL uses observation numbers. |
| cex.id | magnification of point labels. |

## Details

sub.caption - by default the function call-is shown as a subtitle (under the x-axis title) on each plot when plots are on separate pages, or as a subtitle in the outer margin (if any) when there are multiple plots per page.
The "Scale-Location" plot, also called "Spread-Location" or "S-L" plot, takes the square root of the absolute residuals in order to diminish skewness $(\sqrt{|E|}$ is much less skewed than $|E|$ for Gaussian zero-mean $E$ ).

This 'S-L' and the Q-Q plot use standardized residuals which have identical variance (under the hypothesis). They are given as $R_{i} /\left(s \times \sqrt{1-h_{i i}}\right)$ where $h_{i i}$ are the diagonal entries of the hat matrix, lm.influence()\$hat, see also hat.

## Author(s)

John Maindonald and Martin Maechler.

## References

Belsley, D. A., Kuh, E. and Welsch, R. E. (1980) Regression Diagnostics. New York: Wiley. Cook, R. D. and Weisberg, S. (1982) Residuals and Influence in Regression. London: Chapman and Hall.
Hinkley, D. V. (1975) On power transformations to symmetry. Biometrika 62, 101-111.
McCullagh, P. and Nelder, J. A. (1989) Generalized Linear Models. London: Chapman and Hall.

## See Also

```
termplot, lm.influence, cooks.distance.
```


## Examples

```
## Analysis of the life-cycle savings data
## given in Belsley, Kuh and Welsch.
data(LifeCycleSavings)
plot(lm.SR <- lm(sr ~ pop15 + pop75 + dpi + ddpi, data = LifeCycleSavings))
## 4 plots on 1 page; allow room for printing model formula in outer margin:
par(mfrow = c(2, 2), oma = c(0, 0, 2, 0))
plot(lm.SR)
plot(lm.SR, id.n = NULL) # no id's
plot(lm.SR, id.n = 5, labels.id = NULL)# 5 id numbers
## Fit a smmooth curve, where applicable:
plot(lm.SR, panel = panel.smooth)
## Gives a smoother curve
plot(lm.SR, panel = function(x,y) panel.smooth(x, y, span = 1))
par(mfrow=c(2,1))# same oma as above
plot(lm.SR, which = 1:2, sub.caption = "Saving Rates, n=50, p=5")
```

```
plot.table Plot Methods for 'table' Objects
```


## Description

This is a method of the generic plot function for (contingency) table objects. Whereas for two- and more dimensional tables, a mosaicplot is drawn, one-dimensional ones are plotted "bar like".

## Usage

plot.table(x, type $=$ "h", ylim $=c(0, \max (x)), l_{w d}=2$, xlab $=$ NULL, $y l a b=$ deparse(substitute(x)), frame.plot = is.num, ...)

## Arguments

```
\(x \quad\) a table (like) object.
type plotting type.
ylim range of \(y\)-axis.
lwd line width for bars when type \(=\) " \(h\) " is used in the 1D case.
\(x l a b, y l a b \quad x-\) and \(y\)-axis labels.
frame.plot logical indicating if a frame (box) should be drawn in the 1D case. De-
    faults to true when \(x\) has dimnames coerceable to numbers.
... further graphical arguments, see plot.default.
```


## Details

The current implementation ( R 1.2 ) is somewhat experimental and will be improved and extended.

## See Also

plot.factor, the plot method for factors.

## Examples

```
## 1-d tables
(Poiss.tab <- table(N = rpois(200, lam= 5)))
plot(Poiss.tab, main = "plot(table(rpois(200, lam=5)))")
data(state)
plot(table(state.division))
## 4-D :
data(Titanic)
plot(Titanic, main ="plot(Titanic, main= *)")
```

```
plot.ts Plotting Time-Series Objects
```


## Description

Plotting methods for objects of class "ts" or "mts" (multivariate time-series).

## Usage

```
plot(x, y = NULL, type = "l", frame.plot = axes,
        plot.type = c("multiple", "single"),
        xy.labels = n <= 150, xy.lines = do.lab, panel=lines, ...)
    lines(x, ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | es objects, usually of class |
| :---: | :---: |
| type | the type of plot, see plot. When y is present, the default will depend on xy.labels, see below. |
| frame.plot | a function to give the 'frame' for each panel. |
| plot.type | for multivariate time series, should the series by plotted separately (with a common time axis) or on a single plot? |
| xy.labels | logical, indicating if text () labels should be used for an x-y plot. |
| xy.lines | logical, indicating if lines should be drawn for an $x-y$ plot. Default is true, when labels are drawn as well. |
| panel | a function( $\mathrm{x}, \mathrm{col}, \mathrm{bg}, \mathrm{pch}$, type, ...) which gives the action to be carried out in each panel of the display for plot.type="multiple". The default is lines. <br> additional graphical arguments, see plot, plot.default and par. |

## Details

With one principal argument, these functions create time series plots, for multivariate series of two kinds depending on plot.type,
If y is present, both x and y must be univariate, and a "scatter" plot y ~ x will be drawn, enhanced by using text if xy.labels is TRUE or character, and lines if xy.lines is TRUE.

## See Also

ts for basic time series construction and access functionality.

## Examples

```
## Multivariate
z <- ts(matrix(rt(300, df = 3), 100, 3), start=c(1961, 1), frequency=12)
plot(z) # multiple
plot(z, panel=points) # same with points instead of lines
plot(z, plot.type="single", lty=1:3)
```

```
## A phase plot:
data(nhtemp)
plot(nhtemp, c(nhtemp[-1], NA), cex = .8, col="blue",
    main = "Lag plot of New Haven temperatures")
## a clearer way to do this would be
library(ts)
plot(nhtemp, lag(nhtemp, 1), cex = .8, col="blue",
    main = "Lag plot of New Haven temperatures")
library(ts)
data(sunspots)
## xy.lines and xy.labels are FALSE for large series:
plot(lag(sunspots, 1), sunspots, pch = ".")
data(EuStockMarkets)
SMI <- EuStockMarkets[, "SMI"]
plot(lag(SMI, 1), SMI, pch = ".")
plot(lag(SMI, 20), SMI, pch = ".", log = "xy",
    main = "4 weeks lagged SMI stocks -- log scale", xy.lines= TRUE)
detach("package:ts")
```

plot.window Set up World Coordinates for Graphics Window

## Description

This function sets up the world coordinate system for a graphics window. It is called by higher level functions such as plot.default (after plot.new).

## Usage

```
plot.window(xlim, ylim, log = "", asp = NA, ...)
```


## Arguments

xlim, ylim numeric of length 2, giving the x and y coordinates ranges.
log character; indicating which axes should be in log scale.
asp numeric, giving the aspect ratio $y / x$.
... further graphical parameters as in par.

## Details

Note that if asp is a finite positive value then the window is set up so that one data unit in the x direction is equal in length to asp $\times$ one data unit in the y direction.
The special case asp $==1$ produces plots where distances between points are represented accurately on screen. Values with asp > 1 can be used to produce more accurate maps when using latitude and longitude.
Usually, one should rather use the higher level functions such as plot, hist, image, ..., instead and refer to their help pages for explanation of the arguments.

## See Also

xy.coords, plot.xy, plot.default.

## Examples

```
##--- An example for the use of 'asp' :
library(mva)
data(eurodist)
loc <- cmdscale(eurodist)
rx <- range(x <- loc[,1])
ry <- range(y <- -loc[,2])
plot(x, y, type="n", asp=1, xlab="", ylab="")
abline(h=pretty(rx, 10),
    v=pretty(ry, 10), col= "lightgray")
text(x, y, names(eurodist), cex=0.5)
```

```
plot.xy Basic Internal Plot Function
```


## Description

This is the internal function that does the basic plotting of points and lines. Usually, one should rather use the higher level functions instead and refer to their help pages for explanation of the arguments.

## Usage

```
plot.xy(xy, type, pch=1, lty="solid", col=par("fg"), bg=NA, cex=1, ...)
```


## Arguments

| xy | A four-element list as results from xy.coords. |
| :--- | :--- |
| type | 1 character code. |
| pch | character or integer code for kind of points/lines, see points.default. |
| lty | line type code, see lines. |
| col | color code or name, see colors, palette. |
| bg | background ("fill") color for open plot symbols. |
| cex | character expansion. |
| $\ldots$ | further graphical parameters. |

## See Also

```
plot, plot.default, points, lines.
```


## Examples

```
points.default # to see how it calls "plot.xy(xy.coords(x, y), ...)"
```

```
plotmath Mathematical Annotation in R
```


## Description

If the text argument to one of the text-drawing functions (text, mtext, axis) in $R$ is an expression, the argument is interpreted as a mathematical expression and the output will be formatted according to TeX-like rules. Expressions can also be used for titles, subtitles and x - and y -axis labels (but not for axis labels on persp plots).

## Details

A mathematical expression must obey the normal rules of syntax for any $R$ expression, but it is interpreted according to very different rules than for normal $R$ expressions.

It is possible to produce many different mathematical symbols, generate sub- or superscripts, produce fractions, etc.

The output from example (plotmath) includes several tables which show the available features. In these tables, the columns of grey text show sample R expressions, and the columns of black text show the resulting output.
The available features are also described in the tables below:


## Meaning

x plus y
x minus y
juxtapose x and y
x forwardslash y
x plus or minus y
x divided by y
x times y
x subscript i
x superscript 2
juxtapose $\mathrm{x}, \mathrm{y}$, and z
square root of $x$
$y$ th root of $x$
x equals y
x is not equal to y
x is less than y
x is less than or equal to y
$x$ is greater than $y$
$x$ is greater than or equal to $y$
x is approximately equal to y
x and y are congruent
x is defined as y
x is proportional to y
draw x in normal font
draw $x$ in bold font
draw x in italic font
draw x in bolditalic font
comma-separated list
ellipsis (height varies)

| cdots | ellipsis (vertically centred) |
| :---: | :---: |
| ldots | ellipsis (at baseline) |
| x \%subset\% y | $x$ is a proper subset of y |
| x \%subseteq\% y | $x$ is a subset of y |
| x \%notsubset\% y | $x$ is not a subset of y |
| x \%supset\% y | x is a proper superset of y |
| x \%supseteq\% y | $x$ is a superset of y |
| x \%in\% y | $x$ is an element of $y$ |
| $\mathrm{x} \%$ notin\% y | $x$ is not an element of y |
| hat (x) | x with a circumflex |
| tilde(x) | x with a tilde |
| $\operatorname{dot}(\mathrm{x})$ | x with a dot |
| ring(x) | x with a ring |
| bar (xy) | xy with bar |
| widehat (xy) | xy with a wide circumflex |
| widetilde(xy) | xy with a wide tilde |
| x \% <->\% y | x double-arrow y |
| x \% ->\% y | x right-arrow y |
| x \%<-\% y | x left-arrow y |
| x \%up\% y | x up-arrow y |
| x \%down\% y | x down-arrow y |
| x \% <=>\% y | x is equivalent to y |
| x \% $=>\%$ y | x implies y |
| x \% < = \% y | y implies x |
| x \%dblup\% y | x double-up-arrow y |
| x \%dbldown\% y | x double-down-arrow y |
| alpha-omega | Greek symbols |
| Alpha - Omega | uppercase Greek symbols |
| infinity | infinity symbol |
| partialdiff | partial differential symbol |
| $32 *$ degree | 32 degrees |
| $60 *$ minute | 60 minutes of angle |
| $30 *$ second | 30 seconds of angle |
| displaystyle(x) | draw x in normal size (extra spacing) |
| textstyle(x) | draw x in normal size |
| scriptstyle(x) | draw x in small size |
| scriptscriptstyle(x) | draw x in very small size |
| $\mathrm{x} \sim \sim \mathrm{y}$ | put extra space between x and y |
| $\mathrm{x}+\mathrm{phantom}(0)+\mathrm{y}$ | leave gap for "0", but don't draw it |
| $\mathrm{x}+\mathrm{over}(1$, phantom(0)) | leave vertical gap for "0" (don't draw) |
| $\mathrm{frac}(\mathrm{x}, \mathrm{y})$ | x over y |
| over (x, y) | x over y |
| atop (x, y) | $x$ over y (no horizontal bar) |
| $\operatorname{sum}(x[i], i==1, n)$ | sum x[i] for i equals 1 to n |
| $\operatorname{prod}(\mathrm{plain}(\mathrm{P})(\mathrm{X}==\mathrm{x}), \mathrm{x})$ | product of $\mathrm{P}(\mathrm{X}=\mathrm{x})$ for all values of x |
| $\begin{aligned} & \text { integral }(\mathrm{f}(\mathrm{x}) * \mathrm{dx}, \mathrm{a}, \mathrm{~b}) \\ & \text { union }(\mathrm{A}[\mathrm{i}], \mathrm{i}==1, \mathrm{n}) \end{aligned}$ | definite integral of $f(x)$ wrt $x$ union of $\mathrm{A}[\mathrm{i}]$ for i equals 1 to n |
| intersect(A[i], i==1, n ) | intersection of $\mathrm{A}[\mathrm{i}]$ |
| $\begin{aligned} & \lim (f(x), x \%->\% 0) \\ & \min (g(x), x>0) \end{aligned}$ | limit of $f(x)$ as $x$ tends to 0 minimum of $\mathrm{g}(\mathrm{x})$ for x greater than 0 |
| $\inf (\mathrm{S})$ | infimum of $S$ |
| sup (S) | supremum of S |

```
x^y + z normal operator precedence
x^(y + z) visible grouping of operands
x^{y + z} invisible grouping of operands
group("(",list(a, b),"]") specify left and right delimiters
bgroup("(",atop(x,y),")") use scalable delimiters
group(lceil, x, rceil)
```

normal operator precedence visible grouping of operands invisible grouping of operands specify left and right delimiters use scalable delimiters special delimiters

## References

Murrell, P. and Ihaka, R. (2000) An approach to providing mathematical annotation in plots. Journal of Computational and Graphical Statistics, 9, 582-599.

## See Also

axis, mtext, text, title

## Examples

```
x <- seq(-4, 4, len = 101)
y <- cbind(sin(x), cos(x))
matplot(x, y, type = "l", xaxt = "n",
    main = expression(paste(plain(sin) * phi, " and ",
                                    plain(cos) * phi)),
        ylab = expression("sin" * phi, "cos" * phi), # only 1st is taken
        xlab = expression(paste("Phase Angle ", phi)),
        col.main = "blue")
axis(1, at = c(-pi, -pi/2, 0, pi/2, pi),
    lab = expression(-pi, -pi/2, 0, pi/2, pi))
## How to combine "math" and numeric variables :
plot(1:10, type="n", xlab="", ylab="", main = "plot math & numbers")
tt <- 1.23 ; mtext(substitute(hat(theta) == that, list(that= tt)))
for(i in 2:9)
    text(i,i+1, substitute(list(xi,eta) == group("(",list(x,y),")"),
                                    list(x=i, y=i+1)))
```

```
plot(1:10, 1:10)
text(4, 9, expression(hat(beta) == (X^t * X)^{-1} * X^t * y))
text(4, 8.4, "expression(hat(beta) == (X^t * X)^{-1} * X^t * y)",
    cex = .8)
text(4, 7, expression(bar(x) == sum(frac(x[i], n), i==1, n)))
text(4, 6.4, "expression(bar(x) == sum(frac(x[i], n), i==1, n))",
    cex = .8)
text(8, 5, expression(paste(frac(1, sigma*sqrt(2*pi)), " ",
                                    plain(e)^{frac(-(x-mu)^2, 2*sigma^2)})),
    cex= 1.2)
######
# create tables of mathematical annotation functionality
######
make.table <- function(nr, nc) {
    savepar <- par(mar=rep(0, 4), pty="s")
    plot(c(0, nc*2 + 1), c(0, -(nr + 1)),
        type="n", xlab="", ylab="", axes=FALSE)
    savepar
}
```

```
get.r <- function(i, nr) {
    i %% nr + 1
}
get.c <- function(i, nr) {
    i %/%nr + 1
}
draw.title.cell <- function(title, i, nr) {
    r <- get.r(i, nr)
    c <- get.c(i, nr)
    text(2*c - .5, -r, title)
    rect((2*(c - 1) + . 5), -(r - . 5), (2*c + . 5), -(r + . 5))
}
draw.plotmath.cell <- function(expr, i, nr, string = NULL) {
    r <- get.r(i, nr)
    c <- get.c(i, nr)
    if (is.null(string)) {
                string <- deparse(expr)
        string <- substr(string, 12, nchar(string) - 1)
    }
    text((2*(c - 1) + 1), -r, string, col="grey")
    text((2*c), -r, expr, adj=c(.5,.5))
    rect((2*(c - 1) + . 5), -(r - . 5), (2*c + . 5), -(r + . 5), border="grey")
}
nr <- 20
nc <- 2
oldpar <- make.table(nr, nc)
i <- 0
draw.title.cell("Arithmetic Operators", i, nr); i <- i + 1
draw.plotmath.cell(expression(x + y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x - y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x * y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x / y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x %+-% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x %/% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x %*% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(-x), i, nr); i <- i + 1
draw.plotmath.cell(expression(+x), i, nr); i <- i + 1
draw.title.cell("Sub/Superscripts", i, nr); i <- i + 1
draw.plotmath.cell(expression(x[i]), i, nr); i <- i + 1
draw.plotmath.cell(expression(x^2), i, nr); i <- i + 1
draw.title.cell("Juxtaposition", i, nr); i <- i + 1
draw.plotmath.cell(expression(x * y), i, nr); i <- i + 1
draw.plotmath.cell(expression(paste(x, y, z)), i, nr); i <- i + 1
draw.title.cell("Lists", i, nr); i <- i + 1
draw.plotmath.cell(expression(list(x, y, z)), i, nr); i <- i + 1
# even columns up
i <- 20
draw.title.cell("Radicals", i, nr); i <- i + 1
draw.plotmath.cell(expression(sqrt(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(sqrt(x, y)), i, nr); i <- i + 1
draw.title.cell("Relations", i, nr); i <- i + 1
draw.plotmath.cell(expression(x == y), i, nr); i <- i + 1
```

draw. plotmath.cell(expression(x != y), i, nr); i <- i + 1
draw. plotmath. cell (expression(x < y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x <= y), i, nr); i <- i + 1
draw. plotmath.cell (expression(x > y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x >= y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%~~\% y), i, nr); i <- i + 1
draw.plotmath.cell (expression(x \%=~\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x $\%==\%$ y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%prop\% y), i, nr); i <- i + 1
draw.title.cell("Typeface", i, nr); i <- i + 1
draw.plotmath.cell(expression(plain(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(italic(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(bold(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(bolditalic(x)), i, nr); i <- i + 1
\# Need fewer, wider columns for ellipsis ...
$\mathrm{nr}<-20$
nc <- 2
make.table(nr, nc)
i <- 0
draw.title.cell("Ellipsis", i, nr); i <- i + 1
draw.plotmath. cell(expression(list(x[1], ..., x[n])), i, nr); i<-i+1
draw.plotmath.cell(expression(x[1] + ... + x[n]), i, nr); i <- i + 1
draw.plotmath.cell(expression(list(x[1], cdots, $x[n])$ ), i, nr); i <- i + 1
draw.plotmath.cell (expression(x[1] + ldots $+x[n]$ ), i, $n r$ ); i <- i + 1
draw.title.cell("Set Relations", i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%subset\% y), i, nr); i <- i + 1
draw.plotmath.cell (expression(x \%subseteq\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%supset\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%supseteq\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%notsubset\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%in\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%notin\% y), i, nr); i <- i + 1
draw.title.cell("Accents", i, nr); i <- i + 1
draw.plotmath.cell(expression(hat(x)), i, nr); i <- i + 1
draw. plotmath.cell(expression(tilde(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(ring(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(bar(xy)), i, nr); i <- i + 1
draw.plotmath.cell(expression(widehat(xy)), i, nr); i <- i + 1
draw.plotmath.cell(expression(widetilde(xy)), i, nr); i <- i + 1
draw.title.cell("Arrows", i, nr); i <- i + 1
draw.plotmath.cell (expression(x \%<->\% y), i, nr); i <- i + 1
draw.plotmath.cell (expression (x \%->\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%<-\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%up\% y), i, nr) ; i <- i + 1
draw.plotmath.cell(expression(x \%down\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%<=>\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%=>\% y), i, nr); i <- i + 1
draw.plotmath.cell (expression (x \% < = \% y), i, nr) ; i <- i + 1
draw.plotmath.cell(expression(x \%dblup\% y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x \%dbldown\% y), i, nr); i <- i + 1
draw.title.cell("Symbolic Names", i, nr); i <- i + 1
draw.plotmath.cell(expression(Alpha - Omega), i, nr); i <- i + 1
draw.plotmath.cell(expression(alpha - omega), i, nr); i <- i + 1
draw. plotmath.cell(expression(infinity), i, nr); i <- i + 1
draw.plotmath.cell(expression(32 * degree), i, nr); i <- i + 1
draw.plotmath.cell(expression(60 * minute), i, nr); i <- i + 1

```
draw.plotmath.cell(expression(30 * second), i, nr); i <- i + 1
# Need even fewer, wider columns for typeface and style ...
nr <- 20
nc <- 1
make.table(nr, nc)
i <- 0
draw.title.cell("Style", i, nr); i <- i + 1
draw.plotmath.cell(expression(displaystyle(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(textstyle(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(scriptstyle(x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(scriptscriptstyle(x)), i, nr); i <- i + 1
draw.title.cell("Spacing", i, nr); i <- i + 1
draw.plotmath.cell(expression(x ~~ y), i, nr); i <- i + 1
# Need fewer, taller rows for fractions ...
# cheat a bit to save pages
par(new = TRUE)
nr <- 10
nc <- 1
make.table(nr, nc)
i <- 4
draw.plotmath.cell(expression(x + phantom(0) + y), i, nr); i <- i + 1
draw.plotmath.cell(expression(x + over(1, phantom(0))), i, nr); i <- i + 1
draw.title.cell("Fractions", i, nr); i <- i + 1
draw.plotmath.cell(expression(frac(x, y)), i, nr); i <- i + 1
draw.plotmath.cell(expression(over(x, y)), i, nr); i <- i + 1
draw.plotmath.cell(expression(atop(x, y)), i, nr); i <- i + 1
# Need fewer, taller rows and fewer, wider columns for big operators ...
nr <- 10
nc <- 1
make.table(nr, nc)
i <- 0
draw.title.cell("Big Operators", i, nr); i <- i + 1
draw.plotmath.cell(expression(sum(x[i], i=1, n)), i, nr); i <- i + 1
draw.plotmath.cell(expression(prod(plain(P) (X == x), x)), i, nr); i <- i + 1
draw.plotmath.cell(expression(integral(f(x) * dx, a, b)), i, nr); i <- i + 1
draw.plotmath.cell(expression(union(A[i], i==1, n)), i, nr); i <- i + 1
draw.plotmath.cell(expression(intersect(A[i], i==1, n)), i, nr); i <- i + 1
draw.plotmath.cell(expression(lim(f(x), x %->% 0)), i, nr); i <- i + 1
draw.plotmath.cell(expression(min(g(x), x >= 0)), i, nr); i <- i + 1
draw.plotmath.cell(expression(inf(S)), i, nr); i <- i + 1
draw.plotmath.cell(expression(sup(S)), i, nr); i <- i + 1
make.table(nr, nc)
i <- 0
draw.title.cell("Grouping", i, nr); i <- i + 1
draw.plotmath.cell(expression((x + y)*z), i, nr); i <- i + 1
draw.plotmath.cell(expression(x^y + z), i, nr); i <- i + 1
draw.plotmath.cell(expression(x^(y + z)), i, nr); i <- i + 1
# have to do this one by hand
draw.plotmath.cell(expression(x^{y + z}), i, nr, string="x^{y + z}"); i <- i + 1
draw.plotmath.cell(expression(group("(", list(a, b), "]")), i, nr); i <- i + 1
draw.plotmath.cell(expression(bgroup("(", atop(x, y), ")")), i, nr); i <- i + 1
draw.plotmath.cell(expression(group(lceil, x, rceil)), i, nr); i <- i + 1
draw.plotmath.cell(expression(group(lfloor, x, rfloor)), i, nr); i <- i + 1
```

```
draw.plotmath.cell(expression(group("|", x, "|")), i, nr); i <- i + 1
par(oldpar)
```

pmatch Partial String Matching

## Description

pmatch seeks matches for the elements of its first argument among those of its second.

## Usage

```
pmatch(x, table, nomatch = NA, duplicates.ok = FALSE)
```


## Arguments

$\mathrm{x} \quad$ the values to be matched.
table the values to be matched against.
nomatch the value returned at non-matching or multiply partially matching positions.
duplicates.ok should elements be in table be used more than once?

## Details

The behaviour differs by the value of duplicates.ok. Consider first the case if this is true. First exact matches are considered, and the positions of the first exact matches are recorded. Then unique partial matches are considered, and if found recorded. (A partial match occurs if the whole of the element of $x$ matches the beginning of the element of table.) Finally, all remaining elements of $x$ are regarded as unmatched. In addition, an empty string can match nothing, not even an exact match to an empty string. This is the appropriate behaviour for partial matching of character indices, for example.

If duplicates.ok is FALSE, values of table once matched are excluded from the search for subsequent matches. This behaviour is equivalent to the R algorithm for argument matching, except for the consideration of empty strings (which in argument matching are matched after exact and partial matching to any remaining arguments).
charmatch is similar to pmatch with duplicates.ok true, the differences being that it differentiates between no match and an ambiguous partial match, it does match empty strings, and it does not allow multiple exact matches.

## Value

A numeric vector of integers (including NA if nomatch $=N A$ ) of the same length as $x$, giving the indices of the elements in table which matched, or nomatch.

## Note

Versions of R prior to 1.0 .0 had a different behaviour that was seriously incompatible with S (and the current version) when duplicates.ok = TRUE.

## Author(s)

Of this version, B. D. Ripley.

## See Also

match, charmatch and match.arg, match.fun, match.call, for function argument matching etc., grep etc for more general (regexp) matching of strings.

## Examples

```
pmatch("", "") # returns NA
pmatch("m", c("mean", "median", "mode")) # returns NA
pmatch("med", c("mean", "median", "mode")) # returns 2
pmatch(c("", "ab", "ab"), c("abc", "ab"), dup=FALSE)
pmatch(c("", "ab", "ab"), c("abc", "ab"), dup=TRUE)
## compare
charmatch(c("", "ab", "ab"), c("abc", "ab"))
```

png JPEG and PNG graphics devices

## Description

A graphics device for JPEG or PNG format bitmap files.

## Usage

```
jpeg(filename = "Rplot%03d.jpeg", width = 480, height = 480,
    pointsize = 12, quality = 75, bg = "white", ...)
png(filename ="Rplot%03d.png", width = 480, height = 480,
    pointsize = 12, bg = "white", ...)
```


## Arguments

| filename | the name of the output file. The page number is substituted if an integer <br> format is included in the character string. |
| :--- | :--- |
| width | the width of the device in pixels. |
| height | the height of the device in pixels. |
| pointsize | the default pointsize of plotted text. |
| quality | the 'quality' of the JPEG image, as a percentage. Smaller values will give <br> more compression but also more degradation of the image. |
| bg | default background colour. |
| . . | additional arguments to the X11 device. |

## Details

Plots in PNG and JPEG format can easily be converted to many other bitmap formats, and both can be displayed in most modern web browsers. The PNG format is lossless and is best for line diagrams and blocks of solid colour. The JPEG format is lossy, but may be useful for image plots, for example.
png supports transparent backgrounds: use bg = "transparent". Not all PNG viewers render files with transparency correctly. When transparency is in use a very light grey is used as the background and so will appear as transparent if used in the plot. This allows opaque white to be used, as on the example.
$R$ can be compiled without support for either or both of these devices: this will be reported if you attempt to use them on a system where they are not supported. They will not be available if R has been started with --gui=none (and will give a different error message), and they may not be usable unless the X11 display is available to the owner of the R process.

## Value

A plot device is opened: nothing is returned to the R interpreter.

## Warning

If you plot more than one page on one of these devices and do not include somthing like $\% \mathrm{~d}$ for the sequence number in file, the file will contain the last page plotted.

## Note

These are based on the X11 device, so the additional arguments to that device work, but are rarely appropriate. The colour handling will be that of the X11 device in use.

## Author(s)

Guido Masarotto and Brian Ripley

## See Also

Devices, dev.print
capabilities to see if these devices are supported by this build of R.
bitmap provides an alternative way to generate PNG and JPEG plots that does not depend on accessing the X11 display but does depend on having GhostScript installed.

## Examples

```
## these examples will work only if the devices are available
## and the X11 display is available.
## copy current plot to a PNG file
dev.print(png, file="myplot.png", width=480, height=480)
png(file="myplot.png", bg="transparent")
plot(1:10)
rect(1, 5, 3, 7, col="white")
dev.off()
jpeg(file="myplot.jpeg")
example(rect)
```

```
dev.off()
```

```
points Add Points to a Plot
```


## Description

points is a generic function to draw a sequence of points at the specified coordinates. The specified character(s) are plotted, centered at the coordinates.

## Usage

```
points(x, ...)
points.default(x, y=NULL, type="p", pch=par("pch"), col=par("col"),
    \(\mathrm{bg}=\mathrm{NA}, \mathrm{cex}=1, \ldots\) )
```


## Arguments

```
\(x, y \quad\) coordinate vectors of points to plot.
type character indicating the type of plotting; actually any of the types as in
    plot.
pch plotting "character", i.e. symbol to use. pch can either be a character
    or an integer code for a set of graphics symbols. The full set of S symbols
    is available with \(\mathrm{pch}=0: 18\), see the last picture from example(points),
    i.e., the examples below.
    In addition, there is a special set of R plotting symbols which can be ob-
    tained with pch=19:25 and 21:25 can be colored and filled with different
    colors:
    - pch=19: solid circle,
    - pch=20: bullet (smaller circle),
    - pch=21: circle,
    - pch=22: square,
    - pch=23: diamond,
    - pch=24: triangle point-up,
    - pch=25: triangle point down.
col color code or name, see par.
bg background ("fill") color for open plot symbols
cex character expansion
... Further graphical parameters (see plot.xy and par) may also be supplied
    as arguments.
```


## Details

The coordinates can be passed in a plotting structure (a list with x and y components), a two-column matrix, a time series, .... See xy.coords.
Graphical parameters are permitted as arguments to this function.

## See Also

plot, lines, and the underlying "primitive" plot.xy.

## Examples

```
plot(-4:4, -4:4, type = "n")# setting up coord. system
points(rnorm(200), rnorm(200), col = "red")
points(rnorm(100)/2, rnorm(100)/2, col = "blue", cex = 1.5)
op <- par(bg = "light blue")
x <- seq(0,2*pi, len=51)
## something ''between type="b" and type="o" '` :
plot(x, sin(x), type="o", pch=21, bg=par("bg"), col = "blue", cex=.6,
    main='plot(..., type="o", pch=21, bg=par("bg"))')
par(op)
##------- Showing all the extra & some char graphics symbols
Pex <- 3 ## good for both .Device=="postscript" and "x11"
ipch <- 1:(np <- 25+11); k <- floor(sqrt(np)); dd <- c(-1,1)/2
rx <- dd + range(ix <- (ipch-1) %/% k)
ry <- dd + range(iy <- 3 + (k-1)-(ipch-1) %% k)
pch <- as.list(ipch)
pch[25+ 1:11] <- as.list(c("*",".", "०","0","0","+","-",":","|","%","#"))
plot(rx, ry, type="n", axes = FALSE, xlab = "", ylab = "",
    main = paste("plot symbols : points (... pch = *, cex =", Pex,")"))
abline(v = ix, h = iy, col = "lightgray", lty = "dotted")
for(i in 1:np) {
    pc <- pch[[i]]
    points(ix[i], iy[i], pch = pc, col = "red", bg = "yellow", cex = Pex)
    ## red symbols with a yellow interior (where available)
    text(ix[i] - .3, iy[i], pc, col = "brown", cex = 1.2)
}
```


## Poisson The Poisson Distribution

## Description

Density, distribution function, quantile function and random generation for the Poisson distribution with parameter lambda.

## Usage

```
dpois(x, lambda, log = FALSE)
ppois(q, lambda, lower.tail = TRUE, log.p = FALSE)
qpois(p, lambda, lower.tail = TRUE, log.p = FALSE)
rpois(n, lambda)
```


## Arguments

| x | vector of (non-negative integer) quantiles. |
| :--- | :--- |
| q | vector of quantiles. |
| p | vector of probabilities. |

n
lambda vector of positive means.
$\log , \log . \mathrm{p} \quad$ logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

The Poisson distribution has density

$$
p(x)=\frac{\lambda^{x} e^{-\lambda}}{x!}
$$

for $x=0,1,2, \ldots$ The mean and variance are $E(X)=\operatorname{Var}(X)=\lambda$.
If an element of x is not integer, the result of dpois is zero, with a warning. $p(x)$ is computed using Loader's algorithm, see the reference in dbinom.

The quantile is left continuous: qgeom(q, prob) is the largest integer $x$ such that $P(X \leq$ $x)<q$.

Setting lower.tail = FALSE allows to get much more precise results when the default, lower.tail = TRUE would return 1, see the example below.

## Value

dpois gives the (log) density, ppois gives the (log) distribution function, qpois gives the quantile function, and rpois generates random deviates.

## See Also

 dbinom for the binomial and dnbinom for the negative binomial distribution.
## Examples

```
-log(dpois(0:7, lambda=1) * gamma(1+ 0:7)) # == 1
Ni <- rpois(50, lam= 4); table(factor(Ni, 0:max(Ni)))
1 - ppois(10*(15:25), lambda=100) # becomes 0 (cancellation)
    ppois(10*(15:25), lambda=100, lower=FALSE) # no cancellation
par(mfrow = c(2, 1))
x <- seq(-0.01, 5, 0.01)
plot(x, ppois(x, 1), type="s", ylab="F(x)", main="Poisson(1) CDF")
plot(x, pbinom(x, 100, 0.01),type="s", ylab="F(x)",
    main="Binomial(100, 0.01) CDF")
```


## poly <br> Compute Orthogonal Polynomials

## Description

Returns or evaluates orthogonal polynomials of degree 1 to degree over the specified set of points x . These are all orthogonal to the constant polynomial of degree 0.

## Usage

```
poly(x, ..., degree \(=1\), coefs = NULL)
polym(..., degree)
predict(object, newdata, ...)
```


## Arguments

x , newdata a numeric vector at which to evaluate the polynomial. x can also be a matrix.
degree the degree of the polynomial
coefs for prediction, coefficients from a previous fit.
object an object inheriting from class "poly", normally the result of a call to poly with a single vector argument.
... poly, polym: further vectors.
predict.poly: arguments to be passed to or from other methods.

## Details

Although formally degree should be named (as it follows ...), an unnamed second argument of length 1 will be interpreted as the degree.
The orthogonal polynomial is summarized by the coefficients, which can be used to evaluate it via the three-term recursion given in Kennedy \& Gentle (1980, pp. 343-4), and use in the "predict" part of the code.

## Value

For poly with a single vector argument:
A matrix with rows corresponding to points in x and columns corresponding to the degree, with attributes "degree" specifying the degrees of the columns and "coefs" which contains the centring and normalization constants used in constructing the orthogonal polynomials. The matrix is given class c("poly", "matrix") as from R 1.5.0.

Other cases of poly and polym, and predict.poly: a matrix.

## Note

This routine is intended for statistical purposes such as contr.poly: it does not attempt to orthogonalize to machine accuracy.

## Author(s)

B. D. Ripley

## References

Kennedy, W. J. Jr and Gentle, J. E. (1980) Statistical Computing Marcel Dekker.

## See Also

```
contr.poly
```


## Examples

```
(z <- poly(1:10, 3))
predict(z, seq(2, 4, 0.5))
poly(seq(4, 6, 0.5), 3, coefs = attr(z, "coefs"))
polym(1:4, c(1, 4:6), degree=3) # or just poly()
poly(cbind(1:4, c(1, 4:6)), degree=3)
```

polygon Polygon Drawing

## Description

polygon draws the polygons whose vertices are given in x and y .

## Usage

```
polygon(x, y = NULL, density = NULL, angle = 45,
        border = NULL, col = NA, lty = NULL, xpd = NULL, ...)
```


## Arguments

```
\(\mathrm{x}, \mathrm{y} \quad\) vectors containing the coordinates of the vertices of the polygon.
density the density of shading lines, in lines per inch. The default value of NULL
    means that no shading lines are drawn. Non-positive values of density
    also inhibit the drawing of shading lines.
    angle the slope of shading lines, given as an angle in degrees (counter-clockwise).
    col the color for filling the polygon. The default, NA, is to leave polygons
        unfilled.
    border the color to draw the border. The default, NULL, uses par ("fg"). Use
        border \(=\) NA to omit borders.
        For compatibility with S , border can also be logical, it which case FALSE
        is equivalent to NA (borders omitted) and TRUE is equivalent to NULL (use
        the foreground colour),
    lty the line type to be used, as in par.
    xpd (where) should clipping take place? Defaults to par ("xpd").
    graphical parameters can be given as arguments to polygon.
```


## Details

The coordinates can be passed in a plotting structure (a list with x and y components), a two-column matrix, .... See xy. coords.
It is assumed that the polygon is closed by joining the last point to the first point.
The coordinates can contain missing values. The behaviour is similar to that of lines, except that instead of breaking a line into several lines, NA values break the polygon into several complete polygons (including closing the last point to the first point). See the examples below.

When multiple polygons are produced, the values of density, angle, col, border, and lty are recycled in the usual manner.

## Bugs

The present shading algorithm can produce incorrect results for self-intesecting polygons.

## Author(s)

The code implementing polygon shading was donated by Kevin Buhr 〈buhr@stat.wisc.edu〉.

## See Also

segments for even more flexibility, lines, rect, box, abline. par for how to specify colors.

## Examples

```
x <- c(1:9,8:1)
y <- c(1,2*(5:3),2,-1,17,9,8,2:9)
op <- par(mfcol=c(3,1))
for(xpd in c(FALSE,TRUE,NA)) {
    plot(1:10, main=paste("xpd =", xpd)) ; box("figure", col = "pink", lwd=3)
    polygon(x,y, xpd=xpd, col = "orange", lty=2, lwd=2, border = "red")
}
par(op)
n <- 100
xx <- c(0:n, n:0)
yy <- c(c(0,cumsum(rnorm(n))), rev(c(0,cumsum(rnorm(n)))))
plot (xx, yy, type="n", xlab="Time", ylab="Distance")
polygon(xx, yy, col="gray", border = "red")
title("Distance Between Brownian Motions")
# Multiple polygons from NA values
# and recycling of col, border, and lty
op <- par(mfrow=c(2,1))
plot(c(1,9), 1:2, type="n")
polygon(1:9, c(2,1,2,1,1,2,1,2,1),
    col=c("red", "blue"),
    border=c("green", "yellow"),
    lwd=3, lty=c("dashed", "solid"))
plot(c(1,9), 1:2, type="n")
polygon(1:9, c(2,1,2,1,NA,2,1,2,1),
    col=c("red", "blue"),
    border=c("green", "yellow"),
```

```
    lwd=3, lty=c("dashed", "solid"))
par(op)
# Line-shaded polygons
plot(c(1,9), 1:2, type="n")
polygon(1:9, c(2,1,2,1,NA,2,1,2,1),
        density=c(10, 20), angle=c(-45, 45))
```

    polyroot Find Zeros of a Real or Complex Polynomial
    
## Description

Find zeros of a real or complex polynomial.

## Usage

polyroot(z)

## Arguments

z
the vector of polynomial coefficients in increasing order.

## Details

A polynomial of degree $n-1$,

$$
p(x)=z_{1}+z_{2} x+\cdots+z_{n} x^{n-1}
$$

is given by its coefficient vector $\mathbf{z}[1: \mathrm{n}]$. polyroot returns the $n-1$ complex zeros of $p(x)$ using the Jenkins-Traub algorithm.
If the coefficient vector $z$ has zeroes for the highest powers, these are discarded.

## Value

A complex vector of length $n-1$, where $n$ is the position of the largest non-zero element of z .

## References

Jenkins and Traub (1972) TOMS Algorithm 419. Comm. ACM, 15, 97-99.

## See Also

uniroot for numerical root finding of arbitrary functions; complex and the zero example in the demos directory.

## Examples

```
polyroot(c(1, 2, 1))
round(polyroot(choose(8, 0:8)), 11) # guess what!
for (n1 in 1:4) print(polyroot(1:n1), digits = 4)
polyroot(c(1, 2, 1, 0, 0)) # same as the first
```


## pos.to.env Convert Positions in the Search Path to Environments

## Description

Returns the environment at a specified position in the search path.

## Usage

```
pos.to.env(x)
```


## Arguments

$\mathrm{x} \quad$ an integer between 1 and length ( $\operatorname{search}()$ ), the length of the search path.

## Details

Several R functions for manipulating objects in environments (such as get and 1s) allow specifying environments via corresponding positions in the search path. pos.to.env is a convenience function for programmers which converts these positions to corresponding environments; users will typically have no need for it.

## Examples

```
pos.to.env(1) # R_GlobalEnv
# the next returns NULL, which is how package:base is represented.
pos.to.env(length(search()))
```

```
postscript PostScript Graphics
```


## Description

postscript starts the graphics device driver for producing PostScript graphics.
The auxiliary function ps.options can be used to set and view (if called without arguments) default values for the arguments to postscript.

## Usage

```
postscript(file = ifelse(onefile, "Rplots.ps", "Rplot%03d.ps"),
            onefile = TRUE,
            paper, family, encoding, bg, fg,
            width, height, horizontal, pointsize,
            pagecentre, print.it, command)
ps.options(paper, horizontal, width, height, family, encoding,
            pointsize, bg, fg,
            onefile = TRUE, print.it = FALSE, append = FALSE,
            reset = FALSE, override.check = FALSE)
    .PostScript.Options
```


## Arguments

| file | a character string giving the name of the file. If it is "", the output is <br> piped to the command given by the argument command. If it is " $\mid$ cmd", <br> the output is piped to the command given by 'cmd'. |
| :--- | :--- |
| For use with onefile=FALSE give a printf format such as |  |
| "Rplot\%o3d.ps" (the default in that case). |  |

## Details

postscript opens the file file and the PostScript commands needed to plot any graphics requested are stored in that file. This file can then be printed on a suitable device to obtain hard copy.

A postscript plot can be printed via postscript in two ways.

1. Setting print.it $=$ TRUE causes the command given in argument command to be called with argument "file" when the device is closed. Note that the plot file is not deleted unless command arranges to delete it.
2. file="" or file="|cmd" can be used to print using a pipe on systems that support 'popen'. Failure to open the command will probably be reported to the terminal but not to 'popen', in which case close the device by dev. off immediately.

The postscript produced by R is EPS (Encapsulated PostScript) compatible, and can be included into other documents, e.g., into LaTeX, using
includegraphics\{<filename>\}. For use in this way you will probably want to set horizontal = FALSE, onefile = FALSE, paper = "special".

Most of the PostScript prologue used is taken from the R character vector .ps.prolog. This is marked in the output, and can be changed by changing that vector. (This is only advisable for PostScript experts.)
ps.options needs to be called before calling postscript, and the default values it sets can be overridden by supplying arguments to postscript.

## Families

The argument family specifies the font family to be used. In normal use it is one of "AvantGarde", "Bookman", "Courier", "Helvetica", "Helvetica-Narrow", "NewCenturySchoolbook", "Palatino" or "Times", and refers to the standard Adobe PostScript fonts of those names which are included (or cloned) in all common PostScript devices.

Many PostScript emulators (including those based on ghostscript) use the URW equivalents of these fonts, which are "URWGothic", "URWBookman", "NimbusMon", "NimbusSan", "NimbusSanCond", "CenturySch", "URWPalladio" and "NimbusRom" respectively. If your PostScript device is using URW fonts, you will obtain access to more characters and more approriate metrics by using these names. To make these easier to remember, "URWHelvetica" == "NimbusSan" and "URWTimes" == "NimbusRom" are also supported.
It is also possible to specify family="ComputerModern". This is intended to use with the Type 1 versions of the TeX CM fonts. It will normally be possible to include such output in TeX or LaTeX provided it is processed with dvips $-\mathrm{Ppfb}-\mathrm{j} 0$ or the equivalent on your system. (-j0 turns off font subsetting.)
If the second form of argument "family" is used, it should be a character vector of four or five paths to Adobe Font Metric files for the regular, bold, italic, bold italic and (optionally) symbol fonts to be used. If these paths do not contain the file separator, they are taken to refer to files in the R directory 'R_HOME/afm'. Thus the default Helvetica family can be specified by family $=c$ ("hv $\qquad$ .afm", "hvb $\qquad$ .afm", "hvo $\qquad$ .afm", "hvbo____.afm", "sy_____.afm"). It is the user's responsibility to check that suitable fonts are made available, and that they contain the needed characters when re-encoded. The fontnames used are taken from the FontName fields of the afm files. The software including the PostScript plot file should either embed the font outlines (usually from . pfb or .pfa files) or use DSC comments to instruct the print spooler to do so.

## Encodings

Encodings describe which glyphs are used to display the character codes (in the range $0-255$ ). By default R uses ISOLatin1 encoding, and the examples for text are in that encoding. However, the encoding used on machines running R may well be different, and by using the encoding argument the glyphs can be matched to encoding in use.
None of this will matter if only ASCII characters (codes 32-126) are used as all the encodings agree over that range. Some encodings are supersets of ISOLatin1, too. However, if accented and special characters do not come out as you expect, you may need to change the encoding. Three other encodings are supplied with R: "WinAnsi.enc" and "MacRoman.enc"
correspond to the encodings normally used on Windows and MacOS (at least by Adobe), and "PDFDoc.enc" is the first 256 characters of the Unicode encoding, the standard for PDF.
If you change the encoding, it is your responsibility to ensure that the PostScript font contains the glyphs used. One issue here is the Euro symbol which is in the WinAnsi and MacRoman encodings but may well not be in the PostScript fonts. (It is in the URW variants; it is not in the supplied Adobe Font Metric files.)
There is one exception. Character $45("-")$ is always set as minus (its value in Adobe ISOLatin1) even though it is hyphen in the other encodings. Hyphen is available as character 173 (octal 0255) in ISOLatin1.

## Author(s)

Support for Computer Modern fonts is based on a contribution by Brian D'Urso〈durso@hussle.harvard.edu〉.

## See Also

Devices, check.options which is called from both ps.options and postscript.

## Examples

```
# open the file "foo.ps" for graphics output
postscript("foo.ps")
# produce the desired graph(s)
dev.off() # turn off the postscript device
postscript("llp -dlw")
# produce the desired graph(s)
dev.off() # plot will appear on printer
# for URW PostScript devices
postscript("foo.ps", family = "NimbusSan")
## for inclusion in Computer Modern TeX documents, perhaps
postscript("cm_test.eps", width = 4.0, height = 3.0,
    horizontal = FALSE, onefile = FALSE, paper = "special",
    family = "ComputerModern")
## The resultant postscript file can be used by dvips -Ppfb -j0.
## To test out encodings, you can use
TestChars <- function(encoding="ISOLatin1", family="URWHelvetica")
{
    postscript(encoding=encoding, family=family)
    par(pty="s")
    plot(c(0,15), c(0,15), type="n", xlab="", ylab="")
    title(paste("Centred chars in encoding", encoding))
    grid(15, 15, lty=1)
    for(i in c(32:255)) {
        x <- i
        y <- i
        points(x, y, pch=i)
    }
    dev.off()
}
## there will be many warnings. We use URW to get a complete enough
```

```
## set of font metrics.
TestChars()
TestChars("ISOLatin2")
TestChars("WinAnsi")
stopifnot(unlist(ps.options()) == unlist(.PostScript.Options))
ps.options(bg = "pink")
str(ps.options(reset = TRUE))
### ---- error checking of arguments: ----
ps.options(width=0:12, onefile=0, bg=pi)
# override the check for 'onefile', but not the others:
str(ps.options(width=0:12, onefile=1, bg=pi,
    override.check = c(FALSE,TRUE,FALSE)))
```

```
power Create a Power Link Object
```


## Description

Creates a link object based on the link function $\eta=\mu^{\lambda}$.

## Usage

```
power(lambda = 1)
```


## Arguments

lambda a real number.

## Details

If lambda is non-negative, it is taken as zero, and the log link is obtained. The default lambda $=1$ gives the identity link.

## Value

A list with components linkfun, linkinv, mu.eta, and valideta. See make.link for information on their meaning.

See Also
make.link, family

## Examples

```
power()
quasi(link=power(1/3)) [c("linkfun", "linkinv")]
```


## Description

Generates the sequence of "probability" points $(1: m-a) /(m+(1-a)-a)$ where $m$ is either $n$, if length $(n)==1$, or length $(n)$.

## Usage

```
ppoints(n, a = ifelse(n <= 10, 3/8, 1/2))
```


## Arguments

$\mathrm{n} \quad$ either the number of points generate or a vector of observations.
a the offset fraction to be used; typically in $(0,1)$.

## Details

If $0<a<1$, the resulting values are within $(0,1)$ (excluding boundaries). In any case, the resulting sequence is symmetric in $[0,1]$, i.e., $p+\operatorname{rev}(p)==1$.
ppoints() is used in qqplot and qqnorm to generate the set of probabilities at which to evaluate the inverse distribution.

## See Also

qqplot, qqnorm.

## Examples

```
ppoints(4) # the same as ppoints(1:4)
ppoints(10)
ppoints(10, a=1/2)
```

```
precip Annual Precipitation in US Cities
```


## Description

The average amount of precipitation (rainfall) in inches for each of 70 United States (and Puerto Rico) cities.

## Usage

```
data(precip)
```


## Format

A named vector of length 70 .

## Source

Statistical Abstracts of the United States, 1975.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(precip)
dotchart(precip[order(precip)], main = "precip data")
title(sub = "Average annual precipitation (in.)")
```

predict Model Predictions

## Description

predict is a generic function for predictions from the results of various model fitting functions. The function invokes particular methods which depend on the class of the first argument.

The function predict. 1 m makes predictions based on the results produced by lm.

## Usage

predict (object, ...)

## Arguments

```
object a model object for which prediction is desired.
... additional arguments affecting the predictions produced.
```


## Value

The form of the value returned by predict depends on the class of its argument. See the documentation of the particular methods for details of what is produced by that method.

## See Also

predict.lm.

## Examples

```
## All the "predict" methods available in your current search() path:
for(fn in methods("predict"))
    cat(fn,":\n\t",deparse(args(get(fn))),"\n")
```


## Description

Obtains predictions and optionally estimates standard errors of those predictions from a fitted generalized linear model object.

## Usage

```
predict(object, newdata = NULL, type = c("link", "response", "terms"),
        se.fit = FALSE, dispersion = NULL, terms = NULL, ...)
```


## Arguments

| object | a fitted object of class inheriting from "glm". |
| :--- | :--- |
| newdata | optionally, a new data frame from which to make the predictions. If <br> omitted, the fitted linear predictors are used. <br> the type of prediction required. The default is on the scale of the linear <br> predictors; the alternative "response" is on the scale of the response <br> variable. Thus for a default binomial model the default predictions are of <br> log-odds (probabilities on logit scale) and type = "response" gives the <br> predicted probabilities. The "terms" option returns a matrix giving the <br> fitted values of each term in the model formula on the linear predictor <br> scale. |
|  | The value of this argument can be abbreviated. <br> logical switch indicating if standard errors are required. |
| se.fit | the dispersion of the GLM fit to be assumed in computing the standard <br> errors. If omitted, that returned by summary applied to the object is used. |
| dispersion | with type="terms" by default all terms are returned. A character vector |
| terms | specifies which terms are to be returned <br> further arguments passed to or from other methods. |
|  |  |

## Value

If $\mathrm{se}=\mathrm{FALSE}$, a vector or matrix of predictions. If se $=$ TRUE, a list with components

```
fit Predictions
se.fit Estimated standard errors
residual.scale
```

A scalar giving the square root of the dispersion used in computing the standard errors.

## Author(s)

B.D. Ripley

## See Also

glm, SafePrediction

## Examples

```
## example from Venables and Ripley (1997, pp. 231-3.)
ldose <- rep(0:5, 2)
numdead <- c(1, 4, 9, 13, 18, 20, 0, 2, 6, 10, 12, 16)
sex <- factor(rep(c("M", "F"), c(6, 6)))
SF <- cbind(numdead, numalive=20-numdead)
budworm.lg <- glm(SF ~ sex*ldose, family=binomial)
summary(budworm.lg)
plot(c(1,32), c(0,1), type = "n", xlab = "dose",
    ylab = "prob", log = "x")
text(2^ldose, numdead/20, as.character(sex))
ld <- seq(0, 5, 0.1)
lines(2^ld, predict(budworm.lg, data.frame(ldose=ld,
    sex=factor(rep("M", length(ld)), levels=levels(sex))),
    type = "response"))
lines(2^ld, predict(budworm.lg, data.frame(ldose=ld,
    sex=factor(rep("F", length(ld)), levels=levels(sex))),
    type = "response"))
```


## Description

Predicted values based on linear model object

## Usage

```
predict(object, newdata, se.fit = FALSE, scale = NULL, df = Inf,
    interval = c("none", "confidence", "prediction"),
    level = 0.95, type = c("response", "terms"),
    terms = NULL, ...)
```


## Arguments

| object | Object of class inheriting from "lm" |
| :--- | :--- |
| newdata | Data frame in which to predict |
| se.fit | A switch indicating if standard errors are required. |
| scale | Scale parameter for std.err. calculation |
| df | Degrees of freedom for scale |
| interval | Type of interval calculation |
| level | Tolerance/confidence level |
| type | Type of prediction (response or model term) |
| terms | If type="terms", which terms (default is all terms) |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

predict.lm produces predicted values, obtained by evaluating the regression function in the frame newdata (which defaults to model.frame (object). If the logical se.fit is TRUE, standard errors of the predictions are calculated. If the numeric argument scale is set (with optional df), it is used as the residual standard deviation in the computation of the standard errors, otherwise this is extracted from the model fit. Setting intervals specifies computation of confidence or prediction (tolerance) intervals at the specified level.

If the fit is rank-deficient, some of the columns of the design matrix will have been dropped. Prediction from such a fit only makes sense if newdata is contained in the same subspace as the original data. That cannot be checked accurately, so a warning is issued.

## Value

predict.lm produces a vector of predictions or a matrix of predictions and bounds with column names fit, lwr, and upr if interval is set. If se.fit is TRUE, a list with the following components is returned:

```
fit vector or matrix as above
se.fit standard error of predictions
residual.scale
    residual standard deviations
df degrees of freedom for residual
```


## Note

Offsets specified by offset in the fit by lm will not be included in predictions, whereas those specified by an offset term in the formula will be.

## See Also

The model fitting function lm, predict, SafePrediction

## Examples

```
## Predictions
x <- rnorm(15)
y <- x + rnorm(15)
predict(lm(y ~ x))
new <- data.frame(x = seq(-3, 3, 0.5))
predict(lm(y ~ x), new, se.fit = TRUE)
pred.w.plim <- predict(lm(y ~ x), new, interval="prediction")
pred.w.clim <- predict(lm(y ~ x), new, interval="confidence")
matplot(new$x,cbind(pred.w.clim, pred.w.plim[,-1]),
    lty=c(1,2,2,3,3), type="l", ylab="predicted y")
```

```
preplot Pre-computations for a Plotting Objeect
```


## Description

Compute an object to be used for plots relating to the given model object.

## Usage

```
    preplot(object, ...)
```


## Arguments

object a fitted model object.
... additional arguments for specific methods.

## Details

Only the generic function is currently provided in base R, but some add-on packages have methods. Principally here for S compatibility.

## Value

An object set up to make a plot that describes object.

```
presidents Approval Rating of US Presidents
```


## Description

The (approximately) quarterly approval rating for the President of the United states from the first quarter of 1945 to the last quarter of 1974.

## Usage

data(presidents)

## Format

A time series of 120 values.

## Details

The data are actually a fudged version of the approval ratings. See McNeil's book for details.

## Source

The Gallup Organisation.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

data(presidents)
plot(presidents, las = 1, ylab = "Approval rating (\%)", main = "presidents data")
pressure Vapor Pressure of Mercury as a Function of Temperature

## Description

Data on the relation between temperature in degrees Celsius and vapor pressure of mercury in millimeters (of mercury).

## Usage

data(pressure)

## Format

A data frame with 19 observations on 2 variables.

$$
\begin{array}{llll}
{[, 1]} & \text { temperature } & \text { numeric } & \text { temperature }(\operatorname{deg} \mathrm{C}) \\
{[, 2]} & \text { pressure } & \text { numeric } & \text { pressure }(\mathrm{mm})
\end{array}
$$

## Source

Weast, R. C., ed. (1973) Handbook of Chemistry and Physics. CRC Press.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(pressure)
plot(pressure, xlab = "Temperature (deg C)",
    ylab = "Pressure (mm of Hg)",
    main = "pressure data: Vapor Pressure of Mercury")
plot(pressure, xlab = "Temperature (deg C)", log = "y",
    ylab = "Pressure (mm of Hg)",
    main = "pressure data: Vapor Pressure of Mercury")
```

    pretty Pretty Breakpoints
    
## Description

Compute a sequence of about $n+1$ equally spaced nice values which cover the range of the values in x . The values are chosen so that they are 1,2 or 5 times a power of 10 .

## Usage

```
pretty ( \(\mathrm{x}, \mathrm{n}=5\), min. \(\mathrm{n}=\mathrm{n} \% / \% 3\), shrink.sml \(=0.75\),
    high.u.bias \(=1.5\), u5.bias \(=.5+1.5 *\) high.u.bias,
    eps.correct \(=0\) )
```


## Arguments

x numeric vector
n
integer giving the desired number of intervals. Non-integer values are rounded down.
min.n nonnegative integer giving the minimal number of intervals. If min.n == 0 , pretty (.) may return a single value.
shrink.sml positive numeric by a which a default scale is shrunk in the case when range ( x ) is "very small" (usually 0 ).
high.u.bias non-negative numeric, typically $>1$. The interval unit is determined as $\{1,2,5,10\}$ times b , a power of 10. Larger high.u.bias values favor larger units.
u5.bias non-negative numeric multiplier favoring factor 5 over 2. Default and "optimal": u5.bias = $5+1.5 * h i g h . u . b i a s$.
eps.correct integer code, one of $\{0,1,2\}$. If non- 0 , an "epsilon correction" is made at the boundaries such that the result boundaries will be outside range ( x ); in the small case, the correction is only done if eps.correct $>=2$.

## Details

Let $\mathrm{d}<-\max (\mathrm{x})-\min (\mathrm{x}) \geq 0$. If d is not (very close) to 0 , we let $\mathrm{c}<-\mathrm{d} / \mathrm{n}$, otherwise more or less $\mathrm{c}<-\max (\operatorname{abs}($ range (x)) ) $*$ shrink.sml / min.n. Then, the 10 base b is $10^{\left\lfloor\log _{10}(c)\right\rfloor}$ such that $b \leq c<10 b$.

Now determine the basic unit $u$ as one of $\{1,2,5,10\} b$, depending on $c / b \in[1,10)$ and the two "bias" coefficients, $h=$ high.u.bias and $f=$ u5.bias.

## Examples

```
pretty(1:15) # 0 2 4 4 6 8
pretty(1:15, h=2)# 0 5 10 15
pretty(1:15, n=4)# 0 5 10 15
pretty(1:15 * 2) # 0 5 10 15 20 25 30
pretty(1:20) # 0 5 10 15 20
pretty(1:20, n=2) # 0 10 20
pretty(1:20, n=10)# 0 2 4 ... 20
for(k in 5:11) {
    cat("k=",k,": "); print(diff(range(pretty(100 + c(0, pi*10^-k)))))}
##-- more bizarre, when min(x) == max(x):
pretty(pi)
```

```
add.names <- function(v) { names(v) <- paste(v); v}
str(lapply(add.names(-10:20), pretty))
str(lapply(add.names(0:20), pretty, min = 0))
sapply( add.names(0:20), pretty, min = 4)
pretty(1.234e100)
pretty(1001.1001)
pretty(1001.1001, shrink = .2)
for(k in -7:3)
    cat("shrink=",formatC(2^k,wid=9),":",
        formatC(pretty(1001.1001, shrink = 2^k), wid=6),"\n")
```

    Primitive Call a "Primitive" Internal Function
    
## Description

.Primitive returns an entry point to a "primitive" (internally implemented) function.
The advantage of .Primitive over. Internal functions is the potential efficiency of argument passing.

## Usage

.Primitive(name)

## Arguments

name name of the R function.

## See Also

. Internal.

## Examples

```
mysqrt <- .Primitive("sqrt")
c
.Internal # this one *must* be primitive!
get("if") # just 'if' or 'print(if)' are not syntactically ok.
```

```
print Print Values
```


## Description

print prints its argument and returns it invisibly (via invisible(x)). It is a generic function which means that new printing methods can be easily added for new classes.

## Usage

```
print(x, ...)
print.factor(x, quote = FALSE, max.levels = NULL,
    width = getOption("width"), ...)
```


## Arguments

$\mathrm{x} \quad$ an object used to select a method.
... further arguments passed to or from other methods.
quote logical, indicating whether or not strings should be printed with surrounding quotes.
max.levels integer, indicating how many levels should be printed for a factor; if 0 , no extra "Levels" line will be printed. The default, NULL, entails chosing max.levels such that the levels print on one line of width width.
width only used when max.levels is NULL, see above.

## Details

The default method, print.default has its own help page. Use methods("print") to get all the methods for the print generic.
print.factor allows some customization and is used for printing ordered factors as well.
See noquote as an example of a class whose main purpose is a specific print method.

## See Also

The default method print.default, and help for the methods above; further options, noquote.

## Examples

```
ts(1:20)#-- print is the ''Default function'' --> print.ts(.) is called
rr <- for(i in 1:3) print(1:i)
rr
## Printing of factors illustrated for ex
```

```
print.coefmat Print Coefficient Matrices
```


## Description

Utility function to be used in "higher level" print methods, such as print.summary.lm, print.summary.glm and print.anova. The goal is to provide a flexible interface with smart defaults such that often, only x needs to be specified.

## Usage

```
print.coefmat(x, digits=max(3, getOption("digits") - 2),
    signif.stars = getOption("show.signif.stars"),
    dig.tst = max(1, min(5, digits - 1)),
    cs.ind = 1:k, tst.ind = k + 1, zap.ind = integer(0),
    P.values = NULL,
    has.Pvalue = nc >= 4 && substr(colnames(x)[nc],1,3) == "Pr(",
    eps.Pvalue = .Machine$double.eps,
    na.print = "", ...)
```


## Arguments

| x | a numeric matrix like object, to be printed. |
| :--- | :--- |
| digits | minimum number of significant digits to be used for most numbers. |
| signif.stars | logical; if TRUE, P-values are additionally encoded visually as "significance <br> stars" in order to help scanning of long coefficient tables. It defaults to <br> the show.signif.stars slot of options. |
| dig.tst | minimum number of significant digits for the test statistics, see tst.ind. <br> indices (integer) of column numbers which are (like) coefficients and <br> standard errors to be formatted together. |
| tst.ind | indices (integer) of column numbers for test statistics. <br> indices (integer) of column numbers which should be formatted by <br> zapsmall, i.e., by "zapping" values close to 0. |
| zap.ind | logical or NULL; if TRUE, the last column of x is formatted by format.pval <br> as P values. If P.values = NULL, the default, it is set to TRUE only |
| if link\{options\} ("show.coef.Pvalue") is TRUE and x has at least 4 |  |
| columns and the last column name of x starts with "Pr(". |  |

## Details

Despite its name, this is not (yet) a method for the generic print function, because there is no class "coefmat".

## Value

Invisibly returns its argument, x.

## Author(s)

Martin Maechler

## See Also

print.summary.lm, format.pval, format.

## Examples

```
cmat <- cbind(rnorm(3, 10), sqrt(rchisq(3, 12)))
cmat <- cbind(cmat, cmat[,1]/cmat[,2])
cmat <- cbind(cmat, 2*pnorm(-cmat[,3]))
colnames(cmat) <- c("Estimate", "Std.Err", "Z value", "Pr(>z)")
print.coefmat(cmat[,1:3])
print.coefmat(cmat)
options(show.coef.Pvalues = FALSE)
print.coefmat(cmat, digits=2)
print.coefmat(cmat, digits=2, P.values = TRUE)
options(show.coef.Pvalues = TRUE)# revert
```

print.data.frame Printing Data Frames

## Description

These functions create or manipulate data frames, tightly coupled collections of variables which share many of the properties of matrices and of lists, used as the fundamental data structure by most of R's modeling software.

## Usage

```
print(x, ..., digits = NULL, quote = FALSE, right = TRUE)
```


## Arguments

x
object of class data.frame.
... optional arguments to print or plot methods.
digits the minimum number of significant digits to be used.
quote logical, indicating whether or not strings (characters) should be printed with surrounding quotes.
right logical, indicating whether or not strings should be right-aligned. The default is left-alignment.

## Value

For the print method (print.data.frame), see print.matrix.

## See Also

data.frame.

```
print.default Default Printing
```


## Description

print.default is the default method of the generic print function which prints its argument.
print.atomic is almost the same and exists purely for compatibility reasons.

## Usage

```
print.default(x, digits = NULL, quote = TRUE, na.print = NULL,
    print.gap = NULL, right = FALSE, ...)
    print.atomic(x, quote = TRUE, ...)
```


## Arguments

| x | the object to be printed. |
| :--- | :--- |
| digits | a non-null value for digits specifies the minimum number of significant <br> digits to be printed in values. If digits is NULL, the value of digits set <br> by options is used. |
| quote | logical, indicating whether or not strings (characters) should be printed <br> with surrounding quotes. |
| na.print | a character string which is used to indicate NA values in printed output, <br> or NULL (see Details) |
| print.gap | an integer, giving the spacing between adjacent columns in printed ma- <br> trices and arrays, or NULL meaning 1. <br> logical, indicating whether or not strings should be right-aligned. The <br> default is left-alignment. |
| right | (further arguments, currently disregarded) |
| $\ldots$ |  |

## Details

The default for printing NAs is to print NA (without quotes) unless this is a character NA and quote $=$ FALSE, when <NA> is printed.
The same number of decimal places is used throughout a vector, This means that digits specifies the minimum number of significant digits to be used, and that at least one entry will be printed with that minimum number.

## See Also

The generic print, options. The "noquote" class and print method.

## Examples

```
pi
print(pi, digits = 16)
LETTERS[1:16]
print(LETTERS, quote = FALSE)
```


## Description

Pseudo-method for the print generic. Especially useful with the right argument which does not (yet) exist for print. default.

## Usage

```
print.matrix(x, rowlab=character(0), collab=character(0),
    quote=TRUE, right=FALSE, na.print = NULL,
    print.gap = NULL, ...)
```


## Arguments

$x \quad$ numeric or character matrix.
rowlab, collab (optional) character vectors giving row or column names respectively. By default, these are taken from dimnames ( x ).
quote logical; if TRUE and x is of mode "character", quotes (") are used.
right if TRUE and x is of mode "character", the output columns are rightjustified.
na.print how NAs are printed. If this is non-null, its value is used to represent NA.
print.gap not yet used.
... arguments for other methods.

## Details

print.matrix and print. default both print matrices, and each has at least an optional argument that the other lacks. Also, both directly dispatch into. Internal code directly instead of relying on each other. This mainly stems from historic compatibility and similar reasons should be changed in the future.
prmatrix is currently just an alias for print.matrix.

## Value

Invisibly returns its argument, x.

## See Also

print.default, and other print methods.

## Examples

```
print.matrix(m6 <- diag(6), row = rep("",6), coll=rep("",6))
chm <- matrix(scan(system.file("help", "AnIndex", package = "eda"),
    what = ""), , 2, byrow = TRUE)
chm #-> print.default(.) = 'same' as print.matrix(chm)
print.matrix(chm, collab = paste("Column",1:3), right=TRUE, quote=FALSE)
```

```
print.ts Printing Time-Series Objects
```


## Description

Print method for time series objects.

## Usage

print(x, calendar, ...)

## Arguments

x
calendar enable/disable the display of information about month names, quarter names or year when printing. The default is TRUE for a frequency of 4 or 12, FALSE otherwise.
... additional arguments to print.

## Details

This is the print methods for objects inheriting from class "ts".

## See Also

print, ts.

## Examples

```
print(ts(1:10, freq = 7, start = c(12, 2)), calendar = TRUE)
```

proc.time
Running Time of $R$

## Description

proc.time determines how much time (in seconds) the currently running $R$ process already consumed.

## Usage

proc.time()

## Value

A numeric vector of length 5, containing the user, system, and total elapsed times for the currently running $R$ process, and the cumulative sum of user and system times of any child processes spawned by it.
The resolution of the times will be system-specific; it is common for them to be recorded to of the order of $1 / 100$ second, and elapsed time is rounded to the nearest $1 / 100$.
It is most useful for "timing" the evaluation of R expressions, which can be done conveniently with system.time.

## Note

It is possible to compile R without support for proc.time, when the function will not exist.

## See Also

system.time for timing a valid R expression, gc.time for how much of the time was spent in garbage collection.

## Examples

```
ptm <- proc.time()
for (i in 1:50) mad(runif(500))
proc.time() - ptm
```

prod Product of Vector Elements

## Description

prod returns the product of all the values present in its arguments.

## Usage

```
prod(..., na.rm = FALSE)
```


## Arguments

```
... numeric vectors.
na.rm logical. Should missing values be removed?
```


## Details

If na.rm is FALSE an NA value in any of the arguments will cause a value of NA to be returned, otherwise NA values are ignored.

See Also sum, cumprod, cumsum.

## Examples

```
print(prod(1:7)) == print(gamma(8))
```

profile Generic Function for Profiling Models

## Description

Investigates behavior of objective function near the solution represented by fitted.
See documentation on method functions for further details.

## Usage

profile(fitted, ...)

## Arguments

fitted the original fitted model object.
... additional parameters. See documentation on individual methods.

## Value

A list with an element for each parameter being profiled. See the individual methods for further details.

## See Also

```
profile.nls in package nls, profile.glm in package MASS,...
```

proj Projections of Models

## Description

proj returns a matrix or list of matrices giving the projections of the data onto the terms of a linear model. It is most frequently used for aov models.

## Usage

```
proj (object, ...)
proj.aov (object, onedf = FALSE, unweighted.scale = FALSE, ...)
proj.aovlist(object, onedf = FALSE, unweighted.scale = FALSE, ...)
proj.default(object, onedf = TRUE, ...)
proj.lm (object, onedf = FALSE, unweighted.scale = FALSE, ...)
```


## Arguments

object An object of class "lm" or a class inheriting from it, or an object with a similar structure including in particular components qr and effects.
onedf A logical flag. If TRUE, a projection is returned for all the columns of the model matrix. If FALSE, the single-column projections are collapsed by terms of the model (as represented in the analysis of variance table).
unweighted.scale
If the fit producing object used weights, this determines if the projections correspond to weighted or unweighted observations.
... Swallow and ignore any other arguments.

## Details

A projection is given for each stratum of the object, so for aov models with an Error term the result is a list of projections.

## Value

A projection matrix or (for multi-stratum objects) a list of projection matrices.
Each projection is a matrix with a row for each observations and either a column for each term (onedf $=$ FALSE) or for each coefficient (onedf $=$ TRUE). Projection matrices from the default method have orthogonal columns representing the projection of the response onto the column space of the Q matrix from the QR decomposition. The fitted values are the sum of the projections, and the sum of squares for each column is the reduction in sum of squares from fitting that column (after those to the left of it).
The methods for 1 m and aov models add a column to the projection matrix giving the residuals (the projection of the data onto the orthogonal complement of the model space).
Strictly, when onedf = FALSE the result is not a projection, but the columns represent sums of projections onto the columns of the model matrix corresponding to that term. In this case the matrix does not depend on the coding used.

## Author(s)

B.D. Ripley

## See Also

aov, lm, model.tables

## Examples

```
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
        K=factor(K), yield=yield)
npk.aov <- aov(yield ~ block + N*P*K, npk)
proj(npk.aov)
## as a test, not particularly sensible
```

```
options(contrasts=c("contr.helmert", "contr.treatment"))
npk.aovE <- aov(yield ~ N*P*K + Error(block), npk)
proj(npk.aovE)
```

prompt Produce Prototype of an $R$ Documentation File

## Description

Facilitate the constructing of files documenting $R$ functions.

## Usage

```
prompt(object, ...)
prompt.default(object,
    filename = paste(name, ".Rd", sep = ""),
    name = NULL, force.function = FALSE, ...)
prompt.data.frame(object,
    filename = paste(name, ".Rd", sep = ""), name = NULL, ...)
```


## Arguments

| object | an R object, typically a function |
| :--- | :--- |
| name | character string specifying the name of the object. |
| filename | name of the output file |
| force.function |  |
|  | treat object as function in any case |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

An ASCII file filename is produced containing the proper function and argument names of object. You have to edit it before adding the documentation to the source tree, i.e., (currently) to '\$R_HOME/src/library/base/man/'.
When prompt is used in for loops or scripts, the explicit name specification will be useful.

## Note

The documentation file produced by prompt.data.frame does not have the same format as many of the data frame documentation files in the base library. We are trying to settle on a preferred format for the documentation.

## Author(s)

Douglas Bates for prompt.data.frame

## See Also

help and the chapter on "Writing R documentation" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).

## Examples

```
prompt(plot.default)
prompt(interactive, force.function = TRUE)
unlink("plot.default.Rd")
unlink("interactive.Rd")
data(women) # data.frame
prompt(women)
unlink("women.Rd")
data(sunspots) # non-data.frame data
prompt(sunspots)
unlink("sunspots.Rd")
```

prop.table Express table entries as fraction of marginal table

## Description

This is really sweep(x, margin, margin.table(x, margin), "/") for newbies, except that if margin has length zero, then one gets $x / \operatorname{sum}(x)$.

## Usage

prop.table(x, margin=NULL)

## Arguments

| $x$ | table |
| :--- | :--- |
| margin | index, or vector of indices to generate margin for |

## Value

Table like x expressed relative to margin

Author(s)
Peter Dalgaard

See Also
margin.table

## Examples

```
m<-matrix(1:4,2)
m
prop.table(m,1)
```

```
pushBack Push Text Back on to a Connection
```


## Description

Functions to push back text lines onto a connection, and to enquire how many lines are currently pushed back.

## Usage

pushBack(data, connection, newLine = TRUE)
pushBackLength(connection)

## Arguments

```
data a character vector.
connection A connection.
newLine logical. If true, a newline is appended to each string pushed back.
```


## Details

Several character strings can be pushed back on one or more occasions. The occasions form a stack, so the first line to be retrieved will be the first string from the last call to pushBack. Lines which are pushed back are read prior to the normal input from the connection, by the normal text-reading functions such as readLines and scan.

Pushback is only allowed for readable connections.
Not all uses of connections respect pushbacks, in particular the input connection is still wired directly, so for example parsing commands from the console and scan("") ignore pushbacks on stdin.

## Value

pushBack returns nothing.
pushBackLength returns number of lines currently pushed back.

## See Also

```
connections, readLines.
```


## Examples

```
zz <- textConnection(LETTERS)
readLines(zz, 2)
pushBack(c("aa", "bb"), zz)
pushBackLength(zz)
readLines(zz, 1)
pushBackLength(zz)
readLines(zz, 1)
readLines(zz, 1)
close(zz)
```


## Description

qqnorm is a generic functions the default method of which produces a normal QQ plot of the values in y . qqline adds a line to a normal quantile-quantile plot which passes through the first and third quartiles.
qqplot produces a QQ plot of two datasets.
Graphical parameters may be given as arguments to qqnorm, qqplot and qqline.

## Usage

qqnorm(y, ...)
qqnorm(y, ylim, main = "Normal Q-Q Plot", xlab = "Theoretical Quantiles", ylab = "Sample Quantiles", plot.it = TRUE, ...)
qqline (y, ...)
qqplot(x, y, plot.it $=$ TRUE, $x l_{\text {lab }}=$ deparse(substitute( $x$ )), ylab = deparse(substitute(y)), ...)

## Arguments

| x | The first sample for qqplot. |
| :--- | :--- |
| y | The second or only data sample. |
| xlab, ylab, main |  |
|  | plot labels. |
| plot.it | logical. Should the result be plotted? |
| ylim, ... | graphical parameters. |

## Value

For qqnorm and qqplot, a list with components
$\mathrm{x} \quad$ The x coordinates of the points that were/would be plotted
y The corresponding y coordinates

## See Also

ppoints.

## Examples

```
y <- rt(200, df = 5)
qqnorm(y); qqline(y, col = 2)
qqplot(y, rt(300, df = 5))
data(precip)
qqnorm(precip, ylab = "Precipitation [in/yr] for 70 US cities")
```


## Description

qr computes the QR decomposition of a matrix. It provides an interface to the techniques used in the LINPACK routine DQRDC or (for complex matrices) the LAPACK routine ZGEQP3.

## Usage

```
\(\mathrm{qr}(\mathrm{x}, \mathrm{tol}=1 \mathrm{e}-07\) )
qr.coef(qr, y)
qr.qy(qr, \(y\) )
qr.qty(qr, y)
qr.resid(qr, y)
qr.fitted(qr, y, k = qr\$rank)
qr.solve(a, b, tol = 1e-7)
is. \(\mathrm{qr}(\mathrm{x})\)
as.qr(x)
```


## Arguments

## x

a matrix whose QR decomposition is to be computed.
the tolerance for detecting linear dependencies in the columns of x .
qr a QR decomposition of the type computed by qr.
$y, b \quad a \quad v e c t o r ~ o r ~ m a t r i x ~ o f ~ r i g h t-h a n d ~ s i d e s ~ o f ~ e q u a t i o n s . ~$
a A matrix or QR decomposition.
k effective rank.

## Details

The QR decomposition plays an important role in many statistical techniques. In particular it can be used to solve the equation $\boldsymbol{A} \boldsymbol{x}=\boldsymbol{b}$ for given matrix $\boldsymbol{A}$, and vector $\boldsymbol{b}$. It is useful for computing regression coefficients and in applying the Newton-Raphson algorithm.
The functions qr.coef, qr.resid, and qr.fitted return the coefficients, residuals and fitted values obtained when fitting y to the matrix with QR decomposition qr. qr.qy and qr. qty return $\mathrm{Q} \% * \% \mathrm{y}$ and $\mathrm{t}(\mathrm{Q}) \% * \% \mathrm{y}$, where Q is the $\boldsymbol{Q}$ matrix.
All the above functions keep dimnames (and names) of $x$ and $y$ if there are.
qr.solve solves systems of equations via the QR decomposition.
is. qr returns TRUE if x is a list with components named qr , rank and qraux and FALSE otherwise.
It is not possible to coerce objects to mode "qr". Objects either are QR decompositions or they are not.

## Value

The QR decomposition of the matrix as computed by LINPACK or LAPACK. The components in the returned value correspond directly to the values returned by DQRDC/ZGEQP3.
qr a matrix with the same dimensions as x . The upper triangle contains the $\boldsymbol{R}$ of the decomposition and the lower triangle contains information on the $\boldsymbol{Q}$ of the decomposition (stored in compact form).
qraux a vector of length ncol (x) which contains additional information on $\boldsymbol{Q}$.
rank the rank of x as computed by the decomposition: always full rank in the complex case.
pivot information on the pivoting strategy used during the decomposition.

## Note

To compute the determinant of a matrix (do you really need it?), the QR decomposition is much more efficient than using Eigen values (eigen). See det.

The complex case uses column pivoting and does not attempt to detect rank-deficient matrices.

## References

Dongarra, J. J., Bunch, J. R., Moler, C. B. and Stewart, G. W. (1978) LINPACK Users Guide. Philadelphia: SIAM Publications.

Anderson. E. and ten others (1999) LAPACK Users' Guide. Third Edition. SIAM. Available on-line at http://www.netlib.org/lapack/lug/lapack_lug.html.

## See Also

qr.Q, qr.R, qr.X for reconstruction of the matrices. solve.qr, lsfit, eigen, svd. det (using qr) to compute the determinant of a matrix.

## Examples

```
hilbert <- function(n) { i <- 1:n; 1 / outer(i - 1, i, "+") }
h9 <- hilbert(9); h9
qr(h9)$rank #--> only 7
qrh9 <- qr(h9, tol = 1e-10)
qrh9$rank #--> 9
##-- Solve linear equation system H %*% x = y :
y <- 1:9/10
x <- qr.solve(h9, y, tol = 1e-10) # or equivalently :
x <- qr.coef(qrh9, y) #-- is == but much better than
        #-- solve(h9) %*% y
h9 %*% x
                                # = y
```

QR.Auxiliaries Reconstruct the $Q$, R, or X Matrices from a QR Object

## Description

Returns the original matrix from which the object was constructed or the components of the decomposition.

## Usage

```
qr.X(qr, complete = FALSE, ncol =)
qr.Q(qr, complete = FALSE, Dvec = 1)
qr.R(qr, complete = FALSE)
```


## Arguments

qr object representing a QR decomposition. This will typically have come from a previous call to qr or lsfit.
complete logical expression of length 1 . Indicates whether an arbitrary orthogonal completion of the $\boldsymbol{Q}$ or $\boldsymbol{X}$ matrices is to be made, or whether the $\boldsymbol{R}$ matrix is to be completed by binding zero-value rows beneath the square upper triangle.
ncol integer in the range 1:nrow(qr\$qr). The number of columns to be in the reconstructed $\boldsymbol{X}$. The default when complete is FALSE is the first $\min (\mathrm{ncol}(\mathrm{X}), \operatorname{nrow}(\mathrm{X}))$ columns of the original $\boldsymbol{X}$ from which the qr object was constructed. The default when complete is TRUE is a square matrix with the original $\boldsymbol{X}$ in the first ncol (X) columns and an arbitrary orthogonal completion (unitary completion in the complex case) in the remaining columns.

Dvec vector (not matrix) of diagonal values. Each column of the returned $\boldsymbol{Q}$ will be multiplied by the corresponding diagonal value.

## Value

qr. X returns $\boldsymbol{X}$, the original matrix from which the qr object was constructed, provided ncol(X) <= nrow (X). If complete is TRUE or the argument ncol is greater than ncol(X), additional columns from an arbitrary orthogonal (unitary) completion of X are returned.
qr. Q returns $\mathbf{Q}$, the order-nrow $(X)$ orthogonal (unitary) transformation represented by qr. If complete is TRUE, $\mathbf{Q}$ has nrow ( X ) columns. If complete is FALSE, $\mathbf{Q}$ has ncol(X) columns. When Dvec is specified, each column of $\mathbf{Q}$ is multiplied by the corresponding value in Dvec.
qr. R returns $\mathbf{R}$, the upper triangular matrix such that $X==Q \% * \%$. The number of rows of $\mathbf{R}$ is $\operatorname{nrow}(X)$ or $\operatorname{ncol}(X)$, depending on whether complete is TRUE or FALSE.

## See Also

```
qr, qr.qy.
```


## Examples

```
data(LifeCycleSavings)
p <- ncol(x <- LifeCycleSavings[,-1]) # not the 'sr'
qrstr <- qr(x) # dim(x) == c(n,p)
qrstr $ rank # = 4 = p
Q <- qr.Q(qrstr) # dim(Q) == dim(x)
R <- qr.R(qrstr) # dim(R) == ncol(x)
X <- qr.X(qrstr) # X == x
range(X - as.matrix(x))# ~ < 6e-12
## X == Q %*% R :
all((1 - X /( Q %*% R))< 100*.Machine$double.eps)#TRUE
dim(Qc <- qr.Q(qrstr, complete=TRUE)) # Square: dim(Qc) == rep(nrow(x),2)
all((crossprod(Qc) - diag(nrow(x))) < 10*.Machine $double.eps)
QD <- qr.Q(qrstr, D=1:p) # QD == Q %*% diag(1:p)
all(QD - Q %*% diag(1:p) < 8* .Machine$double.eps)
dim(Rc <- qr.R(qrstr, complete=TRUE)) # == dim(x)
dim(Xc <- qr.X(qrstr, complete=TRUE)) # square: nrow(x) ^ 2
all(Xc[,1:p] == X)
```


## quakes

Locations of Earthquakes off Fiji

## Description

The data set give the locations of 1000 seismic events of MB $>4.0$. The events occurred in a cube near Fiji since 1964.

## Usage

data(quakes)

## Format

A data frame with 1000 observations on 5 variables.

| $[, 1]$ | lat | numeric | Latitude of event |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | long | numeric | Longitude |
| $[, 3]$ | depth | numeric | Depth $(\mathrm{km})$ |
| $[, 4]$ | mag | numeric | Richter Magnitude |
| $[, 5]$ | stations | numeric | Number of stations reporting |

## Details

There are two clear planes of seismic activity. One is a major plate junction; the other is a trench off New Zealand. These data constitute a subsample from a larger dataset of containing 5000 observations.

## Source

This is one of the Harvard PRIM-H project data sets. They in turn obtained it from Dr. John Woodhouse, Dept. of Geophysics, Harvard University.

## Examples

data(quakes)
pairs(quakes, main = "Fiji Earthquakes, $N=1000 "$, cex.main=1.2, pch=".")
quantile Sample Quantiles

## Description

The generic function quantile produces sample quantiles corresponding to the given probabilities. The smallest observation corresponds to a probability of 0 and the largest to a probability of 1 .

## Usage

quantile(x, probs $=$ seq( $0,1,0.25$ ), na.rm $=$ FALSE , names = TRUE, ...)

## Arguments

x
probs
na.rm logical; if true, any NA and NaN's are removed from $x$ before the quantiles are computed.
names logical; if true, the result has a names attribute. Set to FALSE for speedup with many probs.
... further arguments passed to or from other methods.

## Details

A vector of length length(probs) is returned; if names = TRUE, it has a names attribute. quantile $(x, p)$ as a function of $p$ linearly interpolates the points ( $(i-1) /(n-1)$, ox $[i]$ ), where ox <- sort( $x$ ) and $n<-$ length $(x)$.

This gives quantile $(x, p)==(1-f) * o x[i]+f * o x[i+1]$, where $r<-1+(n-1) * p$, i <- floor (r), f <- r - i and ox[n+1] := ox[n].

NA and NaN values in probs are propagated to the result.

## See Also

ecdf (in the stepfun package) for empirical distributions of which quantile is the "inverse". boxplot.stats and fivenum for computing "versions" of quartiles, etc.

## Examples

```
quantile(x <- rnorm(1001))# Extremes & Quartiles by default
quantile(x, probs=c(.1,.5,1,2,5,10,50, NA)/100)
n <- length(x) ## the following is exact, because 1/(1001-1) is exact:
stopifnot(sort(x) == quantile(x, probs = ((1:n)-1)/(n-1), names=FALSE))
n <- 777
ox <- sort(x <- round(rnorm(n),1))# round() produces ties
ox <- c(ox, ox[n]) #- such that ox[n+1] := ox[n]
p <- c(0,1,runif(100))
i <- floor(r <- 1 + (n-1)*p)
f <- r - i
all(abs(quantile(x,p) - ((1-f)*ox[i] + f*ox[i+1])) < 20*.Machine$double.eps)
```

```
quartz MacOS X Quartz device
```


## Description

quartz starts a graphics device driver for the MacOS X System. This can only be done on machines that run MacOS X.

## Usage

```
quartz(display = "", width = 6, height = 6, pointsize = 12,
    family = "Helvetica", antialias = TRUE, autorefresh = TRUE)
```


## Arguments

display the display on which the graphics window will appear. The default is to use the value in the user's environment variable DISPLAY.
width the width of the plotting window in inches.
height the height of the plotting window in inches.
pointsize the default pointsize to be used.
family this is the family name of the Postscript font that will be used by the device.
antialias whether to use antialiasing. It is never the case to set it FALSE
autorefresh this sets realtime refreshing. If it is set to FALSE, then the System is charged to refresh the context of the device window.

## Details

Quartz is the graphic engine based on the PDF format. It is used by the graphic interface of MacOS X to render high quality graphics. As PDF it is device independent and can be rescaled without loss of definition.

## See Also

Devices.

## quit $\quad$ Terminate an $R$ Session

## Description

The function quit or its alias q terminate the current R session.

## Usage

```
quit(save = "default", status = 0, runLast = TRUE)
    q(save = "default", status = 0, runLast = TRUE)
.Last <- function(x) { ...... }
```


## Arguments

| save | a character string indicating whether the environment (workspace) should <br> be saved, one of "no", "yes", "ask" or "default". |
| :--- | :--- |
| status | the (numerical) error status to be returned to the operating system, where <br> relevant. Conventionally 0 indicates successful completion. |
| runLast | should .Last() be executed? |

## Details

save must be one of "no", "yes", "ask" or "default". In the first case the workspace is not saved, in the second it is saved and in the third the user is prompted and can also decide not to quit. The default is to ask in interactive use but may be overridden by command-line arguments (which must be supplied in non-interactive use).
Immediately before terminating, the function . Last () is executed if it exists and runLast is true. If in interactive use there are errors in the .Last function, control will be returned to the command prompt, so do test the function thoroughly.
Some error statuses are used by R itself. The default error handler for non-interactive effectively calls q("no", 1, FALSE) and returns error code 1. Error status 2 is used for R 'suicide', that is a catastrophic failure, and other small numbers are used by specific ports for initialization failures. It is recommended that users choose statuses of 10 or more.
Valid values of status are system-dependent, but 0:255 are normally valid.

## See Also

.First for setting things on startup.

## Examples

```
## Unix-flavour example
.Last <- function() {
    cat("Now sending PostScript graphics to the printer:\n")
    system("lpr Rplots.ps")
    cat("bye bye...\n")
}
quit("yes")
```

R.home

Return the R Home Directory

## Description

Return the R home directory.

## Usage

R.home()

## Value

A character string giving the current home directory.
R.Version Version Information

## Description

R.Version() provides detailed information about the version of R running.
R.version is a variable (a list) holding this information (and version is a copy of it for $S$ compatibility), whereas R.version.string is a simple character string, useful for plotting, etc.

## Usage

R.Version()
R.version
R.version.string

## Value

R.Version returns a list with components
platform the platform for which R was built. Under Unix, a triplet of the form CPU-VENDOR-OS, as determined by the configure script. E.g, "i586-unknown-linux".
arch the architecture (CPU) R was built on/for.
os the underlying operating system
system CPU and OS.
status the status of the version (e.g., "Alpha")
status.rev the status revision level
major the major version number
minor the minor version number
year the year the version was released
month the month the version was released
day the day the version was released
language always "R".

## Note

Do not use R.version\$os to test the platform the code is running on: use .Platform\$OS.type instead. Slightly different versions of the OS may report different values of R.version\$os, as may different versions of $R$.

## See Also

.Platform.

## Examples

R.version\$os \# to check how lucky you are ... plot(0) \# any plot
mtext(R.version.string, side=1,line=4,adj=1)\# a useful bottom-right note

Random Random Number Generation

## Description

.Random.seed is an integer vector, containing the random number generator (RNG) state for random number generation in R. It can be saved and restored, but should not be altered by the user.

RNGkind is a more friendly interface to query or set the kind of RNG in use. set.seed is the recommended way to specify seeds.

## Usage

```
.Random.seed <- c(rng.kind, n1, n2, ...)
save.seed <- .Random.seed
RNGkind(kind = NULL, normal.kind = NULL)
set.seed(seed, kind = NULL)
```


## Arguments

kind character or NULL. If kind is a character string, set R's RNG to the kind desired. If it is NULL, return the currently used RNG. Use "default" to return to the R default.
normal.kind character string or NULL. If it is a character string, set the method of Normal generation. Use "default" to return to the R default.
seed a single value, interpreted as an integer.
rng.kind integer code in $0: \mathrm{k}$ for the above kind.
$\mathrm{n} 1, \mathrm{n} 2, \ldots$ integers. See the details for how many are required (which depends on rng.kind).

## Details

The currently available RNG kinds are given below. kind is partially matched to this list. The default is "Marsaglia-Multicarry".
"Wichmann-Hill" The seed, .Random.seed[-1] == r[1:3] is an integer vector of length 3 , where each $r[i]$ is in 1 : $(\mathrm{p}[\mathrm{i}]-1)$, where p is the length 3 vector of primes, $p=(30269,30307,30323)$. The Wichmann-Hill generator has a cycle length of $6.9536 \times 10^{12}(=\operatorname{prod}(\mathrm{p}-1) / 4$, see Applied Statistics (1984) 33, 123 which corrects the original article).
"Marsaglia-Multicarry": A multiply-with-carry RNG is used, as recommended by George Marsaglia in his post to the mailing list 'sci.stat.math'. It has a period of more than $2^{60}$ and has passed all tests (according to Marsaglia). The seed is two integers (all values allowed).
"Super-Duper": Marsaglia's famous Super-Duper from the 70's. This is the original version which does not pass the MTUPLE test of the Diehard battery. It has a period of $\approx 4.6 \times 10^{18}$ for most initial seeds. The seed is two integers (all values allowed for the first seed: the second must be odd).
We use the implementation by Reeds et al. (1982-84).
The two seeds are the Tausworthe and congruence long integers, respectively. A one-to-one mapping to S's .Random.seed[1:12] is possible but we will not publish one, not least as this generator is not exactly the same as that in recent versions of S-PLUS.
"Mersenne-Twister": From Matsumoto and Nishimura (1998). A twisted GFSR with period $2^{19937}-1$ and equidistribution in 623 consecutive dimensions (over the whole period). The "seed" is a 624 -dimensional set of 32 -bit integers plus a current position in that set.
"Knuth-TAOCP": From Knuth (1997). A GFSR using lagged Fibonacci sequences with subtraction. That is, the recurrence used is

$$
X_{j}=\left(X_{j-100}-X_{j-37}\right) \bmod 2^{30}
$$

and the "seed" is the set of the 100 last numbers (actually recorded as 101 numbers, the last being a cyclic shift of the buffer). The period is around $2^{129}$.
"Knuth-TAOCP-2002": The 2002 version which not backwards compatible with the earlier version: the initialization of the GFSR from the seed was altered. R did not allow you to choose consecutive seeds, the reported 'weakness', and already scrambled the seeds.
"user-supplied": Use a user-supplied generator. See Random.user for details.
normal.kind can be "Kinderman-Ramage" (the default), "Ahrens-Dieter", "Box-Muller", "Inversion" or "user-supplied". (For inversion, see the reference in qnorm.)
set. seed uses its single integer argument to set as many seeds as are required. It is intended as a simple way to get quite different seeds by specifying small integer arguments, and also as a way to get valid seed sets for the more complicated methods (especially "MersenneTwister" and "Knuth-TAOCP").

## Value

.Random.seed is an integer vector whose first element codes the kind of RNG and normal generator. The lowest two decimal digits are in in $0:(\mathrm{k}-1)$ where k is the number of available RNGs. The hundreds represent the type of normal generator (starting at 0 ).

In the underlying $C$, .Random.seed [-1] is unsigned; therefore in $R$.Random.seed [-1] can be negative.

RNGkind returns a two-element character vector of the RNG and normal kinds in use before the call, invisibly if either argument is not NULL.
set. seed returns NULL, invisibly.

## Note

Initially, there is no seed; a new one is created from the current time when one is required. Hence, different sessions will give different simulation results, by default.
.Random.seed saves the seed set for the uniform random-number generator, at least for the system generators. It does not necessarily save the state of other generators, and in particular does not save the state of the Box-Muller normal generator. If you want to reproduce work later, call set.seed rather than set. Random.seed.

## Author(s)

of RNGkind: Martin Maechler. Current implementation, B. D. Ripley

## References

Wichmann, B. A. and Hill, I. D. (1982) Algorithm AS 183: An Efficient and Portable Pseudo-random Number Generator, Applied Statistics, 31, 188-190; Remarks: 34, 198 and 35, 89.
De Matteis, A. and Pagnutti, S. (1993) Long-range Correlation Analysis of the WichmannHill Random Number Generator, Statist. Comput., 3, 67-70.
Marsaglia, G. (1997) A random number generator for C. Discussion paper, posting on Usenet newsgroup sci.stat.math on September 29, 1997.

Reeds, J., Hubert, S. and Abrahams, M. (1982-4) C implementation of SuperDuper, University of California at Berkeley. (Personal communication from Jim Reeds to Ross Ihaka.)
Marsaglia, G. and Zaman, A. (1994) Some portable very-long-period random number generators. Computers in Physics, 8, 117-121.
Matsumoto, M. and Nishimura, T. (1998) Mersenne Twister: A 623-dimensionally equidistributed uniform pseudo-random number generator, ACM Transactions on Modeling and Computer Simulation, 8, 3-30.
Source code at http://www.math.keio.ac.jp/~matumoto/emt.html.
Knuth, D. E. (1997) The Art of Computer Programming. Volume 2, third edition. Source code at http://www-cs-faculty.stanford.edu/~knuth/taocp.html.

Knuth, D. E. (2002) The Art of Computer Programming. Volume 2, third edition, ninth printing.
See http://Sunburn.Stanford.EDU/~knuth/news02.html.
Kinderman, A. J. and Ramage, J. G. (1976) Computer generation of normal random variables. Journal of the American Statistical Association 71, 893-896.
Ahrens, J.H. and Dieter, U. (1973) Extensions of Forsythe's method for random sampling from the normal distribution. Mathematics of Computation 27, 927-937.
Box, G.E.P. and Muller, M.E. (1958) A note on the generation of normal random deviates. Annals of Mathmatical Statistics 29, 610-611.

## See Also

runif, rnorm, ....

## Examples

```
runif(1); .Random.seed; runif(1); .Random.seed
## If there is no seed, a ''random'' new one is created:
rm(.Random.seed); runif(1); .Random.seed
RNGkind("Wich")# (partial string matching on 'kind')
## This shows how 'runif(.)' works for Wichmann-Hill,
## using only R functions:
p.WH <- c(30269, 30307, 30323)
a.WH <- c( 171, 172, 170)
next.WHseed <- function(i.seed = .Random.seed[-1])
    { (a.WH * i.seed) %% p.WH }
my.runif1 <- function(i.seed = .Random.seed)
    { ns <- next.WHseed(i.seed[-1]); sum(ns / p.WH) %% 1 }
rs <- .Random.seed
(WHs <- next.WHseed(rs[-1]))
u <- runif(1)
stopifnot(
    next.WHseed(rs[-1]) == .Random.seed[-1],
    all.equal(u, my.runif1(rs))
)
## ----
.Random.seed
ok <- RNGkind()
RNGkind("Super")#matches "Super-Duper"
RNGkind()
.Random.seed # new, corresponding to Super-Duper
## Reset:
RNGkind(ok[1])
```

Random.user User-supplied Random Number Generation

## Description

Function RNGkind allows user-coded uniform and normal random number generators to be supplied. The details are given here.

## Details

A user-specified uniform RNG is called from entry points in dynamically-loaded compiled code. The user must supply the entry point user_unif_rand, which takes no arguments and returns a pointer to a double. The example below will show the general pattern.

Optionally, the user can supply the entry point user_unif_init, which is called with an unsigned int argument when RNGkind (or set.seed) is called, and is intended to be used to initialize the user's RNG code. The argument is intended to be used to set the "seeds"; it is the seed argument to set.seed or an essentially random seed if RNGkind is called.

If only these functions are supplied, no information about the generator's state is recorded in .Random.seed. Optionally, functions user_unif_nseed and user_unif_seedloc can be
supplied which are called with no arguments and should return pointers to the number of "seeds" and to an integer array of "seeds". Calls to GetRNGstate and PutRNGstate will then copy this array to and from .Random.seed.
A user-specified normal RNG is specified by a single entry point user_norm_rand, which takes no arguments and returns a pointer to a double.

## Warning

As with all compiled code, mis-specifying these functions can crash R. Do include the 'R_ext/Random.h' header file for type checking.

## Examples

```
## Marsaglia's conguential PRNG
#include <R_ext/Random.h>
static Int32 seed;
static double res;
static int nseed = 1;
double * user_unif_rand()
{
    seed = 69069 * seed + 1;
    res = seed * 2.32830643653869e-10;
    return &res;
}
void user_unif_init(Int32 seed_in) { seed = seed_in; }
int * user_unif_nseed() { return &nseed; }
int * user_unif_seedloc() { return (int *) &seed; }
/* ratio-of-uniforms for normal */
#include <math.h>
static double x;
double * user_norm_rand()
{
    double u, v, z;
    do {
            u = unif_rand();
            v = 0.857764 * (2. * unif_rand() - 1);
            x = v/u; z = 0.25 * x * x;
            if (z < 1. - u) break;
            if (z > 0.259/u + 0.35) continue;
        } while (z > -log(u));
        return &x;
}
## Use under Unix:
R SHLIB urand.c
R
> dyn.load("urand.so")
 RNGkind("user")
runif(10)
> .Random.seed
R RNGkind(, "user")
```

```
> rnorm(10)
```

> RNGkind()
[1] "user-supplied" "user-supplied"
randu Random Numbers from Congruential Generator

## Description

400 triples of successive random numbers were taken from the VAX FORTRAN function RANDU running under VMS 1.5.

## Usage

data(randu)

## Format

A data frame with 400 observations on 3 variables named $x, y$ and $z$ which give the first, second and third random number in the triple.

## Details

In three dimensional displays it is evident that the triples fall on 15 parallel planes in 3-space. This can be shown theoretically to be true for all triples from the RANDU generator.
These particular 400 triples start 5 apart in the sequence, that is they are ( $(\mathrm{U}[5 \mathrm{i}+1], \mathrm{U}[5 \mathrm{i}+2]$, $\mathrm{U}[5 \mathrm{i}+3]), \mathrm{i}=0, \ldots, 399)$, and they are rounded to 6 decimal places.

Under VMS versions 2.0 and higher, this problem has been fixed.

## Source

David Donoho

## Examples

```
## We could re-generate the dataset by the following R code
seed <- as.double(1)
RANDU <- function() {
    seed <<- ((2^16 + 3) * seed) %% (2^31)
    seed/(2^31)
}
for(i in 1:400) {
    U <- c(RANDU(), RANDU(), RANDU(), RANDU(), RANDU())
    print(round(U[1:3], 6))
}
```

range Range of Values

## Description

range returns a vector containing the minimum and maximum of all the given arguments.

## Usage

```
range(..., na.rm = FALSE)
range.default(..., na.rm = FALSE, finite = FALSE)
```


## Arguments

| $\ldots$. | any numeric objects. |
| :--- | :--- |
| na.rm | logical, indicating if NA's should be omitted. |
| finite | logical, indicating if all non-finite elements should be omitted. |

## Details

This is a generic function; currently, it has only a default method (range.default).
It is also a member of the Summary group of functions, see Methods.
If na.rm is FALSE, NA and NaN values in any of the arguments will cause NA values to be returned, otherwise NA values are ignored.

If finite is TRUE, the minimum and maximum of all finite values is computed, i.e., finite=TRUE includes na.rm=TRUE.

A special situation occurs when there is no (after omission of NAs) nonempty argument left, see min.

## See Also

 min, max, Methods.
## Examples

```
print(r.x <- range(rnorm(100)))
diff(r.x) # the SAMPLE range
x <- c(NA, 1:3, -1:1/0); x
range(x)
range(x, na.rm = TRUE)
range(x, finite = TRUE)
```

```
rank Sample Ranks
```


## Description

Returns the sample ranks of the values in a numeric vector. Ties result in ranks being averaged.

## Usage

rank (x, na.last $=$ TRUE)

## Arguments

x a numeric vector.
na.last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed.

## See Also

order and sort.

## Examples

```
(r1 <- rank(x1 <- c(3,1,4,59,26)))
(r2 <- rank(x2 <- c(3,1,4,5,9,2,6,5,3,5))) # ties
## rank() is "idempotent": rank(rank(x)) == rank(x) :
stopifnot(rank(r1) == r1, rank(r2) == r2)
```

RdUtils Utilities for Processing Rd Files

## Description

Utilities for converting files in R documentation ( Rd ) format to other formats or create indices from them, and for converting documentation in other formats to Rd format.

## Usage

```
R CMD Rdconv [options] file
R CMD Rdindex [options] files
R CMD Rd2dvi [options] files
R CMD Rd2txt [options] file
R CMD Sd2Rd [options] file
```


## Arguments

| file | the path to a file to be processed. |
| :--- | :--- |
| files | a list of file names specifying the $R$ documentation sources to use, by <br> either giving the paths to the files, or the path to a directory with the |
| sources of a package. |  |

## Details

Rdconv converts Rd format to other formats. Currently, plain text, HTML, LaTeX, S version 3 ( Sd ), and S version 4 (.sgml) formats are supported. It can also extract the examples for run-time testing.

Rd2dvi and Rd2txt are user-level programs for producing DVI/PDF output or pretty text output from Rd sources.

Rdindex creates an index table from Rd files.
Sd2Rd converts S (version 3 or 4) documentation formats to Rd format.
Use R CMD foo --help to obtain usage information on utility foo.

## Note

Conversion to S version $3 / 4$ formats is rough: there are some . Rd constructs for which there is no natural analogue. They are intended as a starting point for hand-tuning.

## See Also

The chapter "Processing Rd format" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).
read.00Index
Read 00Index-style Files

## Description

Read item/description information from 00Index-style files. Such files are description lists rendered in tabular form, and currently used for the object, data and demo indices and 'TITLE' files of add-on packages.

## Usage

read.00Index(file)

## Arguments

file the name of a file to read data values from. If the specified file is " ", then input is taken from the keyboard (in this case input can be terminated by a blank line). Alternatively, file can be a connection, which will be opened if necessary, and if so closed at the end of the function call.

## Value

a character matrix with 2 columns named "Item" and "Description" which hold the items and descriptions.

## See Also

formatDL for the inverse operation of creating a 00Index-style file from items and their descriptions.

```
read.ftable Manipulate Flat Contingency Tables
```


## Description

Read, write and coerce "flat" contingency tables.

## Usage

```
read.ftable(file, sep = "", quote = "\"",
    row.var.names, col.vars, skip = 0)
write.ftable(x, file = "", quote = TRUE, digits = getOption("digits"))
as.table(x, ...)
```


## Arguments

file either a character string naming a file or a connection which the data are to be read from or written to. "" indicates input from the console for reading and output to the console for writing.
sep the field separator string. Values on each line of the file are separated by this string.
quote a character string giving the set of quoting characters for read.ftable; to disable quoting altogether, use quote="". For write.table, a logical indicating whether strings in the data will be surrounded by double quotes.
row.var.names a character vector with the names of the row variables, in case these cannot be determined automatically.
col.vars a list giving the names and levels of the column variables, in case these cannot be determined automatically.
skip the number of lines of the data file to skip before beginning to read data.
$x \quad$ an object of class "ftable".
digits an integer giving the number of significant digits to use for (the cell entries of) x .
... further arguments to be passed to or from methods.

## Details

read.ftable reads in a flat-like contingency table from a file. If the file contains the written representation of a flat table (more precisely, a header with all information on names and levels of column variables, followed by a line with the names of the row variables), no further arguments are needed. Similarly, flat tables with only one column variable the name of which is the only entry in the first line are handled automatically. Other variants can be dealt with by skipping all header information using skip, and providing the names of the row variables and the names and levels of the column variable using row.var.names and col.vars, respectively. See the examples below.
Note that flat tables are characterized by their "ragged" display of row (and maybe also column) labels. If the full grid of levels of the row variables is given, one should instead use read.table to read in the data, and create the contingency table from this using xtabs.
write.ftable writes a flat table to a file, which is useful for generating "pretty" ASCII representations of contingency tables.
as.table.ftable converts a contingency table in flat matrix form to one in standard array form. This is a method for the generic function as.table.

## References

Agresti, A. (1990) Categorical data analysis. New York: Wiley.

## See Also

 ftable for more information on flat contingencty tables.```
Examples
## Agresti (1990), page 157, Table 5.8.
## Not in ftable standard format, but o.k.
file <- tempfile()
cat(" Intercourse\n",
    "Race Gender Yes No\n",
    "White Male 43 134\n",
    " Female 26 149\n",
    "Black Male 29 23\n",
    " Female 22 36\n",
    file = file)
file.show(file)
ft <- read.ftable(file)
ft
unlink(file)
## Agresti (1990), page 297, Table 8.16.
## Almost o.k., but misses the name of the row variable.
file <- tempfile()
cat(" \"Tonsil Size\"\n",
    " \"Not Enl.\" \"Enl.\" \"Greatly Enl.\"\n",
    "Noncarriers 497 560 269\n",
    "Carriers 19 29 24\n",
    file = file)
file.show(file)
ft <- read.ftable(file, skip = 2,
    row.var.names = "Status",
    col.vars = list("Tonsil Size" =
```

```
c("Not Enl.", "Enl.", "Greatly Enl.")))
ft
unlink(file)
```

read.fwf Read Fixed Width Format Files

## Description

Read a "table" of fixed width formatted data into a data.frame.

## Usage

```
read.fwf(file, widths, sep = "\t", as.is = FALSE,
    skip = 0, row.names, col.names, n = -1, ...)
```


## Arguments

\(\left.\begin{array}{ll}file \& the name of the file which the data are to be read from. <br>
Alternatively, file can be a connection, which will be opened if neces- <br>

sary, and if so closed at the end of the function call.\end{array}\right]\)| integer vector, giving the widths of the fixed-width fields (of one line). |
| :--- |
| widths |
| sep |
| character; the separator used internally; should be a character that does |
| not occur in the file. |

## Details

Fields that are of zero-width or are wholly beyond the end of the line in file are replaced by NA.

## Value

A data.frame as produced by read.table which is called internally.

## Author(s)

Brian Ripley for R version: original Perl by Kurt Hornik.

## See Also

scan and read.table.

## Examples

```
ff <- tempfile()
cat(file=ff, "123456", "987654", sep="\n")
read.fwf(ff, width=c(1,2,3)) #> 1 23 456 \ 9 87 654
unlink(ff)
cat(file=ff, "123", "987654", sep="\n")
read.fwf(ff, width=c(1,0, 2,3)) #> 1 NA 23 NA \ 9 NA 87 654
unlink(ff)
```

read.socket Read from or Write to a Socket

## Description

read. socket reads a string from the specified socket, write. socket writes to the specified socket. There is very little error checking done by either.

## Usage

```
read.socket(socket, maxlen=256, loop=FALSE)
write.socket(socket, string)
```


## Arguments

| socket | a socket object |
| :--- | :--- |
| maxlen | maximum length of string to read |
| loop | wait for ever if there is nothing to read? |
| string | string to write to socket |

## Value

read.socket returns the string read.

## Author(s)

Thomas Lumley

## See Also

close.socket, make.socket

## Examples

```
finger <- function(user, host = "localhost", port = 79, print = TRUE)
{
    if (!is.character(user))
            stop("user name must be a string")
    user <- paste(user,"\r\n")
    socket <- make.socket(host, port)
    on.exit(close.socket(socket))
    write.socket(socket, user)
    output <- character(0)
    repeat{
```

```
    ss <- read.socket(socket)
    if (ss == "") break
    output <- paste(output, ss)
    }
    close.socket(socket)
    if (print) cat(output)
    invisible(output)
}
finger("root") ## only works if your site provides a finger daemon
```

```
read.table Data Input
```


## Description

Reads a file in table format and creates a data frame from it, with cases corresponding to lines and variables to fields in the file.

## Usage

```
read.table(file, header = FALSE, sep = "", quote = "\")", dec = ".",
            row.names, col.names, as.is = FALSE, na.strings = "NA",
            colClasses = NA, nrows = -1,
            skip = 0, check.names = TRUE, fill = !blank.lines.skip,
            strip.white = FALSE, blank.lines.skip = TRUE,
            comment.char = "\#")
read.csv(file, header = TRUE, sep = ",", quote="\"", dec=".",
    fill = TRUE, ...)
read.csv2(file, header = TRUE, sep = ";", quote="\"", dec=",",
    fill = TRUE, ...)
read.delim(file, header = TRUE, sep = "\t", quote="\"", dec=".",
    fill = TRUE, ...)
read.delim2(file, header = TRUE, sep = "\t", quote="\"", dec=",",
    fill = TRUE, ...)
```


## Arguments

file the name of the file which the data are to be read from. Each row of the table appears as one line of the file. If it does not contain an absolute path, the file name is relative to the current working directory, getwd(). Tilde-expansion is performed where supported.
Alternatively, file can be a connection, which will be opened if necessary, and if so closed at the end of the function call. (If stdin() is used, the prompts for lines may be somewhat confusing. Terminate input with an EOF signal, Ctrl-D on Unix and Ctrl-Z on Windows.)
file can also be a complete URL.

| header | a logical value indicating whether the file contains the names of the variables as its first line. If missing, the value is determined from the file format: header is set to TRUE if and only if the first row contains one fewer field than the number of columns. |
| :---: | :---: |
| sep | the field separator character. Values on each line of the file are separated by this character. If sep $=" "$ (the default for read.table) the separator is "white space", that is one or more spaces, tabs or newlines. |
| quote | the set of quoting characters. To disable quoting altogether, use quote="". See scan for the behaviour on quotes embedded in quotes. |
| dec | the character used in the file for decimal points. |
| row.names | a vector of row names. This can be a vector giving the actual row names, or a single number giving the column of the table which contains the row names, or character string giving the name of the table column containing the row names. |
|  | If there is a header and the first row contains one fewer field than the number of columns, the first column in the input is used for the row names. Otherwise if row.names is missing, the rows are numbered. <br> Using row.names $=$ NULL forces row numbering. |
| col.names | a vector of optional names for the variables. The default is to use "V" followed by the column number. |
| as.is | the default behavior of read.table is to convert character variables (which are not converted to logical, numeric or complex) to factors. The variable as.is controls this conversion. Its value is either a vector of logicals (values are recycled if necessary), or a vector of numeric indices which specify which columns should not be converted to factors. <br> Note: to suppress all conversions including those of numeric columns, set colClasses = "character". |
| na.strings | a vector of strings which are to be interpreted as NA values. Blank fields are also considered to be missing values. |
| colClasses | character. A vector of classes to be assumed for the columns. Recycled as necessary. If this is not one of the atomic vector classes (logical, integer, numeric, complex and character), there needs to be an as method for conversion from "character" to the specified class, or NA when type.convert is used. NB: as is in package methods. |
| nrows | the maximum number of rows to read in. Negative values are ignored. |
| skip | the number of lines of the data file to skip before beginning to read data. |
| check. n ames | logical. If TRUE then the names of the variables in the data frame are checked to ensure that they are syntactically valid variable names. If necessary they are adjusted (by make.names) so that they are, and also to ensure that there are no duplicates. |
| fill | logical. If TRUE then in case the rows have unequal length, blank fields are implicitly added. See Details. |
| strip.white | logical. Used only when sep has been specified, and allows the stripping of leading and trailing white space from character fields (numeric fields are always stripped). See scan for further details, remembering that the columns may include the row names. |
| blank.lines.skip |  |
|  | logical: if TRUE blank lines in the input are ignored |


| comment.char | character: a character vector of length one containing a single character <br> or an empty string. Use "" to turn off the interpretation of comments <br> altogether. |
| :--- | :--- |
| $\ldots$ | Further arguments to read.table. |

## Details

If row.names is not specified and the header line has one less entry than the number of columns, the first column is taken to be the row names. This allows data frames to be read in from the format in which they are printed. If row. names is specified and does not refer to the first column, that column is discarded from such files.
The number of data columns is determined by looking at the first five lines of input (or the whole file if it has less than five lines), or from the length of col.names if it is specified and is longer. This could conceivably be wrong if fill or blank.lines.skip are true.
read.csv and read.csv2 are identical to read.table except for the defaults. They are intended for reading "comma separated value" files ('.csv') or the variant used in countries that use a comma as decimal point and a semicolon as field separator. Similarly, read.delim and read.delim2 are for reading delimited files, defaulting to the TAB character for the delimiter. Notice that header $=$ TRUE and fill $=$ TRUE in these variants.
Comment characters are allowed unless comment.char $=$ " ", and complete comment lines are allowed provided blank.lines.skip = TRUE However, comment lines prior to the header must have the comment character in the first non-blank column.

## Value

A data frame (data.frame) containing a representation of the data in the file. Empty input is an error unless col.names is specified, when a 0 -row data frame is returned: similarly giving just a header line if header $=$ TRUE results in a 0 -row data frame.
This function is the principal means of reading tabular data into R.

## Note

The columns referred to in as.is and colClasses include the column of row names (if any).
Less memory will be used if colClasses is specified as one of the five atomic vector classes.
Using nrows, even as a mild over-estimate, will help memory usage.
Using comment.char = "" will be appreciably faster.
read.table is not the right tool for reading large matrices, especially those with many columns: it is designed to read data frames which may have columns of very different classes. Use scan instead.

## See Also

The $R$ Data Import/Export manual.
scan, type.convert, read.fwf for reading fixed width formatted input; read.table.url for "reading" data from the internet; write.table; data.frame.
count.fields can be useful to determine problems with reading files which result in reports of incorrect record lengths.
readBin Transfer Binary Data To and From Connections

## Description

Read binary data from a connection, or write binary data to a connection.

## Usage

```
readBin(con, what, \(\mathrm{n}=1\), size \(=\mathrm{NA}\), signed \(=\) TRUE,
    endian = .Platform\$endian)
writeBin(object, con, size = NA, endian = .Platform\$endian)
readChar(con, nchars)
writeChar (object, con, nchars = nchar (object), eos = "")
```


## Arguments

| con | A connection object or a character string. |
| :--- | :--- |
| what | Either an object whose mode will give the mode of the vector <br> to be read, or a character vector of length one describing the <br> mode: one of "numeric", "double", "integer", "int", "logical", <br> "complex", "character". <br> integer. The (maximal) number of records to be read. You can use an <br> over-estimate here, but not too large as storage is reserved for n items. <br> integer. The number of bytes per element in the byte stream. The default, |
| n size | NA, uses the natural size. Size changing is not supported for complex <br> vectors. <br> logical. Only used for integers of sizes 1 and 2, when it determines if the <br> quantity on file should be regarded as a signed or unsigned integer. |
| signed | The endian-ness ("big" or "little" of the target system for the file. |
| endian | Using "swap" will force swapping endian-ness. <br> An R object to be written to the connection. |
| nchars | integer, giving the lengths of (unterminated) character strings to be read <br> or written. |
| eos | character. The terminator to be written after each string, followed by an |
|  | ASCII nul; use NULL for no terminator at all. |

## Details

If the con is a character string, the functions call file to obtain an file connection which is opened for the duration of the function call.
If the connection is open it is read/written from its current position. If it is not open, it is opened for the duration of the call and then closed again.
If size is specified and not the natural size of the object, each element of the vector is coerced to an appropriate type before being written or as it is read. Possible sizes are 1, 2,4 and possibly 8 for integer or logical vectors, and 4,8 and possibly $12 / 16$ for numeric vectors. (Note that coercion occurs as signed types except if signed $=$ FALSE when reading
integers of sizes 1 and 2.) Changing sizes is unlikely to preserve NAs, and the extended precision sizes are unlikely to be portable across platforms.
readBin and writeBin read and write C-style zero-terminated character strings. readChar and writeChar allow more flexibility, and can also be used on text-mode connections.

Handling R's missing and special (Inf, -Inf and NaN values is discussed in the $R$ Data Import/Export manual.

## Value

For readBin, a vector of appropriate mode and length the number of items read (which might be less than $n$ ).

For readChar, a character vector of length the number of items read (which might be less than length(nchars)).
For writeBin and writeChar, none.

## Note

Integer read/writes of size 8 will be available if either C type long is of size 8 bytes or C type long long exists and is of size 8 bytes.
Real read/writes of size sizeof(long double) (usually 12 or 16 bytes) will be available only if that type is available and different from double.
Note that as R character strings cannot contain ASCII nul, strings read by readChar which contain such characters will appear to be shorted than requested, but the additional bytes are read from the file.
If the character length requested for readChar is longer than the string, as from version 1.4.0 what is available is returned.

## See Also

The $R$ Data Import/Export manual.
connections, readLines, writeLines.
.Machine for the sizes of long, long long and long double.

## Examples

```
zz <- file("testbin", "wb")
writeBin(1:10, zz)
writeBin(pi, zz, endian="swap")
writeBin(pi, zz, size=4)
writeBin(pi^2, zz, size=4, endian="swap")
writeBin(pi+3i, zz)
writeBin("A test of a connection", zz)
z <- paste("A very long string", 1:100, collapse=" + ")
writeBin(z, zz)
if(.Machine$sizeof.long == 8 || .Machine$sizeof.longlong == 8)
    writeBin(as.integer(5^(1:10)), zz, size = 8)
if((s <-.Machine$sizeof.longdouble) > 8) writeBin((pi/3)^(1:10), zz, size = s)
close(zz)
zz <- file("testbin", "rb")
readBin(zz, integer(), 4)
readBin(zz, integer(), 6)
```

```
readBin(zz, numeric(), 1, endian="swap")
readBin(zz, numeric(), size=4)
readBin(zz, numeric(), size=4, endian="swap")
readBin(zz, complex(), 1)
readBin(zz, character(), 1)
z2 <- readBin(zz, character(), 1)
if(.Machine$sizeof.long == 8 || .Machine$sizeof.longlong == 8)
    readBin(zz, integer(), 10, size = 8)
if((s <-.Machine$sizeof.longdouble) > 8) readBin(zz, numeric(), 10, size = s)
close(zz)
unlink("testbin")
stopifnot(z2 == z)
## test fixed-length strings
zz <- file("testbin", "wb")
x <- c("a", "this will be truncated", "abc")
nc <- c(3, 10, 3)
writeChar(x, zz, nc, eos=NULL)
writeChar(x, zz, eos="\r\n")
close(zz)
zz <- file("testbin", "rb")
readChar(zz, nc)
readChar(zz, nchar(x)+3) # need to read the terminator explicitly
close(zz)
unlink("testbin")
## signed vs unsigned ints
zz <- file("testbin", "wb")
x <- as.integer(seq(0, 255, 32))
writeBin(x, zz, size=1)
writeBin(x, zz, size=1)
x <- as.integer(seq(0, 60000, 10000))
writeBin(x, zz, size=2)
writeBin(x, zz, size=2)
close(zz)
zz <- file("testbin", "rb")
readBin(zz, integer(), 8, size=1)
readBin(zz, integer(), 8, size=1, signed=FALSE)
readBin(zz, integer(), 7, size=2)
readBin(zz, integer(), 7, size=2, signed=FALSE)
close(zz)
unlink("testbin")
```

readline Read a Line from the Terminal

## Description

readline reads a line from the terminal

## Usage

```
readline(prompt="")
```


## Arguments

prompt the string printed when prompting the user for input. Should usually end with a space " ".

## Details

The prompt string will be truncated to a maximum allowed length, normally 256 chars (but can be changed in the source code).

## Value

A character vector of length one.

## Examples

```
fun <- function() {
        ANSWER <- readline("Are you a satisfied R user? ")
        if (substr(ANSWER, 1, 1) == "n")
            cat("This is impossible. YOU LIED!\n")
        else
            cat("I knew it.\n")
}
fun()
```

readLines Read Text Lines from a Connection

## Description

Read text lines from a connection.

## Usage

readLines (con $=\operatorname{stdin}(), \mathrm{n}=-1$, ok $=$ TRUE)

## Arguments

con A connection object or a character string.
$\mathrm{n} \quad$ integer. The (maximal) number of lines to read. Negative values indicate that one should read up to the end of the connection.
ok logical. Is it OK to reach the end of the connection before $\mathrm{n}>0$ lines are read? If not, an error will be generated.

## Details

If the con is a character string, the functions call file to obtain an file connection which is opened for the duration of the function call.

If the connection is open it is read from its current position. If it is not open, it is opened for the duration of the call and then closed again.

If the final line is incomplete (no final EOL marker) the behaviour depends on whether the connection is blocking or not. For a blocking text-mode connection (or a non-textmode connection) the line will be accepted, with a warning. For a non-blocking text-mode connection the incomplete line is pushed back, silently.

## Value

A character vector of length the number of lines read.

## See Also

connections, writeLines, scan

## Examples

```
cat("TITLE extra line", "2 3 5 7", "", "11 13 17", file="ex.data",
    sep="\n")
readLines("ex.data", n=-1)
unlink("ex.data") # tidy up
## difference in blocking
cat("123\nabc", file = "test1")
readLines("test1") # line with a warning
con <- file("test1", "r", blocking = FALSE)
readLines(con) # empty
cat(" def\n", file = "test1", append = TRUE)
readLines(con) # gets both
close(con)
unlink("test1") # tidy up
```

```
real Real Vectors
```


## Description

real creates a double precision vector of the specified length. Each element of the vector is equal to 0 .
as.real attempts to coerce its argument to be of real type.
is.real returns TRUE or FALSE depending on whether its argument is of real type or not.

## Usage

```
real(length = 0)
as.real(x, ...)
is.real(x)
```


## Arguments

| length | desired length. |
| :--- | :--- |
| x | object to be coerced or tested. |
| $\ldots$ | further arguments passed to or from other methods. |

## Note

R has no single precision data type. All real numbers are stored in double precision format.

## Recall Recursive Calling

## Description

Recall is used as a placeholder for the name of the function in which it is called. It allows the definition of recursive functions which still work after being renamed, see example below.

## Usage

```
Recall(...)
```


## Arguments

```
... all the arguments to be passed.
```


## See Also

do.call and call.

## Examples

```
## A trivial (but inefficient!) example:
fib <- function(n) if(n<=2) {if(n>=0) 1 else 0} else Recall(n-1) + Recall(n-2)
fibonacci <- fib; rm(fib)
## renaming wouldn't work without Recall
fibonacci(10) # 55
```

recordPlot Record and Replay Plots

## Description

Functions to save the current plot in an $R$ variable, and to replay it.

## Usage

recordPlot()
replayPlot(x)

## Arguments

X
A saved plot.

## Details

These functions record and replay the displaylist of the current graphics device. The returned object is of class "recordedplot", and replayPlot acts as a print method for that class.
The format of recorded plots was changed in R 1.4.0: plots saved in earlier versions can still be replayed.

## Value

recordPlot returns an object of class "recordedplot", a list with components:
displaylist The saved display list, as a pairlist.
gpar The graphics state, as an integer vector.
replayPlot has no return value.
recover Browsing after an Error

## Description

This function allows the user to browse directly on any of the currently active function calls, and is suitable as an error option. The expression options (error=recover) will make this the error option.

## Usage <br> recover()

## Details

When called, recover prints the list of current calls, and prompts the user to select one of them. The standard R browser is then invoked from the corresponding environment; the user can type ordinary $S$ language expressions to be evaluated in that environment.
When finished browsing in this call, type c to return to recover from the browser. Type another frame number to browse some more, or type 0 to exit recover.
The use of recover largely supersedes dump.frames as an error option, unless you really want to wait to look at the error. If recover is called in non-interactive mode, it behaves like dump.frames. For computations involving large amounts of data, recover has the advantage that it does not need to copy out all the environments in order to browse in them. If you do decide to quit interactive debugging, call dump.frames directly while browsing in any frame (see the examples).
WARNING: The special Q command to go directly from the browser to the prompt level of the evaluator currently interacts with recover to effectively turn off the error option for the next error (on subsequent errors, recover will be called normally).

## Value

Nothing useful is returned. However, you can invoke recover directly from a function, rather than through the error option shown in the examples. In this case, execution continues after you type 0 to exit recover.

## Compatibility Note

The $R$ recover function can be used in the same way as the S-Plus function of the same name; therefore, the error option shown is a compatible way to specify the error action. However, the actual functions are essentially unrelated and interact quite differently with the user. The navigating commands up and down do not exist in the R version; instead, exit the browser and select another frame.

## References

John M. Chambers (1998). Programming with Data; Springer. See the compatibility note above, however.

## See Also

browser for details about the interactive computations; options for setting the error option; dump.frames to save the current environments for later debugging.

## Examples

```
options(error = recover) # setting the error option
### Example of interaction
> myFit <- lm(y ~ x, data = xy, weights = w)
Error in lm.wfit(x, y, w, offset = offset, ...) :
    missing or negative weights not allowed
Enter a frame number, or 0 to exit
1:lm(y ~ x, data = xy, weights = w)
2:lm.wfit(x, y, w, offset = offset, ...)
Selection: 2
Called from: eval(expr, envir, enclos)
Browse[1]> objects() # all the objects in this frame
[1] "method" "n" "ny" "offset" "tol" "w"
[7] "x" "y"
Browse[1]> w
[1] -0.5013844 1.3112515 0.2939348 -0.8983705 -0.1538642
[6] -0.9772989 0.7888790-0.1919154 -0.3026882
Browse[1]> dump.frames() # save for offline debugging
Browse[1]> c # exit the browser
Enter a frame number, or 0 to exit
1:lm(y ~ x, data = xy, weights = w)
2:lm.wfit(x, y, w, offset = offset, ...)
Selection: O # exit recover
>
```

```
rect Draw a Rectangle
```


## Description

rect draws a rectangle (or sequence of rectangles) with the given coordinates, fill and border colors.

## Usage

```
rect(xleft, ybottom, xright, ytop, density \(=\) NULL, angle \(=45\),
    col = NULL, border = NULL, lty = NULL, lwd = par("lwd"),
    xpd = NULL, ...)
```


## Arguments

| xleft | a vector (or scalar) of left x positions. |
| :---: | :---: |
| ybottom | a vector (or scalar) of bottom y positions. |
| xright | a vector (or scalar) of right x positions. |
| ytop | a vector (or scalar) of top y positions. |
| density | the density of shading lines, in lines per inch. The default value of NULL means that no shading lines are drawn. Non-positive values of density also inhibit the drawing of shading lines. |
| angle | angle (in degrees) of the shading lines. |
| col | color(s) to fill or shade the rectangle(s) with. The default NULL, or also NA do not fill, i.e., draw transparent rectangles. |
| border | color for rectangle border(s). |
| lty | line type for borders; defaults to "solid". |
| lwd | width for borders. |
| xpd | logical ("expand"); defaults to par("xpd"). See par(xpd= ). other graphical parameters can be given as arguments. |

## Details

The positions supplied, i.e., xleft, ..., are relative to the current plotting region. If the x-axis goes from 100 to 200 then xleft must be larger than 100 and xright must be less than 200.
It is a primitive function used in hist, barplot, legend, etc.

## See Also

box for the "standard" box around the plot; polygon and segments for flexible line drawing. par for how to specify colors.

## Examples

```
## set up the plot region:
op <- par(bg = "thistle")
plot(c(100, 250), c(300, 450), type = "n", xlab="",
    main = "2 x 11 rectangles; 'rect(100+i,300+i, 150+i,380+i)'")
i <- 4*(0:10)
## draw rectangles with bottom left (100, 300)+i and top right (150, 380)+i
rect(100+i, 300+i, 150+i, 380+i, col=rainbow(11, start=.7,end=.1))
rect(240-i, 320+i, 250-i, 410+i, col=heat.colors(11), lwd=i/5)
## Background alternating ( transparent / "bg" ) :
j <- 10*(0:5)
rect(125+j, 360+j, 141+j, 405+j/2, col = c(NA,0), border = "gold", lwd = 2)
rect(125+j, 296+j/2, 141+j, 331+j/5, col = c(NA,"midnightblue"))
mtext("+ 2 x 6 rect(*, col = c(NA,0)) and col = c(NA,\"m..blue\"))")
par(op)
```


## reg.finalizer Finalization of objects

## Description

Registers an R function to be called upon garbage collection of object.

## Usage

reg.finalizer (e, f)

## Arguments

e Object to finalize. Must be environment or external pointer.
$\mathrm{f} \quad$ Function to call on finalization. Must accept a single argument, which will be the object to finalize.

## Value

NULL.

## Note

The purpose of this function is mainly to allow objects that refer to external items (a temporary file, say) to perform cleanup actions when they are no longer referenced from within R. This only makes sense for objects that are never copied on assignment, hence the restriction to environments and external pointers.

## Examples

```
f <- function(e) print("cleaning....")
g <- function(x){e<-environment(); reg.finalizer(e,f)}
g()
gc() # trigger cleanup
```

relevel Reorder Levels of Factor

## Description

The levels of a factor are re-ordered so that the level specified by ref is first and the others are moved down. This is useful for contr.treatment contrasts which take the first level as the reference.

## Usage

relevel(x, ref, ...)

## Arguments

$x \quad$ An unordered factor.
ref The reference level.
... Additional arguments for future methods.

## Value

A factor of the same length as x .

## Author(s)

B. D. Ripley

## See Also

factor, contr.treatment

## Examples

```
data(warpbreaks)
warpbreaks$tension <- relevel(warpbreaks$tension, ref="M")
summary(lm(breaks ~ wool + tension, data=warpbreaks))
```

REMOVE Remove Add-on Packages

## Description

Utility for removing add-on packages.

## Usage

R CMD REMOVE [options] [-1 lib] pkgs

## Arguments

pkgs a list with the names of the packages to be removed.
lib the path name of the R library tree to remove from. May be absolute or relative.
options further options.

## Details

If used as R CMD REMOVE pkgs without explicitly specifying lib, packages are removed from the library tree rooted at the first directory given in \$R_LIBS if this is set and non-null, and to the default library tree (which is rooted at '\$R_HOME/library') otherwise.
To remove from the library tree lib, use R CMD REMOVE -l lib pkgs.
Use R CMD REMOVE --help for more usage information.

## See Also

INSTALL
remove Remove Objects from a Specified Environment

## Description

remove and rm can be used to remove objects. These can be specified successively as character strings, or in the character vector list, or through a combination of both. All objects thus specified will be removed.
If envir is NULL then the the currently active environment is searched first.
If inherits is TRUE then parents of the supplied directory are searched until a variable with the given name is encountered. A warning is printed for each variable that is not found.

## Usage

```
remove(..., list = character(0), pos = -1, envir = as.environment(pos),
        inherits = FALSE)
rm (..., list = character(0), pos = -1, envir = as.environment(pos),
    inherits = FALSE)
```


## Arguments

|  | the objects to be removed, supplied individually and/or as a character vector |
| :---: | :---: |
| list | a character vector naming objects to be removed. |
| pos | where to do the removal. By default, uses the current environment. See the details for other possibilities. |
| envir | the environment to use. See the details section. |
| inherits | should the enclosing frames of the environment be inspected? |

## Details

The pos argument can specify the environment from which to remove the objects in any of several ways: as an integer (the position in the search list); as the character string name of an element in the search list; or as an environment (including using sys.frame to access the currently active function calls). The envir argument is an alternative way to specify an environment, but is primarily there for back compatibility.

## See Also

ls, objects

## Examples

```
tmp <- 1:4
## work with tmp and cleanup
rm(tmp)
## remove (almost) everything in the working environment.
## You will get no warning, so don't do this unless you are really sure.
rm(list = ls())
```

remove.packages Remove Installed Packages

## Description

Removes installed packages and updates index information as necessary.

## Usage

remove.packages(pkgs, lib)

## Arguments

pkgs a character vector with the names of the packages to be removed.
lib a character string giving the library directory to move the packages from.

## See Also

REMOVE for a command line version; install.packages for installing packages.

```
rep Replicate Elements of Vectors and Lists
```


## Description

rep replicates the values in x according to the values given in times and length. out.

## Usage

rep(x, times, length.out, each)
rep.int(x, times)

## Arguments

| x | a vector (of any mode including a list) or a pairlist. |
| :--- | :--- |
| times | non-negative integer. A vector giving the number of times to repeat each <br> element if of length length $(\mathrm{x})$, or to repeat the whole vector if of length |
|  | 1. |

## Details

If times consists of a single integer, the result consists of the values in x repeated this many times. If times is a vector of the same length as $x$, the result consists of $x[1]$ repeated times [1] times, x[2] repeated times [2] times and so on.
length. out may be given in place of times, in which case x is repeated as many times as is necessary to create a vector of this length. If both length. out and times are specified, times determines the replication, and length. out can be used to truncate the output vector (or extend it by NAs).
Non-integer values of times will be truncated towards zero. If times is a computed quantity it is prudent to add a small fuzz.

## Note

If the original vector has names, these are also replicated and so will almost always contain duplicates.

If length. out is used to extend the vector, the behaviour is different from that of S-PLUS, which recycles the existing vector.
Function rep.int is a simple case handled by internal code, and provided as a separate function purely for $S$ compatibility.

## See Also

```
seq, sequence.
```


## Examples

```
rep(1:4, 2)
rep(1:4, each = 2) # not the same.
rep(1:4, c(2,2,2,2)) # same as second.
rep(1:4, c(2,1,2,1))
rep(1:4, each = 2, len = 4) # first 4 only.
rep(1:4, each = 2, len = 10) # 8 integers plus two NAs
rep(1, 40*(1-.8)) # length 7 on most platforms
rep(1, 40*(1-.8)+1e-7) # better
## replicate a list
fred <- list(happy = 1:10, name = "squash")
rep(fred, 5)
## more esoteric:
stopifnot(identical(rep(letters, 0), character(0)),
    identical(rep.int(1:2, 0), integer(0)))
```

replace Replace Values in a Vector

## Description

replace replaces the values in x with indexes given in list by those given in values. If necessary, the values in values are recycled.

## Usage

```
replace(x, list, values)
```


## Arguments

| x | vector |
| :--- | :--- |
| list | an index vector |
| values | replacement values |

## Value

A vector with the values replaced.

## Note

x is unchanged: remember to assign the result.

```
replications Number of Replications of Terms
```


## Description

Returns a vector or a list of the number of replicates for each term in the formula.

## Usage

replications(formula, data=NULL, na.action)

## Arguments

| formula | a formula or a terms object or a data frame. |
| :--- | :--- |
| data | a data frame used to find the objects in formula. |
| na.action | function for handling missing values. Defaults to a na.action attribute <br> of data, then a setting of the option na.action, or na.fail if that is not <br> set. |

## Details

If formula is a data frame and data is missing, formula is used for data with the formula ~ ..

## Value

A vector or list with one entry for each term in the formula giving the number(s) of replications for each level. If all levels are balanced (have the same number of replications) the result is a vector, otherwise it is a list with a component for each terms, as a vector, matrix or array as required.

A test for balance is !is.list(replications(formula, data)).

## Author(s)

B. D. Ripley

## See Also

model.tables

## Examples

```
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block=gl(6,4), N=factor(N), P=factor(P),
    K=factor(K), yield=yield)
replications(~ . - yield, npk)
```

reshape Reshape Grouped Data

## Description

This function reshapes a dataframe between 'wide' format with repeated measurements in separate columns of the same record and 'long' format with the repeated measurements in separate records.

## Usage

```
reshape(data, varying = NULL, v.names = NULL, timevar = "time",
    idvar = "id", ids = 1:NROW(data),
    times = seq(length = length(varying[[1]])),
    drop = NULL, direction, new.row.names = TRUE,
    split = list(regexp="\\.", include=FALSE))
```


## Arguments

| data | a data frame |
| :--- | :--- |
| varying | names of sets of variables in the wide format that correspond to single <br> variables in long format ('time-varying'). A list of vectors (or optionally <br> a matrix for direction="wide"). See below for more details and options. |
| v.names | names of variables in the long format that correspond to multiple variables <br> in the wide format. |
| timevar | the variable in long format that differentiates multiple records from the <br> same group or individual. |
| idvar | the variable in long format that identifies multiple records from the same <br> group/individual. This variable may also be present in wide format. |
| ids | the values to use for a newly created idvar variable in long format. |

times the values to use for a newly created timevar variable in long format.
drop a vector of names of variables to drop before reshaping
direction "wide" to reshape to wide format, "long" to reshape to long format.
new.row.names logical; if TRUE and direction="wide", create new row names in long format from the values of the id and time variables.
split information for guessing the varying, v.names, and times arguments. See below for details.

## Details

The arguments to this function are described in terms of longitudinal data, as that is the application motivating the functions. A 'wide' longitudinal dataset will have one record for each individual with some time-constant variables that occupy single columns and some time-varying variables that occupy a column for each time point. In 'long' format there will be multiple records for each individual, with some variables being constant across these records and others varying across the records. A 'long' format dataset also needs a 'time' variable identifying which time point each record comes from and an 'id' variable showing which records refer to the same person.

If the data frame resulted from a previous reshape then the operation can be reversed by specifying just the direction argument. The other arguments are stored as attributes on the data frame.

If direction="long" and no varying or v.names arguments are supplied it is assumed that all variables except idvar and timevar are time-varying. They are all expanded into multiple variables in wide format.

If direction="wide" the varying argument can be a vector of column names or column numbers (converted to column names). The function will attempt to guess the v.names and times from these names. The default is variable names like $\mathrm{x} .1, \mathrm{x} .2$, where split=list (regexp="\.", include=FALSE) to specifies to split at the dot and drop it from the name. To have alphabetic followed by numeric times use split=list (regexp=" [A-Za-$z][0-9]$ ", include=TRUE). This splits between the alphabetic and numeric parts of the name and does not drop the regular expression.

## Value

The reshaped data frame with added attributes to simplify reshaping back to the original form.

## See Also

```
stack, aperm
```


## Examples

```
data(Indometh,package="nls")
summary(Indometh)
wide <- reshape(Indometh, v.names="conc", idvar="Subject",
    timevar="time", direction="wide")
wide
reshape(wide, direction="long")
reshape(wide, idvar="Subject", varying=list(names(wide)[2:12]),
        v.names="conc", direction="long")
```

```
## times need not be numeric
df <- data.frame(id=rep(1:4,rep(2,4)), visit=I(rep(c("Before","After"),4)),
    x=rnorm(4), y=runif(4))
df
reshape(df, timevar="visit", idvar="id", direction="wide")
## warns that y is really varying
reshape(df, timevar="visit", idvar="id", direction="wide", v.names="x")
## unbalanced 'long' data leads to NA fill in 'wide' form
df2 <- df[1:7,]
df2
reshape(df2, timevar="visit", idvar="id", direction="wide")
## Alternative regular expressions for guessing names
df3 <- data.frame(id=1:4, age=c (40,50,60,50), dose1=c (1,2,1,2),
    dose2=c(2,1,2,1), dose4=c(3,3,3,3))
reshape(df3, direction="long", varying=3:5,
    split=list(regexp="[a-z] [0-9] ", include=TRUE))
## an example that isn't longitudinal data
data(state)
state.x77 <- as.data.frame(state.x77)
long <- reshape(state.x77, idvar="state", ids=row.names(state.x77),
    times=names(state.x77), timevar="Characteristic",
    varying=list(names(state.x77)), direction="long")
reshape(long, direction="wide")
reshape(long, direction="wide", new.row.names=unique(long$state))
```

residuals Extract Model Residuals

## Description

residuals is a generic function which extracts model residuals from objects returned by modeling functions.

The abbreviated form resid is an alias for residuals. It is intended to encourage users to access object components through an accessor function rather than by directly referencing an object slot.

All object classes which are returned by model fitting functions should provide a residuals method. (Note that the method is 'residuals' and not 'resid'.)

Methods can make use of naresid methods to compensate for the omission of missing values. The default method does.

## Usage

```
residuals(object, ...)
resid(object, ...)
```


## Arguments

object an object for which the extraction of model residuals is meaningful.
... other arguments.

## Value

Residuals extracted from the object object.

## See Also

coefficients, fitted.values, glm, lm.

```
restart-deprecated Restart an Expression
```


## Description

restart performs a type of non-local return.

## Usage

restart(on = TRUE)

## Arguments

on $\quad$ if true a jump point is set; if false the jump point is removed.

## Details

When restart is called with on = TRUE the evaluator marks that function as a return point. Any errors or signals (such as control-C on Unix) cause control to return to the start of the function containing the call to restart. The most recently established function is always entered first.

## Note

The direct use of restart is likely to result in an infinite loop. Use try unless you are sure you know what you are doing.

In fact, as restart is deprecated, don't use it even if you are sure you know what you are doing.

## See Also

try for a safer interface.
options for setting error handlers and suppressing the printing of error messages.

```
rev Reverse a Vector's Elements
```


## Description

rev provides a reversed version of its argument. It can be used in combination with sort to obtain vectors sorted into descending order.

## Usage

rev( x )

## Arguments

X
a vector.

## See Also

seq, sort.

## Examples

```
x <- c(1:5,5:3)
# sort into descending order
rev(sort(x))
stopifnot(rev(1:7) == 7:1)#- don't need 'rev' here
```

rgb $\quad R G B$ Color Specification

## Description

This function creates "colors" corresponding to the given intensities (between 0 and max) of the red, green and blue primaries. The names argument may be used to provide names for the colors.

The values returned by rgb can be used with a col= specification in graphics functions or in par.

## Usage

rgb(red, green, blue, names=NULL, maxColorValue = 1)

## Arguments

red, blue, green
vectors of same length with values in $[0, M]$ where $M$ is maxColorValue. When this is 255 , the red, blue and green values are coerced to integers in $0: 255$ and the result is computed most efficiently.
names character. The names for the resulting vector.
maxColorValue number giving the maximum of the color values range, see above.

See Also
col2rgb the "inverse" for translating $R$ colors to RGB vectors. rainbow, hsv, gray.

## Examples

```
rgb(0,1,0)
(u01 <- seq(0,1, length=11))
stopifnot(rgb(u01,u01,u01) == gray(u01))
reds <- rgb((0:15)/15, g=0,b=0, names=paste("red",0:15,sep="."))
reds
rgb(0, 0:12, 0, max = 255)# integer input
```

RHOME $\quad R$ Home Directory

## Description

Returns the location of the R home directory, which is the root of the installed R tree.

## Usage

R RHOME
rivers Lengths of Major North American Rivers

## Description

This data set gives the lengths (in miles) of 141 "major" rivers in North America, as compiled by the US Geological Survey.

## Usage

data(rivers)

## Format

A vector containing 141 observations.

## Source

World Almanac and Book of Facts, 1975, page 406.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.
rle Run Length Encoding

## Description

Compute the lengths and values of runs of equal values in a vector - or the reverse operation.

## Usage

```
rle(x)
print(x, digits = getOption("digits"), ...)
inverse.rle(x, ...)
```


## Arguments

$\mathrm{x} \quad$ a simple vector for rle() or an object of class "rle" for print() or inverse.rle().
digits, ... potentially further arguments to the corresponding method.

## Value

rle() returns an object of class "rle" which is a list with components

| lengths | an integer vector containing the length of each run. |
| :--- | :--- |
| values | a vector of the same length as lengths with the corresponding values. |

inverse.rle() is the inverse function of rle() .

## Examples

```
x <- rev(rep(6:10, 1:5))
rle(x)
## lengths [1:5] 5 4 3 2 1
## values [1:5] 10 9 8 7 6
z <- c(TRUE,TRUE,FALSE,FALSE,TRUE,FALSE,TRUE,TRUE,TRUE)
rle(z)
rle(as.character(z))
stopifnot(x == inverse.rle(rle(x)),
    z == inverse.rle(rle(z)))
```


## Round

## Description

ceiling takes a single numeric argument x and returns a numeric vector containing the smallest integers not less than the corresponding elements of x .
floor takes a single numeric argument $x$ and returns a numeric vector containing the largest integers not greater than the corresponding elements of x .
round rounds the values in its first argument to the specified number of decimal places (default 0). Note that for rounding off a 5, the IEEE standard is used, "go to the even digit". Therefore round ( 0.5 ) is 0 and round ( -1.5 ) is -2 .
signif rounds the values in its first argument to the specified number of significant digits.
trunc takes a single numeric argument x and returns a numeric vector containing the integers by truncating the values in $x$ toward 0 .
zapsmall determines a digits argument dr for calling round( $x$, digits $=d r$ ) such that values "close to zero" values are "zapped", i.e., treated as 0.

## Usage

```
ceiling(x)
floor(x)
round(x, digits = 0)
signif(x, digits = 6)
trunc(x)
zapsmall(x, digits= getOption("digits"))
```


## Arguments

| x | a numeric vector. |
| :--- | :--- |
| digits | integer indicating the precision to be used. |

## See Also

```
as.integer.
```


## Examples

```
round(.5 + -2:4) # IEEE rounding: -2 0 0 2 2 4 4 4
( x1 <- seq(-2, 4, by = .5) )
round(x1)#-- IEEE rounding !
x1[trunc(x1) != floor(x1)]
x1[round(x1) != floor(x1 + .5)]
(non.int <- ceiling(x1) != floor(x1))
stopifnot(
    trunc(x1) == as.integer(x1),
    non.int == (ceiling(x1) != trunc(x1) | trunc(x1) != floor(x1)),
    (signif(x1, 1) != round(x1,1)) == (non.int & abs(x1) > 1)
)
x2 <- pi * 100^(-1:3)
```

```
round(x2, 3)
signif(x2, 3)
print (x2 / 1000, digits=4)
zapsmall(x2 / 1000, digits=4)
zapsmall(exp(1i*0:4*pi/2))
```

    round.POSIXt Round / Truncate Data-Time Objects
    
## Description

Round or truncate date-time objects.

## Usage

```
round(x, units=c("secs", "mins", "hours", "days"))
trunc.POSIXt(x, units=c("secs", "mins", "hours", "days"))
```


## Arguments

| x | an object inheriting from "POSIXt". |
| :--- | :--- |
| units | one of the units listed. Can be abbreviated. |

## Details

The time is rounded or truncated to the second, minute, hour or day. Timezones are only relevant to days, when midnight in the current timezone is used.

## Value

An object of class "POSIXlt".

## Note

trunc is not generic, so trunc.POSIXt has to be called explicitly.

## See Also

DateTimeClasses

## Examples

```
round(.leap.seconds + 1000, "hour")
trunc.POSIXt(Sys.time(), "day")
```

row Row Indexes

## Description

Returns a matrix of integers indicating their row number in the matrix.

## Usage

```
row(x, as.factor = FALSE)
```


## Arguments

x
a matrix.
as.factor a logical value indicating whether the value should be returned as a factor rather than as numeric.

## Value

An integer matrix with the same dimensions as x and whose ij -th element is equal to i .

## See Also

col to get columns.

## Examples

```
x <- matrix(1:12, 3, 4)
# extract the diagonal of a matrix
dx <- x[row (x) == col(x)]
dx
# create an identity 5-by-5 matrix
x <- matrix(0, nr = 5, nc = 5)
x[row(x) == col(x)] <- 1
x
```

row/colnames Row and Columnn Names

## Description

Retrieve or set the row or column names of an object (the first or second component of its dimnames).

## Usage

```
rownames(x, do.NULL = TRUE, prefix = "row")
rownames(x) <- namevector
colnames(x, do.NULL = TRUE, prefix = "col")
colnames(x) <- namevector
```


## Arguments

x
do.NULL logical. Should this create names if they are NULL?
prefix for created names.

## Details

If do.NULL is FALSE, a character vector (of length $\operatorname{NROW}(x)$ or NCOL ( $x$ ) is returned in any case, prepending prefix to simple numbers, if dimnames ( x ) [ [ i$]$ ] ( $\mathrm{i}=1$ or 2 ) is NULL.

## See Also

dimnames, case.names, variable.names.

## Examples

```
m0 <- matrix(NA, 4, 0)
m2 <- cbind(1,1:4)
rownames(m0)
colnames(m2, do.NULL = FALSE)
colnames(m2) <- c("x","Y")
rownames(m2) <- rownames(m2, do.NULL = FALSE, prefix = "Obs.")
m2
```


## rowsum <br> Give row sums of a matrix or data frame, based on a grouping

 variable
## Description

Compute sums across rows of a matrix-like object for each level of a grouping variable. rowsum is generic, with methods for matrices and data frames.

## Usage

rowsum (x, group, reorder $=$ TRUE, ...)

## Arguments

$x \quad$ a matrix, data frame or vector of numeric data. Missing values are allowed.
group a vector giving the grouping, with one element per row of x . Missing values will be treated as another group and a warning will be given
reorder if TRUE, then the result will be in order of sort (unique(group)), if FALSE, it will be in the order that rows were encountered.
... other arguments for future methods

## Details

The default is to reorder the rows to agree with tapply as in the example below. Reordering should not add noticeably to the time except when there are very many distinct values of group and x has few columns.
The original function was written by Terry Therneau, but this is a new implementation using hashing that is much faster for large matrices.
To add all the rows of a matrix (ie, a single group) use rowSums, which should be even faster.

## Value

a matrix or data frame containing the sums. There will be one row per unique value of group.

## See Also

tapply, aggregate,rowSums

## Examples

```
x <- matrix(runif(100), ncol=5)
group <- sample(1:8, 20, TRUE)
xsum <- rowsum(x, group)
## Slower versions
xsum2 <- tapply(x, list(group[row(x)], col(x)), sum)
xsum3<- aggregate(x,list(group),sum)
```


## Rprof Enable Profiling of R's Execution

## Description

Enable or disable profiling of the execution of R expressions.

## Usage

```
Rprof(filename = "Rprof.out", append = FALSE, interval = 0.02)
```


## Arguments

| filename | The file to be used for recording the profiling results. Set to NULL or "" <br> to disable profiling. |
| :--- | :--- |
| append | logical: should the file be over-written or appended to? <br> interval |
| real: time interval between samples. |  |

## Details

Enabling profiling automatically disables any existing profiling to another or the same file. Profiling works by writing out the call stack every interval seconds, to the file specified. Either the summaryRprof function or the Perl script R CMD Rprof can be used to process the output file to produce a summary of the usage; use R CMD Rprof --help for usage information.

## Note

Profiling is not available on all platforms. By default, it is attempted to compile support for profiling. Configure R with --disable-R-profiling to change this.
As R profiling uses the same mechanisms as C profiling, the two cannot be used together, so do not use Rprof in an executable built for profiling.

## See Also

The chapter on "Tidying and profiling R code" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).

```
summaryRprof
```


## Examples

```
Rprof()
## some code to be profiled
Rprof(NULL)
## some code NOT to be profiled
Rprof (append=TRUE)
## some code to be profiled
Rprof(NULL)
## Post-process the output by
## R CMD Rprof Rprof.out
## at the shell prompt.
```

rug Add a Rug to a Plot

## Description

Adds a rug representation (1-d plot) of the data to the plot.

## Usage

```
rug(x, ticksize=0.03, side=1, lwd=0.5, col,
    quiet = getOption("warn") < 0, ...)
```


## Arguments

| x | A numeric vector |
| :--- | :--- |
| ticksize | The length of the ticks making up the 'rug'. Positive lengths give inwards <br> ticks. |
| side | On which side of the plot box the rug will be plotted. Normally 1 (bottom) <br> or 3 (top). |
| lwd | The line width of the ticks. |
| col | The colour the ticks are plotted in, default is black. |
| quiet | logical indicating if there should be a warning about clipped values. <br> $\ldots$. |
|  | further arguments, passed to axis (...), such as line or pos for specify- <br> ing the location of the rug. | ing the location of the rug.

## Details

Because of the way rug is implemented, only values of $x$ that fall within the plot region are included. There will be a warning if any finite values are omitted, but non-finite values are omitted silently.

Because of the way colours are done the axis itself is coloured the same as the ticks. You can always replot the box in black if you don't like this feature.

## Author(s)

B. D. Ripley

## See Also

jitter which you may want for ties in x .

## Examples

```
data(faithful)
attach(faithful)
plot(density(eruptions, bw=0.15))
rug(eruptions)
rug(jitter(eruptions, amount = .01), side = 3, col = "light blue")
detach("faithful")
```

sample Random Samples and Permutations

## Description

sample takes a sample of the specified size from the elements of x using either with or without replacement.

```
Usage
    sample(x, size, replace = FALSE, prob = NULL)
```


## Arguments

$\mathrm{x} \quad$ Either a (numeric, complex, character or logical) vector of more than one element from which to choose, or a positive integer.
size A positive integer giving the number of items to choose.
replace Should sampling be with replacement?
prob A vector of probability weights for obtaining the elements of the vector being sampled.

## Details

If x has length 1 , sampling takes place from $1: \mathrm{x}$.
By default size is equal to length ( $x$ ) so that sample $(x)$ generates a random permutation of the elements of $x$ (or 1:x).

The optional prob argument can be used to give a vector of weights for obtaining the elements of the vector being sampled. They need not sum to one, but they should be nonnegative and not all zero. If replace is false, these probabilities are applied sequentially, that is the probability of choosing the next item is proportional to the probabilities amongst the remaining items. The number of nonzero weights must be at least size in this case.

## Examples

```
x <- 1:12
# a random permutation
sample(x)
# bootstrap sampling
sample(x,replace=TRUE)
# 100 Bernoulli trials
sample(c(0,1), 100, replace = TRUE)
```

save Save $R$ Objects

## Description

save writes a external representation of R objects to the specified file. The objects can be read back from the file at a later date by using the function load (or data in some cases).
save.image() is just a short-cut for "save my current environment", i.e., save(list = ls(all=TRUE), file = ". RData"). It is what also happens with $q$ ("yes").

## Usage

```
save(..., list = character(0), file = "", ascii = FALSE,
    version = NULL, envir = parent.frame(), compress = FALSE)
save.image(file = ".RData", version = NULL, ascii = FALSE,
            compress = FALSE, safe = TRUE)
sys.load.image(name, quiet)
sys.save.image(name)
```


## Arguments

| $\ldots$. | the names of the objects to be saved. |
| :--- | :--- |
| list | A character vector containing the names of objects to be saved. |
| file | a connection or the name of the file where the data will be saved. Must <br> be a file name for workspace format version 1. |
| ascii | if TRUE, an ASCII representation of the data is written. This is useful <br> for transporting data between machines of different types. The default <br> value of ascii is FALSE which leads to a more compact binary file being <br> written. |


| version | the workspace format version to use. NULL specifies the current default |
| :--- | :--- |
| format. The version used from R 0.99.0 to R 1.3 .1 was version 1 . The |  |
| default format as from R 1.4.0 is version 2. |  | envir | environment to search for objects to be saved. |  |
| :--- | :--- |
| compress | logical specifying whether saving to a named file is to use compression. <br> Ignored when file is a connection and for workspace format version 1. |
| safe | logical. If TRUE, a temporary file is used for creating the saved <br> workspace. The temporary file is renamed to file if the save succeeds. <br> This preserves an existing workspace file if the save fails, but at the cost <br> of using extra disk space during the save. |
| quiet | name of image file to save or load. |

## Details

All R platforms use the XDR representation of binary objects in binary save-d files, and these are portable across all R platforms.

Default values for save.image options can be modified with the save.image.defaults option. This mechanism is experimental and subject to change.
sys.save.image is a system function that is called by q() and its GUI analogs; sys.load.image is called by the startup code. These functions should not be called directly and are subject to change.
sys.save.image closes all connections first, to ensure that it is able to open a connection to save the image. This is appropriate when called from q() and allies, but reinforces the warning that it should not be called directly.

## See Also

```
dput, dump, load, data.
```


## Examples

```
x <- runif(20)
y <- list(a = 1, b = TRUE, c = "oops")
save(x, y, file = "xy.Rdata")
save.image()
unlink("xy.Rdata")
unlink(".RData")
# set save.image defaults using option:
options(save.image.defaults=list(ascii=TRUE, safe=FALSE))
save.image()
unlink(".RData")
```

savehistory Load or Save or Display the Commands History

## Description

Load or save or display the commands history.

## Usage

```
loadhistory(file = ".Rhistory")
savehistory(file = ".Rhistory")
history(max.show = 25, reverse = FALSE)
```


## Arguments

file The name of the file in which to save the history, or from which to load it. The path is relative to the current working directory.
max.show The maximum number of lines to show. Inf will give all of the currently available history.
reverse logical. If true, the lines are shown in reverse order. Note: this is not useful when there are continuation lines.

## Details

This works under the readline and GNOME interfaces, but not if readline is not available.

## Note

If you want to save the history (almost) every session, you can put a call to savehistory () in .Last.

```
scale Scaling and Centering of Matrix-like Objects
```


## Description

scale is generic function whose default method centers and/or scales the columns of a numeric matrix.

## Usage

```
scale(x, center = TRUE, scale = TRUE)
```


## Arguments

$\mathrm{x} \quad$ a numeric matrix (like object).
center either a logical value or a numeric vector of length equal to the number of columns of x .
scale either a logical value or a numeric vector of length equal to the number of columns of x .

## Details

The value of center determines how column centering is performed. If center is a numeric vector with length equal to the number of columns of x , then each column of x has the corresponding value from center subtracted from it. If center is TRUE then centering is done by subtracting the column means (omitting NAs) of x from their corresponding columns, and if center is FALSE, no centering is done.

The value of scale determines how column scaling is performed (after centering). If scale is a numeric vector with length equal to the number of columns of $x$, then each column of x is divided by the corresponding value from scale. If scale is TRUE then scaling is done by dividing the (centered) columns of $x$ by their root-mean-square, and if scale is FALSE, no scaling is done.

The root-mean-square for a column is obtained by computing the square-root of the sum-of-squares of the non-missing values in the column divided by the number of non-missing values minus one.

## Value

For scale.default, the centered, scaled matrix. The numeric centering and scalings used (if any) are returned as attributes "scaled:center" and "scaled:scale"

## See Also

 sweep which allows centering (and scaling) with arbitrary statistics.
## Examples

```
x <- matrix(1:10, nc=2)
(centered.x <- scale(x, scale=FALSE))
cov(centered.scaled.x <- scale(x))# all 1
```


## scan Read Data Values

## Description

Read data into a vector or list from the console or file.

## Usage

```
scan(file = "", what = double(0), nmax = -1, n = -1, sep = "",
    quote = if (sep=="\n") "" else ">\"", dec = ".",
    skip = 0, nlines = 0, na.strings = "NA",
    flush = FALSE, fill = FALSE, strip.white = FALSE, quiet = FALSE,
    blank.lines.skip = TRUE, multi.line = TRUE, comment.char = "")
```


## Arguments

file
what the type of what gives the type of data to be read. If what is a list, it is assumed that the lines of the data file are records each containing length (what) items ("fields").
nmax the maximum number of data values to be read, or if what is a list, the maximum number of records to be read. If omitted (and nlines is not set to a positive value), scan will read to the end of file.
n
sep
the maximum number of data values to be read, defaulting to no limit.
by default, scan expects to read white-space delimited input fields. Alternatively, sep can be used to specify a character which delimits fields. A field is always delimited by a newline unless it is quoted.
quote the set of quoting characters as a single character string.
dec decimal point character.
skip the number of lines of the input file to skip before beginning to read data values.
nlines the maximum number of lines of data to be read.
na.strings character vector. Elements of this vector are to be interpeted as missing (NA) values.
flush logical: if TRUE, scan will flush to the end of the line after reading the last of the fields requested. This allows putting comments after the last field, but precludes putting more that one record on a line.
fill logical: if TRUE, scan will implicitly add empty fields to any lines with fewer fields than implied by what.
strip.white vector of logical value(s) corresponding to items in the what argument. It is used only when sep has been specified, and allows the stripping of leading and trailing white space from character fields (numeric fields are always stripped).
If strip.white is of length 1, it applies to all fields; otherwise, if strip.white[i] is TRUE and the i-th field is of mode character (because what [i] is) then the leading and trailing white space from field $i$ is stripped.
quiet logical: if FALSE (default), scan() will print a line, saying how many items have been read.
blank.lines.skip
logical: if TRUE blank lines in the input are ignored, except when counting skip and nlines.
multi.line logical. Only used if what is a list. If FALSE, all of a record must appear on one line (but more than one record can appear on a single line). Note that using fill $=$ TRUE implies that a record will terminated at the end of a line.
comment.char character: a character vector of length one containing a single character or an empty string. Use "" to turn off the interpretation of comments altogether (the default).

## Details

The value of what can be a list of types, in which case scan returns a list of vectors with the types given by the types of the elements in what. This provides a way of reading columnar data. If any of the types is NULL, the corresponding field is skipped (but a NULL component appears in the result).

The type of what or its components can be one of the five atomic types or NULL,
Empty numeric fields are always regarded as missing values. Empty character fields are scanned as empty character vectors, unless na.strings contains "" when they are regarded as missing values.

If sep is the default (""), the character \in a quoted string escapes the following character, so quotes may included in the string by escaping them.
If sep is non-default, the fields may be quoted in the style of '.csv' files where separators inside quotes ( $\mathbf{\square}$ or "") are ignored and quotes may be put inside strings by doubling them. However, if sep = "
n " it is assumed by default that one wants to read entire lines verbatim.
Note that since sep is a separator and not a terminator, reading a file by scan("foo", sep="
n", blank.lines.skip=FALSE) will give an empty file line if the file ends in a linefeed and not if it does not. This might not be what you expected; see also readLines.

If comment. char occurs (except inside a quoted character field), it signals that the rest of the line should be regarded as a comment and be discarded. Lines beginning with a comment character (possibly after white space) are treated as blank lines.

## Value

if what is a list, a list of the same length and same names (as any) as what.
Otherwise, a vector of the type of what.

## Note

The default for multi.line differs from $S$. To read one record per line, use flush $=$ TRUE and multi.line $=$ FALSE.

If number of items is not specified, the internal mechanism re-allocates memory in powers of two and so could use up to three times as much memory as needed. (It needs both old and new copies.) If you can, specify either $n$ or nmax whenever inputting a large vector, and nmax or nlines when inputting a large list.

## See Also

read.table for more user-friendly reading of data matrices; readLines to read a file a line at a time. write.

## Examples

```
cat("TITLE extra line", "2 3 5 7", "11 13 17", file="ex.data", sep="\n")
pp <- scan("ex.data", skip = 1, quiet= TRUE)
    scan("ex.data", skip = 1)
    scan("ex.data", skip = 1, nlines=1)# only 1 line after the skipped one
str(scan("ex.data", what = list("","",""))) # flush is F -> read "7"
str(scan("ex.data", what = list("","",""), flush = TRUE))
unlink("ex.data") # tidy up
```

```
screen Creating and Controlling Multiple Screens on a Single Device
```


## Description

split.screen defines a number of regions within the current device which can, to some extent, be treated as separate graphics devices. It is useful for generating multiple plots on a single device. Screens can themselves be split, allowing for quite complex arrangements of plots.
screen is used to select which screen to draw in.
erase.screen is used to clear a single screen, which it does by filling with the background colour.
close.screen removes the specified screen definition(s).

## Usage

```
split.screen(figs, screen, erase = TRUE)
screen(n, new = TRUE)
erase.screen(n)
close.screen(n, all.screens = FALSE)
```


## Arguments

| figs | A two-element vector describing the number of rows and the number of <br> columns in a screen matrix or a matrix with 4 columns. If a matrix, then <br> each row describes a screen with values for the left, right, bottom, and <br> top of the screen (in that order) in NDC units. |
| :--- | :--- |
| screen | A number giving the screen to be split. |
| erase | logical: should be selected screen be cleared? <br> n |
| A number indicating which screen to prepare for drawing (screen), erase |  |
| (erase.screen), or close (close.screen). |  |
| new | A logical value indicating whether the screen should be erased as part of <br> the preparation for drawing in the screen. |
| all.screens | A logical value indicating whether all of the screens should be closed. |

## Details

The first call to split.screen places R into split-screen mode. The other split-screen functions only work within this mode. While in this mode, certain other commands should be avoided (see WARNINGS below). Split-screen mode is exited by the command close.screen (all = TRUE)

## Value

split.screen returns a vector of screen numbers for the newly-created screens. With no arguments, split.screen returns a vector of valid screen numbers.
screen invisibly returns the number of the selected screen. With no arguments, screen returns the number of the current screen.
close.screen returns a vector of valid screen numbers.
screen, erase.screen, and close.screen all return FALSE if R is not in split-screen mode.

## Warning

These functions are totally incompatible with the other mechanisms for arranging plots on a device: $\operatorname{par}$ (mfrow), par (mfcol), and layout ().

The functions are also incompatible with some plotting functions, such as coplot, which make use of these other mechanisms.
The functions should not be used with multiple devices.
erase.screen will appear not to work if the background colour is transparent (as it is by default on most devices).

## See Also

```
par, layout, Devices, dev.*
```


## Examples

```
if (interactive()) {
par(bg = "white") # default is likely to be transparent
split.screen(c(2,1)) # split display into two screens
split.screen(c(1,3), screen = 2) # now split the bottom half into 3
screen(1) # prepare screen 1 for output
plot(10:1)
screen(4) # prepare screen 4 for output
plot(10:1)
close.screen(all = TRUE) # exit split-screen mode
split.screen(c(2,1)) # split display into two screens
split.screen(c(1,2),2) # split bottom half in two
plot(1:10)
    # screen 3 is active, draw plot
erase.screen() # forgot label, erase and redraw
plot(1:10, ylab= "ylab 3")
screen(1) # prepare screen 1 for output
plot(1:10)
screen(4) # prepare screen 4 for output
plot(1:10, ylab="ylab 4")
screen(1, FALSE) # return to screen 1, but do not clear
plot(10:1, axes=FALSE, lty=2, ylab="") # overlay second plot
axis(4) # add tic marks to right-hand axis
title("Plot 1")
close.screen(all = TRUE) # exit split-screen mode
}
```


## sd Standard Deviation

## Description

This function computes the standard deviation of the values in $x$. If na.rm is TRUE then missing values are removed before computation proceeds. If x is a matrix or a dataframe, a vector of the standard deviation of the columns is returned.

## Usage

$\operatorname{sd}(\mathrm{x}, \mathrm{na} . \mathrm{rm}=\mathrm{FALSE})$

## Arguments

$$
\begin{array}{ll}
\mathrm{x} & \text { a numeric vector, matrix or data frame. } \\
\text { na.rm } & \text { logical. Should missing values be removed? }
\end{array}
$$

## See Also

var for its square, and mad, the most robust alternative.

## Examples

```
sd(1:2) ^ 2
```

se.aov
Internal Functions Used by model.tables

## Description

Internal function for use by model.tables.

## Usage

```
se.aov(object, n, type = "means")
se.aovlist(object, dn.proj, dn.strata, factors, mf, efficiency,
    n, type = "diff.means", ...)
```


## Author(s)

B. D. Ripley

## See Also

model.tables

```
se.contrast Standard Errors for Contrasts in Model Terms
```


## Description

Returns the standard errors for one or more contrasts in an aov object.

## Usage

```
se.contrast(object, ...)
se.contrast(object, contrast.obj,
    coef = contr.helmert(ncol(contrast))[, 1], data = NULL, ...)
```


## Arguments

| object | A suitable fit, usually from aov. |
| :--- | :--- |
| contrast.obj | The contrasts for which standard errors are requested. This can be speci- <br> fied via a list or via a matrix. A single contrast can be specified by a list of <br> logical vectors giving the cells to be contrasted. Multiple contrasts should <br> be specified by a matrix, each column of which is a numerical contrast <br> vector (summing to zero). |
| coef | Used when \{contrast.obj\} is a list; it should be a vector of the same <br> length as the list with zero sum. The default value is the first Helmert <br> contrast, which contrasts the first and second cell means specified by the <br> list. |
| data | The data frame used to evaluate contrast.obj. |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

Contrasts are usually used to test if certain means are significantly different; it can be easier to use se.contrast than compute them directly from the coefficients.
In multistratum models, the contrasts can appear in more than one stratum; the contrast and standard error are computed in the lowest stratum and adjusted for efficiencies and comparisons between strata.
Suitable matrices for use with coef can be found by calling contrasts and indexing the columns by a factor.

## Value

A vector giving the standard errors for each contrast.

Author(s)
B. D. Ripley

## See Also

contrasts, model.tables

## Examples

```
## From Venables and Ripley (1997) p.210.
N <- c(0,1,0,1,1,1,0,0,0,1,1,0,1,1,0,0,1,0,1,0,1,1,0,0)
P <- c(1,1,0,0,0,1,0,1,1,1,0,0,0,1,0,1,1,0,0,1,0,1,1,0)
K <- c(1,0,0,1,0,1,1,0,0,1,0,1,0,1,1,0,0,0,1,1,1,0,1,0)
yield <- c(49.5,62.8,46.8,57.0,59.8,58.5,55.5,56.0,62.8,55.8,69.5,
55.0, 62.0,48.8,45.5,44.2,52.0,51.5,49.8,48.8,57.2,59.0,53.2,56.0)
npk <- data.frame(block = gl(6,4), N = factor(N), P = factor(P),
    K = factor(K), yield = yield)
options(contrasts=c("contr.treatment", "contr.poly"))
npk.aov1 <- aov(yield ~ block + N + K, npk)
se.contrast(npk.aov1, list(N=="0", N=="1"), data=npk)
# or via a matrix
cont <- matrix(c(-1,1), 2, 1, dimnames=list(NULL, "N"))
se.contrast(npk.aov1, cont[N, , drop=FALSE]/12, data=npk)
## test a multi-stratum model
npk.aov2 <- aov(yield ~ N + K + Error(block/(N + K)), npk)
se.contrast(npk.aov2, list(N == "0", N == "1"))
```


## search

Give Search Path for $R$ Objects

## Description

Gives a list of attached packages (see library), and R objects, usually data.frames.

## Usage

search()
searchpaths()

## Value

A character vector, starting with ".GlobalEnv", and ending with "package:base" which is R's base package required always.
searchpaths gives a similar character vector, with the entries for packages being the path to the package used to load the code.

## See Also

attach and detach to change the search "path", objects to find R objects in there.

## Examples

search ()
searchpaths()

## seek Functions to Reposition Connections

## Description

Functions to re-position connections.

```
Usage
seek(con, where = NA, origin = "start", rw = "", ...)
truncate(con, ...)
isSeekable(con)
```


## Arguments

con a connection.
where integer. A file position (relative to the origin specified by origin), or NA.
rw
character. Empty or "read" or "write", partial matches allowed.
origin character. One of "start", "current", "end".
... further arguments passed to or from other methods.

## Details

seek with where $=$ NA returns the current byte offset of a connection (from the beginning), and with a non-missing where argument the connection is re-positioned (if possible) to the specified position. isSeekable returns whether the connection in principle supports seek: currently only file connections do.
File connections can be open for both writing/appending, in which case R keeps separate positions for reading and writing. Which seek refers to can be set by its rw argument: the default is the last mode (reading or writing) which was used. Most files are only opened for reading or writing and so default to that state. If a file is open for reading and writing but has not been used, the default is to give the reading position (0).
The initial file position for reading is always at the beginning. The initial position for writing is at the beginning of the file for modes " $\mathrm{r}+$ " and " $\mathrm{r}+\mathrm{b}$ ", otherwise at the end of the file. Some platforms only allow writing at the end of the file in the append modes.
truncate truncates a file opened for writing at its current position. It works only for file connections, and is not implemented on all platforms.

## Value

seek returns the current position (before any move), as a byte offset, if relevant, or 0 if not. truncate returns NULL: it stops with an error if it fails (or is not implemented).
isSeekable returns a logical value, whether the connection is support seek.

## See Also

segments Add Line Segments to a Plot

## Description

Draw line segments between pairs of points.

## Usage

```
segments(x0, y0, x1, y1,
    col = par("fg"), lty = par("lty"), lwd = par("lwd"), ...)
```


## Arguments

$\mathrm{x} 0, \mathrm{y} 0 \quad$ coordinates of points from which to draw.
$\mathrm{x} 1, \mathrm{y} 1 \quad$ coordinates of points to which to draw.
col, lty, lwd usual graphical parameters as in par.
... further graphical parameters (from par).

## Details

For each i, a line segment is drawn between the point (x0[i], y0[i]) and the point (x1[i],y1[i]).

The graphical parameters col and lty can be used to specify a color and line texture for the line segments (col may be a vector).

## See Also

arrows, polygon for slightly easier and less flexible line drawing, and lines for the usual polygons.

## Examples

```
x <- runif(12); y <- rnorm(12)
i <- order(x,y); x <- x[i]; y <- y[i]
plot(x,y, main="arrows(.) and segments(.)")
## draw arrows from point to point :
s <- seq(length(x)-1)# one shorter than data
arrows(x[s], y[s], x[s+1], y[s+1], col= 1:3)
s <- s[-length(s)]
segments(x[s], y[s], x[s+2], y[s+2], col= 'pink')
```


## Description

Generate regular sequences.

## Usage

```
from:to
seq(from, to)
seq(from, to, by=)
seq(from, to, length=)
seq(along)
```


## Arguments

| from | starting value of sequence. |
| :--- | :--- |
| to | (maximal) end value of the sequence. |
| by | increment of the sequence. |
| length | desired length of the sequence. |
| along | take the length from the length of this argument. |

## Details

The operator : and the first seq(.) form generate the sequence from, from+1,..., to. seq is a generic function.

The second form generates from, from+by, ..., to.
The third generates a sequence of length equally spaced values from from to to.
The last generates the sequence $1,2, \ldots$, length (along).
If from and to are factors of the same length, then from : to returns the "cross" of the two.

Very small sequences (with from - to of the order of 1e-14 times the larger of the ends) will return from.

## Value

The result is of mode "integer" if from is (numerically equal to an) integer and by is not specified.

## See Also

```
rep, sequence, row, col.
```


## Examples

```
1:4
pi:6 # float
6:pi # integer
seq(0,1, length=11)
str(seq(rnorm(20)))
seq(1,9, by = 2) # match
seq(1,9, by = pi)# stay below
seq(1,6, by = 3)
seq(1.575, 5.125, by=0.05)
stopifnot(
    3 == seq(3,3, by=pi),
    3 == seq(3,3.1,by=pi),
    seq(1,6,by=3) == c(1,4),
    seq}(10,4.05,by=-3)==c(10,7
)
for (x in list(NULL, letters[1:6], list(1,pi)))
    cat("x=",deparse(x),"; seq(along = x):",seq(along = x),"\n")
f1 <- gl(2,3); f1
f2 <- gl(3,2); f2
f1:f2 # a factor, the ''cross'' f1 x f2
```

seq.POSIXt Generate Regular Sequences of Dates

## Description

The method for seq for data-time classes.

## Usage

```
seq(from, to, by, length.out=NULL, along.with=NULL, ...)
```


## Arguments

| from | starting date. Required |
| :--- | :--- |
| to | end date. Optional. If supplied must be after from. |
| by | increment of the sequence. Optional. See Details. |
| length.out | integer, optional. desired length of the sequence. |
| along.with | take the length from the length of this argument. |
| $\ldots$ | arguments passed to or from other methods. |

## Details

by can be specified in several ways.

- A number, taken to be in seconds.
- A object of class difftime
- A character string, containing one of "sec", "min", "hour", "day", "DSTday", "week", "month" or "year". This can optionally be preceded by an integer and a space, or followed by "s".

The difference between "day" and "DSTday" is that the former ignores changes to/from daylight savings time and the latter takes the same clock time each day. ("week" ignores DST, but "7 DSTdays") can be used as an alternative. "month" and "year" allow for DST as from R 1.5.0.)

## Value

A vector of class "POSIXct".

## See Also

DateTimeClasses

## Examples

```
## first days of years
seq(ISOdate(1910,1,1), ISOdate(1999,1,1), "years")
## by month
seq(ISOdate(2000,1,1), by="month", length=12)
## quarters
seq(ISOdate(1990,1,1), ISOdate(2000,1,1), by="3 months")
## days vs DSTdays
seq(ISOdate(2000,3,20), by="day", length = 10)
seq(ISOdate(2000,3,20), by="DSTday", length = 10)
seq(ISOdate(2000,3,20), by="7 DSTdays", length = 4)
```

sequence
Create A Vector of Sequences

## Description

For each element of nvec the sequence seq(nvec[i]) is created. These are appended and the result returned.

## Usage

sequence(nvec)

## Arguments

nvec an integer vector each element of which specifies the upper bound of a sequence.

## See Also

gl, seq, rep.

## Examples

## sets Set Operations

## Description

Performs set union, intersection, (asymmetric!) difference, equality and membership on two vectors.

## Usage

```
union(x, y)
intersect(x, y)
setdiff(x, y)
setequal(x, y)
is.element(el, set)
```


## Arguments

$\mathrm{x}, \mathrm{y}, \mathrm{el}$, set vectors (of the same mode) containing a sequence of items (conceptually) with no duplicated values.

## Details

Each of union, intersect and setdiff will remove any duplicated values in the arguments. is.element ( $\mathrm{x}, \mathrm{y}$ ) is identical to $\mathrm{x} \% \mathrm{in} \% \mathrm{y}$.

## Value

A vector of the same mode as x or y for setdiff and intersect, respectively, and of a common mode for union.

A logical scalar for setequal and a logical of the same length as x for is.element.

## Author(s)

B. D. Ripley

## See Also

\%in\%

## Examples

```
(x <- c(sort(sample(1:20, 9)),NA))
(y <- c(sort(sample(3:23, 7)),NA))
union(x, y)
intersect(x, y)
setdiff(x, y)
setdiff(y, x)
setequal(x, y)
## True for all possible x & y :
setequal( union(x,y),
        c(setdiff (x,y), intersect(x,y), setdiff(y,x)))
```

```
is.element(x, y)# length 10
```

is.element (y, x)\# length 8

SHLIB Build Shared Library for Dynamic Loading

## Description

Compile given source files using R CMD COMPILE, and then link all specified object files into a shared library which can be loaded into $R$ using dyn.load or library.dynam.

## Usage

R CMD SHLIB [options] [-o libname] files

## Arguments

files a list specifying the object files to be included in the shared library. You can also include the name of source files, for which the object files are automagically made from their sources.
libname the full name of the shared library to be built, including the extension (typically '.so' on Unix systems). If not given, the name of the library is taken from the first file.
options Further options to control the processing, or for obtaining information about usage and version of the utility.

## See Also

COMPILE, dyn.load, library.dynam

```
showConnections Display Connections
```


## Description

Display aspects of connections.

## Usage

```
showConnections(all=FALSE)
getConnection(what)
closeAllConnections()
stdin()
stdout()
stderr()
```


## Arguments

all logical: if true all connections, including closed ones and the standard ones are displayed. If false only open user-created connections are included.
what integer: a row number of the table given by showConnections.

## Details

stdin(), stdout() and stderr() are standard connections corresponding to input, output and error on the console respectively (and not necessarily to file streams). They are textmode connections of class "terminal" which cannot be opened or closed, and are read-only, write-only and write-only respectively. The stdout() and stderr() connections can be redirected by sink.
showConnections returns a matrix of information. If a connection object has been lost or forgotten, getConnection will take a row number from the table and return a connection object for that connection, which can be used to close the connection, for example.
closeAllConnections closes (and destroys) all open user connections, restoring all sink diversions as it does so.

## Value

stdin(), stdout() and stderr() return connection objects.
showConnections returns a character matrix of information with a row for each connection, by default only for open non-standard connections.
getConnection returns a connection object, or NULL.

## See Also

connections

## Examples

```
showConnections(all = TRUE)
textConnection(letters)
# oops, I forgot to record that one
showConnections()
# class description mode text isopen can read can write
#3 "letters" "textConnection" "r" "text" "opened" "yes" "no"
close(getConnection(3))
showConnections()
```

sign Sign Function

## Description

sign returns a vector with the signs of the corresponding elements of $x$ (the sign of a real number is 1,0 , or -1 if the number is positive, zero, or negative, respectively).
Note that sign does not operate on complex vectors.

## Usage

$$
\operatorname{sign}(x)
$$

## Arguments

x
a numeric vector

## See Also

abs

## Examples

```
sign(pi) # == 1
sign(-2:3)# -1 -1 0 1 1 1
```

Signals Interrupting Execution of $R$

## Description

On receiving SIGUSR1 R will save the workspace and quit. SIGUSR2 has the same result except that the .Last function and on.exit expressions will not be called.

## Usage

```
kill -USR1 pid
kill -USR2 pid
```


## Arguments

pid The process ID of the R process

## Warning

It is possible that one or more R objects will be undergoing modification at the time the signal is sent. These objects could be saved in a corrupted form.

## SignRank Distribution of the Wilcoxon Signed Rank Statistic

## Description

Density, distribution function, quantile function and random generation for the distribution of the Wilcoxon Signed Rank statistic obtained from a sample with size n.

## Usage

```
dsignrank(x, n, log = FALSE)
psignrank(q, n, lower.tail = TRUE, log.p = FALSE)
qsignrank(p, n, lower.tail = TRUE, log.p = FALSE)
rsignrank(nn, n)
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
$\mathrm{nn} \quad$ number of observations. If length $(\mathrm{nn})>1$, the length is taken to be the number required.
n
numbers of observations in the sample. Must be positive integers less than 50.
$\log$, log.p logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

This distribution is obtained as follows. Let x be a sample of size n from a continuous distribution symmetric about the origin. Then the Wilcoxon signed rank statistic is the sum of the ranks of the absolute values $\mathrm{x}[\mathrm{i}]$ for which $\mathrm{x}[\mathrm{i}]$ is positive. This statistic takes values between 0 and $n(n+1) / 2$, and its mean and variance are $n(n+1) / 4$ and $n(n+1)(2 n+1) / 24$, respectively.

## Value

dsignrank gives the density, psignrank gives the distribution function, qsignrank gives the quantile function, and rsignrank generates random deviates.

## Author(s)

Kurt Hornik 〈hornik@ci.tuwien.ac.at〉

## See Also

dwilcox etc, for the two-sample Wilcoxon rank sum statistic.

## Examples

```
par(mfrow=c (2,2))
for(n in c(4:5,10,40)) {
    x <- seq(0, n*(n+1)/2, length=501)
    plot(x, dsignrank(x,n=n), type='l', main=paste("dsignrank(x,n=",n,")"))
}
```

sink Send $R$ Output to a File

## Description

sink diverts R output to a connection.
sink.number() reports how many diversions are in use.
sink.number (type $=$ "message") reports the number of the connection currently being used for error messages.

## Usage

```
sink(file = NULL, append = FALSE, type = c("output", "message"))
sink.number(type = c("output", "message"))
```


## Arguments

| file | a connection or a character string naming the file to write to, or NULL to <br> stop sink-ing. |
| :--- | :--- |
| append | logical. If TRUE, output will be appended to file; otherwise, it will <br> overwrite the contents of file. |
| type | character. Either the output stream or the messages stream. |

## Details

sink diverts R output to a connection. If file is a character string, a file connection with that name will be established for the duration of the diversion.
Normal R output is diverted by the default type = "output". Only prompts and warning/error messages continue to appear on the terminal. These too can diverted by type $=$ "message" (see below).
sink() or $\operatorname{sink}(f i l e=N U L L) ~ e n d s ~ t h e ~ l a s t ~ d i v e r s i o n ~(o f ~ t h e ~ s p e c i f i e d ~ t y p e) . ~ A s ~ f r o m ~ R ~$ version 1.3.0 there is a stack of diversions for normal output, so output reverts to the previous diversion (if there was one). The stack is of up to 21 connections ( 20 diversions). If file is a connection if will be opened if necessary.
Sink-ing the messages stream should be done only with great care. For that stream file must be an already open connection, and there is no stack of connections.

## Value

For sink.
For sink. number () the number $(0,1,2, \ldots)$ of diversions of output in place.
For sink.number("message") the connection number used for messages, 2 if no diversion has been used.

## Warning

Don't use a connection that is open for sink for any other purpose. The software will stop you closing one such inadvertently.
Do not sink the messages stream unless you understand the source code implementing it and hence the pitfalls.

## Examples

```
sink("sink-examp.txt")
i <- 1:10
outer(i, i, "*")
sink()
unlink("sink-examp.txt")
## capture all the output to a file.
zz <- file("all.Rout", open="wt")
sink(zz)
```

```
sink(zz, type="message")
try (log("a"))
\#\# back to the console
sink (type="message")
sink()
try(log("a"))
```

sleep Students'Sleep Data

## Description

Data which show the effect of two soporific drugs (increase in hours of sleep) on groups consisting of 10 patients each.

## Usage

data(sleep)

## Format

A data frame with 20 observations on 2 variables.

| $[, 1]$ | extra | numeric | increase in hours of sleep |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | group | factor | patient group |

## Source

Student (1908) The probable error of the mean. Biometrika, 6, 20.

## References

Scheffé, Henry (1959) The Analysis of Variance. New York, NY: Wiley.

## Examples

```
data(sleep)
## ANOVA
anova(lm(extra ~ group, data = sleep))
```

$$
\text { slotOp } \quad \text { Extract or Replace Slots }
$$

## Description

Operators to extract or replace tbe contents of a slot in a object with a formal class structure.

## Usage

```
object@name
object@name <- value
```


## Arguments

$$
\begin{array}{ll}
\text { object } & \text { An object from a formally defined class. } \\
\text { name } & \text { The character-string name of the slot. }
\end{array}
$$

## Details

These operators support the formal classes of package methods. See slot for further details.

## See Also

Extract, slot

```
solve
```

Solve a System of Equations

## Description

This generic function solves the equation $\mathrm{a} \% * \% \mathrm{x}=\mathrm{b}$ for x , where b can be either a vector or a matrix.

## Usage

```
solve(a, b, tol = 1e-7, ...)
```


## Arguments

a a numeric matrix containing the coefficients of the linear system.
b a numeric vector or matrix giving the right-hand side(s) of the linear system. If omitted, b is taken to be an identity matrix and solve will return the inverse of $a$.
tol the tolerance for detecting linear dependencies in the columns of a.
... further arguments passed to or from other methods

## Details

As from R version 1.3.0, a or b can be complex, in which case LAPACK routine ZESV is used. This uses double complex arithmetic which might not beavailable on all platforms.

## See Also

backsolve, qr.solve.

## Examples

```
hilbert <- function(n) { i <- 1:n; 1 / outer(i - 1, i, "+") }
h8 <- hilbert(8); h8
solve(h8) # gives error: 'singular'
sh8 <- solve(h8, tol = 1e-10)
round(sh8 %**% h8, 3)
A <- hilbert(4)
A[] <- as.complex(A)
## might not be supported on all platforms
try(solve(A))
```


## Description

Sort (or order) a numeric or complex vector (partially) into ascending (or descending) order.

## Usage

sort $(x$, partial $=$ NULL, na.last $=$ NA, decreasing $=$ FALSE, method = c("shell", "quick"), index.return = FALSE)
is.unsorted(x, na.rm = FALSE)

## Arguments

| x | a numeric or complex vector. |
| :--- | :--- |
| partial | a vector of indices for partial sorting. |
| na.last | for controlling the treatment of NAs. If TRUE, missing values in the data <br> are put last; if FALSE, they are put first; if NA, they are removed. |
| decreasing | logical. Should the sort be increasing or decreasing? <br> character specifying the algorithm used. |
| method | logical indicating if the ordering index vector should be returned as well; <br> index.return <br> logis <br> this is only available for the default na.last = NA. |
| na.rm | logical. Should missing values be removed? |

## Details

If partial is not NULL, it is taken to contain indices of elements of $x$ which are to be placed in their correct positions by partial sorting. After the sort, the values specified in partial are in their correct position in the sorted array. Any values smaller than these values are guaranteed to have a smaller index in the sorted array and any values which are greater are guaranteed to have a bigger index in the sorted array.
The sort order for character vectors will depend on the collating sequence of the locale in use: see Comparison.
is.unsorted returns a logical indicating if $x$ is sorted increasingly, i.e. is.unsorted( $x$ ) is true if any ( $\mathrm{x}!=\operatorname{sort}(\mathrm{x})$ ) (and there are no NAs).
method = "shell" uses Shellsort and was the only method before R version 1.5.x (although improved there to an $O\left(n^{4 / 3}\right)$ variant of Sedgewick (1996)).
Method "quick" uses Singleton's Quicksort implementation and is only available when x is numeric, the sort is increasing and partial is NULL. It is normally somewhat faster than Shellsort (perhaps twice as fast on vectors of length a million) but has poor performance in the rare worst case. (Peto's modification using a pseudo-random midpoint is used to make the worst case rarer.)

## Value

For sort the sorted vector unless index.return is true, when the result is a list with components named x and ix containing the sorted numbers and the ordering index vector. In the latter case, if method == "quick" ties may be reversed in the ordering, unlike sort.list, as quicksort is not stable.

## References

Sedgewick, R. (1986) A new upper bound for Shell sort. J. Algorithms 7, 159-173.
Singleton, R. C. (1969) An efficient algorithm for sorting with minimal storage: Algorithm 347. Communications of the ACM 12, 185-187.

## See Also

order, rank.

## Examples

```
data(swiss)
x <- swiss$Education[1:25]
x; sort(x); sort(x, partial = c(10, 15))
median # shows you another example for 'partial'
stopifnot(!is.unsorted(sort(x)),
    !is.unsorted(LETTERS),
    is.unsorted(c(NA,1:3,2), na.rm = TRUE))
## illustrate 'stable' sorting (of ties):
sort(c(10:3,2:12), method = "sh", index=TRUE) # is stable
```



```
## $ix: 9 8 10 7 11 6 6 12 5 5 13 4 4 14 3 15 2 16 16 1 17 18 19
sort(c(10:3,2:12), method = "qu", index=TRUE) # is not
```



```
## $ix: 9 10 8 7 7 11 6 12 5 5 13 4 4 14 3 15 16 % 2 17 14 18 19
##
## Small speed comparison simulation:
N <- 2000
Sim <- 20
rep <- 50 # << adjust to your CPU
c1 <- c2 <- numeric(Sim)
for(is in 1:Sim){
    x <- rnorm(N)
    gc() ## sort should not have to pay for gc
    c1[is] <- system.time(for(i in 1:rep) sort(x, method = "shell"))[1]
    c2[is] <- system.time(for(i in 1:rep) sort(x, method = "quick"))[1]
    stopifnot(sort(x, meth = "s") == sort(x, meth = "q"))
}
100 * rbind(ShellSort = c1, QuickSort = c2)
cat("Speedup factor of quick sort():\n")
summary({qq <- c1 / c2; qq[is.finite(qq)]})
## A larger test
x <- rnorm(1e6)
gc()
system.time(x1 <- sort(x, method = "shell"))
gc()
system.time(x2 <- sort(x, method = "quick"))
stopifnot(identical(x1, x2))
```

```
source Read R Code from a File or a Connection
```


## Description

source causes $R$ to accept its input from the named file (the name must be quoted). Input is read from that file until the end of the file is reached. parse is used to scan the expressions in, they are then evaluated sequentially in the chosen environment.

## Usage

```
source(file, local = FALSE, echo = verbose, print.eval = echo,
verbose = getOption("verbose"), prompt.echo = getOption("prompt"),
max.deparse.length = 150, chdir = FALSE)
```


## Arguments

file a connection or a character string giving the name of the file or URL to read from.
local if local is FALSE, the statements scanned are evaluated in the user's workspace (the global environment), otherwise in the environment calling source.
echo logical; if TRUE, each expression is printed after parsing, before evaluation.
print.eval logical; if TRUE, the result of eval(i) is printed for each expression $i$; defaults to echo.
verbose if TRUE, more diagnostics (than just echo = TRUE) are printed during parsing and evaluation of input, including extra info for each expression.
prompt.echo character; gives the prompt to be used if echo = TRUE.
max.deparse.length
integer; is used only if echo is TRUE and gives the maximal length of the "echo" of a single expression.
chdir logical; if TRUE, the R working directory is changed to the directory containing file for evaluating.

## Details

All versions of R accept input from a connection with end of line marked by LF (as used on Unix), CRLF (as used on DOS/Windows) or CR (as used on Mac). The final line can be incomplete, that is missing the final EOL marker.
If options("keep.source") is true (the default), the source of functions is keep so they can be listed exactly as input. This imposes a limit of 128 K chars on the function size and a nesting limit of 265 . Use option(keep.source $=$ FALSE) when these limits might take effect: if exceeded they generate an error.

## See Also

demo which uses source; eval, parse and scan; options("keep.source").

## Special Special Functions of Mathematics

## Description

Special mathematical functions related to the beta and gamma functions.

## Usage

```
beta(a, b)
lbeta(a, b)
gamma(x)
lgamma(x)
digamma(x)
trigamma(x)
tetragamma(x)
pentagamma(x)
choose(n, k)
lchoose(n, k)
```


## Arguments

a, b, x
numeric vectors.
n, k
integer vectors.

## Details

The functions beta and lbeta return the beta function and the natural logarithm of the beta function,

$$
B(a, b)=\frac{\Gamma(a) \Gamma(b)}{\Gamma(a+b)}
$$

The functions gamma and lgamma return the gamma function $\Gamma(x)$ and the natural logarithm of the absolute value of the gamma function.
The functions digamma, trigamma, tetragamma and pentagamma return the first, second, third and fourth derivatives of the logarithm of the gamma function.

$$
\operatorname{digamma}(\mathrm{x})=\psi(x)=\frac{d}{d x} \ln \Gamma(x)=\frac{\Gamma^{\prime}(x)}{\Gamma(x)}
$$

The functions choose and lchoose return binomial coefficients and their logarithms.

## References

Abramowitz, M. and Stegun, I. A. (1972) Handbook of Mathematical Functions. New York: Dover. Chapter 6: Gamma and Related Functions.

## See Also

Arithmetic for simple, sqrt for miscellaneous mathematical functions and Bessel for the real Bessel functions.

## Examples

```
choose(5, 2)
for (n in 0:10) print(choose(n, k = 0:n))
curve(gamma(x), -3,4, n=1001, ylim=c(-10,100),
            col="red", lwd=2, main="gamma(x)")
abline(h=0,v=0, lty=3, col="midnightblue")
x <- seq(.1, 4, length = 201); dx <- diff(x)[1]
par(mfrow = c(2, 3))
for (ch in c("", "l","di","tri","tetra","penta")) {
    is.deriv <- nchar(ch) >= 2
    if (is.deriv) dy <- diff(y) / dx
    nm <- paste(ch, "gamma", sep = "")
    y <- get(nm)(x)
    plot(x, y, type = "l", main = nm, col = "red")
    abline(h = 0, col = "lightgray")
    if (is.deriv) lines(x[-1], dy, col = "blue", lty = 2)
}
par(mfrow = c(2, 2))
```

    splinefun Interpolating Splines
    
## Description

Perform cubic spline interpolation of given data points, returning either a list of points obtained by the interpolation or a function performing the interpolation.

## Usage

```
splinefun(x, y = NULL, method = "fmm")
spline(x, y = NULL, n = 3*length(x), method = "fmm",
    xmin = min(x), xmax = max(x))
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | vectors giving the coordinates of the points to be interpolated. Alterna- <br> tively a single plotting structure can be specified: see xy.coords. |
| :--- | :--- |
| method | specifies the type of spline to be used. Possible values are "fmm", <br> "natural" and "periodic". |
| n | interpolation takes place at n equally spaced points spanning the interval <br> [xmin, xmax]. |
| xmin | left-hand endpoint of the interpolation interval. |
| xmax | right-hand endpoint of the interpolation interval. |

## Details

If method $=$ "fmm", the spline used is that of Forsythe, Malcolm and Moler (an exact cubic is fitted through the four points at each end of the data, and this is used to determine the end conditions). Natural splines are used when method = "natural", and periodic splines when method = "periodic".
These interpolation splines can also be used for extrapolation, that is prediction at points outside the range of $x$. Extrapolation makes little sense for method = "fmm"; for natural splines it is linear using the slope of the interpolating curve at the nearest data point.

## Value

spline returns a list containing components x and y which give the ordinates where interpolation took place and the interpolated values.
splinefun returns a function which will perform cubic spline interpolation of the given data points. This is often more useful than spline.

## References

Forsythe, G. E., Malcolm, M. A. and Moler, C. B. (1977) Computer Methods for Mathematical Computations.

## See Also

approx and approxfun for constant and linear interpolation.
Package splines, especially interpSpline and periodicSpline for interpolation splines. That package also generates spline bases that can be used for regression splines. smooth.spline in package modreg for smoothing splines.

## Examples

```
op <- par(mfrow = c(2,1), mgp = c(2,.8,0), mar = .1+c(3,3,3,1))
n <- 9
x <- 1:n
y <- rnorm(n)
plot(x, y, main = paste("spline[fun](.) through", n, "points"))
lines(spline(x, y))
lines(spline(x, y, n = 201), col = 2)
y <- (x-6)^2
plot(x, y, main = "spline(.) -- 3 methods")
lines(spline(x, y, n = 201), col = 2)
lines(spline(x, y, n = 201, method = "natural"), col = 3)
lines(spline(x, y, n = 201, method = "periodic"), col = 4)
legend(6,25, c("fmm","natural","periodic"), col=2:4, lty=1)
f <- splinefun(x, y)
ls(envir = environment(f))
splinecoef <- eval(expression(z), envir = environment(f))
curve(f(x), 1, 10, col = "green", lwd = 1.5)
points(splinecoef, col = "purple", cex = 2)
par(op)
```


## split Divide into Groups

## Description

split divides the data in the vector $x$ into the groups defined by $f$. The assignment forms replace values corresponding to such a division. Unsplit reverses the effect of split.

```
Usage
    split(x, f)
    split.default(x, f)
    split.data.frame(x, f)
    split(x, f) <- value
    split.default(x, f) <- value
    split.data.frame(x, f) <- value
    unsplit(value, f)
```


## Arguments

$\mathrm{x} \quad$ vector or data frame containing values to be divided into groups.
$\mathrm{f} \quad \mathrm{a}$ "factor" such that factor (f) defines the grouping, or a list of such factors in which case their interaction is used for the grouping.
value a list of vectors or data frames compatible with a splitting of x

## Details

f is recycled as necessary and if the length of x is not a multiple of the length of f a warning is printed. unsplit works only with lists of vectors. The data frame method can also be used to split a matrix into a list of matrices, and the assignment form likewise, provided they are invoked explicitly.

## Value

The value returned from split is a list of vectors containing the values for the groups. The components of the list are named by the factor levels given be $f$. If $f$ is longer than $x$ some of these will be of zero length. The assignment forms return their right hand side. unsplit returns a vector for which split ( $x, f$ ) equals value

## See Also

cut

## Examples

```
n <- 10; nn <- 100
g <- factor(round(n * runif(n * nn)))
x <- rnorm(n * nn) + sqrt(as.numeric(g))
xg <- split(x, g)
boxplot(xg, col = "lavender", notch = TRUE, varwidth = TRUE)
sapply(xg, length)
sapply(xg, mean)
```

```
## Calculate z-scores by group
z <- unsplit(lapply(split(x, g), scale), g)
tapply(z, g, mean)
# or
z <- x
split(z, g) <- lapply(split(x, g), scale)
tapply(z, g, sd)
## Split a matrix into a list by columns
ma <- cbind(x = 1:10, y = (-4:5)^2)
split(ma, col(ma))
split(1:10, 1:2)
```

sprintf Use C-style String Formatting Commands

## Description

A wrapper for the C function sprintf, that returns a character vector of length one containing a formatted combination of text and variable values.

```
Usage
    sprintf(fmt, ...)
```


## Arguments

fmt a format string.
... values to be passed into fmt. Only logical, integer, real and character vectors are accepted, and only the first value is read from each vector.

## Details

This is a wrapper for the system's C call. Attempts are made to check that the mode of the values passed match the format supplied, and R's special values (NA, Inf, -Inf and NaN ) are handled correctly.
The following is abstracted from K\&R (see References, below). The string fmt contains normal characters, which are passed through to the output string, and also special characters that operate on the arguments provided through . . . Special characters start with a \% and terminate with one of the letters in the set difeEgGs\%. These letters denote the following types:
d,i Integer value
f Double precision value, in decimal notation of the form "[-]mmm.ddd". The number of decimal places is specified by the precision: the default is 6 ; a precision of 0 suppresses the decimal point.
$\mathrm{e}, \mathrm{E}$ Double precision value, in decimal notation of the form $[-] \mathrm{m}$. ddde $[+-] \mathrm{xx}$ or [-]m.dddE [+-] xx
g , G Double precision value, in \%e or $\% \mathrm{E}$ format if the exponent is less than -4 or greater than or equal to the precision, and \%f format otherwise
s Character string
\% Literal \% (none of the formatting characters given below are permitted in this case)
In addition, between the initial \% and the terminating conversion character there may be, in any order:
m.n Two numbers separated by a period, denoting the field width (m) and the precision (n)

- Left adjustment of converted argument in its field
+ Always print number with sign
a space Prefix a space if the first number is not a sign
0 For numbers, pad to the field width with leading zeros


## Value

A character vector of length one. Character NAs are converted to "NA".

## Author(s)

Original code by Jonathan Rougier, 〈J.C.Rougier@durham.ac.uk〉

## References

Kernighan, B. W. and Ritchie, D. M. (1988) The C Programming Language. Second edition, Prentice Hall. describes the format options in table B-1 in the Appendix.

## See Also

formatC for a way of formatting vectors of numbers in a similar fashion. paste for another way of creating a vector combining text and values.

## Examples

```
## be careful with the format: most things in R are floats
sprintf("%s is %f feet tall\n", "Sven", 7) # OK
try(sprintf("%s is %i feet tall\n", "Sven", 7)) # not OK
sprintf("%s is %i feet tall\n", "Sven", as.integer(7)) # OK again
## use a literal % :
sprintf("%.Of%% said yes (out of a sample of size %.Of)", 66.666, 3)
## various formats of pi :
sprintf("%f", pi)
sprintf("%.3f", pi)
sprintf("%1.0f", pi)
sprintf("%5.1f", pi)
sprintf("%05.1f", pi)
```

```
sprintf("%+f", pi)
sprintf("% f", pi)
sprintf("%-10f", pi)# left justified
sprintf("%e", pi)
sprintf("%E", pi)
sprintf("%g", pi)
sprintf("%g", 1e6 * pi) # -> exponential
sprintf("%.9g", 1e6 * pi) # -> "fixed"
sprintf("%G", 1e-6 * pi)
## no truncation:
sprintf("%1.f",101)
## More sophisticated:
lapply(c("a", "ABC", "and an even longer one"),
    function(ch) sprintf("10-string (%10s'", ch))
sapply(1:18, function(n)
    sprintf(paste("e with %2d digits = %.",n,"g",sep=""),
                n, exp(1)))
```

stack Stack or Unstack Vectors from a Data Frame or List

## Description

Stacking vectors concatenates multiple vectors into a single vector along with a factor indicating where each observation originated. Unstacking reverses this operation.

## Usage

```
stack(x, ...)
stack.default(x, ...)
stack.data.frame(x, select, ...)
unstack(x, ...)
unstack.default(x, form, ...)
unstack.data.frame(x, form = formula(x), ...)
```


## Arguments

| x | object to be stacked or unstacked |
| :--- | :--- |
| select | expression, indicating variables to select from a data frame |
| form | a two-sided formula whose left side evaluates to the vector to be unstacked <br> and whose right side evaluates to the indicator of the groups to create. |
|  | Defaults to formula (x) in unstack. data.frame. |
| $\ldots$ | further arguments passed to or from other methods. |

## Details

The stack function is used to transform data available as separate columns in a data frame or list into a single column that can be used in an analysis of variance model or other linear model. The unstack function reverses this operation.

## Value

unstack produces a list of columns according to the formula form. If all the columns have the same length, the resulting list is coerced to a data frame.
stack produces a data frame with two columns
values the result of concatenating the selected vectors in x
ind a factor indicating from which vector in x the observation originated

## Author(s)

Douglas Bates

## See Also

lm, reshape

## Examples

```
data(PlantGrowth)
formula(PlantGrowth) # check the default formula
pg <- unstack(PlantGrowth) # unstack according to this formula
pg
stack(pg) # now put it back together
stack(pg, select = -ctrl) # omitting one vector
```

```
stackloss Brownlee's Stack Loss Plant Data
```


## Description

Operational data of a plant for the oxidation of ammonia to nitric acid.

## Usage

data(stackloss)

## Format

stackloss is a data frame with 21 observations on 4 variables.

| $[, 1]$ | Air Flow | Flow of cooling air |
| :--- | :--- | :--- |
| $[, 2]$ | Water Temp | Cooling Water Inlet Temperature |
| $[, 3]$ | Acid Conc. | Concentration of acid [per 1000, minus 500] |
| $[, 4]$ | stack.loss | Stack loss |

For compatibility with S-PLUS, the data sets stack.x, a matrix with the first three (independent) variables of the data frame, and stack.loss, the numeric vector giving the fourth (dependent) variable, are provided as well.

## Details

"Obtained from 21 days of operation of a plant for the oxidation of ammonia $\left(\mathrm{NH}_{3}\right)$ to nitric acid $\left(\mathrm{HNO}_{3}\right)$. The nitric oxides produced are absorbed in a countercurrent absorption tower." (Brownlee, cited by Dodge, slightly reformatted by MM.)
Air Flow represents the rate of operation of the plant. Water Temp is the temperature of cooling water circulated through coils in the absorption tower. Acid Conc. is the concentration of the acid circulating, minus 50 , times 10: that is, 89 corresponds to 58.9 per cent acid. stack.loss (the dependent variable) is 10 times the percentage of the ingoing ammonia to the plant that escapes from the absorption column unabsorbed; that is, an (inverse) measure of the over-all efficiency of the plant.

## Source

Brownlee, K. A. (1960, 2nd ed. 1965) Statistical Theory and Methodology in Science and Engineering. New York: Wiley. pp. 491-500.

## References

Dodge, Y. (1996) The guinea pig of multiple regression. In: Robust Statistics, Data Analysis, and Computer Intensive Methods; In Honor of Peter Huber's 60th Birthday, 1996, Lecture Notes in Statistics 109, Springer-Verlag, New York.

## Examples

```
data(stackloss)
summary(lm.stack <- lm(stack.loss ~ stack.x))
```

standardGeneric Formal Method System Placeholders

## Description

These routines are primitives used with the methods package. They should not be used without it and do not need to be called directly in any case.
standardGeneric: dispatch the method defined for generic function named $f$, using the actual arguments in the frame from which standardGeneric is called.
objWithClass: return the result of setting the class of object to value. Defined as a separate primitive function because R types cannot generally be changed in place.
dataClass: returns a single string for the class of object even in the case that the object has an old-style class attribute with multiple strings.
topicName: the string used internally to find documenation of the given type and topic. Called by the ? operator and the special prompt functions in the methods package.

```
Usage
    standardGeneric(f)
    objWithClass(object, value)
    dataClass(object)
    topicName(type, topic)
```


## Author(s)

John Chambers

## stars

Star (Spider/Radar) Plots and Segment Diagrams

## Description

Draw star plots or segment diagrams of a multivariate data set. With one single location, also draws "spider" (or "radar") plots.

## Usage

```
stars(x, full = TRUE, scale = TRUE, radius = TRUE,
    labels \(=\) dimnames \((x)[[1]]\), locations \(=\) NULL,
    nrow \(=\) NULL, ncol \(=\) NULL, len \(=1\),
    key.loc = NULL, key.labels = dimnames(x) [[2]], key.xpd = TRUE,
    xlim = NULL, ylim = NULL, flip.labels = NULL,
    draw.segments = FALSE, col.segments = 1:n.seg, col.stars = NA,
    axes = FALSE, frame.plot = axes,
    main \(=\) NULL, sub \(=\) NULL, xlab \(=\) "", ylab \(=\) "",
    cex \(=0.8\), lwd \(=0.25\), lty = par("lty"), xpd = FALSE,
    mar = pmin(par("mar"),
                            \(1.1+c(2 * a x e s+(x l a b ~!=~ " "), 2 * a x e s+(y l a b ~!=~ " "), 1,0))\),
    add=FALSE, plot=TRUE, ...)
```


## Arguments

X
full logical flag: if TRUE, the segment plots will occupy a full circle. Otherwise, they occupy the (upper) semicircle only.
scale logical flag: if TRUE, the columns of the data matrix are scaled independently so that the maximum value in each column is 1 and the minimum is 0 . If FALSE, the presumption is that the data have been scaled by some other algorithm to the range $[0,1]$.
radius logical flag: in TRUE, the radii corresponding to each variable in the data will be drawn.
labels vector of character strings for labeling the plots. Unlike the S function stars, no attempt is made to construct labels if labels = NULL.
locations Either two column matrix with the x and y coordinates used to place each of the segment plots; or numeric of length 2 when all plots should be superimposed (for a "spider plot"). By default, locations = NULL, the segment plots will be placed in a rectangular grid.
nrow, ncol integers giving the number of rows and columns to use when locations is NULL. By default, nrow == ncol, a square layout will be used.
matrix or data frame of data. One star or segment plot will be produced for each row of x . Missing values (NA) are allowed, but they are treated as if they were 0 (after scaling, if relevant).

len
scale factor for the length of radii or segments.

| key.loc | vector with x and y coordinates of the unit key. |
| :---: | :---: |
| key.labels | vector of character strings for labeling the segments of the unit key. If omitted, the second component of dimnames ( $x$ ) is used, if available. |
| key.xpd | clipping switch for the unit key (drawing and labeling), see par ("xpd"). |
| xlim | vector with the range of x coordinates to plot. |
| ylim | vector with the range of y coordinates to plot. |
| flip.labels | logical indicating if the label locations should flip up and down from diagram to diagram. Defaults to a somewhat smart heuristic. |
| draw.segments | logical. If TRUE draw a segment diagram. |
| col.segments | color vector (integer or character, see par), each specifying a color for one of the segments (variables). Ignored if draw. segments = FALSE. |
| col.stars | color vector (integer or character, see par), each specifying a color for one of the stars (cases). Ignored if draw. segments = TRUE. |
| axes | logical flag: if TRUE axes are added to the plot. |
| frame.plot | logical flag: if TRUE, the plot region is framed. |
| main | a main title for the plot. |
| sub | a sub title for the plot. |
| xlab | a label for the x axis. |
| ylab | a label for the y axis. |
| cex | character expansion factor for the labels. |
| lwd | line width used for drawing. |
| lty | line type used for drawing. |
| xpd | logical or NA indicating if clipping should be done, see par (xpd = . ). |
| mar | argument to par (mar $=*$ ), typically chosing smaller margings than by default. |
|  | further arguments, passed to the first call of plot(), see plot.default and to box() if frame.plot is true. |
| add | logical, if TRUE $a d d$ stars to current plot. |
| plot | logical, if FALSE, nothing is plotted. |

## Details

Missing values are treated as 0 .
Each star plot or segment diagram represents one row of the input x. Variables (columns) start on the right and wind counterclockwise around the circle. The size of the (scaled) column is shown by the distance from the center to the point on the star or the radius of the segment representing the variable.

Only one page of output is produced.

## Note

This code started life as spatial star plots by David A. Andrews. See http://www.udallas. edu:8080/~andrews/software/software.html.

Prior to 1.4.1, scaling only shifted the maximum to 1 , although documented as here.

## Author(s)

Thomas S. Dye

## Examples

```
data(mtcars)
stars(mtcars[, 1:7], key.loc = c(14, 2),
    main = "Motor Trend Cars : stars(*, full = F)", full = FALSE)
stars(mtcars[, 1:7], key.loc = c(14, 1.5),
    main = "Motor Trend Cars : full stars()",flip.labels=FALSE)
## 'Spider' or 'Radar' plot:
stars(mtcars[, 1:7], locations = c(0,0), radius = FALSE,
    key.loc=c(0,0), main="Motor Trend Cars", lty = 2)
## Segment Diagrams:
palette(rainbow(12, s = 0.6, v = 0.75))
stars(mtcars[, 1:7], len = 0.8, key.loc = c(12, 1.5),
    main = "Motor Trend Cars", draw.segments = TRUE)
stars(mtcars[, 1:7], len = 0.6, key.loc = c(1.5, 0),
        main = "Motor Trend Cars", draw.segments = TRUE,
        frame.plot=TRUE, nrow = 4, cex = .7)
data(USJudgeRatings)
## scale linearly (not affinely) to [0, 1]
USJudge <- apply(USJudgeRatings, 2, function(x) x/max(x))
Jnam <- case.names(USJudgeRatings)
Snam <- abbreviate(substring(Jnam,1,regexpr("[,.]",Jnam) - 1), 7)
stars(USJudge, labels = Jnam, scale = FALSE,
    key.loc = c(13, 1.5), main = "Judge not ...", len = 0.8)
stars(USJudge, labels = Snam, scale = FALSE,
    key.loc = c(13, 1.5), radius = FALSE)
loc <- stars(USJudge, labels = NULL, scale = FALSE,
                                    radius = FALSE, frame.plot = TRUE,
                                    key.loc = c(13, 1.5), main = "Judge not ...", len = 1.2)
text(loc, Snam, col = "blue", cex = 0.8, xpd = TRUE)
## 'Segments':
stars(USJudge, draw.segments = TRUE, scale = FALSE, key.loc = c(13,1.5))
## 'Spider':
stars(USJudgeRatings, locations=c(0,0), scale=FALSE,radius = FALSE,
    col.stars=1:10, key.loc = c(0,0), main="US Judges rated")
## 'Radar-Segments'
stars(USJudgeRatings[1:10,], locations = 0:1, scale=FALSE,
    draw.segments = TRUE, col.segments=0, col.stars=1:10,key.loc= 0:1,
        main="US Judges 1-10 ")
palette("default")
stars(cbind(1:16,10*(16:1)),draw.segments=TRUE,
        main = "A Joke -- do *not* use symbols on 2D data!")
```


## Description

Extract and encode the times the first and last observations were taken. Provided only for compatibility with S version 2 .

## Usage

```
    start(x, ...)
```

    end ( \(x, \ldots\) )
    
## Arguments

| x | a univariate or multivariate time-series, or a vector or matrix. |
| :--- | :--- |
| $\ldots$ | extra arguments for future methods. |

## Details

These are generic functions, which will use the tsp attribute of x if it exists. Their default methods decode the start time from the original time units, so that for a monthly series 1995.5 is represented as $c(1995,7)$. For a series of frequency $f$, time $n+i / f$ is presented as $c(n, i+1)($ even for $i=0$ and $f=1)$.

## Warning

The representation used by start and end has no meaning unless the frequency is supplied.

```
See Also
ts, time, tsp.
```

Startup Initialization at Start of an $R$ Session

## Description

In $R$, the startup mechanism is as follows.
Unless --no-environ was given on the command line, R searches for user and site files to process for setting environment variables. The name of the site file is the one pointed to by the environment variable R_ENVIRON; if this is unset or empty, '\$R_HOME/etc/Renviron.site' is used (if it exists, which it does not in a "factory-fresh" installation). The user files searched for are '.Renviron' in the current or in the user's home directory (in that order). See Details for how the files are read.
Then R searches for the site-wide startup profile unless the command line option --no-site-file was given. The name of this file is taken from the value of the R_PROFILE environment variable. If this variable is unset, the default is '\$R_HOME/etc/Rprofile.site', which is used if it exists (which it does not in a "factory-fresh" installation). This code is loaded into package base.

Then, unless --no-init-file was given, R searches for a file called '.Rprofile' in the current directory or in the user's home directory (in that order) and sources it into the user workspace.

It then loads a saved image of the user workspace from '.RData' if there is one (unless --no-restore-data was specified, or --no-restore, on the command line).

Finally, if a function .First is found on the search path, it is executed as .First(). In a "factory-fresh" version of R the .First in the system profile file 'library/base/R/Rprofile' is used. (This requires the ctest library.)

A function.First (and .Last) can be defined in appropriate. Rprofile or Rprofile.site files or have been saved in '.RData'.

The commands history is read from the file specified by the environment variable R_HISTFILE (default .Rhistory) unless --no-restore-history was specified (or --norestore).

The command-line flag --vanilla implies --no-site-file, --no-init-file, --norestore and --no-environ.

## Usage

```
.First <- function() { ...... }
.Rprofile <startup file>
```


## Details

Note that there are two sorts of files used in startup: environment files which contain lists of environment variables to be set, and profile files which contain R code.

Lines in a site or user environment file should be either comment lines starting with \#, or lines of the form name=value. The latter sets the environmental variable name to value, overriding an existing value. If value is of the form $\$\{f 00-\mathrm{bar}\}$, the value is that of the environmental variable foo if that exists and is set to a non-empty value, otherwise bar. This construction can be nested, so bar can be of the same form (as in \$\{foo-\$\{barblah\}\}).

Leading and trailing white space in value are stripped. value is processed in a similar way to a Unix shell. In particular quotes are stripped, and backslashes are removed except inside quotes.

## Note

Prior to R version 1.4.0, the environment files searched were '.Renviron' in the current directory, the file pointed to by R_ENVIRON if set, and '. Renviron' in the user's home directory.

Prior to R version 1.2.1, '.Rprofile' was sourced after '.RData' was loaded, although the documented order was as here.

The format for site and user environment files was changed in version 1.2.0. Older files are quite likely to work but may generate warnings on startup if they contained unnecessary export statements.

Values in environment files were not processed prior to version 1.4.0.

## See Also

. Last for final actions before termination.

## Examples

```
# Example ~/.Renviron on Unix
R_LIBS=~/R/library
PAGER=/usr/local/bin/less
# Example .Renviron on Windows
R_LIBS=C:/R/library
TCL_LIBRARY=c:/packages/Tcl/lib/tcl8.3
# Example of .Rprofile
options(width=65, digits=5)
options(show.signif.stars=FALSE)
ps.options(horizontal=FALSE)
set.seed(1234)
    .First <- function() cat("\n Welcome to R!\n\n")
.Last <- function() cat("\n Goodbye!\n\n")
## if .Renviron contains
FOOBAR="coo\bar"doh\ex"abc\"def'"
## then we get
> cat(Sys.getenv("FOOBAR"), "\n")
coo\bardoh\exabc"def'
```

stat.anova GLM Anova Statistics

## Description

This is a utility function, used in lm and glm methods for anova(..., test != NULL) and should not be used by the average user.

## Usage

```
stat.anova(table, test = c("Chisq", "F", "Cp"), scale, df.scale, n)
```


## Arguments

| table | numeric matrix as results from anova.glm(..., test=NULL). |
| :--- | :--- |
| test | a character string, matching one of "Chisq", "F" or "Cp". |
| scale | a weighted residual sum of squares. |
| df.scale | degrees of freedom corresponding to scale. |
| n | number of observations. |

## Value

A matrix which is the original table, augmented by a column of test statistics, depending on the test argument.

## See Also

anova.lm, anova.glm.

## Examples

```
##-- Continued from ''?glm'':
print(ag <- anova(glm.D93))
stat.anova(ag$table, test = "Cp",
        scale = sum(resid(glm.D93, "pearson")^2)/4, df = 4, n = 9)
```

state States of the U.S.A.

## Description

Data sets related to the 50 states of the United States of America.

## Usage

data(state)

## Details

R currently contains the following "state" data sets. Note that all data are arranged according to alphabetical order of the state names.
state.abb: character vector of 2-letter abbreviations for the state names.
state.area: numeric vector of state areas (in square miles).
state.center: list with components named x and y giving the approximate geographic center of each state in negative longitude and latitude. Alaska and Hawaii are placed just off the West Coast.
state.division: factor giving state divisions (New England, Middle Atlantic, South Atlantic, East South Central, West South Central, East North Central, West North Central, Mountain, and Pacific).
state. name: character vector giving the full state names.
state.region: factor giving the region (Northeast, South, North Central, West) that each state belongs to.
state.x77: matrix with 50 rows and 8 columns giving the following statistics in the respective columns.
Population: population estimate as of July 1, 1975
Income: per capita income (1974)
Illiteracy: illiteracy (1970, percent of population)
Life Exp: life expectancy in years (1969-71)
Murder: murder and non-negligent manslaughter rate per 100,000 population (1976) HS Grad: percent high-school graduates (1970)
Frost: mean number of days with minimum temperature below freezing (1931-1960) in capital or large city
Area: land area in square miles

## Source

U.S. Department of Commerce, Bureau of the Census (1977) Statistical Abstract of the United States.
U.S. Department of Commerce, Bureau of the Census (1977) County and City Data Book.

## stem Stem-and-Leaf Plots

## Description

stem produces a stem-and-leaf plot of the values in $x$. The parameter scale can be used to expand the scale of the plot. A value of scale=2 will cause the plot to be roughly twice as long as the default.

## Usage

```
stem(x, scale = 1, width = 80, atom = 1e-08)
```


## Arguments

$\mathrm{x} \quad$ a numeric vector.
scale This controls the plot length.
width The desired width of plot.
atom a tolerance.

## Examples

```
data(islands)
stem(islands)
stem(log10(islands))
```


## step

Choose a model by AIC in a Stepwise Algorithm

## Description

Select a formula-based model by AIC.

## Usage

```
step(object, scope, scale = 0,
    direction = c("both", "backward", "forward"),
    trace = 1, keep = NULL, steps = 1000, k = 2, ...)
```


## Arguments

object an object representing a model of an appropriate class (mainly "lm" and "glm"). This is used as the initial model in the stepwise search.
scope defines the range of models examined in the stepwise search. This should be either a single formula, or a list containing components upper and lower, both formulae. See the details for how to specify the formulae and how they are used.
scale used in the definition of the AIC statistic for selecting the models, currently only for 1 m , aov and glm models.
direction the mode of stepwise search, can be one of "both", "backward", or "forward", with a default of "both". If the scope argument is missing the default for direction is "backward".
trace if positive, information is printed during the running of step. Larger values may give more detailed information.
keep a filter function whose input is a fitted model object and the associated AIC statistic, and whose output is arbitrary. Typically keep will select a subset of the components of the object and return them. The default is not to keep anything.
steps the maximum number of steps to be considered. The default is 1000 (essentially as many as required). It is typically used to stop the process early.
k
the multiple of the number of degrees of freedom used for the penalty. Only $\mathrm{k}=2$ gives the genuine AIC: $\mathrm{k}=\log (\mathrm{n})$ is sometimes referred to as BIC or SBC.
... any additional arguments to extractAIC.

## Details

step uses add1 and drop1 repeatedly; it will work for any method for which they work, and that is determined by having a valid method for extractAIC. When the additive constant can be chosen so that AIC is equal to Mallows' $C_{p}$, this is done and the tables are labelled appropriately.

The set of models searched is determined by the scope argument. The right-hand-side of its lower component is always included in the model, and right-hand-side of the model is included in the upper component. If scope is a single formula, it specifes the upper component, and the lower model is empty. If scope is missing, the initial model is used as the upper model.

Models specified by scope can be templates to update object as used by update.formula.
There is a potential problem in using glm fits with a variable scale, as in that case the deviance is not simply related to the maximized log-likelihood. The function extractAIC.glm makes the appropriate adjustment for a gaussian family, but may need to be amended for other cases. (The binomial and poisson families have fixed scale by default and do not correspond to a particular maximum-likelihood problem for variable scale.)

## Value

the stepwise-selected model is returned, with up to two additional components. There is an "anova" component corresponding to the steps taken in the search, as well as a "keep" component if the keep= argument was supplied in the call. The "Resid. Dev" column of the analysis of deviance table refers to a constant minus twice the maximized log likelihood: it will be a deviance only in cases where a saturated model is well-defined (thus excluding lm, aov and survreg fits, for example).

## Warning

The model fitting must apply the models to the same dataset. This may be a problem if there are missing values and R's default of na.action $=$ na.omit is used. We suggest you remove the missing values first.

## Note

This function differs considerably from the function in S , which uses a number of approximations and does not compute the correct AIC.
This is a minimal implementation. Use stepAIC for a wider range of object classes.

## Author(s)

B. D. Ripley

## See Also

stepAIC, add1, drop1

## Examples

```
example(lm)
step(lm.D9)
data(swiss)
summary(lm1 <- lm(Fertility ~ ., data = swiss))
slm1 <- step(lm1)
summary(slm1)
slm1$anova
```

stop Stop Function Execution

## Description

stop stops execution of the current expression, prints the message given as its argument, then executes an error action.
geterrmessage gives the last error message.

## Usage

stop(..., call. = TRUE)
geterrmessage()

## Arguments

... character vectors (which are pasted together with no separator) or NULL.
call. logical, indicating if the call should become part of the error message.

## Details

The error action is controlled by the current error handler set by options (error=). The default behaviour (the NULL error-handler) in interactive use is to return to the top level prompt, and in non-interactive use to (effectively) call q("no", status=1, runLast=FALSE).
Errors will be truncated to getOption("warning.length") characters, default 1000.

## Value

geterrmessage gives the last error message, as character string ending in " n".

## See Also

warning, try to catch errors and retry, and options for setting error handlers. stopifnot for validity testing.

## Examples

```
options(error = expression(NULL))# don't stop on stop(.) << Use with CARE! >>
iter <- 12
if(iter > 10) stop("too many iterations")
tst1 <- function(...) stop("dummy error")
tst1(1:10,long,calling, expression)
tst2 <- function(...) stop("dummy error", call. = FALSE)
tst2(1:10,long,calling,expression,but.not.seen.in.Error)
options(error = NULL)# revert to default
```

```
stopifnot Ensure the 'Truth' of R Expressions
```


## Description

If any of the expressions in . . . are not all TRUE, stop is called, producing an error message indicating the first element of ... which was not true.

## Usage

stopifnot(...)

## Arguments

... any number of (logical) R expressions which should evaluate to TRUE.

## Details

stopifnot(A, B) is conceptually equivalent to \{if(!all(A)) stop(...) ; if(!all(B)) stop(...) \}.

## Value

(NULL if all statements in . . . are TRUE.)

## See Also

stop, warning.

## Examples

```
stopifnot(1 == 1, all.equal(pi, 3.14159265), 1 < 2) # all TRUE
m <- matrix(c(1,3,3,1), 2,2)
stopifnot(m == t(m), diag(m) == rep(1,2)) # all(.) |=> TRUE
options(error = expression(NULL))# "disable stop(.)" << Use with CARE! >>
stopifnot(all.equal(pi, 3.141593), 2 < 2, all(1:10 < 12), "a" < "b")
stopifnot(all.equal(pi, 3.1415927), 2 < 2, all(1:10 < 12), "a" < "b")
options(error = NULL)# revert to default error handler
```

str Compactly Display the Structure of an Arbitrary $R$ Object

## Description

Compactly display the internal structure of an R object, a "diagnostic" function and an alternative to summary (and to some extent, dput). Ideally, only one line for each "basic" structure is displayed. It is especially well suited to compactly display the (abbreviated) contents of (possibly nested) lists. The idea is to give reasonable output for any R object. It calls args for (non-primitive) function objects.
ls.str and lsf.str are useful "versions" of ls, calling str on each object. They are not foolproof and should rather not be used for programming, but are provided for their usefulness.

## Usage

```
str(object, ...)
str.data.frame(object, ...)
str.default(object, max.level = 0, vec.len = 4, digits.d = 3,
    nchar.max = 128, give.attr = TRUE, give.length = TRUE,
    wid = getOption("width"), nest.lev = 0,
    indent.str = paste(rep(" ", max(0, nest.lev + 1)), collapse = ".."),
    ...)
    ls.str(pos = 1, pattern, ..., envir = as.environment(pos), mode = "any",
    max.level = 1, give.attr = FALSE)
lsf.str(pos = 1, ..., envir = as.environment(pos))
```


## Arguments

| object | any R object about which you want to have some information. |
| :--- | :--- |
| max.level | maximal level of nesting which is applied for displaying nested structures, <br> e.g., a list containing sub lists. Default 0: Display all nesting levels. |
| vec.len | numeric $(>=0)$ indicating how many "first few" elements are displayed <br> of each vector. The number is multiplied by different factors (from . 5 to |
| 3) depending on the kind of vector. Default 4. |  |
| digits.d | number of digits for numerical components (as for print). |


| nchar.max | maximal number of characters to show for character strings. Longer <br> strings are truncated, see longch example below. |
| :--- | :--- |
| give.attr | logical; if TRUE (default), show attributes as sub structures. <br> give.length <br> wid |
| logical; if TRUE (default), indicate length (as [1:...]). <br> the page width to be used. The default is the currently active <br> options("width"). |  |
| nest.lev | current nesting level in the recursive calls to str. |
| indent.str | the indentation string to use. |
| f. | potential further arguments (required for Method/Generic reasons). <br> pos <br> envir |
| integer indicating search path position. |  |
| pattern | environment to use, see ls. <br> regular expression passed to ls. Only names matching pattern are con- <br> sidered. |
| mode | character specifying the mode of objects to consider. Passed to exists <br> and get. |

## Value

str does not return anything, for efficiency reasons. The obvious side effect is output to the terminal.
ls.str and lsf.str invisibly return a character vector of the matching names, similarly to ls.

## Author(s)

Martin Maechler 〈maechler@stat.math.ethz.ch〉 since 1990.

## See Also

summary, args.

## Examples

```
## The following examples show some of 'str' capabilities
str(1:12)
str(ls)
str(args)#- more useful than args(args) !
data(freeny); str(freeny)
str(str)
str(.Machine, digits = 20)
str( lsfit(1:9,1:9))
str( lsfit(1:9,1:9), max =1)
op <- options(); str(op)#- save first; otherwise internal options() is used.
need.dev <- !exists(".Device") || is.null(.Device)
if(need.dev) postscript()
str(par()); if(need.dev) graphics.off()
nchar(longch <- paste(rep(letters,100), collapse=""))
str(longch)
str(longch, nchar.max = 52)
lsf.str()#- how do the functions look like which I am using?
```

```
ls.str(mode = "list")#- what are the structured objects I have defined?
## which base functions have "file" in their name ?
lsf.str(pos = length(search()), pattern = "file")
```

```
stripchart 1-D Scatter Plots
```


## Description

stripchart produces one dimensional scatter plots (or dot plots) of the given data. These plots are are good alternative to boxplots when sample sizes are small.

## Usage

```
stripchart(x, method="overplot", jitter=0.1, offset=1/3,
    vertical=FALSE, group.names,
    xlim=NULL, ylim=NULL, main="", ylab="", xlab="",
    pch=0, col=par("fg"), cex=par("cex"))
```


## Arguments

x
method the method to be used to separate coincident points. The default method "overplot" causes such points to be overplotted, but it is also possible to specify "jitter" to jitter the points, or "stack" have coincident points stacked. The last method only makes sense for very granular data.
jitter when jittering is used, jitter gives the amount of jittering applied.
offset when stacking is used, points are stacked this many line-heights (symbol widths) apart.
vertical when vertical is TRUE the plots are drawn vertically rather than the default horizontal.
group.names group labels which will be printed alongside (or underneath) each plot.
xlim, ylim, main, ylab, xlab, pch, col, cex
Graphical parameters.

## Details

Extensive examples of the use of this kind of plot can be found in Box, Hunter and Hunter or Seber and Wild.

## Examples

```
x <- round(rnorm(50), 1)
stripchart(x)
```

```
strptime Date-time Conversion Functions to and from Character
```


## Description

Functions to convert between character representations and objects of classes "POSIXlt" and "POSIXct" representing calendar dates and times.

## Usage

```
format.POSIXct(x, format = "", tz = "", usetz = FALSE, ...)
format.POSIXlt(x, format = "", usetz = FALSE, ...)
as.character(x, ...)
strftime(x, format="%Y-%m-%d %X", usetz = FALSE, ...)
strptime(x, format)
ISOdatetime(year, month, day, hour, min, sec, tz = "")
ISOdate(year, month, day, hour = 12, min = 0, sec = 0, tz = "GMT")
```


## Arguments

```
x An object to be converted.
tz A timezone specification to be used for the conversion. System-specific,
    but "" is the current time zone, and "GMT" is UTC.
format A character vector. The default is "%Y-%m-%d %H:%M:%S" if any com-
    ponent has a time component which is not midnight, and "%Y-%m-%d"
    otherwise.
... Further arguments to be passed from or to other methods.
usetz logical. Should the timezone be appended to the output? This is used
    in printing time, and as a workaround for problems with using "%Z" on
    most Linux systems.
year, month, day
    numerical values to specify a day.
hour, min, sec
    numerical values for a time within a day.
```


## Details

strftime is an alias for format.POSIXlt, and format.POSIXct first converts to class "POSIXct" by calling as.POSIXct. Note that only that conversion depends on the time zone.
The usual vector re-cycling rules are applied to x and format so the answer will be of length that of the longer of the vectors.

Locale-specific conversions to and from character strings are used where appropriate and available. This affects the names of the days and months, the AM/PM indicator (if used) and the separators in formats such as $\% \mathrm{x}$ and $\% \mathrm{X}$.

The details of the formats are system-specific, but the following are defined by the POSIX standard for strftime and are likely to be widely available. Any character in the format string other than the \% escapes is interpreted literally (and \%\% gives \%).
\%a Abbreviated weekday name.
\%A Full weekday name.
\%b Abbreviated month name.
\%B Full month name.
$\%$ c Date and time, locale-specific.
\%d Day of the month as decimal number (01-31).
$\%$ H Hours as decimal number (00-23).
\%I Hours as decimal number (01-12).
$\% j$ Day of year as decimal number (001-366).
\%m Month as decimal number (01-12).
\%M Minute as decimal number ( $00-59$ ).
$\%$ p AM/PM indicator in the locale. Used in conjuction with \%I and not with \%H.
$\%$ S Second as decimal number (00-61), allowing for up to two leap-seconds.
\%U Week of the year as decimal number ( $00-53$ ) using the first Sunday as day 1 of week 1.
\%w Weekday as decimal number ( $0-6$, Sunday is 0 ).
$\%$ Week of the year as decimal number $(00-53)$ using the first Monday as day 1 of week 1 .
$\%$ x Date, locale-specific.
\%X Time, locale-specific.
\%y Year without century (00-99). If you use this on input, which century you get is systemspecific. So don't! Often values up to 69 are prefixed by 20 and $70-99$ by 19 .
\%Y Year with century.
\%Z (output only.) Time zone as a character string (empty if not available). Note: do not use this on Linux unless the TZ environment variable is set.

Where leading zeros are shown they will be used on output but are optional on input.
ISOdatetime and ISOdate are convenience wrappers for strptime, that differ only in their defaults.

## Value

The format methods and strftime return character vectors representing the time. strptime turns character representations into an object of class "POSIXlt". ISOdatetime and ISOdate return an object of class "POSIXct".

## Note

The default formats follow the rules of the ISO 8601 international standard which expresses a day as "2001-02-03" and a time as "14:01:02" using leading zeroes as here. The ISO form uses no space to separate dates and times.
If the date string does not specify the date completely, the returned answer may be systemspecific. The most common behaviour is to assume that unspecified seconds, minutes or hours are zero, and a missing year, month or day is the current one.
If the timezone specified is invalid on your system, what happens is system-specific but it will probably be ignored.
OS facilities will probably not print years before 1CE (aka 1AD) correctly.

## References

International Organization for Standardization $(1988,1997, \ldots)$ ISO 8601. Data elements and interchange formats - Information interchange - Representation of dates and times. The 1997 version is available on-line at ftp://ftp.qsl.net/pub/g1smd/8601v03.pdf

## See Also

DateTimeClasses for details of the date-time classes; locales to query or set a locale.
Your system's help pages on strftime and strptime to see how to specify their formats.

## Examples

```
## locale-specific version of date()
format(Sys.time(), "%a %b %d %X %Y")
## we would include the timezone as in
## format(Sys.time(), "%a %b %d %X %Y %Z")
## but this crashes some Linux systems
## read in date info in format 'ddmmmyyyy'
## This will give NA(s) in some locales; setting the C locale
## as in the commented lines will overcome this on most systems.
## lct <- Sys.getlocale("LC_TIME"); Sys.setlocale("LC_TIME", "C")
x <- c("1jan1960", "2jan1960", "31mar1960", "30jul1960")
z <- strptime(x, "%d%%b%Y")
## Sys.setlocale("LC_TIME", lct)
z
## read in date/time info in format 'm/d/y h:m:s'
dates <- c("02/27/92", "02/27/92", "01/14/92",
    "02/28/92", "02/01/92")
times <- c("23:03:20", "22:29:56", "01:03:30",
    "18:21:03", "16:56:26")
x <- paste(dates, times)
z <- strptime(x, "%m/%d/%y %H:%M:%S")
z
```

    strsplit Split the Elements of a Character Vector
    
## Description

Split the elements of a character vector x into substrings according to the presence of substring split within them.

## Usage

```
strsplit(x, split, extended = TRUE)
```


## Arguments

$\mathrm{x} \quad$ character vector, to be split.
split character vector containing a regular expression to use as "split". If empty matches occur, in particular if split has length 0 , x is split into single characters. If split has length greater than 1 , it is re-cycled along x .
extended if TRUE, extended regular expression matching is used, and if FALSE basic regular expressions are used.

## Value

A list of length length ( x ) the i-th element of which contains the vector of splits of $\mathrm{x}[\mathrm{i}]$.

## See Also

paste for the reverse, grep and sub for string search and manipulation; further nchar, substr.

## Examples

```
noquote(strsplit("A text I want to display with spaces", NULL)[[1]])
x <- c("asfef", "qwerty", "yuiop[", "b", "stuff.blah.yech")
# split x on the letter e
strsplit(x,"e")
unlist(strsplit("a.b.c", "."))
## [1] "" "" "" "" ""
## Note that 'split' is a regexp!
## If you really want to split on '.', use
unlist(strsplit("a.b.c", "\\."))
## [1] "a" "b" "c"
## a useful function: rev() for strings
strReverse <- function(x)
    sapply(lapply(strsplit(x,NULL), rev), paste, collapse="")
strReverse(c("abc", "Statistics"))
a <- readLines(file.path(R.home(),"AUTHORS")) [-(1:8)]
a <- a[0:1-length(a)]
sub("\t.*","", a)
strReverse(sub(" .*","", a))
```

structure Attribute Specification

## Description

structure returns the given object with its attributes set.

## Usage

```
structure(.Data, ...)
```


## Arguments

| . Data an object which will have various attributes attached to it. |  |
| :--- | :--- |
| $\ldots$. | attributes, specified in tag=value form, which will be attached to data. |

## Examples

```
structure(1:6, dim = 2:3)
```

strwidth Plotting Dimensions of Character Strings and Math Expressions

## Description

These functions compute the width or height, respectively, of the given strings or mathematical expressions s[i] on the current plotting device in user coordinates, inches or as fraction of the figure width par("fin").

## Usage

```
strwidth(s, units = "user", cex = NULL)
strheight(s, units = "user", cex = NULL)
```


## Arguments

s character vector or expressions whose string widths in plotting units are to be determined. An attempt is made to coerce other vectors to character, and other language objects to expressions.
units character indicating in which units $s$ is measured; should be one of "user", "inches", "figure"; partial matching is performed.
cex character expansion to which is applies. By default, the current par("cex") is used.

## Value

Numeric vector with the same length as s, giving the width or height for each s[i]. NA strings are given width and height 0 (as they are not plotted).

## See Also

```
text, nchar
```


## Examples

```
str.ex <- c("W","w","I",".","WwI.")
op <- par(pty='s'); plot(1:100,1:100); par('usr')
sw <- strwidth(str.ex); sw
sum(sw[1:4] == sw[5])#- since the last string contains the others
sw / strwidth(str.ex, cex = .5)
# between 1.5 and 4.2 (!), font dependent
sw.i <- strwidth(str.ex, "inches"); 25.4 * sw.i # width in [mm]
unique(sw / sw.i)
```

```
# constant factor: 1 value
mean(sw.i / strwidth(str.ex, "fig")) / par('fin')[1] # = 1: are the same
## See how letters fall in classes -- depending on graphics device and font!
all.lett <- c(letters, LETTERS)
shL <- strheight(all.lett, units = "inches")
table(shL)# all have same heights ...
mean(shL) / par("cin")[2] # should be 1 (exactly?)
swL <- strwidth(all.lett)
swL <- 3 * swL / min(swL)
all(swL == round(swL))#- TRUE !
swL <- as.integer(swL)
n.classes <- length(tL <- table(swL)); tL
iL <- order(swL)
structure(swL[iL], names = all.lett[iL])
lett.classes <- structure(vector("list", n.classes), names= names(tL))
for(i in 1:n.classes)
    lett.classes[[i]] <- all.lett[swL == as.numeric(names(tL)[i])]
lett.classes
sumex <- expression(sum(x[i], i=1,n), e^{i * pi} == -1)
strwidth(sumex)
strheight(sumex)
rm(sumex); par(op)#- reset to previous setting
```

```
strwrap Wrap Character Strings to Format Paragraphs
```


## Description

Each character string in the input is first split into paragraphs (on lines containing whitespace only). The paragraphs are then formatted by breaking lines at word boundaries. The target columns for wrapping lines and the indentation of the first and all subsequent lines of a paragraph can be controlled independently.

## Usage

```
strwrap(x, width = 0.9 * getOption("width"), indent = 0, exdent = 0,
    prefix = "", simplify = TRUE)
```


## Arguments

| x | a character vector |
| :--- | :--- |
| width | a positive integer giving the target column for wrapping lines in the out- <br> put. |
| indent | a non-negative integer giving the indentation of the first line in a para- <br> graph. |
| exdent | a non-negative integer specifying the indentation of subsequent lines in <br> paragraphs. |
| prefix | a character string to be used as prefix for each line. |

simplify a logical. If TRUE, the result is a single character vector of line text; otherwise, it is a list of the same length as x the elements of which are character vectors of line text obtained from the corresponding element of x . (Hence, the result in the former case is obtained by unlisting that of the latter.)

## Details

Whitespace in the input is destroyed. Double spaces after periods (thought as representing sentence ends) are preserved. Currently, it possible sentence ends at line breaks are not considerd specially.

Indentation is relative to the number of characters in the prefix string.

## Examples

```
## Read in file 'THANKS'.
x <- paste(readLines(file.path(R.home(), "THANKS")), collapse = "\n")
## Split into paragraphs and remove the first three ones
x <- unlist(strsplit(x, "\n[ \t\n]*\n"))[-(1:3)]
## Join the rest
x <- paste(x, collapse = "\n\n")
## Now for some fun:
writeLines(strwrap(x, width = 60))
writeLines(strwrap(x, width = 60, indent = 5))
writeLines(strwrap(x, width = 60, exdent = 5))
writeLines(strwrap(x, prefix = "THANKS> "))
```

subset

Subsetting Vectors and Data Frames

## Description

Return subsets of vectors or data frames which meet conditions.

## Usage

```
subset(x, ...)
subset.default(x, subset, ...)
subset.data.frame(x, subset, select, ...)
```


## Arguments

| x | object to be subsetted |
| :--- | :--- |
| $\ldots$ | how to subset, depends on object |
| subset | logical expression |
| select | expression, indicating variables to select from a data frame |

## Details

For ordinary vectors, the result is simply $x[$ subset \& !is.na(subset)].
For dataframes, the subset argument works similarly on the rows. Note that subset will be evaluated in the dataframe.

The select argument exists only for dataframes. It works by first replacing variable names in the selection expression with the corresponding column numbers in the dataframe and then using the resulting integer vector to index the columns. This allows the use of the standard indexing conventions so that for examples ranges of variables can be specified easily.

## Value

Selected rows and columns of the object x .

## Author(s)

Peter Dalgaard

## See Also

[, transform

## Examples

```
data(airquality)
subset(airquality, Temp > 80, select = c(Ozone, Temp))
subset(airquality, Day == 1, select = -Temp)
subset(airquality, select = Ozone:Wind)
attach(airquality)
subset(Ozone, Temp > 80)
```

```
substitute Substituting and Quoting Expressions
```


## Description

substitute returns the parse tree for the (unevaluated) expression expr, substituting any variables bound in env.
quote simply returns its argument. The argument is not evaluated and can be any R expression.

## Usage

```
substitute(expr, env=<<see below>>)
quote(expr)
```


## Arguments

expr Any syntactically valid $R$ expression
env An environment or a list object. Defaults to the current evaluation environment.

## Details

The typical use of substitute is to create informative labels for data sets and plots. The myplot example below shows a simple use of this facility. It uses the functions deparse and substitute to create labels for a plot which are character string versions of the actual arguments to the function myplot.
Substitution takes place by examining each component of the parse tree as follows: If it is not a bound symbol in env, it is unchanged. If it is a promise object, i.e. a formal argument to a function or explicitly created using delay (), the expression slot of the promise replaces the symbol. If it is an ordinary variable, its value is substituted, unless env is .GlobalEnv in which case the symbol is left unchanged.

## Value

The mode of the result is generally "call" but may in principle be any type. In particular, single-variable expressions have mode "name" and constants have the appropriate base mode.

## Note

Substitute works on a purely lexical basis. There is no guarantee that the resulting expression makes any sense.
Substituting and quoting often causes confusion when the argument is expression(...). The result is a call to the expression constructor function and needs to be evaluated with eval to give the actual expression object.

## See Also

missing for argument "missingness".

## Examples

```
(s.e <- substitute(expression(a + b), list(a = 1))) #> expression(1 + b)
(s.s <- substitute( a + b, list(a = 1))) #> 1 + b
c(mode(s.e), typeof(s.e)) # "call", "language"
c(mode(s.s), typeof(s.s)) # (the same)
# but:
(e.s.e <- eval(s.e)) #> expression(1 + b)
c(mode(e.s.e), typeof(e.s.e)) # "expression", "expression"
substitute(x <- x + 1, list(x=1)) # nonsense
myplot <- function(x, y)
    plot(x, y, xlab=deparse(substitute(x)),
        ylab=deparse(substitute(y)))
## Simple examples about lazy evaluation, etc:
f1<- function(x, y = x) { x <- x + 1; y }
s1 <- function(x, y = substitute(x)) { x <- x + 1; y }
s2 <- function(x, y) { if(missing(y)) y <- substitute(x); x <- x + 1; y }
a <- 10
f1(a)# 11
s1(a)# 11
s2(a)# a
typeof(s2(a))# "symbol"
```

```
substr Substrings of a Character Vector
```


## Description

Extract or replace substrings in a character vector.

## Usage

```
substr(x, start, stop)
substring(text, first, last = 1000000)
substr(x, start, stop) <- value
substring(text, first, last = 1000000) <- value
```


## Arguments

| x, text | a character vector |
| :--- | :--- |
| start, first | integer. The first element to be replaced. |
| stop, last | integer. The last element to be replaced. |
| value | a character vector, recycled if necessary. |

## Details

substring is compatible with S, with first and last instead of start and stop. For vector arguments, it expands the arguments cyclically to the length of the longest.

When extracting, if start is larger than the string length then "" is returned.
For the replacement functions, if start is larger than the string length then no replacement is done. If the portion to be replaced is longer than the replacement string, then only the portion the length of the string is replaced.

## Value

For substr, a character vector of the same length as x .
For substring, a character vector of length the longest of the arguments.

## Note

The S 4 version of substring<- ignores last; this version does not.

## See Also

 strsplit, paste, nchar.
## Examples

```
substr("abcdef",2,4)
print(ss <- substring("abcdef",1:6,1:6))
stopifnot(ss == strsplit ("abcdef",NULL)[[1]])
## strsplit is more efficient ...
substr(rep("abcdef",4),1:4,4:5)
```

```
    x <- c("asfef", "qwerty", "yuiop[", "b", "stuff.blah.yech")
    stopifnot(substr(x, 2, 5) == substring(x, 2, 5))
    substr(x, 2, 5)
    substring(x, 2, 4:6)
    substring(x, 2) <- c("..", "+++")
    x
```

    sum Sum of Vector Elements
    
## Description

sum returns the sum of all the values present in its arguments. If na.rm is FALSE an NA value in any of the arguments will cause a value of NA to be returned, otherwise NA values are ignored.

## Usage

sum(..., na.rm=FALSE)

## Arguments

... numeric or complex vectors.
na.rm logical. Should missing values be removed?

## Value

The sum. If all of . . . are of type integer, then so is the sum, and in that case the result will be NA (with a warning) if integer overflow occurs.

NB: the sum of an empty set is zero, by definition.

```
summary
Object Summaries
```


## Description

summary is a generic function used to produce result summaries of the results of various model fitting functions. The function invokes particular methods which depend on the class of the first argument.

## Usage

```
summary(object, ...)
summary.default (object, ..., digits = max(3, getOption("digits")-3))
summary.data.frame(object, maxsum = 7,
    digits = max(3, getOption("digits")-3), ...)
    summary.factor (object, maxsum = 100, ...)
    summary.matrix (object, ...)
```


## Arguments

object an object for which a summary is desired.
maxsum integer, indicating how many levels should be shown for factors.
digits integer, used for number formatting with signif() (for summary.default) or format() (for summary.data.frame).
... additional arguments affecting the summary produced.

## Details

For factors, the frequency of the first maxsum - 1 most frequent levels is shown, where the less frequent levels are summarized in "(Others)" (resulting in maxsum frequencies).
The functions summary.lm and summary.glm are examples of particular methods which summarise the results produced by 1 m and glm .

## Value

The form of the value returned by summary depends on the class of its argument. See the documentation of the particular methods for details of what is produced by that method.

## See Also

anova, summary.glm, summary.lm.

## Examples

```
data(attenu)
summary(attenu, digits = 4) #-> summary.data.frame(...), default precision
summary(attenu $ station, maxsum = 20) #-> summary.factor(...)
```


## summary.manova Summary Method for Multivariate Analysis of Variance

## Description

A summary method for class "manova".

## Usage

```
summary (object,
test \(=\) c("Pillai", "Wilks", "Hotelling-Lawley", "Roy"),
intercept \(=\) FALSE, ....)
```


## Arguments

object An object of class "manova" or an aov object with multiple responses.
test The name of the test statistic to be used. Partial matching is used so the name can be abbreviated.
intercept logical. If TRUE, the intercept term is included in the table.
... further arguments passed to or from other methods.

## Details

The summary.manova method uses a multivariate test statistic for the summary table. Wilks' statistic is most popular in the literature, but the default Pillai-Bartlett statistic is recommended by Hand and Taylor (1987).

## Value

A list with components

SS A names list of sums of squares and product matrices.
Eigenvalues A matrix of eigenvalues,
stats A matrix of the statistics, approximate F value and degrees of freedom.

## Author(s)

B.D. Ripley

## References

Krzanowski, W. J. (1988) Principles of Multivariate Analysis. A User's Perspective. Oxford.

Hand, D. J. and Taylor, C. C. (1987) Multivariate Analysis of Variance and Repeated Measures. Chapman and Hall.

## See Also

 aov
## Examples

```
## Example on producing plastic filem from Krzanowski (1998, p. 381)
tear <- c(6.5, 6.2, 5.8, 6.5, 6.5, 6.9, 7.2, 6.9, 6.1, 6.3,
    6.7, 6.6, 7.2, 7.1, 6.8, 7.1, 7.0, 7.2, 7.5, 7.6)
gloss <- c(9.5, 9.9, 9.6, 9.6, 9.2, 9.1, 10.0, 9.9, 9.5, 9.4,
    9.1, 9.3, 8.3, 8.4, 8.5, 9.2, 8.8, 9.7, 10.1, 9.2)
opacity <- c(4.4, 6.4, 3.0, 4.1, 0.8, 5.7, 2.0, 3.9, 1.9, 5.7,
    2.8, 4.1, 3.8, 1.6, 3.4, 8.4, 5.2, 6.9, 2.7, 1.9)
Y <- cbind(tear, gloss, opacity)
rate <- factor(gl(2,10), labels=c("Low", "High"))
additive <- factor(gl(2, 5, len=20), labels=c("Low", "High"))
fit <- manova(Y ~ rate * additive)
summary.aov(fit) # univariate ANOVA tables
summary(fit, test="Wilks") # ANOVA table of Wilks' lambda
```

summaryRprof Summarise Output of $R$ Profiler

## Description

Summarise the output of the Rprof function to show the amount of time used by different $R$ functions.

## Usage

```
summaryRprof(filename = "Rprof.out", chunksize = 5000)
```


## Arguments

filename Name of a file produced by Rprof ()

## Details

This function is an alternative to $R$ CMD Rprof. It provides the convenience of an all-R implementation but will be slower for large files.
As the profiling output file could be larger than available memory, it is read in blocks of chunksize lines. Increasing chunksize will make the function run faster if sufficient memory is available.

## Value

A list with components

| by.self | Timings sorted by 'self' time |
| :--- | :--- |
| by.total | Timings sorted by 'total' time |
| sampling.time | Total length of profiling run |

## See Also

The chapter on "Tidying and profiling R code" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).
Rprof

## Examples

```
## Rprof() is not available on all platforms
Rprof(tmp <- tempfile())
example(glm)
Rprof()
summaryRprof(tmp)
unlink(tmp)
```

```
sunflowerplot Produce a Sunflower Scatter Plot
```


## Description

Multiple points are plotted as "sunflowers" with multiple leaves ("petals") such that overplotting is visualized instead of accidental and invisible.

## Usage

```
sunflowerplot(x, y = NULL, number, log = "", digits = 6,
    xlab = NULL, ylab = NULL, xlim = NULL, ylim = NULL,
    add = FALSE, rotate = FALSE,
    pch = 16, cex = 0.8, cex.fact = 1.5,
    size = 1/8, seg.col = 2, seg.lwd = 1.5, ...)
```


## Arguments

$\mathrm{x} \quad$ numeric vector of x -coordinates of length n , say, or another valid plotting structure, as for plot.default, see also xy. coords.
$y \quad$ numeric vector of y -coordinates of length n .
number integer vector of length n . number [i] $=$ number of replicates for ( $\mathrm{x}[\mathrm{i}], \mathrm{y}[\mathrm{i}]$ ), may be 0 .
Default: compute the exact multiplicity of the points $\mathrm{x}[\mathrm{]}, \mathrm{y}[]$.
$\log \quad$ character indicating log coordinate scale, see plot.default.
digits when number is computed (i.e., not specified), $x$ and $y$ are rounded to digits significant digits before multiplicities are computes.

xlim,ylim numeric(2) limiting the extents of the x -, or y -axis.
add logical; should the plot be added on a previous one? Default is FALSE.
rotate logical; if TRUE, randomly rotate the sunflowers (preventing artefacts).
pch plotting character to be used for points (number [i]==1) and center of sunflowers.
cex numeric; character size expansion of center points (s. pch).
cex.fact numeric shrinking factor to be used for the center points when there are flower leaves, i.e. cex / cex.fact is used for these.
size of sunflower leaves in inches, $1[\mathrm{in}]:=2.54[\mathrm{~cm}]$. Default: $1 / 8$; approximately 3.2 mm .
seg.col color to be used for the segments which make the sunflowers leaves, see $\operatorname{par}(\mathrm{col}=)$; col = "gold" reminds of real sunflowers.
seg.lwd numeric; the line width for the leaves' segments.
... further arguments to plot [if add=FALSE].

## Details

For number $[i]==1$, a (slightly enlarged) usual plotting symbol (pch) is drawn. For number [i] > 1, a small plotting symbol is drawn and number [i] equi-angular "rays" emanate from it.
If rotate=TRUE and number [i] >= 2, a random direction is chosen (instead of the $y$-axis) for the first ray. The goal is to jitter the orientations of the sunflowers in order to prevent artefactual visual impressions.

## Value

A list with three components of same length,

| $x$ | $x$ coordinates |
| :--- | :--- |
| $y$ | $y$ coordinates |
| number | number |

## Side Effects

A scatter plot is drawn with "sunflowers" as symbols.

## Author(s)

Andreas Ruckstuhl, Werner Stahel, Martin Maechler, Tim Hesterberg, 1989-1993. Port to R by Martin Maechler <maechler@stat.math.ethz.ch〉.

## References

Chambers, J. M., Cleveland, W. S., Kleiner, B. and Tukey, P. A. (1983) Graphical Methods for Data Analysis. Wadsworth.
Schilling, M. F. and Watkins, A. E. (1994) A suggestion for sunflower plots. The American Statistician, 48, 303-305.

## See Also

```
density
```


## Examples

```
data(iris)
## 'number' is computed automatically:
sunflowerplot(iris[, 3:4])
## Imitating Chambers et al., p.109, closely:
sunflowerplot(iris[, 3:4],cex=.2, cex.f=1, size=.035, seg.lwd=.8)
sunflowerplot(x=sort(2*round(rnorm(100))), y= round(rnorm(100),0),
    main = "Sunflower Plot of Rounded N(0,1)")
## A 'point process' {explicit 'number' argument}:
sunflowerplot(rnorm(100),rnorm(100), number=rpois(n=100,lambda=2),
    rotate=TRUE, main="Sunflower plot")
```


## Description

Monthly mean relative sunspot numbers from 1749 to 1983. Collected at Swiss Federal Observatory, Zurich until 1960, then Tokyo Astronomical Observatory.

## Usage

data(sunspots)

## Format

A time series of monthly data from 1749 to 1983.

## Source

Andrews, D. F. and Herzberg, A. M. (1985) Data: A Collection of Problems from Many Fields for the Student and Research Worker. New York: Springer-Verlag.

## See Also

sunspot.month (package ts) has a longer (and a bit different) series.

## Examples

```
data(sunspots)
plot(sunspots, main = "sunspots data", xlab = "Year",
    ylab = "Monthly sunspot numbers")
```

    svd Singular Value Decomposition of a Matrix
    
## Description

Compute the singular-value decomposition of a rectangular matrix.

## Usage

$\operatorname{svd}(\mathrm{x}, \mathrm{nu}=\min (\mathrm{n}, \mathrm{p}), \mathrm{nv}=\min (\mathrm{n}, \mathrm{p}))$
La.svd(x, nu $=\min (n, p), n v=\min (n, p), m e t h o d=c(" d g e s d d ", ~ " d g e s v d "))$

## Arguments

$\mathrm{x} \quad$ a matrix whose SVD decomposition is to be computed.
nu the number of left singular vectors to be computed. This must be one of 0 , $\operatorname{nrow}(x)$ and $n c o l(x)$, except for method $=$ "dgesdd".
nv the number of right singular vectors to be computed. This must be one of 0 and $\mathrm{ncol}(\mathrm{x})$.
method The LAPACK routine to use in the real case.

## Details

The singular value decomposition plays an important role in many statistical techniques.
svd provides an interface to the LINPACK routine DSVDC. La.svd provides an interface to the LAPACK routines DGESVD and DGESDD. The latter is usually substantially faster if singular vectors are required: see http://www.cs.berkeley.edu/~demmel/ DOE2000/Report0100.html. Most benefit is seen with an optimized BLAS system.
La.svd is preferred to svd for new projects, but it is not an exact replacement as it returns the transpose of the right singular vector matrix, and the signs of the singular vectors may differ from those given by svd. (They may also differ between methods and between platforms.)
Both functions handle complex matrices via LAPACK routine ZGESVD.
Computing the singular vectors is the slow part for large matrices.
Using method="dgesdd" requires IEEE 754 arithmetic. Should this not be supported on your platform, method="dgesvd" is used, with a warning.

## Value

The SVD decomposition of the matrix as computed by LINPACK,

$$
\boldsymbol{X}=\boldsymbol{U} \boldsymbol{D} \boldsymbol{V}^{\prime}
$$

where $\boldsymbol{U}$ and $\boldsymbol{V}$ are orthogonal, $\boldsymbol{V}^{\prime}$ means $V$ transposed, and $\boldsymbol{D}$ is a diagonal matrix with the singular values $D_{i i}$. Equivalently, $\boldsymbol{D}=\boldsymbol{U}^{\prime} \boldsymbol{X} \boldsymbol{V}$, which is verified in the examples, below. The components in the returned value correspond directly to the values returned by DSVDC.
$\mathrm{d} \quad$ a vector containing the singular values of x .
$u \quad a \quad$ matrix whose columns contain the left singular vectors of $x$.
$v \quad$ a matrix whose columns contain the right singular vectors of x .
For La.svd the return value replaces v by vt, the (conjugated if complex) transpose of v .

## References

Dongarra, J. J., Bunch, J. R., Moler, C. B. and Stewart, G. W. (1978) LINPACK Users Guide. Philadelphia: SIAM Publications.
Anderson. E. and ten others (1999) LAPACK Users' Guide. Third Edition. SIAM.
Available on-line at http://www.netlib.org/lapack/lug/lapack_lug.html.

## See Also

eigen, qr.
capabilities to test for IEEE 754 arithmetic.

## Examples

```
hilbert <- function(n) { i <- 1:n; 1 / outer(i - 1, i, "+") }
str(X <- hilbert(9)[,1:6])
str(s <- svd(X))
Eps <- 100 * .Machine$double.eps
D <- diag(s$d)
```

```
stopifnot (abs(X - s\$u \%*\% D \% *\% t(s\$v)) < Eps) \# X = U D V'
stopifnot(abs(D - t(s\$u) \%*\% X \% \% \% s\$v) < Eps) \# D = U' X V
\(\mathrm{X}<-\operatorname{cbind}(1,1: 7)\)
str (s <- svd(X)) ; D <- diag(s\$d)
stopifnot(abs(X - s\$u \%*\% D \%*\% t(s\$v)) < Eps) \# X = U D V'
stopifnot (abs(D - t(s\$u) \(\% * \%\) X \(\% * \%\) s\$v) < Eps) \# D = U' X V
```


## sweep Sweep out Array Summaries

## Description

Return an array obtained from an input array by sweeping out a summary statistic.

## Usage

sweep(x, MARGIN, STATS, FUN="-", ...)

## Arguments

x
MARGIN
STATS
FUN
... optional arguments to FUN.

## Value

An array with the same shape as x , but with the summary statistics swept out.

## See Also

apply on which sweep is based; scale for centering and scaling.

## Examples

```
data(attitude)
med.att <- apply(attitude, 2, median)
sweep(data.matrix(attitude), 2, med.att)# subtract the column medians
```


## Description

Standardized fertility measure and socio-economic indicators for each of 47 French-speaking provinces of Switzerland at about 1888.

## Usage

data(swiss)

## Format

A data frame with 47 observations on 6 variables, each of which is in percent, i.e., in $[0,100]$.

| $[, 1]$ | Fertility | $I_{g}$, "common standardized fertility measure" |
| :--- | :--- | :--- |
| $[, 2]$ | Agriculture | \% of males involved in agriculture as occupation |
| $[, 3]$ | Examination | \% "draftees" receiving highest mark on army examination |
| $[, 4]$ | Education | \% education beyond primary school for "draftees". |
| $[, 5]$ | Catholic | \% catholic (as opposed to "protestant"). |
| $[, 6]$ | Infant.Mortality | live births who live less than 1 year. |

All variables but 'Fertility' give proportions of the population.

## Details

(paraphrasing Mosteller and Tukey):
Switzerland, in 1888, was entering a period known as the "demographic transition"; i.e., its fertility was beginning to fall from the high level typical of underdeveloped countries.
The data collected are for 47 French-speaking "provinces" at about 1888.
Here, all variables are scaled to $[0,100]$, where in the original, all but "Catholic" were scaled to $[0,1]$.

## Note

Files for all 182 districts in 1888 and other years are available at http://opr.princeton. edu/archive/eufert/switz.html.
They state that variables Examination and Education are averages for 1887, 1888 and 1889.

## Source

Project "16P5", pages 549-551 in
Mosteller, F. and Tukey, J. W. (1977) Data Analysis and Regression: A Second Course in Statistics. Addison-Wesley, Reading Mass.
indicating their source as "Data used by permission of Franice van de Walle. Office of Population Research, Princeton University, 1976. Unpublished data assembled under NICHD contract number No 1-HD-O-2077."

## Examples

```
data(swiss)
pairs(swiss, panel = panel.smooth, main = "swiss data",
    col = 3 + (swiss$Catholic > 50))
summary(lm(Fertility ~ . , data = swiss))
```

switch Select One of a List of Alternatives

## Description

switch evaluates EXPR and accordingly chooses one of the further arguments (in ...).

## Usage

```
switch(EXPR, ...)
```


## Arguments

EXPR an expression evaluating to a number or a character string.
... the list of alternatives, given explicitly.

## Details

If the value of EXPR is an integer between 1 and nargs()-1 then the corresponding element of . . . is evaluated and the result returned.
If EXPR returns a character string then that string is used to match the names of the elements in .... If there is an exact match then that element is evaluated and returned if there is one, otherwise the next element is chosen, e.g., switch("cc", $a=1, c c=, d=2$ ) evaluates to 2.
In the case of no match, if there's a further argument in switch that one is returned, otherwise NULL.

## Warning

Beware of partial matching: an alternative $E=f o o$ will match the first argument EXPR unless that is named. See the examples for good practice in naming the first argument.

## Examples

```
centre <- function(x, type) {
    switch(type,
        mean = mean(x),
        median = median(x),
        trimmed = mean(x, trim = .1))
}
x <- rcauchy(10)
centre(x, "mean")
centre(x, "median")
centre(x, "trimmed")
ccc <- c("b","QQ","a","A","bb")
```

```
for(ch in ccc) cat(ch,":",switch(EXPR = ch, a=1, b=2:3), "\n")
for(ch in ccc) cat(ch,":",switch(EXPR = ch, a=,A=1, b=2:3, "Otherwise: last"),"\n")
## Numeric EXPR don't allow an 'otherwise':
for(i in c(-1:3,9)) print(switch(i, 1,2,3,4))
```

symbols
Draw symbols on a plot

## Description

This function draws symbols on a plot. One of six symbols; circles, squares, rectangles, stars, thermometers, and boxplots, can be plotted at a specified set of x and y coordinates. Specific aspects of the symbols, such as relative size, can be customized by additional parameters.

## Usage

```
symbols(x, y, circles, squares, rectangles, stars,
    thermometers, boxplots, inches=TRUE, add=FALSE,
    fg=1, bg=NA, xlab = NULL, ylab = NULL, main = NULL,
    xlim = NULL, ylim = NULL, ...)
```


## Arguments

|  | a |
| :---: | :---: |
| y | a vector giving the y coordinates of the symbols. |
| circles | a vector giving the radii of the circles. |
| squares | a vector giving the length of the sides of the squares. |
| rectangles | a matrix with two columns. The first column gives widths and the second the heights of rectangle symbols. |
| stars | a matrix with three or more columns giving the lengths of the rays from the center of the stars. NA values are replaced by zeroes. |
| thermometers | a matrix with three or four columns. The first two columns give the width and height of the thermometer symbols. If there are three columns, the third is taken as a proportion. The thermometers are filled from their base to this proportion of their height. If there are four columns, the third and fourth columns are taken as proportions. The thermometers are filled between these two proportions of their heights. |
| boxplots | a matrix with five columns. The first two columns give the width and height of the boxes, the next two columns give the lengths of the lower and upper whiskers and the fifth the proportion (with a warning if not in $[0,1]$ ) of the way up the box that the median line is drawn. |
| inches | If inches is FALSE, the units are taken to be those of the x axis. If inches is TRUE, the symbols are scaled so that the largest symbol is one inch in height. If a number is given the symbols are scaled to make largest symbol this height in inches. |
| add | if add is TRUE, the symbols are added to an existing plot, otherwise a new plot is created. |


| fg | colors the symbols are to be drawn in (the default is the value of the col <br> graphics parameter). <br> if specified, the symbols are filled with this color. The default is to leave <br> the symbols unflled. |
| :--- | :--- |
| bg | the x label of the plot if add is not true; this applies to the following <br> arguments as well. Defaults to the deparsed expression used for x. |
| xlab | the y label of the plot. |
| ylab | a main title for the plot. |
| main | numeric of length 2 giving the x limits for the plot. |
| ylim | numeric of length 2 giving the y limits for the plot. |
| $\ldots$ | graphics parameters can also be passed to this function. |

## Details

Observations which have missing coordinates or missing size parameters are not plotted. The exception to this is stars. In that case, the length of any rays which are NA is reset to zero.

Circles of radius zero are plotted at radius one pixel (which is device-dependent).

## References

W. S. Cleveland (1985) The Elements of Graphing Data. Monterey, California: Wadsworth.

## See Also

 stars for drawing stars with a bit more flexibility; sunflowerplot.
## Examples

```
x <- 1:10
y <- sort(10*runif(10))
z <- runif(10)
z3 <- cbind(z, 2*runif(10), runif(10))
symbols(x, y, thermometers=cbind(.5, 1, z), inches=.5, fg = 1:10)
symbols(x, y, thermometers = z3, inches=FALSE)
text(x,y, apply(format(round(z3, dig=2)), 1, paste, collapse = ","),
    adj = c(-.2,0), cex = .75, col = "purple", xpd=NA)
data(trees)
## Note that example(trees) shows more sensible plots!
N <- nrow(trees)
attach(trees)
## Girth is diameter in inches
symbols(Height, Volume, circles=Girth/24, inches=FALSE,
    main="Trees' Girth")# xlab and ylab automatically
## Colors too:
palette(rainbow(N, end = 0.9))
symbols(Height, Volume, circles=Girth/16, inches=FALSE, bg = 1:N,
    fg="gray30", main="symbols(*, circles=Girth/16, bg = 1:N)")
palette("default"); detach()
```

symnum
Symbolic Number Coding

## Description

Symbolically encode a given numeric or logical vector or array.

## Usage

symnum(x, cutpoints=c(0.3, 0.6, 0.8, 0.9, 0.95),
symbols=c(" ", ".", ",", "+", "*", "B"),
legend $=$ length (symbols) >= 3 ,
na="?", eps=1e-5,
corr = missing(cutpoints), show.max = if(corr) "1", show.min = NULL,
lower.triangular $=$ corr \& is.matrix(x),
diag.lower.tri $=$ corr \& !is.null(show.max))

## Arguments

x
cutpoints
symbols character vector, one shorter than (the augmented, see corr below) cutpoints. symbols $[\mathrm{j}]=s_{j}$ are used as "code" for the (half open) inter$\operatorname{val}\left(c_{j}, c_{j+1}\right]$.
For logical argument $x$, the default is $c(" . ",| | ")$ (graphical $0 / 1 \mathrm{~s}$ ).
legend logical indicating if a "legend" attribute is desired.
na character or logical. How NAs are coded. If na == FALSE, NAs are coded invisibly, including the "legend" attribute below, which otherwise mentions NA coding.
eps absolute precision to be used at left and right boundary.
corr logical. If TRUE, $x$ contains correlations. The cutpoints are augmented by 0 and 1 and abs ( x ) is coded.
show.max If TRUE, or of mode character, the maximal cutpoint is coded especially.
show.min If TRUE, or of mode character, the minmal cutpoint is coded especially.
lower.triangular logical. If TRUE and x is a matrix, only the lower triangular part of the matrix is coded as non-blank.
diag.lower.tri
logical. If lower.triangular and this are TRUE, the diagonal part of the matrix is shown.

## Value

An atomic character object of class noquote and the same dimensions as x .
If legend (TRUE by default when there more than 2 classes), it has an attribute "legend" containing a legend of the returned character codes, in the form

$$
c_{1} s_{1} c_{2} s_{2} \ldots s_{n} c_{n+1}
$$

where $c_{j}=$ cutpoints[j] and $s_{j}=$ symbols[j].

## Author(s)

Martin Maechler 〈maechler@stat.math.ethz.ch〉

## See Also

as.character

## Examples

```
ii <- 0:8; names(ii) <- ii
symnum(ii, cut= 2*(0:4), sym = c(".", "-", "+", "$"))
symnum(ii, cut= 2*(0:4), sym = c(".", "-", "+", "$"), show.max=TRUE)
symnum(1:12 %% 3 == 0)# use for logical
##-- Symbolic correlation matrices:
data(attitude)
symnum(cor(attitude), diag = FALSE)
symnum(cor(rbind(1, rnorm(25), rnorm(25)^2)))
symnum(cor(matrix(rexp(30, 1), 5, 18))) # <<-- PATTERN ! --
symnum(cm1 <- cor(matrix(rnorm(90) , 5, 18))) # < White Noise SMALL n
symnum(cm1, diag=FALSE)
symnum(cm2 <- cor(matrix(rnorm(900), 50, 18))) # < White Noise "BIG" n
symnum(cm2, lower=FALSE)
## NA's:
Cm <- cor(matrix(rnorm(60), 10, 6)); Cm[c(3,6), 2] <- NA
symnum(Cm, show.max=NULL)
## Graphical P-values (aka "significance stars"):
pval <- rev(sort(c(outer(1:6, 10^-(1:3)))))
symp <- symnum(pval, corr=FALSE,
    cutpoints = c(0, .001,.01,.05, .1, 1),
    symbols = c("***","**","*","."," "))
noquote(cbind(P.val = format(pval), Signif= symp))
```

Syntax Operator Syntax

## Description

Outlines R syntax and gives the precedence of operators

## Details

The following unary and binary operators are defined. They are listed in precedence groups, from highest to lowest.

| $[$ [ $[$ | indexing |
| :--- | :--- |
| $::$ | name space/variable name separator |
| \$ @ | component / slot extraction |
| - | exponentiation (right to left) |
| -+ | unary minus and plus |


| : | sequence operator |
| :---: | :---: |
| \%any\% | special operators |
| * / | multiply, divide |
| + - | (binary) add, subtract |
| < > <= >= == != | ordering and comparison |
| ! | negation |
| \& \&\& | and |
| 1 \|| | or |
| ~ | as in formulae |
| -> ->> | rightwards assignment |
| = | assignment (right to left) |
| <- _ <<- | assignment (right to left) |
| ? | help (unary and binary) |

Within an expression operators of equal precedence are evaluated from left to right except where indicated.

The links in the See Also section covers most other aspects of the basic syntax.

## Note

There are substantial precedence differences between $R$ and $S$. In particular, in $S$ ? has the same precedence as + - and \& \&\& | \| have equal precedence.

## See Also

Arithmetic, Comparison, Control, Extract, Logic, Paren
The $R$ Language Definition manual.

Sys.getenv
Get Environment Variables

## Description

Sys.getenv obtains the values of the environment variables named by x .

## Usage

Sys.getenv(x)

## Arguments

x
a character vector, or missing

## Value

A vector of the same length as x , with the variable names as its names attribute. Each element holds the value of the environment variable named by the corresponding component of x (or " " if no environment variable with that name was found).

On most platforms Sys.getenv() will return a named vector giving the values of all the environment variables.

## See Also

Sys.putenv, getwd for the working directory.

## Examples

```
Sys.getenv(c("R_HOME", "R_PAPERSIZE", "R_PRINTCMD", "HOST"))
```

```
Sys.info Extract System and User Information
```


## Description

Reports system and user information.

## Usage

Sys.info()

## Details

This function is not implemented on all R platforms, and returns NULL when not available. Where possible it is based on POSIX system calls.
Sys.info() returns details of the platform $R$ is running on, whereas R.version gives details of the platform R was built on: they may well be different.

## Value

A character vector with fields
sysname The operating system.
release The OS release.
version The OS version.
nodename A name by which the machine is known on the network (if any).
machine A concise description of the hardware.
login The user's login name, or "unknown" if it cannot be ascertained.
user $\quad$ The name of the real user ID, or "unknown" if it cannot be ascertained.
The first five fields come from the uname(2) system call. The login name comes from getlogin(2), and the user name from getpwuid(getuid())

## Note

The meaning of OS "release" and "version" is highly system-dependent and there is no guarantee that the node or login or user names will be what you might reasonably expect. (In particular on some Linux distributions the login name is unknown from sessions with re-directed inputs.)

Author(s)
B. D. Ripley

## See Also

.Platform, and R.version.

## Examples

```
Sys.info()
## An alternative (and probably better) way to get the login name on Unix
Sys.getenv("LOGNAME")
```

sys.parent Functions to Access the Function Call Stack

## Description

These functions provide access to environments ("frames" in S terminology) associated with functions further up the calling stack.

## Usage

```
sys.call(which = 0)
sys.frame(which \(=0\) )
sys.nframe()
sys.function( \(\mathrm{n}=0\) )
sys.parent(n = 1)
sys.calls()
sys.frames()
sys.parents()
sys.on.exit()
sys.status()
parent.frame(n = 1)
```


## Arguments

which the frame number if non-negative, the number of generations to go back if negative. (See the Details section.)
n the number of frame generations to go back.

## Details

.GlobalEnv is given number 0 in the list of frames. Each subsequent function evaluation increases the frame stack by 1 and the environment for evaluation of that function is returned by sys.frame with the appropriate index.
The parent of a function evaluation is the environment in which the function was called. It is not necessarily numbered one less than the frame number of the current evaluation, nor is it the environment within which the function was defined. sys.parent returns the number of the parent frame if n is 1 (the default), the grandparent if n is 2 , and so on. sys.frame returns the environment associated with a given frame number.
sys.call and sys.frame both accept integer values for the argument which. Non-negative values of which are normal frame numbers whereas negative values are counted back from the frame number of the current evaluation.
sys.nframe returns the number of the current frame in that list. sys.function gives the definition of the function curently being evaluated in the frame n generations back.
sys.frames gives a list of all the active frames and sys.parents gives the indices of the parent frames of each of the frames.
Notice that even though the sys. $x x x$ functions (except sys.status) are interpreted, their contexts are not counted nor are they reported. There is no access to them.
sys.status() returns a list with components sys.calls, sys.parents and sys.frames.
sys.on.exit() retrieves the expression stored for use by on.exit in the function currently being evaluated. (Note that this differs from S, which returns a list of expressions for the current frame and its parents.)
parent.frame( $n$ ) is a convenient shorthand for sys.frame(sys.parent(n)) (implemented slightly more efficiently).

## See Also

eval for the usage of sys.frame and parent.frame.

## Examples

```
ff <- function(x) gg(x)
gg <- function(y) sys.status()
str(ff(1))
gg <- function(y) {
    ggg <- function() {
            cat("current frame is", sys.nframe(), "\n")
            cat("parents are", sys.parents(), "\n")
            print(sys.function(0)) # ggg
            print(sys.function(2)) # gg
    }
    if(y > 0) gg(y-1) else ggg()
}
gg(3)
t1 <- function() {
    aa <- "here"
    t2 <- function() {
        ## in frame 2 here
        cat("current frame is", sys.nframe(), "\n")
        str(sys.calls()) ## list with two components t1() and t2()
        cat("parents are frame nos", sys.parents(), "\n") ## 0 1
        print(ls(envir=sys.frame(-1))) ## [1] "aa" "t2"
        invisible()
    }
    t2()
}
t1()
test.sys.on.exit <- function() {
    on.exit(print(1))
    ex <- sys.on.exit()
    str(ex)
    cat("exiting...\n")
}
```

```
test.sys.on.exit()
## gives 'language print(1)', prints 1 on exit
```

```
Sys.putenv Set Environment Variables
```


## Description

putenv sets environment variables (for other processes called from within R or future calls to Sys.getenv from this R process).

## Usage

Sys.putenv(...)

## Arguments

$\ldots \quad$ arguments in name=value form, with value coercible to a character string.

## Details

Non-standard R names must be quoted: see the Examples section.

## Value

A logical vector of the same length as x , with elements being true if setting the corresponding variable succeeded.

## Note

Not all systems need support Sys.putenv.

## See Also

Sys.getenv, setwd for the working directory.

## Examples

```
print(Sys.putenv("R_TEST"="testit", ABC=123))
Sys.getenv("R_TEST")
```

Sys.sleep Suspend Execution for a Time Interval

## Description

Suspend execution of R expressions for a given number of seconds

## Usage

```
Sys.sleep(time)
```


## Arguments

time The time interval to suspend execution for, in seconds.

## Details

Using this function allows R to be given very low priority and hence not to interfere with more important foreground tasks. A typical use is to allow a process lauched from R to set itself up and read its input files before R execution is resumed.

The intention is that this function suspends execution of $R$ expressions but wakes the process up often enough to respond to GUI events, typically every 0.5 seconds.

There is no guarantee that the process will sleep for the whole of the specified interval, and it may well take slightly longer in real time to resume execution. The resolution of the time interval is system-dependent, but will normally be down to 0.02 secs or better.

## Value

Invisible NULL.

## Note

This function is not implemented on all systems.

## Author(s)

B. D. Ripley

## Examples

```
testit <- function(x)
{
    p1 <- proc.time()
    Sys.sleep(x)
    proc.time() - p1 # The cpu usage should be negligible
}
testit(3.7)
```


## Description

Parses expressions in the given file, and then successively evaluates them in the specified environment.

## Usage

```
sys.source(file, envir = NULL, chdir = FALSE,
    keep.source = getOption("keep.source.pkgs"))
```


## Arguments

| file | a character string naming the file to be read from |
| :--- | :--- |
| envir | an $R$ object specifying the environment in which the expressions are to <br> be evaluated. May also be a list or an integer. The default value NULL <br> corresponds to evaluation in the base environment. This is probably not <br> what you want; you should typically supply an explicit envir argument. |
| chdir | logical; if TRUE, the R working directory is changed to the directory con- <br> taining file for evaluating. |
| keep.source | logical. If TRUE, functions "keep their source" including comments, see <br> options (keep.source $=*)$ for more details. |

## Details

For large files, keep.source = FALSE may save quite a bit of memory.

## See Also

source, and library which uses sys.source.
Sys.time Get Current Time and Timezone

## Description

Sys.time returns the system's idea of the current time and Sys.timezone returns the current time zone.

## Usage

Sys.time()
Sys.timezone()

## Value

Sys.time returns an object of class "POSIXct" (see DateTimeClasses).
Sys.timezone returns an OS-specific character string, possibly an empty string.

## See Also

date for the system time in a fixed-format character string.

## Examples

```
Sys.time()
## locale-specific version of date()
format(Sys.time(), "%a %b %d %X %Y")
Sys.timezone()
```

```
system Invoke a System Command
```


## Description

system invokes the OS command specified by command.

## Usage

system(command, intern = FALSE, ignore.stderr = FALSE)

## Arguments

command the system command to be invoked, as a string.
intern a logical, indicates whether to make the output of the command an $R$ object.
ignore.stderr a logical indicating whether error messages (written to 'stderr') should be ignored.

## Details

If intern is TRUE then popen is used to invoke the command and the output collected, line by line, into an $R$ character vector which is returned as the value of system.

If intern is FALSE then the $C$ function system is used to invoke the command and the value returned by system is the exit status of this function.
unix is a deprecated alternative, available for backwards compatibility.

## Value

If intern=TRUE, a character vector giving the output of the command, one line per character string. If the command could not be run or gives an error a R error is generated.

If intern=FALSE, the return value is an error code.

## See Also

.Platform for platform specific variables.

## Examples

```
# list all files in the current directory using the -F flag
system("ls -F")
# t1 is a character vector, each one
# representing a separate line of output from who
t1 <- system("who", TRUE)
system("ls fizzlipuzzli", TRUE, TRUE)# empty since file doesn't exist
```

```
system.file Find Names of R System Files
```


## Description

Finds the full file names of files in packages etc.

## Usage

system.file(..., package = "base", lib.loc = NULL, pkg, lib)

## Arguments

| $\ldots$. | character strings, specifying subdirectory and file(s) within some package. <br> The default, none, returns the root of the package. Wildcards are not <br> supported. |
| :--- | :--- |
| package | a character string with the name of a single package. An error occurs if <br> more than one package name is given. |
| lib.loc | a character vector with path names of R libraries, or NULL. The default <br> value of NULL corresponds to all libraries currently known. If the default <br> is used, the loaded packages are searched before the libraries. |
| pkg | previous name for argument package. Deprecated. |
| lib | previous name for argument lib.loc. Deprecated. |

## Value

A character vector of positive length, containing the file names that matched ..., or the empty string, " ", if none matched. If matching the root of a package, there is no trailing separator.

As a special case, system.file() gives the root of the base package only.

## See Also

```
list.files
```


## Examples

```
system.file() # The root of the 'base' package
system.file(package = "lqs") # The root of package 'lqs'
system.file("INDEX")
system.file("help", "AnIndex", package = "stepfun")
```

```
system.time CPU Time Used
```


## Description

Return CPU (and other) times that expr used.

## Usage

system.time(expr)
unix.time (expr)

## Arguments

expr Valid R expression to be "timed"

## Details

system.time calls the builtin proc.time, evaluates expr, and then calls proc.time once more, returning the difference between the two proc.time calls.
The values returned by the proc.time are (on Unix) those returned by the C library function times $(3 \mathrm{v})$, if available.
unix.time is an alias of system.time, for compatibility reasons.

## Value

A numeric vector of length 5 containing the user cpu, system cpu, elapsed, subproc1, subproc2 times. The subproc times are the user and system cpu time used by child processes (and so are usually zero).
The resolution of the times will be system-specific; it is common for them to be recorded to of the order of $1 / 100$ second, and elapsed time is rounded to the nearest $1 / 100$.

## Note

It is possible to compile R without support for system.time, when all the values will be NA.

## See Also

proc.time, time which is for time series.

## Examples

```
system.time(for(i in 1:50) mad(runif(500)))
exT <- function(n = 100) {
    # Purpose: Test if system.time works ok; n: loop size
    system.time(for(i in 1:n) x <- mean(rt(1000, df=4)))
}
#-- Try to interrupt one of the following (using Ctrl-C):
exT() #- '1.4' on -0-optimized Ultra1
system.time(exT()) #~ +/- same
```

t
Matrix Transpose

## Description

Given a matrix or data.frame x , t returns the transpose of x .

## Usage

$t(x)$

## Arguments

x
a matrix or data frame, typically.

## Details

A data frame is first coerced to a matrix: see as.matrix. When $x$ is a vector, it is treated as "column", i.e., the result is a 1 -row matrix.

## See Also

aperm for permuting the dimensions of arrays.

## Examples

```
a <- matrix(1:30, 5,6)
ta <- t(a) ##-- i.e., a[i, j] == ta[j, i] for all i,j :
for(j in seq(ncol(a)))
    if(! a[, j] == ta[j, ]) stop("wrong transpose")
```

table Cross Tabulation

## Description

table uses the cross-classifying factors to build a contingency table of the counts at each combination of factor levels.

## Usage

```
table(..., exclude = c(NA, NaN), dnn, deparse.level = 1)
as.table(x, ...)
is.table(x)
as.data.frame.table(x, row.names = NULL, optional = FALSE, ...)
```


## Arguments

```
... objects which can be interpreted as factors (including character strings),
    or a list (or data frame) whose components can be so interpreted
exclude values to use in the exclude argument of factor when interpreting non-
    factor objects
dnn the names to be given to the dimensions in the result ('the dimname
    names').
deparse.level controls how the default dnn is constructed. See details.
x an arbitrary R object.
row.names a character vector giving the row names for the data frame.
optional a logical controlling whether row names are set. Currently not used
```


## Details

If the argument dnn is not supplied, the internal function list.names is called to compute the 'dimname names'. If the arguments in ... are named, those names are used. For the remaining arguments, deparse.level = 0 gives an empty name, deparse.level = 1 uses the supplied argument if it is a symbol, and deparse.level $=2$ will deparse the argument.
There is a summary method for contingency table objects created by table or xtabs, which gives basic information and performs a chi-squared test for independence of factors (note that the function chisq.test in package ctest currently only handles 2-d tables).
as.table and is.table coerce to and test for contingency table, respectively.
as.data.frame.table is a method for the generic function as.data.frame to convert the array-based representation of a contingency table to a data frame containing the classifying factors and the corresponding counts (the latter as component Freq). This is the inverse of xtabs.

## Examples

```
## Simple frequency distribution
table(rpois(100,5))
data(warpbreaks)
attach(warpbreaks)
## Check the design:
table(wool, tension)
data(state)
table(state.division, state.region)
data(airquality)
attach(airquality)
# simple two-way contingency table
table(cut(Temp, quantile(Temp)), Month)
a <- letters[1:3]
table(a, sample(a)) # dnn is c("a", "")
table(a, sample(a), deparse.level = 0) # dnn is c("", "")
table(a, sample(a), deparse.level = 2) # dnn is c("a", "sample(a)")
## xtabs() <-> as.data.frame.table() :
data(UCBAdmissions) ## already a contingency table
DF <- as.data.frame(UCBAdmissions)
class(tab <- xtabs(Freq ~ ., DF))# xtabs & table
```

```
## tab *is* ''the same') as the original table:
all(tab == UCBAdmissions)
all.equal(dimnames(tab), dimnames(UCBAdmissions))
```

tabulate Tabulation for Vectors

## Description

tabulate takes the integer valued vector bin and counts the number of times each integer occurs in it. tabulate is used as the basis of the table function.

## Usage

tabulate(bin, nbins $=\max (1$, bin))

## Arguments

| bin | a vector of integers, or a factor. |
| :--- | :--- |
| nbins | the number of bins to be used. |

## Details

If bin is a factor, its internal integer representation is tabulated. If the elements of bin are not integers, they are rounded to the nearest integer. Elements outside the range 1, ... , nbin are (silently) ignored in the tabulation.

## See Also

factor, table.

## Examples

```
tabulate(c(2,3,5))
tabulate(c(2,3,3,5), nb = 10)
tabulate(c(-2,0,2,3,3,5), nb = 3)
tabulate(factor(letters[1:10]))
```

tapply Apply a Function Over a "Ragged" Array

## Description

Apply a function to each cell of a ragged array, that is to each (non-empty) group of values given by a unique combination of the levels of certain factors.

## Usage

```
tapply(X, INDEX, FUN = NULL, ..., simplify = TRUE)
```


## Arguments

$$
\begin{array}{ll}
\text { X } & \text { an atomic object, typically a vector. } \\
\text { INDEX } & \text { list of factors, each of same length as X. } \\
\text { FUN } & \begin{array}{l}
\text { the function to be applied. In the case of functions like }+, \% * \%, \text { etc., the } \\
\text { function name must be quoted. If FUN is NULL, tapply returns a vector } \\
\text { which can be used to subscript the multi-way array tapply normally } \\
\text { produces. }
\end{array} \\
\ldots & \begin{array}{l}
\text { optional arguments to FUN. }
\end{array} \\
\text { simplify } & \begin{array}{l}
\text { If FALSE, tapply always returns an array of mode "list". If TRUE (the } \\
\text { default), then if FUN always returns a scalar, tapply returns an array with }
\end{array} \\
& \text { the mode of the scalar. }
\end{array}
$$

## Value

When FUN is present, tapply calls FUN for each cell that has any data in it. If FUN returns a single atomic value for each cell (e.g., functions mean or var) and when simplify is TRUE, tapply returns a multi-way array containing the values. The array has the same number of dimensions as INDEX has components; the number of levels in a dimension is the number of levels (nlevels()) in the corresponding component of INDEX.
Note that contrary to S, simplify = TRUE always returns an array, possibly 1-dimensional. If FUN does not return a single atomic value, tapply returns an array of mode list whose components are the values of the individual calls to FUN, i.e., the result is a list with a dim attribute.

## See Also

the convenience function aggregate (using tapply); apply, lapply with its version sapply.

## Examples

```
groups <- as.factor(rbinom(32, n = 5, p = .4))
tapply(groups, groups, length) #- is almost the same as
table(groups)
data(warpbreaks)
## contingency table from data.frame : array with named dimnames
tapply(warpbreaks$breaks, warpbreaks[,-1], sum)
tapply(warpbreaks$breaks, warpbreaks[, 3, drop = FALSE], sum)
n <- 17; fac <- factor(rep(1:3, len = n), levels = 1:5)
table(fac)
tapply(1:n, fac, sum)
tapply(1:n, fac, sum, simplify = FALSE)
tapply(1:n, fac, range)
tapply(1:n, fac, quantile)
## example of ... argument: find quarterly means
data(presidents)
tapply(presidents, cycle(presidents), mean, na.rm = TRUE)
ind <- list(c(1, 2, 2), c("A", "A", "B"))
table(ind)
tapply(1:3, ind) #-> the split vector
tapply(1:3, ind, sum)
```

taskCallback Add or remove a top-level task callback

## Description

addTaskCallback registers an R function that is to be called each time a top-level task is completed.
removeTaskCallback un-registers a function that was registered earlier via addTaskCallback.

These provide low-level access to the internal/native mechanism for managing taskcompletion actions. One can use taskCallbackManager at the S-language level to manage S functions that are called at the completion of each task. This is easier and more direct.

## Usage

addTaskCallback(f, data $=$ NULL, name $=$ character $(0)$ )
removeTaskCallback(id)

## Arguments

$\mathrm{f} \quad$ the function that is to be invoked each time a top-level task is successfully completed. This is called with 5 or 4 arguments depending on whether data is specified or not, respectively. The return value should be a logical value indicating whether to keep the callback in the list of active callbacks or discard it.
data if specified, this is the 5-th argument in the call to the callback function f.
id a string or an integer identifying the element in the internal callback list to be removed. Integer indices are 1-based, i.e the first element is 1 . The names of currently registered handlers is available using getTaskCallbackNames and is also returned in a call to addTaskCallback.
name character: names to be used.

## Details

Top-level tasks are individual expressions rather than entire lines of input. Thus an input line of the form expression1 ; expression2 will give rise to 2 top-level tasks.

A top-level task callback is called with the expression for the top-level task, the result of the top-level task, a logical value indicating whether it was successfully completed or not (always TRUE at present), and a logical value indicating whether the result was printed or not. If the data argument was specified in the call to addTaskCallback, that value is given as the fifth argument.
The callback function should return a logical value. If the value is FALSE, the callback is removed from the task list and will not be called again by this mechanism. If the function returns TRUE, it is kept in the list and will be called on the completion of the next top-level task.

## Value

addTaskCallback returns an integer value giving the position in the list of task callbacks that this new callback occupies. This is only the current position of the callback. It can be used to remove the entry as long as no other values are removed from earlier positions in the list first.
removeTaskCallback returns a logical value indicating whether the specified element was removed. This can fail (i.e. return FALSE) if an incorrect name or index is given that does not correspond to the name or position of an element in the list.

## Note

This is an experimental feature and the interface may be changed in the future.
There is also C-level access to top-level task callbacks to allow C routines rather than R functions be used.

## See Also

```
getTaskCallbackNames taskCallbackManager http://developer.r-project.org/
```

TaskHandlers.pdf

## Examples

```
times <- function(total = 3, str="Task a") {
    ctr <- 0
    function(expr, value, ok, visible) {
        ctr <<- ctr + 1
        cat(str, ctr, "\n")
        if(ctr == total) {
            cat("handler removing itself\n")
        }
        return(ctr < total)
    }
}
# add the callback that will work for
# 4 top-level tasks and then remove itself.
n <- addTaskCallback(times(4))
# now remove it, assuming it is still first in the list.
removeTaskCallback(n)
```

\# There is no point in running this
\# as
addTaskCallback(times(4))
sum (1:10)
$\operatorname{sum}(1: 10)$
sum ( $1: 10$ )
sum (1:10)
sum (1:10)
taskCallbackManager Create an $R$-level task callback manager

## Description

This provides an entirely S-language mechanism for managing callbacks or actions that are invoked at the conclusion of each top-level task. Essentially, we register a single R function from this manager with the underlying, native task-callback mechanism and this function handles invoking the other $R$ callbacks under the control of the manager. The manager consists of a collection of functions that access shared variables to manage the list of user-level callbacks.

```
Usage
taskCallbackManager(handlers = list(), registered = FALSE, verbose = FALSE)
```


## Arguments

handlers this can be a list of callbacks in which each element is a list with an element named " f " which is a callback function, and an optional element named "data" which is the 5 -th argument to be supplied to the callback when it is invoked. Typically this argument is not specified, and one uses add to register callbacks after the manager is created.
registered a logical value indicating whether the evaluate function has already been registered with the internal task callback mechanism. This is usually FALSE and the first time a callback is added via the add function, the evaluate function is automatically registered. One can control when the function is registered by specifying TRUE for this argument and calling addTaskCallback manually.
verbose a logical value, which if TRUE, causes information to be printed to the console about certain activities this dispatch manager performs. This is useful for debugging callbacks and the handler itself.

## Value

A list containing 6 functions:
add register a callback with this manager, giving the function, an optional 5-th argument, an optional name by which the the callback is stored in the list, and a register argument which controls whether the evaluate function is registered with the internal C-level dispatch mechanism if necessary.
remove remove an element from the manager's collection of callbacks, either by name or position/index.
evaluate the 'real' callback function that is registered with the C-level dispatch mechanism and which invokes each of the R-leve callbacks within this manager's control.
suspend a function to set the suspend state of the manager. If it is suspended, none of the callbacks will be invoked when a task is completed. One sets the state by specifying a logical value for the status argument.

```
register a function to register the evaluate function with the internal C-level
    dispatch mechanism. This is done automatically by the add function, but
    can be called manually.
callbacks returns the list of callbacks being maintained by this manager.
```


## Note

This is an experimental feature and the interface may be changed in the future.

```
See Also
addTaskCallback removeTaskCallback getTaskCallbackNames http://developer.
r-project.org/TaskHandlers.pdf
```


## Examples

```
# create the manager
h <- taskCallbackManager()
    # add a callback
h$add(function(expr, value, ok, visible) {
                                    cat("In handler\n")
                                    return(TRUE)
                            }, name = "simpleHandler")
# look at the internal callbacks.
getTaskCallbackNames()
    # look at the R-level callbacks
names(h$callback())
#
getTaskCallbackNames()
removeTaskCallback("R-taskCallbackManager")
```

taskCallbackNames Query the names of the current internal top-level task callbacks

## Description

This provides a way to get the names (or identifiers) for the currently registered task callbacks that are invoked at the conclusion of each top-level task. These identifies can be used to remove a callback.

## Usage

getTaskCallbackNames()

## Arguments

## Value

A character vector giving the name for each of the registered callbacks which are invoked when a top-level task is completed successfully. Each name is the one used when registering the callbacks and returned as the in the call to addTaskCallback.

## Note

One can use taskCallbackManager to manage user-level task callbacks, i.e. S-language functions, entirely within the S language and access the names more directly.

## See Also

```
addTaskCallback removeTaskCallback taskCallbackManager http://developer.
r-project.org/TaskHandlers.pdf
```


## Examples

```
n <- addTaskCallback(function(expr, value, ok, visible) {
                        cat("In handler\n")
                        return(TRUE)
    }, name = "simpleHandler")
getTaskCallbackNames()
    # now remove it by name
removeTaskCallback("simpleHandler")
h <- taskCallbackManager()
h$add(function(expr, value, ok, visible) {
                        cat("In handler\n")
        return(TRUE)
    }, name = "simpleHandler")
getTaskCallbackNames()
removeTaskCallback("R-taskCallbackManager")
```

TDist The Student $t$ Distribution

## Description

Density, distribution function, quantile function and random generation for the $t$ distribution with df degrees of freedom (and optional noncentrality parameter ncp).

## Usage

```
\(\mathrm{dt}(\mathrm{x}, \mathrm{df}, \log =\mathrm{FALSE})\)
pt(q, df, ncp=0, lower.tail = TRUE, log.p = FALSE)
qt ( \(p\), df, lower.tail = TRUE, log. \(p=\) FALSE \()\)
rt(n, df)
```


## Arguments

```
\(x, q \quad\) vector of quantiles.
\(p \quad\) vector of probabilities.
n number of observations. If length \((\mathrm{n})>1\), the length is taken to be the
    number required.
df degrees of freedom ( \(>0\), maybe non-integer).
ncp non-centrality parameter \(\delta\); currently ncp <= 37.62.
\(\log , \log . \mathrm{p} \quad \operatorname{logical}\); if TRUE, probabilities p are given as \(\log (\mathrm{p})\).
lower.tail logical; if TRUE (default), probabilities are \(P[X \leq x]\), otherwise, \(P[X>\)
    \(x]\).
```


## Details

The $t$ distribution with $\mathrm{df}=\nu$ degrees of freedom has density

$$
f(x)=\frac{\Gamma((\nu+1) / 2)}{\sqrt{\pi \nu} \Gamma(\nu / 2)}\left(1+x^{2} / \nu\right)^{-(\nu+1) / 2}
$$

for all real $x$. It has mean 0 (for $\nu>1$ ) and variance $\frac{\nu}{\nu-2}$ (for $\nu>2$ ).
The general non-central $t$ with parameters $(\nu, \delta)=$ (df, ncp) is defined as a the distribution of $T_{\nu}(\delta):=\frac{U+\delta}{\chi_{\nu} / \sqrt{\nu}}$ where $U$ and $\chi_{\nu}$ are independent random variables, $U \sim \mathcal{N}(0,1)$, and $\chi_{\nu}^{2}$ is chi-squared, see pchisq.
The most used applications are power calculations for $t$-tests:
Let $T=\frac{\bar{X}-\mu_{0}}{S / \sqrt{n}}$ where $\bar{X}$ is the mean and $S$ the sample standard deviation (sd) of $X_{1}, X_{2}, \ldots, X_{n}$ which are i.i.d. $N\left(\mu, \sigma^{2}\right)$. Then $T$ is distributed as non-centrally $t$ with $\mathrm{df}=n-1$ degrees of freedom and non-centrality parameter $\mathrm{ncp}=\left(\mu-\mu_{0}\right) \sqrt{n} / \sigma$.

## Value

dt gives the density, pt gives the distribution function, qt gives the quantile function, and rt generates random deviates.

## References

Lenth, R. V. (1989). Algorithm AS 243 - Cumulative distribution function of the noncentral $t$ distribution, Appl. Statist. 38, 185-189.

## See Also

df for the F distribution.

## Examples

```
1 - pt(1:5, df = 1)
qt(.975, df = c(1:10, 20,50,100,1000))
tt <- seq(0,10, len=21)
ncp <- seq(0,6, len=31)
ptn <- outer(tt,ncp, function(t,d) pt(t, df = 3, ncp=d))
image(tt,ncp,ptn, zlim=c(0,1),main=t.tit <- "Non-central t - Probabilities")
persp(tt,ncp,ptn, zlim=0:1, r=2, phi=20, theta=200, main=t.tit,
    xlab = "t", ylab = "noncentrality parameter", zlab = "Pr(T <= t)")
```

```
tempfile Create Names for Temporary Files
```


## Description

tempfile returns a vector of character strings which can be used as names for temporary files in the directory given by tempdir.

## Usage

```
tempfile(pattern = "file")
tempdir()
```


## Arguments

pattern a non-empty character vector giving the initial part of the name.

## Details

If pattern has length greater than one then the result is of the same length giving a temporary file name for each component of pattern.

The names are very likely to be unique among calls to tempfile in an $R$ session and across simultaneous R sessions. The filenames are guaranteed not to be currently in use.

The file name is made of the pattern, the process number in hex and a random suffix in hex. The filenames will be in the directory given by tempdir(). This will be a subdirectory of the directory given TMPDIR if set, otherwise "/tmp".

## Value

For tempfile a character vector giving the names of possible (temporary) files. Note that no files are generated by tempfile.

For tempdir, the path of the per-session temporary directory.

## See Also

unlink for deleting files.

## Examples

```
tempfile(c("ab", "a b c")) # give file name with spaces in!
```

```
termplot Plot regression terms
```


## Description

Plots regression terms against their predictors, optionally with standard errors and partial residuals added.

## Usage

```
termplot(model, data=NULL, envir=environment(formula(model)),
    partial.resid=FALSE, rug=FALSE,
    terms=NULL, se=FALSE, xlabs=NULL, ylabs=NULL, main = NULL,
    col.term = 2, lwd.term = 1.5,
    col.se = "orange", lty.se = 2, lwd.se = 1,
    col.res = "gray", cex = 1, pch = par("pch"),
    ask = interactive() && nb.fig < n.tms && .Device !="postscript",
    use.factor.levels=TRUE,
    ...)
```


## Arguments

model fitted model object
data data frame in which variables in model can be found
envir environment in which variables in model can be found
partial.resid logical; should partial residuals be plotted?
rug add rugplots (jittered 1-d histograms) to the axes?
terms which terms to plot (default NULL means all terms)
se plot pointwise standard errors?
xlabs vector of labels for the x axes
ylabs vector of labels for the $y$ axes
main logical, or vector of main titles; if TRUE, the model's call is taken as main title, NULL or FALSE mean no titles.
col.term, lwd.term
color and line width for the "term curve", see lines.
col.se, lty.se, lwd.se
color, line type and line width for the "twice-standard-error curve" when
se $=$ TRUE.
col.res, cex, pch
color, plotting character expansion and type for partial residuals, when partial.resid = TRUE, see points.
ask logical; if TRUE, the user is asked before each plot, see par (ask=.).
use.factor.levels
Should x -axis ticks use factor levels or numbers for factor terms?
... other graphical parameters

## Details

The model object must have a predict method that accepts type=terms, eg glm in the base package, coxph and survreg in the survival package.
For the partial.resid=TRUE option it must have a residuals method that accepts type="partial", which lm and glm do.
The data argument should rarely be needed. One exception is that models with missing data using na.action=na.omit will need to specify a data argument. A work-around is to use na.action=na.exclude instead.

Nothing sensible happens for interaction terms.

## See Also

For (generalized) linear models, plot.lm and predict.glm.

## Examples

```
rs <- require(splines)
x <- 1:100
z <- factor(rep(LETTERS[1:4],25))
y <- rnorm(100, sin(x/10)+as.numeric(z))
model <- glm(y ~ ns(x,6) + z)
par(mfrow=c(2,2)) ## 2 x 2 plots for same model :
termplot(model, main = paste("termplot( ", deparse(model$call)," ...)"))
termplot(model, rug=TRUE)
termplot(model, partial=TRUE, rug= TRUE,
    main="termplot(..., partial = TRUE, rug = TRUE)")
termplot(model, partial=TRUE, se = TRUE, main = TRUE)
if(rs) detach("package:splines")
```

```
terms Model Terms
```


## Description

The function terms is a generic function which can be used to extract terms objects from various kinds of $R$ data objects.

## Usage

```
terms(x, ...)
```


## Arguments

x object used to select a method to dispatch. further arguments passed to or from other methods.

## Details

There are methods for classes "aovlist", and "terms" "formula" (see terms.formula): the default method just extracts the terms component of the object (if any).

## Value

An object of class c("terms", "formula") which contains the terms representation of a symbolic model. See terms.object for its structure.

## See Also

```
terms.object, terms.formula, lm, glm, formula.
```

terms.formula $\quad A$ function to construct a terms object from a formula.

## Description

This function takes a formula and some optional arguments and constructs a terms object. The terms object can then be used to construct a model.matrix.

## Usage

```
terms.formula(x, specials=NULL, abb=NULL, data=NULL, neg.out=TRUE,
                        keep.order=FALSE, ...)
```


## Arguments

x
specials What functions in the formula should be marked as special in the terms object.
abb
data A data frame from which the meaning of the special symbol . can be inferred. It is unused if there is no . in the formula.
neg.out TRUE if terms with a minus, - should be removed. If FALSE these are kept in and indicate a negative order (for fractionate?).
keep.order A logical value indicating whether the terms should keep their positions. If FALSE the terms are reordered so that main effects come first.
... further arguments passed to or from other methods.

## Details

Not all of the options work in the same way that they do in $S$ and not all are implemented.

## Value

A terms object is returned.

## See Also

```
terms.object, terms.default
```

terms.object Description of Terms Objects

## Description

An object of class terms holds information about a model. Usually the model was specified in terms of a formula and that formula was used to determine the terms object.

The object itself is simply the formula supplied to the call of terms.formula. The object has a number of attributes and they are used to construct the model frame.

## Value

An object with the following attributes:
factors A matrix of variables by terms showing which variables appear in which terms. The entries are 0 if the variable does not occur in the term, 1 if it does occur and should be coded by contrasts, and 2 if it occurs and should be coded via dummy variables for all levels (as when an intercept or lower-order term is missing).
term.labels A character vector containing the labels for each of the terms in the model.
variables A list of the variables in the model
intercept Either 0, indicating no intercept is to be fit, or 1 indicating that an intercept is to be fit.
order A vector of the same length as term.labels indicating the order of interaction for each term
response The index of the variable (in variables) of the response (the left hand side of the formula).
offset If the model contains offset terms there is an offset attribute indicating which terms are offsets
specials If the specials argument was given to terms.formula there is a specials attribute, a list of vectors indicating the terms that contain these special functions.

The object has class c("terms", "formula").

## Note

These objects are different from those found in S. In particular there is no formula attribute, instead the object is itself a formula. Thus, the mode of a terms object is different as well.
An example of the specials argument can be seen in the aov function.

## See Also

terms, terms.default, formula.
text
Add Text to a Plot

## Description

text draws the strings given in the vector labels at the coordinates given by x and y . y may be missing since xy . coords ( $\mathrm{x}, \mathrm{y}$ ) is used for construction of the coordinates.

## Usage

```
text (x, ...)
text.default (x, y = NULL, labels = seq(along = x), adj = NULL,
        pos = NULL, offset = 0.5, vfont = NULL,
        cex = 1, col = NULL, font = NULL, xpd = NULL, ...)
```


## Arguments

$x, y \quad$ numeric vectors of coordinates where the text labels should be written. If the length of x and y differs, the shorter one is recycled.
labels one or more character strings or expressions specifying the text to be written. An attempt is made to coerce other vectors to character, and other language objects to expressions.
adj one or two values in $[0,1]$ which specify the x (and optionally y) adjustment of the labels. On most devices values outside that interval will also work.
pos a position specifier for the text. If specified this overrides any adj value given. Values of 1, 2, 3 and 4, respectively indicate positions below, to the left of, above and to the right of the specified coordinates.
offset when pos is specified, this value gives the offset of the label from the specified coordinate in fractions of a character width.
vfont if a character vector of length 2 is specified, then Hershey vector fonts are used. The first element of the vector selects a typeface and the second element selects a style.
cex numeric character expansion factor; multiplied by par("cex") yields the final character size.
col, font the color and font to be used; these default to the values of the global graphical parameters in par().
xpd (where) should clipping take place? Defaults to par ("xpd").
... further graphical parameters (from par).

## Details

labels must be of type character or expression (or be coercible to such a type). In the latter case, quite a bit of mathematical notation is available such as sub- and superscripts, greek letters, fractions, etc.
adj allows adjustment of the text with respect to ( $\mathrm{x}, \mathrm{y}$ ). Values of $0,0.5$, and 1 specify left/bottom, middle and right/top, respectively. The default is for centered text, i.e., adj $=$ $c(0.5,0.5)$. Accurate vertical centering needs character metric information on individual characters, which is only available on some devices.

The pos and offset arguments can be used in conjunction with values returned by identify to recreate an interactively labelled plot.
Text can be rotated by using graphical parameters srt (see par); this rotates about the centre set by adj.
Graphical parameters col, cex and font can be vectors and will then be applied cyclically to the labels (and extra values will be ignored).

## See Also

mtext, title, Hershey for details on Hershey vector fonts, plotmath for details and more examples on mathematical annotation.

## Examples

```
plot(-1:1,-1:1, type = "n", xlab = "Re", ylab = "Im")
K <- 16; text(exp(1i * 2 * pi * (1:K) / K), col = 2)
## The following two examples use latin1 characters: these may not
## appear correctly (or be omitted entirely).
plot(1:10, 1:10, main = "text(...) examples\n~~~~~~~~~~~~~"',
    sub = "R is GNU l', but not ■ ...")
mtext("ñISO-accents■: ś éè ф\emptyset å<Å æ<\mathbb{E", side=3)}
points(c(6,2), c(2,1), pch = 3, cex = 4, col = "red")
text(6, 2, "the text is CENTERED around (x,y) = (6,2) by default",
    cex = .8)
text(2, 1, "or Left/Bottom - JUSTIFIED at (2,1) by 'adj = c(0,0)'",
    adj = c(0,0))
text(4, 9, expression(hat(beta) == (X^t * X)^{-1} * X^t * y))
text(4, 8.4, "expression(hat(beta) == (X^t * X)^{-1} * X^t * y)", cex = .75)
text(4, 7, expression(bar(x) == sum(frac(x[i], n), i==1, n)))
## Two more latin1 examples
text(5,10.2,
    "Le français, c’est façile: Règles, Liberté, Egalité, Fraternité...")
text(5,9.8, "Jetz no chli züritüütsch: (noch ein biSSchen Zürcher deutsch)")
```

```
textConnection Text Connections
```


## Description

Input and output text connections.

## Usage

textConnection(object, open = "r")

## Arguments

object character. A description of the connection. For an input is an R character vector object, and for an output connection the name for the $R$ character vector to receive the output.
open character. Either "r" (or equivalently "") for an input connection or "w" or "a"for an output connection.

## Details

An input text connection is opened and the character vector is copied at time the connection object is created, and close destroys the copy.
An output text connection is opened and creates an $R$ character vector of the given name in the user's workspace. This object will at all times hold the completed lines of output to the connection, and isIncomplete will indicate if there is an incomplete final line. Closing the connection will output the final line, complete or not.
Opening a text connection with mode $=$ " a " will attempt to append to an existing character vector with the given name in the user's workspace. If none is found (even if an object exists of the right name but the wrong type) a new character vector wil be created, with a warning.

You cannot seek on a text connection, and seek will always return zero as the position.

## Value

A connection object of class "textConnection" which inherits from class "connection".

## See Also

connections, showConnections, pushBack

## Examples

```
zz <- textConnection(LETTERS)
readLines(zz, 2)
scan(zz, "", 4)
pushBack(c("aa", "bb"), zz)
scan(zz, "", 4)
close(zz)
zz <- textConnection("foo", "w")
writeLines(c("testit1", "testit2"), zz)
cat("testit3 ", file=zz)
isIncomplete(zz)
cat("testit4\n", file=zz)
isIncomplete(zz)
close(zz)
foo
# capture R output: use part of example from help(lm)
zz <- textConnection("foo", "w")
ctl <- c(4.17, 5.58, 5.18, 6.11, 4.5, 4.61, 5.17, 4.53, 5.33, 5.14)
trt <- c(4.81, 4.17, 4.41, 3.59, 5.87, 3.83, 6.03, 4.89, 4.32, 4.69)
group <- gl(2, 10, 20, labels = c("Ctl", "Trt"))
weight <- c(ctl, trt)
sink(zz)
anova(lm.D9 <- lm(weight ~ group))
cat("\nSummary of Residuals:\n\n")
summary(resid(lm.D9))
sink()
close(zz)
cat(foo, sep = "\n")
```

time Sampling Times of Time Series

## Description

time creates the vector of times at which a time series was sampled.
cycle gives the positions in the cycle of each observation.
frequency returns the number of samples per unit time and deltat the time interval between observations (see ts).

## Usage

```
time(x, offset=0, ...)
cycle(x, ...)
frequency(x, ...)
deltat(x, ...)
```


## Arguments

x
a univariate or multivariate time-series, or a vector or matrix.
offset can be used to indicate when sampling took place in the time unit. 0 (the default) indicates the start of the unit, 0.5 the middle and 1 the end of the interval.
... extra arguments for future methods.

## Details

These are all generic functions, which will use the tsp attribute of x if it exists. time and cycle have methods for class ts that coerce the result to that class.

## See Also

ts, start, tsp, window.
date for clock time, system.time for CPU usage.

## Examples

```
data(presidents)
cycle(presidents)
# a simple series plot: c() makes the x and y arguments into vectors
plot(c(time(presidents)), c(presidents), type="l")
```

Titanic Survival of passengers on the Titanic

## Description

This data set provides information on the fate of passengers on the fatal maiden voyage of the ocean liner 'Titanic', summarized according to economic status (class), sex, age and survival.

## Usage

data(Titanic)

## Format

A 4-dimensional array resulting from cross-tabulating 2201 observations on 4 variables. The variables and their levels are as follows:

| No | Name | Levels |
| ---: | :--- | :--- |
| 1 | Class | 1st, 2nd, 3rd, Crew |
| 2 | Sex | Male, Female |
| 3 | Age | Child, Adult |
| 4 | Survived | No, Yes |

## Details

The sinking of the Titanic is a famous event, and new books are still being published about it. Many well-known facts-from the proportions of first-class passengers to the "women and children first" policy, and the fact that that policy was not entirely successful in saving the women and children in the third class-are reflected in the survival rates for various classes of passenger.
These data were originally collected by the British Board of Trade in their investigation of the sinking. Note that there is not complete agreement among primary sources as to the exact numbers on board, rescued, or lost.

Due in particular to the very successful film 'Titanic', the last years saw a rise in public interest in the Titanic. Very detailed data about the passengers is now available on the Internet, at sites such as Encyclopedia Titanica (http://www.rmplc.co.uk/eduweb/sites/phind).

## Source

Dawson, Robert J. MacG. (1995), The 'Unusual Episode' Data Revisited. Journal of Statistics Education, 3. http://www.amstat.org/publications/jse/v3n3/datasets.dawson. html

The source provides a data set recording class, sex, age, and survival status for each person on board of the Titanic, and is based on data originally collected by the British Board of Trade and reprinted in:
British Board of Trade (1990), Report on the Loss of the 'Titanic' (S.S.). British Board of Trade Inquiry Report (reprint). Gloucester, UK: Allan Sutton Publishing.

## Examples

```
data(Titanic)
mosaicplot(Titanic, main = "Survival on the Titanic")
## Higher survival rates in children?
apply(Titanic, c(3, 4), sum)
## Higher survival rates in females?
apply(Titanic, c(2, 4), sum)
## Use loglm() in package 'MASS' for further analysis ...
```


## title Plot Annotation

## Description

This function can be used to add labels to a plot. Its first four principal arguments can also be used as arguments in most high-level plotting functions. They must be of type character or expression. In the latter case, quite a bit of mathematical notation is available such as sub- and superscripts, greek letters, fractions, etc.

## Usage

```
title(main \(=\) NULL, sub \(=\) NULL, xlab \(=\) NULL, ylab \(=\) NULL,
            line \(=\) NA, outer \(=\) FALSE, ...)
```


## Arguments

| main | The main title (on top) using font and size (character expansion) <br> par("font.main") and color par("col.main"). |
| :--- | :--- |
| sub | Sub-title (at bottom) using font and size par ("font.sub") and color <br> par("col.sub"). |
| xlab | X axis label using font and character expansion par("font.axis") and <br> color par("col.axis"). |
| ylab | Y axis label, same font attributes as xlab. |
| line | specifying a value for line overrides the default placement of labels, and <br> places them this many lines from the plot. <br> a logical value. If TRUE, the titles are placed in the outer margins of the |
| outer | plot. |
| $\ldots$ | further graphical parameters (from par). |

## Details

The labels passed to title can be simple strings or expressions, or they can be a list containing the string to be plotted, and a selection of the optional modifying graphical parameters cex=, col=, font=.

## See Also

mtext, text; plotmath for details on mathematical annotation.

## Examples

```
data(cars)
plot(cars, main = "") # here, could use main directly
title(main = "Stopping Distance versus Speed")
plot(cars, main = "")
title(main = list("Stopping Distance versus Speed", cex=1.5,
    col="red", font=3))
x <- seq(-4, 4, len = 101)
y <- cbind(sin(x), cos(x))
matplot(x, y, type = "l", xaxt = "n",
    main = expression(paste(plain(sin) * phi, " and ",
                plain(cos) * phi)),
    ylab = expression("sin" * phi, "cos" * phi), # only 1st is taken
    xlab = expression(paste("Phase Angle ", phi)),
    col.main = "blue")
axis(1, at = c(-pi, -pi/2, 0, pi/2, pi),
    lab = expression(-pi, -pi/2, 0, pi/2, pi))
abline(h = 0, v = pi/2 * c(-1,1), lty = 2, lwd = .1, col = "gray70")
```


## ToothGrowth The Effect of Vitamin $C$ on Tooth Growth in Guinea Pigs

## Description

The response is the length of odontoblasts (teeth) in each of 10 guinea pigs at each of three dose levels of Vitamin C ( $0.5,1$, and 2 mg ) with each of two delivery methods (orange juice or ascorbic acid).

## Usage

data(ToothGrowth)

## Format

A data frame with 60 observations on 3 variables.

| $[, 1]$ | len | numeric | Tooth length |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | supp | factor | Supplement type (VC or OJ). |
| $[, 3]$ | dose | numeric | Dose in milligrams. |

## Source

C. I. Bliss (1952) The Statistics of Bioassay. Academic Press.

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(ToothGrowth)
```

```
coplot(len ~ dose | supp, data = ToothGrowth, panel = panel.smooth,
        xlab = "ToothGrowth data: length vs dose, given type of supplement")
```

    toString toString Converts its Argument to a Character String
    
## Description

This is a helper function for format. It converts its argument to a string. If the argument is a vector then its elements are concatenated with a , as a separtor. Most methods should honor the width argument. The minimum value for width is six.

## Usage

```
toString(x, ...)
toString.default(x, width, ...)
```


## Arguments

$\mathrm{x} \quad$ The object to be converted.
width The returned value is at most the first width characters.
... Optional arguments for methods.

## Value

A character vector of length 1 is returned.

## Author(s)

Robert Gentleman

## See Also

format

## Examples

```
x <- c("a", "b", "aaaaaaaaaaa")
toString(x)
toString(x, width=8)
```

| traceInteractive Tracing and Debugging of Calls to a Function or <br> Method |
| :--- | :--- |

## Description

A call to trace allows you to insert debugging code (e.g., a call to browser or recover) at chosen places in any function. A call to untrace cancels the tracing. Specified methods can be traced the same way, without tracing all calls to the function. Trace code can be any R expression. Tracing can be temporarily turned on or off globally by calling tracingState.

## Usage

```
trace(what, tracer, exit, at, print = TRUE, signature = NULL)
untrace(what, signature = NULL)
tracingState(on)
```


## Arguments

| what | The name (quoted or not) of a function to be traced or untraced. More <br> than one name can be given in the quoted form, and the same action will <br> be applied to each one. |
| :--- | :--- |
| tracer | Either a function or an unevaluated expression. The function will be <br> called or the expression will be evaluated either at the beginning of the <br> call, or before those steps in the call specified by the argument at. See <br> the details section. |
| exit | Either a function or an unevaluated expression. The function will be <br> called or the expression will be evaluated on exiting the function. See the <br> details section. |
| at | An optional numeric vector. If supplied, tracer will be called just before <br> the corresponding step in the body of the function. See the details section. |
| print | If TRUE, a descriptive line is printed before any trace expression is evalu- <br> ated. |
| signature | If this argument is supplied, it should be a signature for a method for <br> function what. In this case, the method, and not the function itself, is |
| traced. |  |

## Details

The trace function operates by constructing a revised version of the function (or of the method, if signature is supplied), and assigning the new object back where the original was found. If only the what argument is given, a line of trace printing is produced for each call to the function (back compatible with the earlier version of trace).

The object constructed by trace is from a class that extends "function" and which contains the original, untraced version. A call to untrace re-assigns this version.

If the argument tracer or exit is the name of a function, the tracing expression will be a call to that function, with no arguments. This is the easiest and most common case, with the functions browser and recover the likeliest candidates; the former browses in the frame of the function being traced, and the latter allows browsing in any of the currently active calls.
The tracer or exit argument can also be an unevaluated expression (such as returned by a call to quote or substitute). This expression itself is inserted in the traced function, so it will typically involve arguments or local objects in the traced function. An expression of this form is useful if you only want to interact when certain conditions apply (and in this case you probably want to supply print=FALSE in the call to trace also).
When the at argument is supplied, it should be a vector of integers referring to the substeps of the body of the function (this only works if the body of the function is enclosed in \{ ...\}. In this case tracer is not called on entry, but instead just before evauating each of the steps listed in at. (Hint: you don't want to try to count the steps in the printed version of a function; instead, look at as.list(body (f)) to get the numbers associated with the steps in function f.)
An intrinsic limitation in the exit argument is that it won't work if the function itself uses on.exit, since the existing calls will override the one supplied by trace.
Tracing does not nest. Any call to trace replaces previously traced versions of that function or method, and untrace always restores an untraced version. (Allowing nested tracing has too many potentials for confusion and for accidentally leaving traced versions behind.)
Tracing primitive functions (builtins and specials) from the base package works, but only by a special mechanism and not very informatively. Tracing a primitive causes the primitive to be replaced by a function with argument ... (only). You can get a bit of information out, but not much. A warning message is issued when trace is used on a primitive.
The practice of saving the traced version of the function back where the function came from means that tracing carries over from one session to another, if the traced function is saved in the session image. (In the next session, untrace will remove the tracing.) On the other hand, functions that were in a package, not in the global environment, are not saved in the image, so tracing expires with the session for such functions.
Tracing a method is basically just like tracing a function, with the exception that the traced version is stored by a call to setMethod rather than by direct assignment, and so is the untraced version after a call to untrace.

The version of trace described here is largely compatible with the version in S-Plus, although the two work by entirely different mechanisms. The S-Plus trace uses the session frame, with the result that tracing never carries over from one session to another ( R does not have a session frame). Another relevant distinction has nothing directly to do with trace: The browser in S-Plus allows changes to be made to the frame being browsed, and the changes will persist after exiting the browser. The R browser allows changes, but they disappear when the browser exits. This may be relevant in that the S-Plus version allows you to experiment with code changes interactively, but the $R$ version does not. (A future revision may include a "destructive" browser for R.)

## Value

The traced function(s) name(s). The relevant consequence is the assignment that takes place.

## Note

The version of function tracing that includes any of the arguments except for the function name requires the methods package (because it uses special classes of objects to store and restore versions of the traced functions).

In rare cases, this may change some behavior. In particular, although the methods package generally supports older styles of classes and methods, it is stricter in treating class attributes. Just occasionally, older code may stop on an error with the methods package attached, but not otherwise.

## See Also

browser and recover, the likeliest tracing functions; also, quote and substitute for constructing general expressions.

## Examples

```
f <- function(x, y) {
    y <- pmax(y, .001)
    x - y
}
## arrange to call the browser on entering and exiting
## function f
trace("f", browser, exit = browser)
## instead, conditionally assign some data, and then browse
## on exit, but only then. Don't bother me otherwise
trace("f", quote(if(any(y < 0)) yOrig <- y),
    exit = quote(if(exists("yOrig")) browser()),
    print = FALSE)
## trace a utility function, with recover so we
## can browse in the calling functions as well.
trace("as.matrix", recover)
## turn off the tracing
untrace(c("f", "as.matrix"))
```


## Description

traceback() prints the call stack of the last error, i.e., the sequence of calls that lead to the error. This is useful when an error occurs with an unidentifiable error message. This stack is stored as a list in .Traceback, which traceback prints in a user-friendly format.

## Usage

traceback()

## Value

traceback() returns nothing, but prints the deparsed call stack deepest call first. The calls may print on more that one line, and the first line is labelled by the frame number.

## Examples

```
foo <- function(x) { print(1); bar(2) }
bar <- function(x) { x + a.variable.which.does.not.exist }
foo(2) # gives a strange error
traceback()
## 2: bar(2)
## 1: foo(2)
bar
## Ah, this is the culprit ...
```

transform Transform an Object, e.g. a Data Frame

## Description

transform is a generic function, which-at least currently-only does anything useful with dataframes. transform. default converts its first argument to a dataframe if possible and calls transform.data.frame.

## Usage

```
transform(x, ...)
transform.default(x, ...)
transform.data.frame(x, ...)
```


## Arguments

$\begin{array}{ll}\mathrm{x} & \text { The object to be transformed } \\ \ldots . & \text { Further arguments of the form tag=value }\end{array}$

## Details

The ... arguments to transform.data.frame are tagged vector expressions, which are evaluated in the dataframe x . The tags are matched against names ( x ), and for those that match, the value replace the corresponding variable in x , and the others are appended to x .
trees

## Value

The modified value of x .

## Note

If some of the values are not vectors of the appropriate length, you deserve whatever you get!

## Author(s)

Peter Dalgaard

## See Also

```
subset, list, data.frame
```


## Examples

```
data(airquality)
transform(airquality, Ozone = -Ozone)
transform(airquality, new = -Ozone, Temp = (Temp-32)/1.8)
attach(airquality)
transform(Ozone, logOzone = log(Ozone)) # marginally interesting ...
detach(airquality)
```

trees Girth, Height and Volume for Black Cherry Trees

## Description

This data set provides measurements of the girth, height and volume of timber in 31 felled black cherry trees. Note that girth is the diameter of the tree (in inches) measured at 4 ft 6 in above the ground.

## Usage

data(trees)

## Format

A data frame with 31 observations on 3 variables.

| $[, 1]$ | Girth | numeric | Tree diameter in inches |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | Height | numeric | Height in ft |
| $[, 3]$ | Volume | numeric | Volume of timber in cubic ft |

## Source

Ryan, T. A., Joiner, B. L. and Ryan, B. F. (1976) The Minitab Student Handbook. Duxbury Press.

## References

Atkinson, A. C. (1985) Plots, Transformations and Regression. Oxford University Press.

## Examples

```
data(trees)
pairs(trees, panel = panel.smooth, main = "trees data")
plot(Volume ~ Girth, data = trees, log = "xy")
coplot(log(Volume) ~ log(Girth) | Height, data = trees,
    panel = panel.smooth)
summary(fm1 <- lm(log(Volume) ~ log(Girth), data = trees))
summary(fm2 <- update(fm1, ~ . + log(Height), data = trees))
step(fm2)
## i.e. Volume ~= c * Height * Girth^2 seems reasonable
```

Trig Trigonometric Functions

## Description

These functions give the obvious trigonometric functions. They respectively compute the cosine, sine, tangent, arc-cosine, arc-sine, arc-tangent, and the two-argument arc-tangent.

## Usage

```
cos(x)
sin(x)
tan(x)
acos(x)
asin(x)
atan(x)
atan2(y, x)
```


## Arguments

$\mathrm{x}, \mathrm{y}$ numeric vector

## Details

The arc-tangent of two arguments atan2 $(\mathrm{y}, \mathrm{x})$ returns the angle between the x -axis and the vector from the origin to $(x, y)$, i.e., for positive $\operatorname{arguments} \operatorname{atan} 2(y, x)==\operatorname{atan}(y / x)$.
Angles are in radians, not degrees (i.e. a right angle is $\pi / 2$ ).

## Examples

```
cos(0) == 1
sin(3*pi/2) == cos(pi)
x <- rnorm(99)
all.equal( }\operatorname{sin}(-x),-\operatorname{sin}(x)
all.equal( cos(-x), cos(x))
x <- abs(x); y <- abs(rnorm(x))
all(abs(atan2(y, x) - atan(y/x)) <= .Machine$double.eps)# TRUE
table(abs(atan2(y, x) - atan(y/x)) / .Machine$double.eps) # depends!
```

```
x <- 1:99/100
all(Mod(1 - (cos(x) + 1i*sin(x)) / exp(1i*x)) < 1.1 * .Machine$double.eps)
    2* abs(1 - x / acos(cos(x))) / .Machine$double.eps #-- depends ?
all(abs(1 - x / asin(\operatorname{sin}(x))) <= .Machine$double.eps) # TRUE
all(abs(1 - x / atan(tan(x))) <= .Machine$double.eps) # TRUE
```

try
Try an Expression Allowing Error Recovery.

## Description

try is a wrapper to run an expression that might fail and allow the user's code to handle error-recovery.

## Usage

```
try (expr, first = TRUE)
```


## Arguments

```
expr an R expression to try
first not for user use!
```


## Details

try is a user-friendly wrapper to restart. The argument first is used to record if restart has already been used, and so ensure that restart is called only once.

## Value

The value of the expression if expr is evaluated without error, but an invisible object of class "try-error" containing the error message if it if fails. The normal error handling will print the same message unless options("show.error.messages") is false.

## See Also

options for setting error handlers and suppressing the printing of error messages; geterrmessage for retrieving the last error message.

## Examples

```
## this example will not work correctly in example(try), but
## it does work correctly if pasted in
options(show.error.messages = FALSE)
try(log("a"))
print(.Last.value)
options(show.error.messages = TRUE)
## run a simulation, keep only the results that worked.
set.seed(123)
x <- rnorm(50)
doit <- function(x)
{
```

```
    x <- sample(x, replace=TRUE)
    if(length(unique(x)) > 30) mean(x)
    else stop("too few unique points")
}
options(show.error.messages = FALSE)
## alternative 1
res <- lapply(1:100, function(i) try(doit(x)))
## alternative 2
res <- vector("list", 100)
for(i in 1:100) res[[i]] <- try(doit(x))
options(show.error.messages = TRUE)
unlist(res[sapply(res, function(x) !inherits(x, "try-error"))])
```

ts Time-Series Objects

## Description

The function ts is used to create time-series objects.
as.ts and is.ts coerce an object to a time-series and test whether an object is a time series.

## Usage

```
ts (data \(=N A\), start \(=1\), end \(=\) numeric ( 0 ) , frequency \(=1\),
    deltat \(=1\), ts.eps = getOption("ts.eps"), class, names)
as.ts(x)
is.ts(x)
```


## Arguments

| data | a numeric vector or matrix of the observed time-series values. A data <br> frame will be coerced to a numeric matrix via data.matrix. |
| :--- | :--- |
| start | the time of the first observation. Either a single number or a vector of <br> two integers, which specify a natural time unit and a (1-based) number <br> of samples into the time unit. See the examples for the use of the second <br> form. <br> the time of the last observation, specified in the same way as start. |
| end | the number of observations per unit of time. |
| frequency | the fraction of the sampling period between successive observations; e.g., <br> deltat <br> 1/12 for monthly data. Only one of frequency or deltat should be <br> provided. <br> time series comparison tolerance. Frequencies are considered equal if their |
| ts.eps | absolute difference is less than ts.eps. <br> class |
| class to be given to the result, or none if NULL or "none". The default is |  |
| names | "ts" for a single series, c("mts", "ts") for multiple series. |
| a character vector of names for the series in a multiple series: defaults to |  |
| the colnames of data, or Series 1, Series 2,.... |  |

## Details

The function ts is used to create time-series objects. These are vector or matrices with class of "ts" (and additional attributes) which represent data which has been sampled at equispaced points in time. In the matrix case, each column of the matrix data is assumed to contain a single (univariate) time series.
Class "ts" has a number of methods. In particular arithmetic will attempt to align time axes, and subsetting to extract subsets of series can be used (e.g. EuStockMarkets[, "DAX"]). However, subsetting the first (or only) dimension will return a matrix or vector, as will matrix subsetting.
The value of argument frequency is used when the series is sampled an integral number of times in each unit time interval. For example, one could use a value of 7 for frequency when the data are sampled daily, and the natural time period is a week, or 12 when the data are sampled monthly and the natural time period is a year. Values of 4 and 12 are assumed in (e.g.) print methods to imply a quarterly and monthly series respectively.
as.ts will use the tsp attribute of the object if it has one to set the start and end times and frequency.

## See Also

tsp, frequency, start, end, time, window; print.ts, the print method for time series objects; plot.ts, the plot method for time series objects. Standard package ts for many additional time-series functions.

## Examples

```
ts(1:10, frequency = 4, start = c(1959, 2)) # 2nd Quarter of 1959
print( ts(1:10, freq = 7, start = c(12, 2)), calendar = TRUE) # print.ts(.)
## Using July 1954 as start date:
gnp <- ts(cumsum(1 + round(rnorm(100), 2)),
            start = c(1954, 7), frequency = 12)
plot(gnp) # using 'plot.ts' for time-series plot
## Multivariate
z <- ts(matrix(rnorm(300), 100, 3), start=c(1961, 1), frequency=12)
class(z)
plot(z)
plot(z, plot.type="single", lty=1:3)
# Ensure working arithmetic for 'ts' objects :
stopifnot(z == z)
stopifnot(z-z == 0)
## A phase plot:
data(nhtemp)
plot(nhtemp, c(nhtemp[-1], NA), cex = .8, col="blue",
    main = "Lag plot of New Haven temperatures")
## a clearer way to do this would be
library(ts)
plot(nhtemp, lag(nhtemp, 1), cex = .8, col="blue",
    main = "Lag plot of New Haven temperatures")
```


## Description

Methods for objects of class "ts", typically the result of ts.

## Usage

diff(x, lag=1, differences=1, ...)
na.omit(object, ...)

## Arguments

$x \quad$ an object of class "ts" containing the values to be differenced.
lag an integer indicating which lag to use.
differences an integer indicating the order of the difference.
object a univariate or multivariate time series.
... further arguments to be passed to or from methods.

## Details

The na.omit method omits initial and final segments with missing values in one or more of the series. 'Internal' missing values will lead to failure.

## Value

For the na.omit method, a time series without missing values. The class of object will be preserved.

## Author(s)

B. D. Ripley

## See Also

diff; na.omit, na.fail, na.contiguous.
tsp Tsp Attribute of Time-Series-like Objects

## Description

tsp returns the tsp attribute (or NULL). It is included for compatibility with S version 2 . tsp<- sets the tsp attribute. hasTsp ensures x has a tsp attribute, by adding one if needed.

## Usage

```
tsp(x)
tsp(x) <- value
hasTsp(x)
```


## Arguments

$x \quad a \quad$ vector or matrix or univariate or multivariate time-series.
value a numeric vector of length 3 or NULL.

## Details

The tsp attribute was previously described here as c(start(x), end(x), frequency(x)), but this is incorrect. It gives the start time in time units, the end time and the frequency.

Assignments are checked for consistency.
Assigning NULL which removes the tsp attribute and any "ts" class of x .

## See Also

ts, time, start.

Tukey
The Studentized Range Distribution

## Description

Functions on the distribution of the studentized range, $R / s$, where $R$ is the range of a standard normal sample of size $n$ and $s^{2}$ is independently distributed as chi-squared with $d f$ degrees of freedom, see pchisq.

## Usage

ptukey (q, nmeans, df, nranges = 1, lower.tail = TRUE, log. $\mathrm{p}=$ FALSE)
qtukey ( p , nmeans, df , nranges $=1$, lower.tail = TRUE, log.p = FALSE)

## Arguments

q vector of quantiles.
$p \quad$ vector of probabilities.
nmeans sample size for range (same for each group).
df degrees of freedom for $s$ (see below).
nranges number of groups whose maximum range is considered.
$\log , \log \cdot \mathrm{p} \quad \operatorname{logical}$; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

If $n_{g}=$ nranges is greater than one, $R$ is the maximum of $n_{g}$ groups of nmeans observations each.

## Value

ptukey gives the distribution function and qtukey its inverse, the quantile function.

## Note

A Legendre 16-point formula is used for the integral of ptukey. The computations are relatively expensive, especially for qtukey which uses a simple secant method for finding the inverse of ptukey. qtukey will be accurate to the 4th decimal place.

## References

Copenhaver, Margaret Diponzio and Holland, Burt S. (1988) Multiple comparisons of simple effects in the two-way analysis of variance with fixed effects. Journal of Statistical Computation and Simulation, 30, 1-15.

## See Also

pnorm and qnorm for the corresponding functions for the normal distribution.

## Examples

```
system.time(curve(ptukey(x, nm=6, df=5), from=-1, to=8, n=101))
(ptt <- ptukey(0:10, 2, df= 5))
(qtt <- qtukey(.95, 2, df= 2:11))
## The precision may be not much more than about 8 digits:
summary(abs(.95- ptukey(qtt,2, df = 2:11)))
```

```
TukeyHSD Compute Tukey Honest Significant Differences
```


## Description

Create a set of confidence intervals on the differences between the means of the levels of a factor with the specified family-wise probability of coverage. The intervals are based on the Studentized range statistic, Tukey's 'Honest Significant Difference' method. There is a plot method.

## Usage

```
TukeyHSD(x, which = seq(along=tabs), ordered = FALSE, conf.level = 0.95, ...)
```


## Arguments

## x

A fitted model object, usually an aov fit.
which A list of terms in the fitted model for which the intervals should be calculated. Defaults to all the terms.
ordered A logical value indicating if the levels of the factor should be ordered according to increasing average in the sample before taking differences. If ordered is true then the calculated differences in the means will all be positive. The significant differences will be those for which the lwr end point is positive.
conf.level A numeric value between zero and one giving the family-wise confidence level to use.
... Optional additional arguments. None are used at present.

## Details

When comparing the means for the levels of a factor in an analysis of variance, a simple comparison using t-tests will inflate the probability of declaring a significant difference when it is not in fact present. This because the intervals are calculated with a given coverage probability for each interval but the interpretation of the coverage is usually with respect to the entire family of intervals.

John Tukey introduced intervals based on the range of the sample means rather than the individual differences. The intervals returned by this function are based on this Studentized range statistics.
Technically the intervals constructed in this way would only apply to balanced designs where there are the same number of observations made at each level of the factor. This function incorporates an adjustment for sample size that produces sensible intervals for mildly unbalanced designs.

## Value

A list with one component for each term requested in which. Each component is a matrix with columns diff giving the difference in the observed means, lwr giving the lower end point of the interval, and upr giving the upper end point.

## Author(s)

Douglas Bates

## References

Miller, R. G. (1981) Simultaneous Statistical Inference. Springer.
Yandell, B. S. (1997) Practical Data Analysis for Designed Experiments. Chapman \& Hall.

## See Also

aov, qtukey, model.tables

## Examples

```
data(warpbreaks)
summary(fm1 <- aov(breaks ~ wool + tension, data = warpbreaks))
TukeyHSD(fm1, "tension", ordered = TRUE)
plot(TukeyHSD(fm1, "tension"))
```

type.convert Type Conversion on Character Variables

## Description

Convert a character vector to logical, integer, numeric, complex or factor as appropriate.

## Usage

type.convert(x, na.strings = "NA", as.is = FALSE, dec = ".")

## Arguments

X
na.strings a vector of strings which are to be interpreted as NA values. Blank fields are also considered to be missing values.
as.is logical. See Details.
dec the character to be assumed for decimal points.

## Details

This is principally a helper function for read.table. Given a character vector, it attempts to convert it to logical, integer, numeric or complex, and failing that converts it to factor unless as.is = TRUE. The first type that can accept all the non-missing values is chosen.
Vectors which are entirely missing values are converted to logical, since NA is primarily logical.

## Value

A vector of the selected class, or a factor.

## See Also

read.table
typeof
The Type of an Object

## Description

typeof determines the (R internal) type or storage mode of any object

## Usage

typeof(x)

## Arguments

$\mathrm{x} \quad$ any R object.

## Value

A character string.

## See Also

```
mode, storage.mode.
```


## Examples

```
typeof(2)
mode(2)
```


## UCBAdmissions Student Admissions at UC Berkeley

## Description

Aggregate data on applicants to graduate school at Berkeley for the six largest departments in 1973 classified by admission and sex.

## Usage

data(UCBAdmissions)

## Format

A 3-dimensional array resulting from cross-tabulating 4526 observations on 3 variables. The variables and their levels are as follows:

| No | Name | Levels |
| ---: | :--- | :--- |
| 1 | Admit | Admitted, Rejected |
| 2 | Gender | Male, Female |
| 3 | Dept | A, B, C, D, E, F |

## Details

This data set is frequently used for illustrating Simpson's paradox, see Bickel et al. (1975). At issue is whether the data show evidence of sex bias in admission practices. There were 2691 male applicants, of whom $1198(44.5 \%)$ were admitted, compared with 1835 female applicants of whom 557 (30.4\%) were admitted. This gives a sample odds ratio of 1.83 , indicating that males were almost twice as likely to be admitted. In fact, graphical methods (as in the example below) or log-linear modelling show that the apparent association between admission and sex stems from differences in the tendency of males and females to apply to the individual departments (females used to apply "more" to departments with higher rejection rates).

This data set can also be used for illustrating methods for graphical display of categorical data, such as the general-purpose mosaic plot or the "fourfold display" for 2-by-2-by- $k$ tables. See the home page of Michael Friendly (http://www.math.yorku.ca/SCS/friendly.html) for further information.

## References

Bickel, P. J., Hammel, E. A., and O'Connell, J. W. (1975) Sex bias in graduate admissions: Data from Berkeley. Science, 187, 398-403.

## Examples

```
data(UCBAdmissions)
## Data aggregated over departments
apply(UCBAdmissions, c(1, 2), sum)
mosaicplot(apply(UCBAdmissions, c(1, 2), sum),
    main = "Student admissions at UC Berkeley")
## Data for individual departments
opar <- par(mfrow = c(2, 3), oma = c(0, 0, 2, 0))
```

```
for(i in 1:6)
        mosaicplot(UCBAdmissions[,,i],
        xlab = "Admit", ylab = "Sex",
        main = paste("Department", LETTERS[i]))
mtext(expression(bold("Student admissions at UC Berkeley")),
            outer = TRUE, cex = 1.5)
par(opar)
```

Uniform The Uniform Distribution

## Description

These functions provide information about the uniform distribution on the interval from min to max. dunif gives the density, punif gives the distribution function qunif gives the quantile function and runif generates random deviates.

## Usage

```
dunif(x, min=0, max=1, log = FALSE)
punif(q, min=0, max=1, lower.tail = TRUE, log.p = FALSE)
qunif(p, min=0, max=1, lower.tail = TRUE, log.p = FALSE)
runif(n, min=0, max=1)
```


## Arguments

```
x,q vector of quantiles.
p vector of probabilities.
n number of observations. If length(n) > 1, the length is taken to be the
    number required.
min,max lower and upper limits of the distribution.
log, log.p logical; if TRUE, probabilities p are given as log(p).
lower.tail logical; if TRUE (default), probabilities are P[X\leqx], otherwise, P[X>
    x].
```


## Details

If min or max are not specified they assume the default values of 0 and 1 respectively.
The uniform distribution has density

$$
f(x)=\frac{1}{\max -\min }
$$

for $\min \leq x \leq \max$.
For the case of $u:=\min ==\max$, the limit case of $X \equiv u$ is assumed.

## See Also

.Random.seed about random number generation, rnorm, etc for other distributions.

## Examples

```
u <- runif(20)
## The following relations always hold :
    punif(u) == u
    dunif(u) == 1
    runif(10, 2,2) == 2
var(runif(10000))#- ~ = 1/12 = .08333
```

unique Extract Unique Elements

## Description

unique returns a vector, data frame or array like x but with duplicate elements removed.

## Usage

unique (x, incomparables = FALSE, ...)
unique. $\operatorname{array(x,~incomparables~=~FALSE,~MARGIN~}=1, \ldots$ )

## Arguments

x
an atomic vector or a data frame or an array.
incomparables a vector of values that cannot be compared. Currently, FALSE is the only possible value, meaning that all values can be compared.
... arguments for particular methods.
MARGIN the array margin to be held fixed: a single integer.

## Details

This is a generic function with methods for vectors, data frames and arrays (including matrices).

The array method calculates for each element of the dimension specified by MARGIN if the remaining dimensions are identical to those for an earlier element (in row-major order). This would most commonly be used to find unique rows (the default) or columns (with MARGIN $=2$ ).

## Value

An object of the same type of $x$. but if an element is equal to one with a smaller index, it is removed. Dimensions of arrays are not dropped.

## See Also

duplicated which gives the indices of duplicated elements.

## Examples

```
unique(c(3:5, 11:8, 8 + 0:5))
length(unique(sample(100, 100, replace=TRUE)))
## approximately 100(1 - 1/e) = 63.21
my.unique <- function(x) x[!duplicated(x)]
for(i in 1:4)
    { x <- rpois(100, pi); stopifnot(unique(x) == my.unique(x)) }
data(iris)
unique(iris)
stopifnot(dim(unique(iris)) == c(149, 5))
```

uniroot One Dimensional Root (Zero) Finding

## Description

The function uniroot searches the interval from lower to upper for a root (i.e. zero) of the function $f$ with respect to its first argument.

## Usage

```
uniroot(f, interval, lower = min(interval), upper = max(interval),
    tol = .Machine$double.eps^0.25, maxiter = 1000, ...)
```


## Arguments

| f | the function for which the root is sought. |
| :--- | :--- |
| interval | a vector containing the end-points of the interval to be searched for the <br> root. |
| lower | the lower end point of the interval to be searched. |
| upper | the upper end point of the interval to be searched. |
| tol | the desired accuracy (convergence tolerance). |
| maxiter | the maximum number of iterations. |
| $\ldots$ | additional arguments to $f$. |

## Details

Either interval or both lower and upper must be specified. The function uses Fortran subroutine "zeroin"" (from Netlib) based on algorithms given in the reference below.

If the algorithm does not converge in maxiter steps, a warning is printed and the current approximation is returned.

## Value

A list with four components: root and f.root give the location of the root and the value of the function evaluated at that point. iter and estim. prec give the number of iterations used and an approximate estimated precision for root.

## References

Brent, R. (1973) Algorithms for Minimization without Derivatives. Englewood Cliffs, NJ: Prentice-Hall.

## See Also

polyroot for all complex roots of a polynomial; optimize, nlm.

## Examples

```
f <- function (x,a) x - a
str(xmin <- uniroot(f, c(0, 1), tol = 0.0001, a = 1/3))
str(uniroot(function(x) x*(x^2-1) + .5, low = -2, up = 2, tol = 0.0001),
    dig = 10)
str(uniroot(function(x) x*(x^2-1) + .5, low = -2, up =2, tol = 1e-10 ),
    dig = 10)
## Find the smallest value x for which exp(x) > 0 (numerically):
r <- uniroot(function(x) 1e80*exp(x) -1e-300, ,-1000,0, tol=1e-20)
str(r, digits= 15)##> around -745.1332191
exp(r$r) # = 0, but not for r$r * 0.999...
minexp <- r$r * (1 - .Machine$double.eps)
exp(minexp) # typically denormalized
```

```
units Graphical Units
```


## Description

xinch and yinch convert the specified number of inches given as their arguments into the correct units for plotting with graphics functions. Usually, this only makes sense when normal coordinates are used, i.e., no log scale (see the log argument to par).
xyinch does the same for a pair of numbers xy, simultaneously.
cm translates inches in to cm (centimeters).

## Usage

```
xinch (x = 1, warn.log = TRUE)
yinch (y = 1, warn.log = TRUE)
xyinch (xy = 1, warn.log = TRUE)
cm ( x )
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | numeric vector |
| :--- | :--- |
| xy | numeric of length 1 or 2. |
| warn.log | logical; if TRUE, a warning is printed in case of active log scale. |

## Examples

```
all(c(xinch(),yinch()) == xyinch()) # TRUE
xyinch()
xyinch #- to see that is really delta{"usr"} / "pin"
cm(1)# = 2.54
## plot labels offset 0.12 inches to the right
## of plotted symbols in a plot
data(mtcars)
attach(mtcars)
plot(mpg, disp, pch=19, main= "Motor Trend Cars")
text(mpg + xinch(0.12), disp, rownames(mtcars),adj=0, cex = .7, col='blue')
detach(mtcars)
```

```
unlink Delete Files and Directories
```


## Description

unlink deletes the file(s) or directories specified by x .

## Usage

unlink(x, recursive = FALSE)

## Arguments

$x \quad a \quad$ character vector with the names of the file(s) or directories to be deleted. Wildcards (normally '*' and '?') are allowed.
recursive logical. Should directories be deleted recusively?

## Details

If recusive $=$ FALSE directories are not deleted, not even empty ones.
file.remove can only remove files, but gives more detailed error information.

## Value

The return value of the corresponding system command, rm -f, normally 0 for success, 1 for failure. Not deleting a non-existent file is not a failure.

## Note

Prior to $R$ version 1.2.0 the default on Unix was recursive $=$ TRUE, and on Windows empty directories could be deleted.

## See Also

file.remove.

```
unlist Flatten Lists
```


## Description

Given a list structure x , unlist simplifies it to produce a vector which contains all the atomic components which occur in x .

## Usage

unlist(x, recursive $=$ TRUE, use.names $=$ TRUE)

## Arguments

$\mathrm{x} \quad$ A list or vector.
recursive logical. Should unlisting be applied to list components of x ?
use.names logical. Should names be preserved?

## Details

If recursive $=$ FALSE, the function will not recurse beyond the first level items in x . x can be a vector, but then unlist does nothing useful, not even drop names.
By default, unlist tries to retain the naming information present in x . If use.names $=$ FALSE all naming information is dropped.
Where possible the list elements are coerced to a common mode during the unlisting, and so the result often ends up as a character vector.
A list is a (generic) vector, and the simplified vector might still be a list (and might be unchanged). Non-vector elements of the list (for example language elements such as names, formulas and calls) are not coerced, and so a list containing one or more of these remains a list. (The effect of unlisting an 1 m fit is a list which has individual residuals as components,)

## Value

A vector of an appropriate mode to hold the list components.

## See Also

c, as.list.

## Examples

```
unlist(options())
unlist(options(), use.names=FALSE)
l.ex <- list(a = list(1:5, LETTERS[1:5]), b = "Z", c = NA)
unlist(l.ex, recursive = FALSE)
unlist(l.ex, recursive = TRUE)
11 <- list(a="a", b=2, c=pi+2i)
unlist(l1) # a character vector
12 <- list(a="a", b=as.name("b"), c=pi+2i)
unlist(l2) # remains a list
```

```
unname Remove 'names' or 'dimnames'
```


## Description

Remove the names or dimnames attribute of an R object.

## Usage

unname(obj, force = FALSE)

## Arguments

obj the R object which is wanted "nameless".
force logical; if true, the dimnames are even removed from data.frames. This argument is currently experimental and hence might change!

## Value

Object as obj but without names or dimnames.

## Examples

```
## Answering a question on R-help (14 Oct 1999):
col3 <- 750+ 100* rt(1500, df = 3)
breaks <- factor(cut(col3,breaks=360+5*(0:155)))
str(table(breaks)) # The names are quite larger than the data ...
barplot(unname(table(breaks)), axes= FALSE)
```

```
update
Update and Re-fit a Model Call
```


## Description

update will update and (by default) re-fit a model. It does this by extracting the call stored in the object, updating the call and (by default) evaluating that call. Sometimes it is useful to call update with only one argument, for example if the data frame has been corrected.

## Usage

update(object, ...)
update.default(object, formula., ..., evaluate = TRUE)

## Arguments

object An existing fit from a model function such as 1 m , glm and many others.
formula. Changes to the formula - see update.formula for details.
... Additional arguments to the call, or arguments with changed values. Use name=NULL to remove the argument name.
evaluate If true evaluate the new call else return the call.

## Value

If evaluate $=$ TRUE the fitted object, otherwise the updated call.

## Author(s)

B.D. Ripley

## See Also

update.formula

## Examples

```
oldcon <- options(contrasts = c("contr.treatment", "contr.poly"))
## Annette Dobson (1990) "An Introduction to Generalized Linear Models".
## Page 9: Plant Weight Data.
ctl <- c(4.17,5.58,5.18,6.11,4.50,4.61,5.17,4.53,5.33,5.14)
trt <- c(4.81,4.17,4.41,3.59,5.87,3.83,6.03,4.89,4.32,4.69)
group <- gl(2, 10, 20, labels = c("Ctl", "Trt"))
weight <- c(ctl, trt)
lm.D9 <- lm(weight ~ group)
lm.D9
summary(lm.D90 <- update(lm.D9, . ~ . - 1))
options(contrasts = c("contr.helmert", "contr.poly"))
update(lm.D9)
options(oldcon)
```

update.formula

Model Updating

## Description

update.formula is used to update model formulae. This typically involves adding or dropping terms, but updates can be more general.

## Usage

update.formula(old, new, ...)

## Arguments

old a model formula to be updated.
new a formula giving a template which specifies how to update.
... further arguments passed to or from other methods.

## Details

The function works by first identifying the left-hand side and right-hand side of the old formula. It then examines the new formula and substitutes the $l h s$ of the old formula for any occurence of "." on the left of new, and substitutes the rhs of the old formula for any occurence of "." on the right of new.

## Value

The updated formula is returned.

## See Also

```
terms, model.matrix.
```


## Examples

```
update.formula(y ~ x, ~ . + x2) #> y ~ x + x2
update.formula(y ~ x, log(.) ~ . ) #> log(y) ~ x
```

update.packages Download Packages from CRAN

## Description

These functions can be used to automatically compare the version numbers of installed packages with the newest available version on CRAN and update outdated packages on the fly.

## Usage

```
update.packages(lib.loc = NULL, CRAN = getOption("CRAN"),
                    contriburl = contrib.url(CRAN),
                        method, instlib = NULL,
                            ask=TRUE, available=NULL, destdir=NULL)
installed.packages(lib.loc = NULL, priority = NULL)
CRAN.packages(CRAN = getOption("CRAN"), method = "auto",
    contriburl = contrib.url(CRAN))
old.packages(lib.loc = NULL, CRAN = getOption("CRAN"),
    contriburl = contrib.url(CRAN),
    method, available = NULL)
download.packages(pkgs, destdir, available = NULL,
            CRAN = getOption("CRAN"),
            contriburl = contrib.url(CRAN), method = "auto")
install.packages(pkgs, lib, CRAN = getOption("CRAN"),
    contriburl = contrib.url(CRAN),
        method, available = NULL, destdir = NULL)
```


## Arguments

lib.loc character vector describing the location of R library trees to search through (and update packages therein).
CRAN character, the base URL of the CRAN mirror to use, i.e., the URL of a CRAN root such as "http://cran.r-project.org" (the default) or its Statlib mirror, "http://lib.stat.cmu.edu/R/CRAN".
contriburl URL of the contrib section of CRAN. Use this argument only if your CRAN mirror is incomplete, e.g., because you burned only the contrib section on a CD. Overrides argument CRAN.

| method | Download method, see download.file. |
| :--- | :--- |
| pkgs | character vector of the short names of packages whose current versions <br> should be downloaded from CRAN. |
| destdir | directory where downloaded packages are stored. <br> character vector or NULL (default). If non-null, used to select packages; <br> priority <br> "high" is equivalent to c("base", "recommended"). |
| available | list of packages available at CRAN as returned by CRAN.packages. <br> lib,instlib |
| character string giving the library directory where to install the packages. |  |
| ask | logical indicating to ask before packages are actually downloaded and <br> installed. |

## Details

installed.packages scans the 'DESCRIPTION' files of each package found along lib.loc and returns a list of package names, library paths and version numbers. CRAN. packages returns a similar list, but corresponding to packages currently available in the contrib section of CRAN, the comprehensive R archive network. The current list of packages is downloaded over the internet (or copied from a local CRAN mirror). Both functions use read.dcf for parsing the description files. old.packages compares the two lists and reports installed packages that have newer versions on CRAN.
download.packages takes a list of package names and a destination directory, downloads the newest versions of the package sources and saves the in destdir. If the list of available packages is not given as argument, it is also directly obtained from CRAN. If CRAN is local, i.e., the URL starts with "file:", then the packages are not downloaded but used directly.

The main function of the bundle is update.packages. First a list of all packages found in lib.loc is created and compared with the packages available on CRAN. Outdated packages are reported and for each outdated package the user can specify if it should be automatically updated. If so, the package sources are downloaded from CRAN and installed in the respective library path (or instlib if specified) using the R INSTALL mechanism.
install.packages can be used to install new packages, it takes a vector of package names and a destination library, downloads the packages from CRAN and installs them. If the library is omitted it defaults to the first directory in .libPaths(), with a warning if there is more than one.

For install.packages and update.packages, destdir is the directory to which packages will be downloaded. If it is NULL (the default) a temporary directory is used, and the user will be given the option of deleting the temporary files once the packages are installed. (They will always be deleted at the end of the R session.)

## See Also

See download.file for how to handle proxies and other options to monitor file transfers.

```
INSTALL, REMOVE, library, .packages, read.dcf
```


## Examples

```
str(ip <- installed.packages(priority = "high"))
ip[, c(1,3:5)]
```

```
url.show Display a text URL
```


## Description

Extension of file.show to display text files on a remote server.

## Usage

```
url.show(url, title = url, file = tempfile(),
            delete.file = TRUE, method, ...)
```


## Arguments

| url | The URL to read from. |
| :--- | :--- |
| title | Title for the browser. |
| file | File to copy to. |
| delete.file | Delete the file afterwards? |
| method | File transfer method: see download.file |
| $\ldots$ | Arguments to pass to file.show. |

## See Also

```
url, file.show,download.file
```


## Examples

```
url.show("http://lib.stat.cmu.edu/datasets/csb/ch3a.txt")
```

USArrests Violent Crime Rates by US State

## Description

This data set contains statistics, in arrests per 100,000 residents for assault, murder, and rape in each of the 50 US states in 1973. Also given is the percent of the population living in urban areas.

## Usage

data(USArrests)

## Format

A data frame with 50 observations on 5 variables.

| $[, 1]$ | Murder | numeric | Murder arrests (per 100,000) |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | Assault | numeric | Assault arrests (per 100,000) |
| $[, 3]$ | UrbanPop | numeric | Percent urban population |
| $[, 4]$ | Rape | numeric | Rape arrests (per 100,000) |

## Source

World Almanac and Book of facts 1975. (Crime rates).
Statistical Abstracts of the United States 1975. (Urban rates).

## References

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## See Also

The state data sets.

## Examples

```
data(USArrests)
pairs(USArrests, panel = panel.smooth, main = "USArrests data")
```

USJudgeRatings Lawyers' Ratings of State Judges in the US Superior Court

## Description

Lawyers' ratings of state judges in the US Superior Court

## Usage

data(USJudgeRatings)

## Format

A data frame containing 43 observations on 12 numeric variables.

| $[, 1]$ | CONT | Number of contacts of lawyer with judge. |
| :--- | :--- | :--- |
| $[, 2]$ | INTG | Judicial integrity. |
| $[, 3]$ | DMNR | Demeanor. |
| $[, 4]$ | DILG | Diligence. |
| $[, 5]$ | CFMG | Case flow managing. |
| $[, 6]$ | DECI | Prompt decisions. |
| $[, 7]$ | PREP | Preparation for trial. |
| $[, 8]$ | FAMI | Familiarity with law. |
| $[, 9]$ | ORAL | Sound oral rulings. |
| $[, 10]$ | WRIT | Sound written rulings. |
| $[, 11]$ | PHYS | Physical ability. |
| $[, 12]$ | RTEN | Worthy of retention. |

## Source

New Haven Register, 14 January, 1977 (from John Hartigan).

## Examples

data(USJudgeRatings)

```
pairs(USJudgeRatings, main = "USJudgeRatings data")
```

USPersonalExpenditure

Personal Expenditure Data

## Description

This data set consists of United States personal expenditures (in billions of dollars) in the categories; food and tobacco, household operation, medical and health, personal care, and private education for the years 1940, 1945, 1950, and 1960.

## Usage

data(USPersonalExpenditure)

## Format

A matrix with 5 rows and 5 columns.

## Source

The World Almanac and Book of Facts, 1962, page 756.

## References

Tukey, J. W. (1977) Exploratory Data Analysis. Addison-Wesley.
McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(USPersonalExpenditure)
USPersonalExpenditure
require(eda)
medpolish(log10(USPersonalExpenditure))
```


## uspop The Population of the United States

## Description

This data set gives the population of the United States (in millions) as recorded by the decennial census for the period 1790-1970.

## Usage

data(uspop)

## Format

A time series of 19 values.

## Source

McNeil, D. R. (1977) Interactive Data Analysis. New York: Wiley.

## Examples

```
data(uspop)
plot(uspop, log = "y", main = "uspop data", xlab = "Year",
    ylab = "U.S. Population (millions)")
```


## VADeaths Death Rate Data

## Description

Death rates per 100 in Virginia in 1940.

## Usage

data(VADeaths)

## Format

A matrix with 5 rows and 5 columns.

## Details

The death rates are cross-classified by age group (rows) and population group (columns). The age groups are: $50-54,55-59,60-64,65-69,70-74$ and the population groups are Rural/Male, Rural/Female, Urban/Male and Urban/Female.

This provides a rather nice 3 -way analysis of variance example.

## Source

Moyneau, L., Gilliam, S. K., and Florant, L. C.(1947) Differences in Virginia death rates by color, sex, age, and rural or urban residence. American Sociological Review, 12, 525-535.

## References

McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(VADeaths)
n <- length(dr <- c(VADeaths))
nam <- names(VADeaths)
d.VAD <- data.frame(
    Drate = dr,
age = rep(ordered(rownames(VADeaths)),length=n),
gender= gl(2,5,n, labels= c("M", "F")),
site = gl(2,10, labels= c("rural", "urban")))
coplot(Drate ~ as.numeric(age) | gender * site, data = d.VAD,
    panel = panel.smooth, xlab = "VADeaths data - Given: gender")
summary(aov.VAD <- aov(Drate ~ . 2 2, data = d.VAD))
```

```
opar <- par(mfrow = c(2,2), oma = c(0, 0, 1.1, 0))
plot(aov.VAD)
par(opar)
```

    vcov Calculate Variance-Covariance Matrix for a Fitted Model Object
    
## Description

Returns the variance-covariance matrix of the main parameters of a fitted model object.

## Usage

vcov(object, ...)

## Arguments

object a fitted model object.
... additional arguments for method functions. For the glm method this can be used to pass a dispersion parameter.

## Details

This is a generic function. Functions with names beginning in vcov. will be methods for this function. Classes with methods for this function include: lm, glm, nls, lme, gls, coxph and survreg

## Value

A matrix of the estimated covariances between the parameter estimates in the linear or non-linear predictor of the model.

| vector Vectors |
| :--- | :--- |

## Description

vector produces a vector of the given length and mode.
as.vector, a generic, attempts to coerce its argument into a vector of mode mode (the default is to coerce to whichever mode is most convenient). The attributes of x are removed.
is.vector returns TRUE if $x$ is a vector (of mode logical, integer, real, complex, character or list if not specified) and FALSE otherwise.

## Usage

```
vector(mode = "logical", length = 0)
as.vector(x, mode = "any")
is.vector(x, mode = "any")
```


## Arguments

| mode | A character string giving an atomic mode, or "any". |
| :--- | :--- |
| length | A non-negative integer specifying the desired length. |
| x | An object. |

## Details

is.vector returns FALSE if x has any attributes except names. (This is incompatible with S.) On the other hand, as.vector removes all attributes including names.

Note that factors are not vectors; is.vector returns FALSE and as.vector converts to character mode.

## Value

For vector, a vector of the given length and mode. Logical vector elements are initialized to FALSE, numeric vector elements to 0 and character vector elements to "".

## See Also

c, is.numeric, is.list, etc.

## Examples

```
df <- data.frame(x=1:3, y=5:7)
## Error:
    as.vector(data.frame(x=1:3, y=5:7), mode="numeric")
x <- c(a = 1, b = 2)
is.vector(x)
as.vector(x)
all.equal(x, as.vector(x)) ## FALSE
###-- All the following are TRUE:
is.list(df)
! is.vector(df)
! is.vector(df, mode="list")
is.vector(list(), mode="list")
is.vector(NULL, mode="NULL")
```

```
volcano
```

Topographic Information for the Maunga Whau Volcano

## Description

Maunga Whau (Mt Eden) is one of about 50 volcanos in the Auckland volcanic field. This data set gives topographic information for Maunga Whau on a 10 m by 10 m grid.

## Usage

data(volcano)

## Format

A matrix with 87 rows and 61 columns, rows corresponding to grid lines running east to west and columns to grid lines running south to north.

## Source

Digitized from a topographic map by Ross Ihaka. These data should not be regarded as accurate.

## See Also

filled.contour for a nice plot.

## Examples

```
data(volcano)
filled.contour(volcano, color = terrain.colors, asp = 1)
title(main = "volcano data: filled contour map")
```


## warning Warning Messages

## Description

Generates a warning message that corresponds to its argument(s) and (optionally) the expression or function from which it was called.

## Usage

warning(..., call. = TRUE)

## Arguments

... character vectors (which are pasted together with no separator) or NULL.
call. logical, indicating if the call should become part of the warning message.

## Details

The result depends on the value of options("warn").
If warn is negative warnings are ignored; if it is zero they are stored and printed after the top-level function has completed; if it is one they are printed as they occur and if it is 2 (or larger) warnings are turned into errors.

If warn is zero (the default), a top-level variable last.warning is created. It contains the warnings which can be printed via a call to warnings.
Warnings will be truncated to getOption("warning.length") characters, default 1000.

## See Also

stop for fatal errors, warnings, and options with argument warn=.

## Examples

```
testit <- function() warning("testit")
testit() ## shows call
testit <- function() warning("problem in testit", call. = FALSE)
testit() ## no call
```

warnings
Print Warning Messages

## Description

warnings prints the top-level variable last.warning in a pleasing form.

## Usage

warnings(...)

## Arguments

... arguments to be passed to cat.

## See Also

warning.

## Examples

```
ow <- options("warn")
for(w in -1:1) {
    options(warn = w); cat("\n warn =",w,"\n")
    for(i in 1:3) { cat(i,"..\n"); m <- matrix(1:7, 3,4) }
}
warnings()
options(ow) # reset
```

warpbreaks The Number of Breaks in Yarn during Weaving

## Description

This data set gives the number of warp breaks per loom, where a loom corresponds to a fixed length of yarn.

## Usage

data(warpbreaks)

## Format

A data frame with 54 observations on 3 variables.

| $[, 1]$ | breaks | numeric | The number of breaks |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | wool | factor | The type of wool (A or B) |
| $[, 3]$ | tension | factor | The level of tension (L, M, H) |

There are measurements on 9 looms for each of the six types of warp (AL, AM, AH, BL, BM, BH).

## Source

Tippett, L. H. C. (1950) Technological Applications of Statistics. Wiley. Page 106.

## References

Tukey, J. W. (1977) Exploratory Data Analysis. Addison-Wesley.
McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## See Also

xtabs for ways to display these data as a table.

## Examples

```
data(warpbreaks)
summary(warpbreaks)
opar <- par(mfrow = c(1,2), oma = c(0, 0, 1.1, 0))
plot(breaks ~ tension, data = warpbreaks, col = "lightgray",
    varwidth = TRUE, subset = wool == "A", main = "Wool A")
plot(breaks ~ tension, data = warpbreaks, col = "lightgray",
    varwidth = TRUE, subset = wool == "B", main = "Wool B")
mtext("warpbreaks data", side = 3, outer = TRUE)
par(opar)
summary(fm1 <- lm(breaks ~ wool*tension, data = warpbreaks))
anova(fm1)
```

weekdays
Extract Parts of a POSIXt Object

## Description

Extract the weekday, month or quarter, or the Julian time (days since some origin). These are generic functions: the methods for the internal date-time classes are documented here.

## Usage

```
weekdays(x, abbreviate = FALSE)
months(x, abbreviate = FALSE)
quarters(x, ...)
julian(x, origin = as.POSIXct("1970-01-01", tz="GMT"), ...)
```


## Arguments

```
x an object inheriting from class "POSIXt".
abbreviate logical. Should the names be abbreviated?
origin an length-one object inheriting from class "POSIXt".
... arguments for other methods.
```


## Value

weekdays and months return a character vector of names in the locale in use.
quarters returns a character vector of "Q1" to "Q4".
julian returns the number of days (possibly fractional) since the origin, with the origin as a "origin" attribute.

## Note

Other components such as the day of the month or the year are very easy to compute: just use as.POSIXIt and extract the relevant component.

See Also

DateTimeClasses

## Examples

```
weekdays(.leap.seconds)
months(.leap.seconds)
quarters(.leap.seconds)
```

Weibull The Weibull Distribution

## Description

Density, distribution function, quantile function and random generation for the Weibull distribution with parameters shape and scale.

## Usage

```
dweibull(x, shape, scale = 1, log = FALSE)
pweibull(q, shape, scale = 1, lower.tail = TRUE, log.p = FALSE)
qweibull(p, shape, scale = 1, lower.tail = TRUE, log.p = FALSE)
rweibull(n, shape, scale = 1)
```


## Arguments

| $\mathrm{x}, \mathrm{q}$ | vector of quantiles. |
| :--- | :--- |
| p | vector of probabilities. |
| n | number of observations. If length $(\mathrm{n})>1$, the length is taken to be the <br> number required. |
| shape, scale | shape and scale parameters, the latter defaulting to 1. |
| log, log.p | logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$. <br> lower.tail |
| logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ <br> $x]$. |  |

## Details

The Weibull distribution with shape parameter $a$ and scale parameter $\sigma$ has density given by

$$
f(x)=(a / \sigma)(x / \sigma)^{a-1} \exp \left(-(x / \sigma)^{a}\right)
$$

for $x>0$. The cumulative is $F(x)=1-\exp \left(-(x / \sigma)^{a}\right)$, the mean is $E(X)=\sigma \Gamma(1+1 / a)$, and the $\operatorname{Var}(X)=\sigma^{2}\left(\Gamma(1+2 / a)-(\Gamma(1+1 / a))^{2}\right)$.

## Value

dweibull gives the density, pweibull gives the distribution function, qweibull gives the quantile function, and rweibull generates random deviates.

## Note

The cumulative hazard $H(t)=-\log (1-F(t))$ is -pweibull(t, a, b, lower = FALSE, $\log =$ TRUE $)$ which is just $H(t)=(t / b)^{a}$.

## See Also

dexp for the Exponential which is a special case of a Weibull distribution.

## Examples

```
x <- c(0,rlnorm(50))
all.equal(dweibull(x, shape = 1), dexp(x))
all.equal(pweibull(x, shape = 1, scale = pi), pexp(x, rate = 1/pi))
## Cumulative hazard H():
all.equal(pweibull(x, 2.5, pi, lower=FALSE, log=TRUE), -(x/pi)^2.5, tol=1e-15)
all.equal(qweibull(x/11, shape = 1, scale = pi), qexp(x/11, rate = 1/pi))
```

```
weighted.mean Weighted Arithmetic Mean
```


## Description

Compute a weighted mean of a numeric vector.

## Usage

```
weighted.mean(x, w, na.rm=FALSE)
```


## Arguments

x
w a vector of weights the same length as x giving the weights to use for each element of x .
na.rm a logical value indicating whether NA values in $x$ should be stripped before the computation proceeds.

## Details

If w is missing then all elements of x are given the same weight.
Missing values in w are not handled.

## See Also

mean

## Examples

```
## GPA from Siegel }199
wt <- c(5, 5, 4, 1)/15
x <- c(3.7,3.3,3.5,2.8)
xm <- weighted.mean(x,wt)
```

weighted.residuals Compute Weighted Residuals

## Description

Computed weighted residuals from a linear model fit.

## Usage

weighted.residuals(obj, drop0 = TRUE)

## Arguments

obj $\quad$ R object, typically of class lm or glm.
drop0 logical. If TRUE, drop all cases with weights $==0$.

## Details

Weighted residuals are the usual residuals $R_{i}$, multiplied by $\sqrt{w_{i}}$, where $w_{i}$ are the weights as specified in lm's call.
Dropping cases with weights zero is compatible with lm.influence and related functions.

## Value

Numeric vector of length $n^{\prime}$, where $n^{\prime}$ is the number of of non- 0 weights (drop0 $=$ TRUE) or the number of observations, otherwise.

## See Also

```
residuals,lm.influence, etc.
```


## Examples

```
example("lm")
all.equal(weighted.residuals(lm.D9),
    residuals(lm.D9))
x <- 1:10
w <- 0:9
y <- rnorm(x)
weighted.residuals(lmxy <- lm(y ~ x, weights = w))
weighted.residuals(lmxy, drop0 = FALSE)
```

which
Which indices are TRUE?

## Description

Give the TRUE indices of a logical object, allowing for array indices.

## Usage

which ( x , arr. ind = FALSE)

## Arguments

| x | a logical vector or array. NAs are allowed and omitted (treated as if |
| :--- | :--- |
| FALSE). |  |
| arr.ind | logical; should array indices be returned when x is an array? |

## Value

If arr.ind == FALSE (the default), an integer vector with length equal to sum(x), i.e., to the number of TRUEs in x ; Basically, the result is (1:length $(\mathrm{x})$ ) [ x ].
If arr.ind == TRUE and $x$ is an array (has a dim attribute), the result is a matrix who's rows each are the indices of one element of x ; see Examples below.

## Author(s)

Werner Stahel and Peter Holzer 〈holzer@stat.math.ethz.ch〉, for the array case.

## See Also

Logic, which.min for the index of the minimum or maximum.

## Examples

```
which(LETTERS == "R")
which(ll <- c(TRUE,FALSE,TRUE,NA,FALSE,FALSE,TRUE))#> 1 3 7
names(ll) <- letters[seq(ll)]
which(11)
which((1:12)%%2 == 0) # which are even?
str(which(1:10 > 3, arr.ind=TRUE))
( m <- matrix(1:12,3,4) )
which(m %% 3 == 0)
which(m %% 3 == 0, arr.ind=TRUE)
rownames(m) <- paste("Case",1:3, sep="_")
which(m %% 5 == 0, arr.ind=TRUE)
dim(m) <- c(2,2,3); m
which(m %% 3 == 0, arr.ind=FALSE)
which(m %% 3 == 0, arr.ind=TRUE)
vm <- c(m)
dim(vm) <- length(vm) #-- funny thing with length(dim(...)) == 1
which(vm %% 3 == 0, arr.ind=TRUE)
```

```
which.min Where is the Min() or Max() ?
```


## Description

Determines the location, i.e., index of the (first) minimum or maximum of a numeric vector.

## Usage

which.min(x)
which.max (x)

## Arguments

x
numeric vector, whose min or max is searched.

## Value

an integer of length 1 or 0 (iff x has no non-NAs), giving the index of the first minimum or maximum respectively of x .
If this extremum is unique (or empty), the result is the same (but more efficient) as which (x $==\min (x))$ or which $(x==\max (x))$ respectively.

## Author(s)

Martin Maechler

## See Also

which, max.col, max, etc.

## Examples

```
x <- c(1:4,0:5,11)
which.min(x)
which.max(x)
data(presidents)
presidents[1:30]
range(presidents, na.rm = TRUE)
which.min(presidents)# 28
which.max(presidents)# 2
```


## Wilcoxon Distribution of the Wilcoxon Rank Sum Statistic

## Description

Density, distribution function, quantile function and random generation for the distribution of the Wilcoxon rank sum statistic obtained from samples with size $m$ and $n$, respectively.

## Usage

```
dwilcox(x, m, n, log = FALSE)
pwilcox(q, m, n, lower.tail = TRUE, log.p = FALSE)
qwilcox(p, m, n, lower.tail = TRUE, log.p = FALSE)
rwilcox(nn, m, n)
```


## Arguments

$x, q \quad$ vector of quantiles.
$\mathrm{p} \quad$ vector of probabilities.
nn number of observations. If length (nn) > 1, the length is taken to be the number required.
$m, n \quad$ numbers of observations in the first and second sample, respectively.
$\log$, log.p logical; if TRUE, probabilities p are given as $\log (\mathrm{p})$.
lower.tail logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X>$ $x]$.

## Details

This distribution is obtained as follows. Let x and y be two random, independent samples of size $m$ and $n$. Then the Wilcoxon rank sum statistic is the number of all pairs ( $x[i], y[j]$ ) for which $y[j]$ is not greater than $x[i]$. This statistic takes values between 0 and $m * n$, and its mean and variance are $m * n / 2$ and $m * n *(m+n+1) / 12$, respectively.

## Value

dwilcox gives the density, pwilcox gives the distribution function, qwilcox gives the quantile function, and rwilcox generates random deviates.

## Author(s)

Kurt Hornik 〈hornik@ci.tuwien.ac.at〉

## See Also

dsignrank etc, for the one-sample Wilcoxon rank statistic.

## Examples

```
x <- -1:(4*6 + 1)
fx <- dwilcox(x, 4, 6)
all(fx == dwilcox(x, 6, 4))
Fx <- pwilcox(x, 4, 6)
all(abs(Fx - cumsum(fx)) < 10 * .Machine$double.eps)
layout(rbind(1,2),width=1,heights=c(3,2))
plot(x, fx,type='h', col="violet",
    main= "Probabilities (density) of Wilcoxon-Statist.(n=6,m=4)")
plot(x, Fx,type="s", col="blue",
    main= "Distribution of Wilcoxon-Statist.(n=6,m=4)")
abline(h=0:1, col="gray20",lty=2)
layout(1)# set back
N <- 200
hist(U <- rwilcox(N, m=4,n=6), breaks=0:25 - 1/2, border="red", col="pink",
    sub = paste("N =",N))
mtext("N * f(x), f() = true ''density')", side=3, col="blue")
    lines(x, N*fx, type='h', col='blue', lwd=2)
points(x, N*fx, cex=2)
## Better is a Quantile-Quantile Plot
qqplot(U, qw <- qwilcox((1:N - 1/2)/N, m=4,n=6),
        main = paste("Q-Q-Plot of empirical and theoretical quantiles",
            "Wilcoxon Statistic, (m=4, n=6)",sep="\n"))
n <- as.numeric(names(print(tU <- table(U))))
text(n+.2, n+.5, labels=tU, col="red")
```

window

Time Windows

## Description

window is a generic function which extracts the subset of the object x observed between the times start and end. If a frequency is specified, the series is then re-sampled at the new frequency.

## Usage

```
window(x, ...)
window.default(x, start = NULL, end = NULL,
frequency \(=\) NULL, deltat \(=\) NULL, extend \(=\) FALSE,... )
window.ts \((x\), start \(=\) NULL, end \(=\) NULL,
                frequency \(=\) NULL, deltat \(=\) NULL, extend \(=\) FALSE,.. )
```


## Arguments

$x \quad$ a time-series or other object.
start the start time of the period of interest.
end the end time of the period of interest.
frequency, deltat
the new frequency can be specified by either (or both if they are consistent).
extend logical. If true, the start and end values are allowed to extend the series. If false, attempts to extend the series give a warning and are ignored.
... further arguments passed to or from other methods.

## Details

The start and end times can be specified as for ts. If there is no observation at the new start or end, the immediately following (start) or preceding (end) observation time is used.

## Value

The value depends on the method. window. default will return a vector or matrix with an appropriate tsp attribute.
window.ts differs from window. default only in ensuring the result is a ts object.
If extend $=$ TRUE the series will be padded with NA if needed.

## See Also

time, ts.

## Examples

```
data(presidents)
window(presidents, 1960, c(1969,4)) # values in the 1960's
window(presidents, deltat=1) # All Qtr1s
window(presidents, start=c(1945,3), deltat=1) # All Qtr3s
window(presidents, 1944, c(1979,2), extend=TRUE)
```

with Evaluate an Expression in a Data Environment

## Description

Evaluate an R expression in an environment constructed from data.

## Usage

with(data, expr, ...)

## Arguments

data data to use for constructing an environment. For the default method this may be an environment, a list, a data frame, or an integer as in sys.call.
expr expression to evaluate.
... arguments to be passed to future methods.

## Details

with is a generic function that evaluates expr in a local environment constructed from data. The environment has the caller's environment as its parent. This is useful for simplifying calls to modeling functions.

```
See Also
evalq.
```


## Examples

```
#examples from glm:
library(MASS)
data(anorexia)
with(anorexia, {
    anorex.1 <- glm(Postwt ~ Prewt + Treat + offset(Prewt),
                    family = gaussian)
    summary(anorex.1)
})
with(data.frame(u = c(5,10,15,20,30,40,60,80,100),
    lot1 = c(118,58,42,35,27,25,21,19,18),
    lot2 = c(69,35,26,21,18,16,13,12,12)),
        list(summary(glm(lot1 ~ log(u), family=Gamma)),
            summary(glm(lot2 ~ log(u), family=Gamma))))
# example from boxplot:
data(ToothGrowth)
with(ToothGrowth, {
        boxplot(len ~ dose, boxwex = 0.25, at = 1:3 - 0.2,
            subset= supp == "VC", col="yellow",
            main="Guinea Pigs' Tooth Growth",
            xlab="Vitamin C dose mg",
            ylab="tooth length", ylim=c(0,35))
        boxplot(len ~ dose, add = TRUE, boxwex = 0.25, at = 1:3 + 0.2,
            subset= supp == "OJ", col="orange")
        legend(2, 9, c("Ascorbic acid", "Orange juice"),
                fill = c("yellow", "orange"))
})
# alternate form that avoids subset argument:
with(subset(ToothGrowth, supp == "VC"),
    boxplot(len ~ dose, boxwex = 0.25, at = 1:3 - 0.2,
            col="yellow", main="Guinea Pigs' Tooth Growth",
            xlab="Vitamin C dose mg",
            ylab="tooth length", ylim=c(0,35)))
```

```
with(subset(ToothGrowth, supp == "OJ"),
    boxplot(len ~ dose, add = TRUE, boxwex = 0.25, at = 1:3 + 0.2,
            col="orange"))
legend(2, 9, c("Ascorbic acid", "Orange juice"),
        fill = c("yellow", "orange"))
```

    women Average Heights and Weights for American Women
    
## Description

This data set gives the average heights and weights for American women aged 30-39.

## Usage

data(women)

## Format

A data frame with 15 observations on 2 variables.

| $[, 1]$ | height | numeric | Height (in) |
| :--- | :--- | :--- | :--- |
| $[, 2]$ | weight | numeric | Weight (lbs) |

## Details

The data set appears to have been taken from the American Society of Actuaries Build and Blood Pressure Study for some (unknown to us) earlier year.
The World Almanac notes: "The figures represent weights in ordinary indoor clothing and shoes, and heights with shoes.

## Source

The World Almanac and Book of Facts, 1975.

## References

McNeil, D. R. (1977) Interactive Data Analysis. Wiley.

## Examples

```
data(women)
plot(women, xlab = "Height (in)", ylab = "Weight (lb)",
    main = "women data: American women aged 30-39")
```

[^2]
## Description

The data (usually a matrix) $x$ are written to file file. If $x$ is a two-dimensional matrix you need to transpose it to get the columns in file the same as those in the internal representation.

## Usage

```
write(x, file = "data",
    ncolumns = if(is.character(x)) 1 else 5,
    append = FALSE)
```


## Arguments

x
file A connection, or a character string naming the file to write to. If "", print to the standard output connection. If it is " $\mid$ cmd", the output is piped to the command given by 'cmd'.
ncolumns the number of columns to write the data in.
append if TRUE the data $x$ is appended to file file.

## See Also

save for writing any R objects, write.table for data frames, and scan for reading data.

## Examples

```
# create a 2 by 5 matrix
x <- matrix(1:10,ncol=5)
# the file data contains x, two rows, five cols
# 1 3 5 6 9 will form the first row
write(t(x))
# the file data now contains the data in x,
# two rows, five cols but the first row is 1 2 3 4 5
write(x)
unlink("data") # tidy up
```

```
write.table Data Output
```


## Description

write.table prints its required argument x (after converting it to a data frame if it is not one already) to file. The entries in each line (row) are separated by the value of sep.

## Usage

```
write.table(x, file = "", append = FALSE, quote = TRUE, sep = " ",
    eol \(=\) "\n", na \(=\) "NA", dec \(=" . "\), row.names = TRUE,
    col.names \(=\) TRUE, qmethod \(=c(" e s c a p e ", ~ " d o u b l e "))\)
```


## Arguments

X
file either a character string naming a file or a connection. " " indicates output to the console.
append
quote
sep the field separator string. Values within each row of $x$ are separated by
eol the character(s) to print at the end of each line (row).
na the string to use for missing values in the data.
dec the string to use for decimal points.
row. names either a logical value indicating whether the row names of $x$ are to be
col.names either a logical value indicating whether the column names of $x$ are to be
the object to be written, typically a data frame. If not, it is attempted to coerce x to a data frame.
logical. If TRUE, the output is appended to the file. If FALSE, any existing file of the name is destroyed.
a logical or a numeric vector. If TRUE, any character or factor columns will be surrounded by double quotes. If a numeric vector, its elements are taken as the indices of the variable (columns) to quote. In both cases, row and columns names are quoted if they are written, but not if quote is FALSE. this string. written along with x , or a character vector of row names to be written. written along with x , or a character vector of column names to be written.
qmethod a character string specifying how to deal with embedded double quote characters when quoting strings. Must be one of "escape" (default), in which case the quote character is escaped in C style by a backslash, or "double", in which case it is doubled. You can specify just the initial letter.

## Details

Normally there is no column name for a column of row names. If col.names=NA a blank column name is added. This can be used to write CSV files for input to spreadsheets.
write. table can be slow for data frames with large numbers (hundreds or more) of columns: this is inevitable as each column could be of a different class and so must be handled separately. Function write.matrix in package MASS may be much more efficient if x is a matrix or can be represented in a numeric matrix.

## See Also

The ' $R$ Data Import/Export' manual.

```
read.table, write.
```

write.matrix.

## Examples

```
## To write a CSV file for input to Excel one might use
write.table(x, file = "foo.csv", sep = ",", col.names = NA)
## and to read this file back into R one needs
read.table("file.csv", header = TRUE, sep = ",", row.names=1)
```

```
writeLines Write Lines to a Connection
```


## Description

Write text lines to a connection.

## Usage

```
writeLines(text, con = stdout(), sep = "\n")
```


## Arguments

| text | A character vector |
| :--- | :--- |
| con | A connection object or a character string. |
| sep | character. A string to be written to the connection after each line of text. |

## Details

If the con is a character string, the functions call file to obtain an file connection which is opened for the duration of the function call.

If the connection is open it is written from its current position. If it is not open, it is opened for the duration of the call and then closed again.

Normally writeLines is used with a text connection, and the default separator is converted to the normal separator for that platform (LF on Unix/Linux, CRLF on Windows, LF on Macintosh). For more control, open a binary connection and specify the precise value you want written to the file in sep. For even more control, use writeChar on a binary connection.

## See Also

```
connections, writeChar, readLines, cat
```

```
x11 X Window System Graphics
```


## Description

X11 starts a graphics device driver for the X Window System (version 11). This can only be done on machines that run $\mathrm{X} . \mathrm{x} 11$ is recognized as a synonym for X11.

## Usage

```
X11(display = "", width = 7, height = 7, pointsize = 12,
    gamma = 1, colortype = getOption("X11colortype"),
    maxcubesize = 256, canvas = "white")
```


## Arguments

| display | the display on which the graphics window will appear. The default is to <br> use the value in the user's environment variable DISPLAY. |
| :--- | :--- |
| width | the width of the plotting window in inches. |
| height | the height of the plotting window in inches. |
| pointsize | the default pointsize to be used. |
| gamma | the gamma correction factor. This value is used to ensure that the colors <br> displayed are linearly related to RGB values. A value of around 0.5 is <br> appropriate for many PC displays. A value of 1.0 (no correction) is usually <br> appropriate for high-end displays or Macintoshs. |
| colortype | the kind of color model to be used. The possibilities are "mono", "gray", <br> "pseudo", "pseudo. cube" and "true". Ignored if an X11 is already open. |
| maxcubesize | can be used to limit the size of color cube allocated for pseudocolor de- <br> vices. |
| canvas | color. The color of the canvas, which is visible only when the background <br> color is transparent. |

## Details

By default, an X11 device will use the best color rendering strategy that it can. The choice can be overriden with the colortype parameter. A value of "mono" results in black and white graphics, "gray" in grayscale and "true" in truecolor graphics (if this is possible). The values "pseudo" and "pseudo.cube" provide color strategies for pseudocolor displays. The first strategy provides on-demand color allocation which produces exact colors until the color resources of the display are exhausted. The second causes a standard color cube to be set up, and requested colors are approximated by the closest value in the cube. The default strategy for pseudocolor displays is "pseudo".
Note: All X11 devices share a colortype which is set by the first device to be opened. To change the colortype you need to close all open X11 devices then open one with the desired colortype.

With colortype equal to "pseudo.cube" or "gray" successively smaller palettes are tried until one is completely allocated. If allocation of the smallest attempt fails the device will revert to "mono".

## See Also

Devices.

## xfig XFig Graphics Device

## Description

xfig starts the graphics device driver for producing XFig (version 3.2) graphics.
The auxiliary function ps.options can be used to set and view (if called without arguments) default values for the arguments to xfig and postscript.

## Usage

```
xfig(file = "Rplots.fig", onefile = FALSE, ...)
```


## Arguments

$\left.\left.\begin{array}{ll}\text { file } & \begin{array}{l}\text { a character string giving the name of the file. If it is "", the output } \\ \text { is piped to the command given by the argument command. For use with } \\ \text { onefile=FALSE give a printf format such as "Rplot\%d.fig" (the default } \\ \text { in that case). }\end{array} \\ \text { onefile } \\ \text { logical: if true allow multiple figures in one file. If false, assume only one } \\ \text { page per file and generate a file number containing the page number. } \\ \text { further options for xfig(), such as: }\end{array}\right] \begin{array}{l}\text { the size of paper in the printer. The choices are "A4", "Letter" and } \\ \text { paper } \\ \text { "Legal" (and these can be lowercase). A further choice is "default", } \\ \text { which is the default. If this is selected, the papersize is taken from the } \\ \text { option "papersize" if that is set and to "A4" if it is unset or empty. }\end{array}\right\}$

## Details

Although xfig can produce multiple plots in one file, the XFig format does not say how to separate or view them. So onefile=FALSE is the default.

## Note

On some line textures ( 0 <= lty > 4) are used. Eventually this will be partially remedied, but the XFig file format does not allow as general line textures as the R model. Unimplemented line textures are displayed as dash-double-dotted.

There is a limit of 512 colours (plus white and black) per file.

## See Also

Devices, postscript.

```
xtabs
```


## Cross Tabulation

## Description

Create a contingency table from cross-classifying factors, usually contained in a data frame, using a formula interface.

## Usage

```
xtabs(formula \(=\sim\)., data \(=\) parent.frame(), subset, na.action,
            exclude \(=c(N A, N a N), ~ d r o p . u n u s e d . l e v e l s=F A L S E)\)
```


## Arguments

formula a formula object with the cross-classifying variables, separated by + , on the right hand side. Interactions are not allowed. On the left hand side, one may optionally give a vector or a matrix of counts; in the latter case, the columns are interpreted as corresponding to the levels of a variable. This is useful if the data has already been tabulated, see the examples below.
data a data frame, list or environment containing the variables to be crosstabulated.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs.
exclude a vector of values to be excluded when forming the set of levels of the classifying factors.
drop.unused.levels
a logical indicating whether to drop unused levels in the classifying factors. If this is FALSE and there are unused levels, the table will contain zero marginals, and a subsequent chi-squared test for independence of the factors will not work.

## Details

There is a summary method for contingency table objects created by table or xtabs, which gives basic information and performs a chi-squared test for independence of factors (note that the function chisq. test in package ctest currently only handles 2-d tables).
If a left hand side is given in formula, its entries are simply summed over the cells corresponding to the right hand side; this also works if the lhs does not give counts.

## Value

A contingency table in array representation of class c("xtabs", "table"), with a "call" attribute storing the matched call.

## See Also

table for "traditional" cross-tabulation, and as.data.frame.table which is the inverse operation of xtabs (see the DF example below).

## Examples

```
data(esoph)
## 'esoph' has the frequencies of cases and controls for all levels of
## the variables 'agegp', 'alcgp', and 'tobgp'.
xtabs(cbind(ncases, ncontrols) ~ ., data = esoph)
## Output is not really helpful ... flat tables are better:
ftable(xtabs(cbind(ncases, ncontrols) ~ ., data = esoph))
## In particular if we have fewer factors ...
ftable(xtabs(cbind(ncases, ncontrols) ~ agegp, data = esoph))
data(UCBAdmissions)
## This is already a contingency table in array form.
DF <- as.data.frame(UCBAdmissions)
## Now 'DF' is a data frame with a grid of the factors and the counts
## in variable 'Freq'.
DF
## Nice for taking margins ...
xtabs(Freq ~ Gender + Admit, DF)
## And for testing independece ...
summary(xtabs(Freq ~ ., DF))
data(warpbreaks)
## Create a nice display for the warp break data.
warpbreaks$replicate <- rep(1:9, len = 54)
ftable(xtabs(breaks ~ wool + tension + replicate, data = warpbreaks))
```

xy.coords

Extracting Plotting Structures

## Description

xy. coords is used by many function to obtain $x$ and $y$ coordinates for plotting. The use of this common mechanism across all R functions produces a measure of consistency.
plot.default and lowess are examples of functions which use this mechanism.

## Usage

xy.coords(x, y, xlab=NULL, ylab=NULL, log=NULL, recycle = FALSE)

## Arguments

$\mathrm{x}, \mathrm{y}$
the x and y coordinates of a set of points. Alternatively, a single argument
x can be be provided. In this case, an attempt is made to interpret the
argument in a way suitable for plotting. If the argument is a formula yvar
$\sim \sim$ xvar, xvar and yvar are used as x and y variables; if the argument is a
list containing components x and y , these are used are assumed to define
plotting coordinates; if the argument contains a time series, the x values
are taken to be time and the y values to be the time series; if the argument
is a matrix with two columns, the first is assumed to contain the x values
and the second the y values; in any other case, the argument is coerced
to a vector and the values plotted against their indices.
$\mathrm{xlab}, \mathrm{ylab} \quad$ names for the x and y variables to be extracted.
log character, "x", "y" or both, as for plot. Sets negative values to NA and gives a warning.
recycle logical; if TRUE, recycle (rep) the shorter of $x$ or $y$ if their lengths differ.

## Value

A list with the components

```
x numeric (i.e. "double") vector of abscissa values.
y numeric vector of the same length as x.
xlab character(1) or NULL, the 'label' of x.
ylab character(1) or NULL, the 'label' of y.
```


## Examples

```
xy.coords(fft(c(1:10)), NULL)
data(cars) ; attach(cars)
xy.coords(dist ~ speed, NULL)$xlab # = "speed"
str(xy.coords(1:3, 1:2, recycle=TRUE))
str(xy.coords(-2:10,NULL, log="y"))
##> warning: 3 y values <=0 omitted ..
```

xyz.coords

Extracting Plotting Structures

## Description

Utility for obtaining consistent $\mathrm{x}, \mathrm{y}$ and z coordinates and labels for three dimensional (3D) plots.

## Usage

```
xyz.coords(x, y, z, xlab=NULL, ylab=NULL, zlab=NULL, log=NULL,
    recycle=FALSE)
```


## Arguments

$$
\begin{aligned}
& \mathrm{x}, \mathrm{y}, \mathrm{z} \\
& \text { the } \mathrm{x}, \mathrm{y} \text { and } \mathrm{z} \text { coordinates of a set of points. Alternatively, a single } \\
& \text { argument } \mathrm{x} \text { can be be provided. In this case, an attempt is made to } \\
& \text { interpret the argument in a way suitable for plotting. } \\
& \text { If the argument is a formula } \mathrm{zvar} \sim \mathrm{xvar}+\mathrm{yvar} \text {, xvar, yvar and zvar } \\
& \text { are used as } \mathrm{x}, \mathrm{y} \text { and } \mathrm{z} \text { variables; if the argument is a list containing } \\
& \text { components } \mathrm{x}, \mathrm{y} \text { and } \mathrm{z} \text {, these are assumed to define plotting coordinates; } \\
& \text { if the argument is a matrix with three columns, the first is assumed to } \\
& \text { contain the } \mathrm{x} \text { values, etc. } \\
& \text { Alternatively, two arguments } \mathrm{x} \text { and } \mathrm{y} \text { can be be provided. One may be } \\
& \text { real, the other complex; in any other case, the arguments are coerced to } \\
& \text { vectors and the values plotted against their indices. } \\
& \text { xlab, ylab, zlab } \\
& \text { names for the } \mathrm{x}, \mathrm{y} \text { and } \mathrm{z} \text { variables to be extracted. }
\end{aligned}
$$

log character, "x", "y", "z" or combinations. Sets negative values to NA and gives a warning.
recycle logical; if TRUE, recycle (rep) the shorter ones of $x, y$ or $z$ if their lengths differ.

## Value

A list with the components
$x \quad$ numeric (i.e. double) vector of abscissa values.
$\mathrm{y} \quad$ numeric vector of the same length as x .
z numeric vector of the same length as x .
xlab character(1) or NULL, the axis label of $x$.
ylab character (1) or NULL, the axis label of $y$.
zlab character(1) or NULL, the axis label of $z$.

## Author(s)

Uwe Ligges and Martin Maechler

## See Also

xy. coords for 2D.

## Examples

```
str(xyz.coords(data.frame(10*1:9, -4),y=NULL, z=NULL))
str(xyz.coords(1:6, fft(1:6),z=NULL,xlab="X", ylab="Y"))
y <- 2 * (x2 <- 10 + (x1 <- 1:10))
str(xyz.coords(y ~ x1 + x2,y=NULL, z=NULL))
str(xyz.coords(data.frame(x=-1:9,y=2:12,z=3:13), y=NULL, z=NULL,
    log="xy"))
##> Warning message: 2 x values <= 0 omitted ...
```

```
zcbind Bind Two or More Time Series
```


## Description

Bind Two or More Time Series which have common frequency.

## Usage

.cbind.ts(sers, nmsers, dframe $=$ FALSE, union $=$ TRUE)

## Arguments

$$
\begin{array}{ll}
\text { sers } & \begin{array}{l}
\text { a list of two or more univariate or multivariate time series, or objects } \\
\text { which can coerced to time series. }
\end{array} \\
\text { nmsers } & \begin{array}{l}
\text { a character vector of the same length as sers with the names for the time } \\
\text { series. }
\end{array} \\
\text { dframe } & \begin{array}{l}
\text { logical; if TRUE return the result as a data frame. } \\
\text { union }
\end{array} \\
\text { logical; if TRUE, act as ts.union or ts.intersect. }
\end{array}
$$

## Details

This is an internal function which is not to be called by the user.

```
zip.file.extract Extract File from a Zip Archive
```


## Description

This will extract the file named file from the zip archive, if possible, and write it in a temporary location.

## Usage

```
zip.file.extract(file, zipname = "R.zip")
```


## Arguments

| file | A file name. |
| :--- | :--- |
| zipname | The file name of a zip archive, including the ". zip" extension if required. |

## Details

The method used is selected by options(unzip=). All platforms support an "internal" unzip: this is the default under Windows and MacOS, and the fall-back under Unix if no unzip program was found during configuration and R_UNZIPCMD is not set.
The file will be extracted if it is in the archive and any required unzip utility is available. It will probably be extracted to the directory given by tempdir, overwriting an existing file of that name.

## Value

The name of the original or extracted file. Success is indicated by returning a different name.

## Note

The "internal" method is very simple, and will not set file dates.

## Author(s)

B. D. Ripley

## Chapter 2

## The ctest package

```
ansari.test Ansari-Bradley Test
```


## Description

Performs the Ansari-Bradley two-sample test for a difference in scale parameters.

## Usage

ansari.test(x, y, alternative = c("two.sided", "less", "greater"),
exact $=$ NULL, conf.int $=$ FALSE, conf.level $=0.95, \ldots$ )
ansari.test(formula, data, subset, na.action, ...)

## Arguments

$x \quad$ numeric vector of data values.
$\mathrm{y} \quad$ numeric vector of data values.
alternative indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter.
exact a logical indicating whether an exact p-value should be computed.
conf.int a logical,indicating whether a confidence interval should be computed.
conf.level confidence level of the interval.
formula a formula of the form lhs ~ rhs where lhs is a numeric variable giving the data values and rhs a factor with two levels giving the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

Suppose that x and y are independent samples from distributions with densities $f((t-$ $m) / s) / s$ and $f(t-m)$, respectively, where $m$ is an unknown nuisance parameter and $s$, the ratio of scales, is the parameter of interest. The Ansari-Bradley test is used for testing the null that $s$ equals 1 , the two-sided alternative being that $s \neq 1$ (the distributions differ only in variance), and the one-sided alternatives being $s>1$ (the distribution underlying x has a larger variance, "greater") or $s<1$ ("less").

By default (if exact is not specified), an exact p-value is computed if both samples contain less than 50 finite values and there are no ties. Otherwise, a normal approximation is used.
Optionally, a nonparametric confidence interval and an estimator for $s$ are computed. If exact p-values are available, an exact confidence interval is obtained by the algorithm described in Bauer (1972), and the Hodges-Lehmann estimator is employed. Otherwise, the returned confidence interval and point estimate are based on normal approximations.

## Value

A list with class "htest" containing the following components:
statistic the value of the Ansari-Bradley test statistic.
p.value the p-value of the test.
null.value the ratio of scales $s$ under the null, 1.
alternative a character string describing the alternative hypothesis.
method the string "Ansari-Bradley test".
data.name a character string giving the names of the data.
conf.int a confidence interval for the scale parameter. (Only present if argument conf.int = TRUE.)
estimate an estimate of the ratio of scales. (Only present if argument conf.int $=$ TRUE.)

## Note

To compare results of the Ansari-Bradley test to those of the F test to compare two variances (under the assumption of normality), observe that $s$ is the ratio of scales and hence $s^{2}$ is the ratio of variances (provided they exist), whereas for the F test the ratio of variances itself is the parameter of interest. In particular, confidence intervals are for $s$ in the Ansari-Bradley test but for $s^{2}$ in the F test.

## References

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 83-92.
David F. Bauer (1972), Constructing confidence sets using rank statistics. Journal of the American Statistical Association 67, 687-690.

## See Also

fligner.test for a rank-based (nonparametric) $k$-sample test for homogeneity of variances; mood.test for another rank-based two-sample test for a difference in scale parameters; var.test and bartlett.test for parametric tests for the homogeneity in variance.

## Examples

```
## Hollander & Wolfe (1973, p. 86f):
## Serum iron determination using Hyland control sera
ramsay <- c(111, 107, 100, 99, 102, 106, 109, 108, 104, 99,
    101, 96, 97, 102, 107, 113, 116, 113, 110, 98)
jung.parekh <- c(107, 108, 106, 98, 105, 103, 110, 105, 104,
    100, 96, 108, 103, 104, 114, 114, 113, 108, 106, 99)
ansari.test(ramsay, jung.parekh)
ansari.test(rnorm(10), rnorm(10, 0, 2), conf.int = TRUE)
```

```
bartlett.test Bartlett Test for Homogeneity of Variances
```


## Description

Performs Bartlett's test of the null that the variances in each of the groups (samples) are the same.

## Usage

```
bartlett.test(x, g, ...)
bartlett.test(formula, data, subset, na.action, ...)
```


## Arguments

x
g
formula a formula of the form lhs ~ rhs where lhs gives the data values and rhs the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

If x is a list, its elements are taken as the samples or fitted linear models to be compared for homogeneity of variances. In this case, the elements must either all be numeric data vectors or fitted linear model objects, $g$ is ignored, and one can simply use bartlett.test ( x ) to perform the test. If the samples are not yet contained in a list, use bartlett.test (list ( x , ...)).
Otherwise, x must be a numeric data vector, and g must be a vector or factor object of the same length as x giving the group for the corresponding elements of x .

## Value

A list of class "htest" containing the following components:
statistic Bartlett's K-squared test statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic.
p.value the p-value of the test.
method the character string "Bartlett test for homogeneity of variances".
data.name a character string giving the names of the data.

## References

Bartlett, M. S. (1937). Properties of sufficiency and statistical tests. Proceedings of the Royal Statistical Society Series A 160, 268-282.

## See Also

var.test for the special case of comparing variances in two samples from normal distributions; fligner.test for a rank-based (nonparametric) $k$-sample test for homogeneity of variances; ansari.test and mood.test for two rank based two-sample tests for difference in scale.

## Examples

```
data(InsectSprays)
plot(count ~ spray, data = InsectSprays)
bartlett.test(InsectSprays$count, InsectSprays$spray)
bartlett.test(count ~ spray, data = InsectSprays)
```

binom.test Exact Binomial Test

## Description

Performs an exact test of a simple null hypothesis about the probability of success in a Bernoulli experiment.

## Usage

```
binom.test(x, n, p = 0.5,
    alternative = c("two.sided", "less", "greater"),
    conf.level = 0.95)
```


## Arguments

\(\left.$$
\begin{array}{ll}\mathrm{x} & \begin{array}{l}\text { number of successes, or a vector of length } 2 \text { giving the numbers of successes } \\
\text { and failures, respectively. }\end{array} \\
\mathrm{n} & \begin{array}{l}\text { number of trials; ignored if } \mathrm{x} \text { has length } 2 . \\
\mathrm{p}\end{array}
$$ <br>

hypothesized probability of success.\end{array}\right]\)| indicates the alternative hypothesis and must be one of "two.sided", |
| :--- |
| conf.level | | "greater" or "less". You can specify just the initial letter. |
| :--- |
| confidence level for the returned confidence interval. |

## Details

Confidence intervals are obtained by a procedure first given in Clopper and Pearson (1934). This guarantees that the confidence level is at least conf.level, but in general does not give the shortest-length confidence intervals.

## Value

A list with class "htest" containing the following components:
statistic the number of successes.
parameter the number of trials.
p.value the p-value of the test.
conf.int a confidence interval for the probability of success.
estimate the estimated probability of success.
null.value the probability of success under the null, $p$.
alternative a character string describing the alternative hypothesis.
method the character string "Exact binomial test".
data.name a character string giving the names of the data.

## References

Clopper, C. J. \& Pearson, E. S. (1934). The use of confidence or fiducial limits illustrated in the case of the binomial. Biometrika, 26, 404-413.

Conover, W. J. (1971), Practical nonparametric statistics. New York: John Wiley \& Sons. Pages 97-104.

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 15-22.

## See Also

prop.test for a general (approximate) test for equal or given proportions.

## Examples

```
## Conover (1971), p. 97f.
## Under (the assumption of) simple Mendelian inheritance, a cross
## between plants of two particular genotypes produces progeny 1/4 of
## which are ''dwarf"' and 3/4 of which are ''giant'', respectively.
## In an experiment to determine if this assumption is reasonable, a
## cross results in progeny having 243 dwarf and 682 giant plants.
## If ''giant'' is taken as success, the null hypothesis is that p =
## 3/4 and the alternative that p != 3/4.
binom.test(c(682, 243), p = 3/4)
binom.test(682, 682 + 243, p = 3/4) # The same.
## => Data are in agreement with the null hypothesis.
```


## Description

chisq.test performs chi-squared tests on contingency tables.

```
Usage
chisq.test(x, y = NULL, correct = TRUE,
    p = rep(1/length(x), length(x)),
    simulate.p.value = FALSE, B = 2000)
```


## Arguments

$\mathrm{x} \quad$ a vector or matrix.
$\mathrm{y} \quad$ a vector; ignored if x is a matrix.
correct a logical indicating whether to apply continuity correction when computing the test statistic.
$p \quad a$ vector of probabilities of the same length of $x$.
simulate.p.value
a logical indicating whether to compute p-values by Monte Carlo simulation.

B
an integer specifying the number of replicates used in the Monte Carlo simulation.

## Details

If x is a matrix with one row or column, or if x is a vector and y is not given, x is treated as a one-dimensional contingency table. In this case, the hypothesis tested is whether the population probabilities equal those in $p$, or are all equal if $p$ is not given.
If x is a matrix with at least two rows and columns, it is taken as a two-dimensional contingency table, and hence its entries should be nonnegative integers. Otherwise, $x$ and y must be vectors or factors of the same length; incomplete cases are removed, the objects are coerced into factor objects, and the contingency table is computed from these. Then, Pearson's chi-squared test of the null that the joint distribution of the cell counts in a 2 dimensional contingency table is the product of the row and column marginals is performed. If simulate.p.value is FALSE, the p-value is computed from the asymptotic chi-squared distribution of the test statistic; continuity correction is only used in the 2-by-2 case if correct is TRUE. Otherwise, if simulate.p.value is TRUE, the p-value is computed by Monte Carlo simulation with B replicates. This is done by random sampling from the set of all contingency tables with given marginals, and works only if the marginals are positive.

## Value

A list with class "htest" containing the following components:
statistic the value the chi-squared test statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic, NA if the p-value is computed by Monte Carlo simulation.

| p.value | the p-value for the test. |
| :--- | :--- |
| method | a character string indicating the type of test performed, and whether <br> Monte Carlo simulation or continuity correction was used. |
| data.name | a character string giving the name(s) of the data. |
| observed | the observed counts. |
| expected | the expected counts under the null hypothesis. |
| residuals | the Pearson residuals, (observed - expected) / sqrt (expected). |

## Examples

```
data(InsectSprays) # Not really a good example
chisq.test(InsectSprays$count > 7, InsectSprays$spray)
                                    # Prints test summary
chisq.test(InsectSprays$count > 7, InsectSprays$spray)$obs
                                    # Counts observed
chisq.test(InsectSprays$count > 7, InsectSprays$spray)$exp
                                    # Counts expected under the null
## Effect of simulating p-values
x <- matrix(c(12, 5, 7, 7), nc = 2)
chisq.test(x)$p.value # 0.4233
chisq.test(x, simulate.p.value = TRUE, B = 10000)$p.value
                                    # around 0.29!
## Testing for population probabilities
## Case A. Tabulated data
x <- c(A = 20, B = 15, C = 25)
chisq.test(x)
chisq.test(as.table(x)) # the same
## Case B. Raw data
x <- trunc(5 * runif(100))
chisq.test(table(x)) # NOT 'chisq.test(x)'!
```

cor.test Test for Association Between Paired Samples

## Description

Test for association between paired samples, using one of Pearson's product moment correlation coefficient, Kendall's $\tau$ or Spearman's $\rho$.

## Usage

```
cor.test(x, y,
    alternative = c("two.sided", "less", "greater"),
    method = c("pearson", "kendall", "spearman"),
    exact = NULL, conf.level = 0.95, ...)
cor.test(formula, data, subset, na.action, ...)
```


## Arguments

| $\mathrm{x}, \mathrm{y}$ | numeric vectors of data values. x and y must have the same length. |
| :---: | :---: |
| alternative | indicates the alternative hypothesis and must be one of "two.sided" "greater" or "less". You can specify just the initial letter. "greater" corresponds to positive association, "less" to negative association. |
| method | a character string indicating which correlation coefficient is to be used for the test. One of "pearson", "kendall", or "spearman", can be abbreviated. |
| exact | a logical indicating whether an exact p-value should be computed. Only used for Kendall's $\tau$. See the Details for the meaning of NULL (the default) |
| conf.level | confidence level for the returned confidence interval. Currently only used for the Pearson product moment correlation coefficient if there are at least 4 complete pairs of observations. |
| formula | a formula of the form $\sim u+v$, where each of $u$ and $v$ are numeric variables giving the data values for one sample. The samples must be of the same length. |
| data | an optional data frame containing the variables in the model formula. |
| subset | an optional vector specifying a subset of observations to be used. |
| na.action | a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action"). |
|  | further arguments to be passed to or from methods. |

## Details

The three methods each estimate the association between paired samples and compute a test of the value being zero. They use different measures of association, all in the range $[-1,1]$ with 0 indicating no association. These are sometimes referred to as tests of no correlation, but that term is often confined to the default method.
If method is "pearson", the test statistic is based on Pearson's product moment correlation coefficient cor ( $\mathrm{x}, \mathrm{y}$ ) and follows a t distribution with length $(\mathrm{x})-2$ degrees of freedom if the samples follow independent normal distributions. If there are at least 4 complete pairs of observation, an asymptotic confidence interval is given based on Fisher's Z transform.
If method is "kendall" or "spearman", Kendall's $\tau$ or Spearman's $\rho$ statistic is used to estimate a rank-based measure of association. These tests may be used if the data do not necessarily come from a bivariate normal distribution.
For Kendall's test, by default (if exact is NULL), an exact p-value is computed if there are less than 50 paired samples containing finite values and there are no ties. Otherwise, the test statistic is the estimate scaled to zero mean and unit variance, and is approximately normally distributed.
For Spearman's test, p-values are computed using algorithm AS 89.

## Value

A list with class "htest" containing the following components:
statistic the value of the test statistic.
parameter the degrees of freedom of the test statistic in the case that it follows a $t$ distribution.

| p.value <br> estimate | the p-value of the test. <br> the estimated measure of association, with name "cor", "tau", or "rho" <br> correspoding to the method employed. |
| :--- | :--- |
| null.value | the value of the association measure under the null hypothesis, always 0. |
| alternative | a character string describing the alternative hypothesis. |
| method | a character string indicating how the association was measured. |
| data.name | a character string giving the names of the data. |
| conf.int | a confidence interval for the measure of association. Currently only given <br> for Pearson's product moment correlation coefficient in case of at least 4 <br> complete pairs of observations. |

## References

D. J. Best \& D. E. Roberts (1975), Algorithm AS 89: The Upper Tail Probabilities of Spearman's $\rho$. Applied Statistics, 24, 377-379.

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 185-194 (Kendall and Spearman tests).

## Examples

```
## Hollander & Wolfe (1973), p. 187f.
## Assessment of tuna quality. We compare the Hunter L measure of
## lightness to the averages of consumer panel scores (recoded as
## integer values from 1 to 6 and averaged over 80 such values) in
## 9 lots of canned tuna.
x <- c(44.4, 45.9, 41.9, 53.3, 44.7, 44.1, 50.7, 45.2, 60.1)
y <- c( 2.6, 3.1, 2.5, 5.0, 3.6, 4.0, 5.2, 2.8, 3.8)
## The alternative hypothesis of interest is that the
## Hunter L value is positively associated with the panel score.
cor.test(x, y, method = "kendall", alternative = "greater")
## => p=0.05972
cor.test(x, y, method = "kendall", alternative = "greater",
    exact = FALSE) # using large sample approximation
## => p=0.04765
## Compare this to
cor.test(x, y, method = "spearm", alternative = "g")
cor.test(x, y, alternative = "g")
## Formula interface.
data(USJudgeRatings)
pairs(USJudgeRatings)
cor.test(~ CONT + INTG, data = USJudgeRatings)
```

fisher.test Fisher's Exact Test for Count Data

## Description

Performs Fisher's exact test for testing the null of independence of rows and columns in a contingency table with fixed marginals.

## Usage

```
fisher.test(x, y = NULL, workspace = 200000, hybrid = FALSE,
    or = 1, alternative = "two.sided", conf.level = 0.95)
```


## Arguments

$\mathrm{x} \quad$ either a two-dimensional contingency table in matrix form, or a factor object.
$\mathrm{y} \quad$ a factor object; ignored if x is a matrix.
workspace an integer specifying the size of the workspace used in the network algorithm.
hybrid a logical indicating whether the exact probabilities (default) or a hybrid approximation thereof should be computed. In the hybrid case, asymptotic chi-squared probabilities are only used provided that the "Cochran" conditions are satisfied.
or the hypothesized odds ratio. Only used in the 2 by 2 case.
alternative indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less". You can specify just the initial letter. Only used in the 2 by 2 case.
conf.level confidence level for the returned confidence interval. Only used in the 2 by 2 case.

## Details

If x is a matrix, it is taken as a two-dimensional contingency table, and hence its entries should be nonnegative integers. Otherwise, both x and y must be vectors of the same length. Incomplete cases are removed, the vectors are coerced into factor objects, and the contingency table is computed from these.

In the one-sided 2 by 2 cases, p-values are obtained directly using the hypergeometric distribution. Otherwise, computations are based on a C version of the FORTRAN subroutine FEXACT which implements the network developed by Mehta and Patel (1986) and improved by Clarkson, Fan \& Joe (1993). The FORTRAN code can be obtained from http://www.netlib. org/toms/643.

In the 2 by 2 case, the null of conditional independence is equivalent to the hypothesis that the odds ratio equals one. Exact inference can be based on observing that in general, given all marginal totals fixed, the first element of the contingency table has a non-central hypergeometric distribution with non-centrality parameter given by the odds ratio (Fisher, 1935).

## Value

A list with class "htest" containing the following components:
p .value the p -value of the test.
conf.int a confidence interval for the odds ratio. Only present in the 2 by 2 case.
estimate an estimate of the odds ratio. Note that the conditional Maximum Likelihood Estimate (MLE) rather than the unconditional MLE (the sample odds ratio) is used. Only present in the 2 by 2 case.
null.value the odds ratio under the null, or. Only present in the 2 by 2 case.
alternative a character string describing the alternative hypothesis.
method the character string "Fisher's Exact Test for Count Data".
data.name a character string giving the names of the data.

## References

Alan Agresti (1990). Categorical data analysis. New York: Wiley. Pages 59-66.
Fisher, R. A. (1935). The logic of inductive inference. Journal of the Royal Statistical Society Series A 98, 39-54.
Fisher, R. A. (1962). Confidence limits for a cross-product ratio. Australian Journal of Statistics 4, 41.

Cyrus R. Mehta \& Nitin R. Patel (1986). Algorithm 643. FEXACT: A Fortran subroutine for Fisher's exact test on unordered $r * c$ contingency tables. ACM Transactions on Mathematical Software, 12, 154-161.
Douglas B. Clarkson, Yuan-an Fan \& Harry Joe (1993). A Remark on Algorithm 643: FEXACT: An Algorithm for Performing Fisher's Exact Test in $r \times c$ Contingency Tables. ACM Transactions on Mathematical Software, 19, 484-488.

## See Also

```
chisq.test
```


## Examples

```
## Agresti (1990), p. 61f, Fisher's Tea Drinker
## A British woman claimed to be able to distinguish whether milk or
## tea was added to the cup first. To test, she was given 8 cups of
## tea, in four of which milk was added first. The null hypothesis
## is that there is no association between the true order of pouring
## and the women's guess, the alternative that there is a positive
## association (that the odds ratio is greater than 1).
TeaTasting <-
matrix(c(3, 1, 1, 3),
    nr = 2,
    dimnames = list(Guess = c("Milk", "Tea"),
        Truth = c("Milk", "Tea")))
fisher.test(TeaTasting, alternative = "greater")
## => p=0.2429, association could not be established
## Fisher (1962), Convictions of like-sex twins in criminals
Convictions <-
matrix(c(2, 10, 15, 3),
    nr = 2,
```

```
    dimnames =
    list(c("Dizygotic", "Monozygotic"),
        c("Convicted", "Not convicted")))
Convictions
fisher.test(Convictions, alternative = "less")
fisher.test(Convictions, conf.level = 0.95)$conf.int
fisher.test(Convictions, conf.level = 0.99)$conf.int
```

fligner.test Fligner-Killeen Test for Homogeneity of Variances

## Description

Performs a Fligner-Killeen (median) test of the null that the variances in each of the groups (samples) are the same.

## Usage

fligner.test(x, g, ...)
fligner.test(formula, data, subset, na.action, ...)

## Arguments

$\mathrm{x} \quad$ a numeric vector of data values, or a list of numeric data vectors.
$\mathrm{g} \quad \mathrm{a}$ vector or factor object giving the group for the corresponding elements of x . Ignored if x is a list.
formula a formula of the form lhs ~ rhs where lhs gives the data values and rhs the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

If x is a list, its elements are taken as the samples to be compared for homogeneity of variances, and hence have to be numeric data vectors. In this case, $g$ is ignored, and one can simply use fligner.test ( x ) to perform the test. If the samples are not yet contained in a list, use fligner.test(list(x, ...)).

Otherwise, x must be a numeric data vector, and g must be a vector or factor object of the same length as x giving the group for the corresponding elements of x .
The Fligner-Killeen (median) test has been determined in a simulation study as one of the many tests for homogeneity of variances which is most robust against departures from normality, see Conover, Johnson \& Johnson (1981). It is a $k$-sample simple linear rank which uses the ranks of the absolute values of the centered samples and weights $a(i)=$ qnorm $((1+i /(n+1)) / 2)$. The version implemented here uses median centering in each of the samples (F-K:med $X^{2}$ in the reference).

## Value

A list of class "htest" containing the following components:
statistic the Fligner-Killeen:med $X^{2}$ test statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic.
p.value the p-value of the test.
method the character string "Fligner-Killeen test for homogeneity of variances".
data.name a character string giving the names of the data.

## References

W. J. Conover \& Mark E. Johnson \& Myrie M. Johnson (1981). A comparative study of tests for homogeneity of variances, with applications to the outer continental shelf bidding data. Technometrics 23, 351-361.

## See Also

ansari.test and mood.test for rank-based two-sample test for a difference in scale parameters; var.test and bartlett.test for parametric tests for the homogeneity of variances.

## Examples

```
data(InsectSprays)
plot(count ~ spray, data = InsectSprays)
fligner.test(InsectSprays$count, InsectSprays$spray)
fligner.test(count ~ spray, data = InsectSprays)
## Compare this to bartlett.test()
```

friedman.test Friedman Rank Sum Test

## Description

Performs a Friedman rank sum test with unreplicated blocked data.

## Usage

friedman.test(y, groups, blocks, ...)
friedman.test(formula, data, subset, na.action, ...)

## Arguments

y either a numeric vector of data values, or a data matrix.
groups a vector giving the group for the corresponding elements of y if this is a vector; ignored if y is a matrix. If not a factor object, it is coerced to one.
blocks a vector giving the block for the corresponding elements of y if this is a vector; ignored if y is a matrix. If not a factor object, it is coerced to one.
formula $\quad a$ formula of the form $a \sim b \mid c$, where $a, b$ and $c$ give the data values and corresponding groups and blocks, respectively.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

friedman.test can be used for analyzing unreplicated complete block designs (i.e., there is exactly one observation in y for each combination of levels of groups and blocks) where the normality assumption may be violated.
The null hypothesis is that apart from an effect of blocks, the location parameter of y is the same in each of the groups.
If $y$ is a matrix, groups and blocks are obtained from the column and row indices, respectively. NA's are not allowed in groups or blocks; if y contains NA's, corresponding blocks are removed.

## Value

A list with class "htest" containing the following components:

| statistic | the value of Friedman's chi-squared statistic. |
| :--- | :--- |
| parameter | the degrees of freedom of the approximate chi-squared distribution of the <br> test statistic. |
| p.value the p-value of the test. <br> method the character string "Friedman rank sum test". <br> data.name a character string giving the names of the data. |  |

## References

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 139-146.

## See Also

quade.test.

## Examples

```
## Hollander & Wolfe (1973), p. 140ff.
## Comparison of three methods (''round out'), ''narrow angle'', and
## '(wide angle')) for rounding first base. For each of 18 players
## and the three method, the average time of two runs from a point on
## the first base line 35ft from home plate to a point 15ft short of
## second base is recorded.
RoundingTimes <-
matrix(c(5.40, 5.50, 5.55,
    5.85, 5.70, 5.75,
    5.20, 5.60, 5.50,
    5.55, 5.50, 5.40,
    5.90, 5.85, 5.70,
    5.45, 5.55, 5.60,
    5.40, 5.40, 5.35,
```

```
    5.45, 5.50, 5.35,
    5.25, 5.15, 5.00,
    5.85, 5.80, 5.70,
    5.25, 5.20, 5.10,
    5.65, 5.55, 5.45,
    5.60, 5.35, 5.45,
    5.05, 5.00, 4.95,
    5.50, 5.50, 5.40,
    5.45, 5.55, 5.50,
    5.55, 5.55, 5.35,
    5.45, 5.50, 5.55,
    5.50, 5.45, 5.25,
    5.65, 5.60, 5.40,
    5.70, 5.65, 5.55,
    6.30, 6.30, 6.25),
    nr = 22,
    byrow = TRUE,
    dimnames = list(1 : 22,
                            c("Round Out", "Narrow Angle", "Wide Angle")))
friedman.test(RoundingTimes)
## => strong evidence against the null that the methods are equivalent
## with respect to speed
data(warpbreaks)
wb <- aggregate(warpbreaks$breaks,
    by = list(w = warpbreaks$wool,
                        t = warpbreaks$tension),
    FUN = mean)
wb
friedman.test(wb$x, wb$w, wb$t)
friedman.test(x ~ w | t, data = wb)
```


## kruskal.test Kruskal-Wallis Rank Sum Test

## Description

Performs a Kruskal-Wallis rank sum test.

## Usage

kruskal.test(x, g, ...)
kruskal.test(formula, data, subset, na.action, ...)

## Arguments

$\mathrm{x} \quad$ a numeric vector of data values, or a list of numeric data vectors.
g a vector or factor object giving the group for the corresponding elements of x . Ignored if x is a list.
formula a formula of the form lhs ~ rhs where lhs gives the data values and rhs the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

kruskal.test performs a Kruskal-Wallis rank sum test of the null that the location parameters of the distribution of x are the same in each group (sample). The alternative is that they differ in at least one.
If x is a list, its elements are taken as the samples to be compared, and hence have to be numeric data vectors. In this case, $g$ is ignored, and one can simply use kruskal.test ( x ) to perform the test. If the samples are not yet contained in a list, use kruskal.test (list ( x , ...)).
Otherwise, $x$ must be a numeric data vector, and $g$ must be a vector or factor object of the same length as x giving the group for the corresponding elements of x .

## Value

A list with class "htest" containing the following components:

```
statistic the Kruskal-Wallis rank sum statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the
    test statistic.
p.value the p-value of the test.
method the character string "Kruskal-Wallis rank sum test".
data.name a character string giving the names of the data.
```


## References

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 115-120.

## See Also

The Wilcoxon rank sum test (wilcox.test) as the special case for two samples; lm together with anova for performing one-way location analysis under normality assumptions; with Student's t test (t.test) as the special case for two samples.

## Examples

```
## Hollander & Wolfe (1973), 116.
## Mucociliary efficiency from the rate of removal of dust in normal
## subjects, subjects with obstructive airway disease, and subjects
## with asbestosis.
x <- c(2.9, 3.0, 2.5, 2.6, 3.2) # normal subjects
y <- c(3.8, 2.7, 4.0, 2.4) # with obstructive airway disease
z <- c(2.8, 3.4, 3.7, 2.2, 2.0) # with asbestosis
kruskal.test(list(x, y, z))
## Equivalently,
x <- c(x, y, z)
g<- factor(rep(1:3, c(5, 4, 5)),
    labels = c("Normal subjects",
                                "Subjects with obstructive airway disease",
```

```
    "Subjects with asbestosis"))
kruskal.test(x, g)
## Formula interface.
data(airquality)
boxplot(Ozone ~ Month, data = airquality)
kruskal.test(Ozone ~ Month, data = airquality)
```

ks.test

Kolmogorov-Smirnov Tests

## Description

Performs one or two sample Kolmogorov-Smirnov tests.

## Usage

```
ks.test(x, y, ..., alternative = c("two.sided", "less", "greater"),
    exact \(=\) NULL)
```


## Arguments

| x | a numeric vector of data values. |
| :--- | :--- |
| y | either a numeric vector of data values, or a character string naming a <br> distribution function. |
| $\ldots$ | parameters of the distribution specified by y. |
| alternative | indicates the alternative hypothesis and must be one of "two.sided" <br> (default), "less", or "greater". You can specify just the initial letter. |
| exact | NULL or a logical indicating whether an exact p-value should be computed. |
|  | See Details for the meaning of NULL. Only used in the two-sided two- <br> sample case. |

## Details

If y is numeric, a two-sample test of the null hypothesis that x and y were drawn from the same continuous distribution is performed.
Alternatively, y can be a character string naming a continuous distribution function. In this case, a one sample test of the null that the distribution function underlying x is y with parameters specified by ... is carried out.

The presence of ties generates a warning, since continuous distributions do not generate them.

The possible values "two.sided", "less" and "greater" of alternative specify the null hypothesis that the true distribution function of x is equal to, not less than or not greater than the hypothesized distribution function (one-sample case) or the distribution function of y (two-sample case), respectively.

Exact p-values are only available for the two-sided two-sample test with no ties. In that case, if exact = NULL (the default) an exact p-value is computed if the product of the sample sizes is less than 10000. Otherwise, asymptotic distributions are used whose approximations may be inaccurate in small samples.

## Value

A list with class "htest" containing the following components:
statistic the value of the test statistic.
p.value the p-value of the test.
alternative a character string describing the alternative hypothesis.
method a character string indicating what type of test was performed.
data.name a character string giving the name(s) of the data.

## References

Conover, W. J. (1971), Practical nonparametric statistics. New York: John Wiley \& Sons. Pages 295-301 (one-sample "Kolmogorov" test), 309-314 (two-sample "Smirnov" test).

## See Also

shapiro.test which performs the Shapiro-Wilk test for normality.

## Examples

```
x <- rnorm(50)
y <- runif(30)
# Do x and y come from the same distribution?
ks.test(x, y)
# Does x come from a shifted gamma distribution with shape 3 and scale 2?
ks.test(x+2, "pgamma", 3, 2) # two-sided
ks.test(x+2, "pgamma", 3, 2, alternative = "gr")
```

```
mantelhaen.test Cochran-Mantel-Haenszel Chi-Squared Test for Count Data
```


## Description

Performs a Cochran-Mantel-Haenszel chi-squared test of the null that two nominal variables are conditionally independent in each stratum, assuming that there is no three-way interaction.

## Usage

```
mantelhaen.test(x, y = NULL, z = NULL,
    alternative = c("two.sided", "less", "greater"),
    correct = TRUE, exact = FALSE, conf.level = 0.95)
```


## Arguments

x either a 3-dimensional contingency table in array form where each dimension is at least 2 and the last dimension corresponds to the strata, or a factor object with at least 2 levels.
y
a factor object with at least 2 levels; ignored if x is an array.
z
a factor object with at least 2 levels identifying to which stratum the corresponding elements in x and y belong; ignored if x is an array.

| alternative | indicates the alternative hypothesis and must be one of "two.sided", <br> "greater" or "less". You can specify just the initial letter. Only used <br> in the 2 by 2 by $K$ case. |
| :--- | :--- |
| correct | a logical indicating whether to apply continuity correction when comput- <br> ing the test statistic. Only used in the 2 by 2 by $K$ case. |
| exact | a logical indicating whether the Mantel-Haenszel test or the exact condi- <br> tional test (given the strata margins) should be computed. Only used in <br> the 2 by 2 by $K$ case. |
| conf.level | confidence level for the returned confidence interval. Only used in the 2 <br> by 2 by $K$ case. |

## Details

If x is an array, each dimension must be at least 2 , and the entries should be nonnegative integers. NA's are not allowed. Otherwise, $x$, $y$ and $z$ must have the same length. Triples containing NA's are removed. All variables must take at least two different values.

## Value

A list with class "htest" containing the following components:
statistic Only present if no exact test is performed. In the classical case of a 2 by 2 by $K$ table (i.e., of dichotomous underlying variables), the Mantel-Haenszel chi-squared statistic; otherwise, the generalized Cochran-Mantel-Haenszel statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic (1 in the classical case). Only present if no exact test is performed.
p.value the p-value of the test.
conf.int a confidence interval for the common odds ratio. Only present in the 2 by 2 by $K$ case.
estimate an estimate of the common odds ratio. If an exact test is performed, the conditional Maximum Likelihood Estimate is given; otherwise, the Mantel-Haenszel estimate. Only present in the 2 by 2 by $K$ case.
null.value the common odds ratio under the null of independence, 1. Only present in the 2 by 2 by $K$ case.
alternative a character string describing the alternative hypothesis. Only present in the 2 by 2 by $K$ case.
method a character string indicating the method employed, and whether or not continuity correction was used.
data.name a character string giving the names of the data.

## Note

The asymptotic distribution is only valid if there is no three-way interaction. In the classical 2 by 2 by $K$ case, this is equivalent to the conditional odds ratios in each stratum being identical. Currently, no inference on homogeneity of the odds ratios is performed.
See also the example below.

## References

Alan Agresti (1990). Categorical data analysis. New York: Wiley. Pages 230-235.

## Examples

```
## Agresti (1990), pages 231--237, Penicillin and Rabbits
## Investigation of the effectiveness of immediately injected or 1.5
## hours delayed penicillin in protecting rabbits against a lethal
## injection with beta-hemolytic streptococci.
Rabbits <-
array(c(0, 0, 6, 5,
    3, 0, 3, 6,
    6, 2, 0, 4,
    5, 1, 6, 0,
    2, 5, 0, 0),
    dim = c(2, 2, 5),
    dimnames = list(
        Delay = c("None", "1.5h"),
        Response = c("Cured", "Died"),
        Penicillin.Level = c("1/8", "1/4", "1/2", "1", "4")))
Rabbits
## Classical Mantel-Haenszel test
mantelhaen.test(Rabbits)
## => p = 0.047, some evidence for higher cure rate of immediate
## injection
## Exact conditional test
mantelhaen.test(Rabbits, exact = TRUE)
## => p - 0.040
## Exact conditional test for one-sided alternative of a higher
## cure rate for immediate injection
mantelhaen.test(Rabbits, exact = TRUE, alternative = "greater")
## => p = 0.020
## UC Berkeley Student Admissions
data(UCBAdmissions)
mantelhaen.test(UCBAdmissions)
## No evidence for association between admission and gender
## when adjusted for department. However,
apply(UCBAdmissions, 3, function(x) (x[1,1]*x[2, 2])/(x[1, 2]*x[2,1]))
## This suggests that the assumption of homogeneous (conditional)
## odds ratios may be violated. The traditional approach would be
## using the Woolf test for interaction:
woolf <- function(x) {
    x <- x + 1 / 2
    k <- dim(x) [3]
    or <- apply(x, 3, function(x) (x[1,1]*x[2,2])/(x[1,2]*x[2,1]))
    w <- apply(x, 3, function(x) 1 / sum(1 / x))
    1 - pchisq(sum(w * (log(or) - weighted.mean(log(or), w)) ~ 2), k - 1)
}
woolf(UCBAdmissions)
## => p = 0.003, indicating that there is significant heterogeneity.
## (And hence the Mantel-Haenszel test cannot be used.)
```

```
mcnemar.test McNemar's Chi-squared Test for Count Data
```


## Description

Performs McNemar's chi-squared test for symmetry of rows and columns in a twodimensional contingency table.

## Usage

mcnemar.test (x, y = NULL, correct = TRUE)

## Arguments

| x | either a two-dimensional contingency table in matrix form, or a factor <br> object. |
| :--- | :--- |
| y | a factor object; ignored if x is a matrix. |
| correct | a logical indicating whether to apply continuity correction when comput- <br> ing the test statistic. |

## Details

The null is that the probabilities of being classified into cells [ $i, j$ ] and [ $j, i$ ] are the same.
If x is a matrix, it is taken as a two-dimensional contingency table, and hence its entries should be nonnegative integers. Otherwise, both $x$ and $y$ must be vectors of the same length. Incomplete cases are removed, the vectors are coerced into factor objects, and the contingency table is computed from these.

Continuity correction is only used in the 2-by-2 case if correct is TRUE.

## Value

A list with class "htest" containing the following components:
statistic the value of McNemar's statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic.
p .value the p -value of the test.
method a character string indicating the type of test performed, and whether continuity correction was used.
data.name a character string giving the name(s) of the data.

## References

Alan Agresti (1990). Categorical data analysis. New York: Wiley. Pages 350-354.

## Examples

```
## Agresti (1990), p. 350.
## Presidential Approval Ratings.
## Approval of the President's performance in office in two surveys,
## one month apart, for a random sample of 1600 voting-age Americans.
Performance <-
matrix(c(794, 86, 150, 570),
    nr = 2,
    dimnames = list("1st Survey" = c("Approve", "Disapprove"),
        "2nd Survey" = c("Approve", "Disapprove")))
Performance
mcnemar.test(Performance)
## => very strong association between the two successive ratings
```

```
mood.test Mood Two-Sample Test of Scale
```


## Description

Performs Mood's two-sample test for a difference in scale parameters.

## Usage

```
mood.test(x, y, alternative = c("two.sided", "less", "greater"), ...)
mood.test(formula, data, subset, na.action, ...)
```


## Arguments

$\mathrm{x}, \mathrm{y} \quad$ numeric vectors of data values.
alternative indicates the alternative hypothesis and must be one of "two.sided" (default), "greater" or "less" all of which can be abbreviated.
formula a formula of the form lhs ~ rhs where lhs is a numeric variable giving the data values and rhs a factor with two levels giving the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

The underlying model is that the two samples are drawn from $f(x-l)$ and $f((x-l) / s) / s$, respectively, where $l$ is a common location parameter and $s$ is a scale parameter.

The null hypothesis is $s=1$.
There are more useful tests for this problem.

## Value

A list with class "htest" containing the following components:
statistic the value of the test statistic.
p.value the p-value of the test.
alternative a character string describing the alternative hypothesis.
method the character string "Mood two-sample test of scale".
data.name a character string giving the names of the data.

## References

Conover, W. J. (1971), Practical nonparametric statistics. New York: John Wiley \& Sons. Pages 234f.

## See Also

fligner.test for a rank-based (nonparametric) k-sample test for homogeneity of variances; ansari.test for another rank-based two-sample test for a difference in scale parameters; var.test and bartlett.test for parametric tests for the homogeneity in variance.

## Examples

```
## Same data as for the Ansari-Bradley test:
## Serum iron determination using Hyland control sera
ramsay <- c(111, 107, 100, 99, 102, 106, 109, 108, 104, 99,
    101, 96, 97, 102, 107, 113, 116, 113, 110, 98)
jung.parekh <- c(107, 108, 106, 98, 105, 103, 110, 105, 104,
    100, 96, 108, 103, 104, 114, 114, 113, 108, 106, 99)
mood.test(ramsay, jung.parekh)
## Compare this to ansari.test(ramsay, jung.parekh)
```

oneway.test

Test for Equal Means in a One-Way Layout

## Description

Test whether two or more samples from normal distributions have the same means. The variances are not necessarily assumed to be equal.

## Usage

oneway.test(formula, data, subset, na.action, var.equal = FALSE)

## Arguments

| formula | a formula of the form $\mathrm{lhs} \sim$ <br> rhs the corresponding groups. |
| :--- | :--- |
| data | an optional data frame containing the variables in the model formula. |
| subset | an optional vector specifying a subset of observations to be used. |
| na.action | a function which indicates what should happen when the data contain <br> NAs. Defaults to getOption("na.action"). |

var.equal a logical variable indicating whether to treat the variances in the samples as equal. If TRUE, then a simple F test for the equality of means in a oneway analysis of variance is preformed. If FALSE, an approximate method of Welch (1951) is used, which generalizes the commonly known 2-sample Welch test to the case of arbitrarily many samples.

## Value

A list with class "htest" containing the following components:
statistic the value of the test statistic.
parameter the degrees of freedom of the exact or approximate $F$ distribution of the test statistic.
p.value the p-value of the test.
method a character string indicating the test performed.
data.name a character string giving the names of the data.

## References

B. L. Welch (1951), On the comparison of several mean values: an alternative approach. Biometrika, 38, 330-336.

## See Also

The standard t test ( t .test) as the special case for two samples; the Kruskal-Wallis test kruskal.test for a nonparametric test for equal location parameters in a one-way layout.

## Examples

```
data(sleep)
## Not assuming equal variances
oneway.test(extra ~ group, data = sleep)
## Assuming equal variances
oneway.test(extra ~ group, data = sleep, var.equal = TRUE)
## which gives the same result as
anova(lm(extra ~ group, data = sleep))
```

pairwise.prop.test Pairwise comparisons of proportions

## Description

Calculate pairwise comparisons between pairs of proportions with correction for multiple testing

## Usage

pairwise.prop.test(x, n, p.adjust.method=p.adjust.methods, ...)

## Arguments



## Value

Object of class "pairwise.htest"

## See Also

```
prop.test, p.adjust
```


## Examples

```
smokers <- c( 83, 90, 129, 70 )
patients <- c( 86, 93, 136, 82 )
pairwise.prop.test(smokers, patients)
```

```
pairwise.t.test Pairwise t tests
```


## Description

Calculate pairwise comparisons between group levels with corrections for multiple testing

## Usage

pairwise.t.test(x, g, p.adjust.method=p.adjust.methods, pool.sd=TRUE, ...)

## Arguments

| x | Response vector |
| :--- | :--- |
| g | Grouping vector or factor |
| p.adjust.method |  |
|  | Method for adjusting p values (see p.adjust) |
| pool.sd | Switch to allow/disallow the use of a pooled SD |
| $\ldots$ | Additional arguments to pass to t.test |

## Value

Object of class "pairwise.htest"

## See Also

```
t.test, p.adjust
```


## Examples

```
data(airquality)
attach(airquality)
Month <- factor(Month, labels = month.abb[5:9])
pairwise.t.test(Ozone, Month)
pairwise.t.test(Ozone, Month, p.adj = "bonf")
pairwise.t.test(Ozone, Month, pool.sd = FALSE)
detach()
```

pairwise.table Tabulate $p$ values for pairwise comparisons

## Description

Creates table of p values for pairwise comparisons with corrections for multiple testing.

## Usage

pairwise.table(compare.levels, level.names, p.adjust.method)

## Arguments

compare.levels
Function to compute (raw) p value given indices i and j
level.names Names of the group levels
p.adjust.method

Method for multiple testing adjustment

## Details

Functions that do multiple group comparisons create separate compare.levels functions (assumed to be symmetrical in i and j ) and passes them to this function.

## Value

Table of p values in lower triangular form.

See Also
pairwise.t.test, et al.

```
pairwise.wilcox.test Pairwise Wilcoxon rank sum tests
```


## Description

Calculate pairwise comparisons between group levels with corrections for multiple testing

## Usage

pairwise.wilcox.test(x, g, p.adjust.method=p.adjust.methods, ...)

## Arguments

x Response vector
g Grouping vector or factor
p.adjust.method
Method for adjusting p values (see p.adjust)
... Additional arguments to pass to t.test

## Value

Object of class "pairwise.htest"

## See Also

wilcox.test, p.adjust

## Examples

```
data(airquality)
attach(airquality)
Month <- factor(Month, labels = month.abb[5:9])
pairwise.wilcox.test(Ozone, Month)
pairwise.wilcox.test(Ozone, Month, p.adj = "bonf")
detach()
```


## power.prop.test Power calculations two sample test for of proportions

## Description

Compute power of test, or determine parameters to obtain target power.

## Usage

```
power.prop.test(n=NULL, p1=NULL, p2=NULL, sig.level=0.05,
    power=NULL,
    alternative=c("two.sided", "one.sided"))
```


## Arguments

| n | Number of observations (per group) |
| :--- | :--- |
| p1 | probability in one group |
| p2 | probability in other group |
| sig.level | Significance level (Type I error probability) |
| power | Power of test (1 minus Type II error probability) |
| alternative | One- or two-sided test |

## Details

Exactly one of the parameters n, p1, p2, power, and sig.level must be passed as NULL, and that parameter is determined from the others. Notice that sig.level has a non-NULL default so NULL must be explicitly passed if you want it computed.

## Value

Object of class "power.htest", a list of the arguments (including the computed one) augmented with method and note elements.

## Note

uniroot is used to solve power equation for unknowns, so you may see errors from it, notably about inability to bracket the root when invalid arguments are given. If one of them is computed p1 < p2 will hold, although this is not enforced when both are specified.

## Author(s)

Peter Dalgaard. Based on previous work by Claus Ekstrøm

## See Also

```
prop.test, uniroot
```


## Examples

```
power.prop.test(n=50, p1=.50, p2=.75)
power.prop.test(p1=.50, p2=.75, power=.90)
power.prop.test(n=50, p1=.5, power=.90)
```

power.t.test Power calculations for one and two sample t tests

## Description

Compute power of test, or determine parameters to obtain target power.

## Usage

```
power.t.test(n=NULL, delta=NULL, sd=1, sig.level=0.05, power=NULL,
type=c("two.sample", "one.sample", "paired"),
alternative=c("two.sided", "one.sided"))
```


## Arguments

| n | Number of observations (per group) |
| :--- | :--- |
| delta | True difference in means |
| sd | Standard deviation |
| sig.level | Significance level (Type I error probability) |
| power | Power of test (1 minus Type II error probability) |
| type | Type of $t$ test |
| alternative | One- or two-sided test |

## Details

Exactly one of the parameters n, delta, power, sd, and sig.level must be passed as NULL, and that parameter is determined from the others. Notice that the last two have non-NULL defaults so NULL must be explicitly passed if you want to compute them.

## Value

Object of class "power.htest", a list of the arguments (including the computed one) augmented with method and note elements.

## Note

uniroot is used to solve power equation for unknowns, so you may see errors from it, notably about inability to bracket the root when invalid arguments are given.

## Author(s)

Peter Dalgaard. Based on previous work by Claus Ekstrøm

## See Also

```
t.test, uniroot
```


## Examples

```
power.t.test(n=20, delta=1)
power.t.test(power=.90, delta=1)
power.t.test(power=.90, delta=1, alt="one.sided")
```

```
print.pairwise.htest Print method for pairwise tests
```


## Description

Display results of pairwise comparison procedures

## Usage

print(x, ...)

## Arguments

$\begin{array}{ll}\mathrm{x} & \text { Object of class "pairwise.htest" } \\ \ldots & \text { further arguments to be passed to or from methods. }\end{array}$

## Value

None

## See Also

```
    pairwise.t.test, et al.
```

print.power.htest Print method for power calculation object

## Description

Print object of class "power.htest" in nice layout.

## Usage

print (x, ...)

## Arguments

x
Object of class "power.htest".
... further arguments to be passed to or from methods.

## Details

A power.htest object is just a named list of numbers and character strings, supplemented with method and note elements. The method is displayed as a title, the note as a footnote, and the remaining elements are given in an aligned 'name $=$ value' format.

## Value

none

Author(s)
Peter Dalgaard

See Also

```
power.t.test, power.prop.test
```

prop.test Test for Equal or Given Proportions

## Description

prop.test can be used for testing the null that the proportions (probabilities of success) in several groups are the same, or that they equal certain given values.

```
Usage
    prop.test(x, n, p = NULL,
        alternative = c("two.sided", "less", "greater"),
        conf.level = 0.95, correct = TRUE)
```


## Arguments

x
$\mathrm{n} \quad$ a vector of counts of trials; ignored if x is a matrix.
$p \quad a \quad$ vector of probabilities of success. The length of $p$ must be the same as the number of groups specified by $x$, and its elements must be greater than 0 and less than 1.
alternative a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter. Only used for testing the null that a single proportion equals a given value, or that two proportions are equal; ignored otherwise.
conf.level confidence level of the returned confidence interval. Must be a single number between 0 and 1 . Only used when testing the null that a single proportion equals a given value, or that two proportions are equal; ignored otherwise.
correct a logical indicating whether Yates' continuity correction should be applied.

## Details

Only groups with finite numbers of successes and failures are used. Counts of successes and failures must be nonnegative and hence not greater than the corresponding numbers of trials which must be positive. All finite counts should be integers.
If p is NULL and there is more than one group, the null tested is that the proportions in each group are the same. If there are two groups, the alternatives are that the probability of success in the first group is less than, not equal to, or greater than the probability of success in the second group, as specified by alternative. A confidence interval for the difference of proportions with confidence level as specified by conf.level and clipped to $[-1,1]$ is returned. Continuity correction is used only if it does not exceed the difference of the sample proportions in absolute value. Otherwise, if there are more than 2 groups, the alternative is always "two.sided", the returned confidence interval is NULL, and continuity correction is never used.

If there is only one group, then the null tested is that the underlying probability of success is p , or .5 if p is not given. The alternative is that the probability of success if less than, not equal to, or greater than $p$ or 0.5 , respectively, as specified by alternative. A confidence
interval for the underlying proportion with confidence level as specified by conf.level and clipped to $[0,1]$ is returned. Continuity correction is used only if it does not exceed the difference between sample and null proportions in absolute value.

Finally, if p is given and there are more than 2 groups, the null tested is that the underlying probabilities of success are those given by p. The alternative is always "two.sided", the returned confidence interval is NULL, and continuity correction is never used.

## Value

A list with class "htest" containing the following components:

| statistic | the value of Pearson's chi-squared test statistic. <br> parameter |
| :--- | :--- |
| the degrees of freedom of the approximate chi-squared distribution of the <br> test statistic. |  |
| estimate | the p-value of the test. |
| a vector with the sample proportions $\mathrm{x} / \mathrm{n}$. |  |

## See Also

binom.test for an exact test of a binomial hypothesis.

## Examples

```
heads <- rbinom(1, size=100, pr = .5)
prop.test(heads, 100) # continuity correction TRUE by default
prop.test(heads, 100, correct = FALSE)
## Data from Fleiss (1981), p. 139.
## H0: The null hypothesis is that the four populations from which
## the patients were drawn have the same true proportion of smokers.
## A: The alternative is that this proportion is different in at
## least one of the populations.
smokers <- c( 83, 90, 129, 70 )
patients <- c( 86, 93, 136, 82 )
prop.test(smokers, patients)
```


## Description

Performs chi-squared test for trend in proportions, i.e., a test asymptotically optimal for local alternatives where the log odds vary in proportion with score. By default, score is chosen as the group numbers.

## Usage

prop.trend.test(x, n, score = 1:length(x))

## Arguments

| x | Number of events |
| :--- | :--- |
| n | Number of trials |
| score | Group score |

## Value

An object of class "htest" with title, test statistic, p-value, etc.

## Note

This really should get integrated with prop.test

## Author(s)

Peter Dalgaard

## See Also

prop.test

## Examples

```
smokers <- c( 83, 90, 129, 70 )
patients <- c( 86, 93, 136, 82 )
prop.test(smokers, patients)
prop.trend.test(smokers, patients)
prop.trend.test(smokers, patients,c(0,0,0,1))
```

```
quade.test Quade Test
```


## Description

Performs a Quade test with unreplicated blocked data.

## Usage

```
quade.test(y, groups, blocks, ...)
quade.test(formula, data, subset, na.action, ...)
```


## Arguments

| y | either a numeric vector of data values, or a data matrix. |
| :--- | :--- |
| groups | a vector giving the group for the corresponding elements of y if this is a <br> vector; ignored if y is a matrix. If not a factor object, it is coerced to one. |
| blocks | a vector giving the block for the corresponding elements of y if this is a <br> vector; ignored if y is a matrix. If not a factor object, it is coerced to one. |
| formula | a formula of the form a $\sim \mathrm{b} \mid \mathrm{c}$, where $\mathrm{a}, \mathrm{b}$ and c give the data values <br> and corresponding groups and blocks, respectively. |
| data | an optional data frame containing the variables in the model formula. |
| subset | an optional vector specifying a subset of observations to be used. |
| na.action | a function which indicates what should happen when the data contain <br> NAs. Defaults to getOption("na. action"). |
| $\ldots$ | further arguments to be passed to or from methods. |

## Details

quade.test can be used for analyzing unreplicated complete block designs (i.e., there is exactly one observation in y for each combination of levels of groups and blocks) where the normality assumption may be violated.
The null hypothesis is that apart from an effect of blocks, the location parameter of y is the same in each of the groups.
If $y$ is a matrix, groups and blocks are obtained from the column and row indices, respectively. NA's are not allowed in groups or blocks; if y contains NA's, corresponding blocks are removed.

## Value

A list with class "htest" containing the following components:

```
statistic the value of Quade's F statistic.
parameter a vector with the numerator and denominator degrees of freedom of the
    approximate F distribution of the test statistic.
p.value the p-value of the test.
method the character string "Quade test".
data.name a character string giving the names of the data.
```


## References

D. Quade (1979), Using weighted rankings in the analysis of complete blocks with additive block effects. Journal of the American Statistical Association, 74, 680-683.
W. J. Conover (1999), Practical nonparametric statistics. New York: John Wiley \& Sons. Pages 373-380.

## See Also

friedman.test.

## Examples

```
## Conover (1999, p. 375f):
## Numbers of five brands of a new hand lotion sold in seven stores
## during one week.
y <- matrix(c( 5, 4, 7, 10, 12,
                    1, 3, 1, 0, 2,
                    16, 12, 22, 22, 35,
            5, 4, 3, 5, 4,
            10, 9, 7, 13, 10,
            19, 18, 28, 37, 58,
            10, 7, 6, 8, 7),
            nr = 7, byrow = TRUE,
            dimnames =
            list(Store = as.character(1:7),
            Brand = LETTERS[1:5]))
y
quade.test(y)
```

shapiro.test Shapiro-Wilk Normality Test

## Description

Performs the Shapiro-Wilk test for normality.

## Usage

```
shapiro.test(x)
```


## Arguments

$\mathrm{x} \quad$ a numeric vector of data values, the number of which must be between 3 and 5000. Missing values are allowed.

## Value

A list with class "htest" containing the following components:
statistic the value of the Shapiro-Wilk statistic.
p.value the p-value for the test.
method the character string "Shapiro-Wilk normality test".
data.name a character string giving the name(s) of the data.

## References

Patrick Royston (1982) An Extension of Shapiro and Wilk's $W$ Test for Normality to Large Samples. Applied Statistics, 31, 115-124.
Patrick Royston (1982) Algorithm AS 181: The $W$ Test for Normality. Applied Statistics, 31, 176-180.
Patrick Royston (1995) A Remark on Algorithm AS 181: The $W$ Test for Normality. Applied Statistics, 44, 547-551.

## See Also

qqnorm for producing a normal quantile-quantile plot.

## Examples

```
shapiro.test(rnorm(100, mean = 5, sd = 3))
shapiro.test(runif(100, min = 2, max = 4))
```

t.test Student's $t$-Test

## Description

Performs one and two sample t-tests on vectors of data.

## Usage

```
t.test(x, y = NULL, alternative = c("two.sided", "less", "greater"),
            mu = 0, paired = FALSE, var.equal = FALSE,
            conf.level = 0.95, ...)
t.test(formula, data, subset, na.action, ...)
```


## Arguments

x
$\mathrm{y} \quad$ an optional numeric vector data values.
alternative a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
mu a number indicating the true value of the mean (or difference in means if you are performing a two sample test).
paired a logical indicating whether you want a paired t-test.
var.equal a logical variable indicating whether to treat the two variances as being equal. If TRUE then the pooled variance is used to estimate the variance otherwise the Welch approximation to the degrees of freedom is used.
conf.level confidence level of the interval.
formula a formula of the form $1 \mathrm{hs} \sim \mathrm{rhs}$ where lh h is a numeric variable giving the data values and rhs a factor with two levels giving the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

The formula interface is only applicable for the 2 -sample tests.
If paired is TRUE then both $x$ and $y$ must be specified and they must be the same length. Missing values are removed (in pairs if paired is TRUE). If var. equal is TRUE then the pooled estimate of the variance is used. By default, if var. equal is FALSE then the variance is estimated separately for both groups and the Welch modification to the degrees of freedom is used.

## Value

A list with class "htest" containing the following components:

```
statistic the value of the t-statistic.
parameter the degrees of freedom for the t-statistic.
p.value the p-value for the test.
conf.int a confidence interval for the mean appropriate to the specified alternative
    hypothesis.
estimate the estimated mean or difference in means depending on whether it was
    a one-sample test or a two-sample test.
null.value the specified hypothesized value of the mean or mean difference depending
    on whether it was a one-sample test or a two-sample test.
alternative a character string describing the alternative hypothesis.
method a character string indicating what type of t-test was performed.
data.name a character string giving the name(s) of the data.
```


## See Also

prop.test

## Examples

```
t.test(1:10,y=c(7:20)) # P = .00001855
t.test(1:10,y=c(7:20, 200)) # P = . 1245 -- NOT significant anymore
## Classical example: Student's sleep data
data(sleep)
plot(extra ~ group, data = sleep)
## Traditional interface
attach(sleep)
t.test(extra[group == 1], extra[group == 2])
detach()
## Formula interface
t.test(extra ~ group, data = sleep)
```

```
var.test F Test to Compare Two Variances
```


## Description

Performs an F test to compare the variances of two samples from normal populations.

## Usage

```
var.test(x, y, ratio = 1, alternative = c("two.sided", "less", "greater"),
    conf.level = 0.95, ...)
var.test(formula, data, subset, na.action, ...)
```


## Arguments

$\mathrm{x}, \mathrm{y}$ numeric vectors of data values, or fitted linear model objects (inheriting from class "lm").
ratio the hypothesized ratio of the population variances of x and y .
alternative a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
conf.level confidence level for the returned confidence interval.
formula a formula of the form lhs ~ $r$ hs where lh is a numeric variable giving the data values and rhs a factor with two levels giving the corresponding groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.

## Details

The null hypothesis is that the ratio of the variances of the populations from which x and y were drawn, or in the data to which the linear models x and y were fitted, is equal to ratio.

## Value

A list with class "htest" containing the following components:

| statistic | the value of the F test statistic. |
| :--- | :--- |
| parameter | the degrees of the freedom of the F distribtion of the test statistic. |
| p.value | the p-value of the test. |
| conf.int | a confidence interval for the ratio of the population variances. |
| estimate | the ratio of the sample variances of x and y. |
| null.value | the ratio of population variances under the null. |
| alternative | a character string describing the alternative hypothesis. |
| method | the character string "F test to compare two variances". |
| data.name | a character string giving the names of the data. |

## See Also

bartlett.test for testing homogeneity of variances in more than two samples from normal distributions; ansari.test and mood.test for two rank based (nonparametric) two-sample tests for difference in scale.

## Examples

```
x <- rnorm(50, mean = 0, sd = 2)
y <- rnorm(30, mean = 1, sd = 1)
var.test(x, y) # Do x and y have the same variance?
var.test(lm(x ~ 1), lm(y ~ 1)) # The same.
```

wilcox.test Wilcoxon Rank Sum and Signed Rank Tests

## Description

Performs one and two sample Wilcoxon tests on vectors of data.

## Usage

```
wilcox.test(x, y = NULL, alternative = c("two.sided", "less", "greater"),
    mu = 0, paired = FALSE, exact = NULL, correct = TRUE,
    conf.int = FALSE, conf.level = 0.95, ...)
wilcox.test(formula, data, subset, na.action, ...)
```


## Arguments

```
x numeric vector of data values.
y an optional numeric vector of data values.
alternative a character string specifying the alternative hypothesis, must be one of
    "two.sided" (default), "greater" or "less". You can specify just the
    initial letter.
mu a number specifying an optional location parameter.
paired a logical indicating whether you want a paired test.
exact a logical indicating whether an exact p-value should be computed.
correct a logical indicating whether to apply continuity correction in the normal
    approximation for the p-value.
conf.int a logical indicating whether a confidence interval should be computed.
conf.level confidence level of the interval.
formula a formula of the form lhs ~ rhs where lhs is a numeric variable giving
    the data values and rhs a factor with two levels giving the corresponding
    groups.
data an optional data frame containing the variables in the model formula.
subset an optional vector specifying a subset of observations to be used.
na.action a function which indicates what should happen when the data contain
    NAs. Defaults to getOption("na.action").
... further arguments to be passed to or from methods.
```


## Details

The formula interface is only applicable for the 2 -sample tests.
If only $x$ is given, or if both $x$ and $y$ are given and paired is TRUE, a Wilcoxon signed rank test of the null that the distribution of $x$ (in the one sample case) or of $x-y$ (in the paired two sample case) is symmetric about mu is performed.

Otherwise, if both x and y are given and paired is FALSE, a Wilcoxon rank sum test (equivalent to the Mann-Whitney test) is carried out. In this case, the null hypothesis is that the location of the distributions of x and y differ by mu.
By default (if exact is not specified), an exact p-value is computed if the samples contain less than 50 finite values and there are no ties. Otherwise, a normal approximation is used.
Optionally (if argument conf.int is true), a nonparametric confidence interval and an estimator for the pseudomedian (one-sample case) or for the difference of the location parameters $\mathrm{x}-\mathrm{y}$ is computed. (The pseudomedian of a distribution $F$ is the median of the distribution of $(u+v) / 2$, where $u$ and $v$ are independent, each with distribution $F$. If $F$ is symmetric, then the pseudomedian and median coincide. See Hollander \& Wolfe (1973), page 34.) If exact p-values are available, an exact confidence interval is obtained by the algorithm described in Bauer (1972), and the Hodges-Lehmann estimator is employed. Otherwise, the returned confidence interval and point estimate are based on normal approximations.

## Value

A list with class "htest" containing the following components:

```
statistic the value of the test statistic with a name describing it.
parameter the parameter(s) for the exact distribution of the test statistic.
p.value the p-value for the test.
null.value the location parameter mu.
alternative a character string describing the alternative hypothesis.
method the type of test applied.
data.name a character string giving the names of the data.
conf.int a confidence interval for the location parameter. (Only present if argu-
    ment conf.int = TRUE.)
estimate an estimate of the location parameter. (Only present if argument
    conf.int = TRUE.)
```


## References

Myles Hollander \& Douglas A. Wolfe (1973), Nonparametric statistical inference. New York: John Wiley \& Sons. Pages 27-33 (one-sample), 68-75 (two-sample).
David F. Bauer (1972), Constructing confidence sets using rank statistics. Journal of the American Statistical Association 67, 687-690.

## See Also

kruskal.test for testing homogeneity in location parameters in the case of two or more samples; t.test for a parametric alternative under normality assumptions.

## Examples

```
## One-sample test.
## Hollander & Wolfe (1973), 29f.
## Hamilton depression scale factor measurements in 9 patients with
## mixed anxiety and depression, taken at the first (x) and second
## (y) visit after initiation of a therapy (administration of a
## tranquilizer).
x <- c(1.83, 0.50, 1.62, 2.48, 1.68, 1.88, 1.55, 3.06, 1.30)
y <- c(0.878, 0.647, 0.598, 2.05, 1.06, 1.29, 1.06, 3.14, 1.29)
wilcox.test(x, y, paired = TRUE, alternative = "greater")
wilcox.test(y - x, alternative = "less") # The same.
wilcox.test(y - x, alternative = "less",
exact = FALSE, correct = FALSE) # H&W large sample
                                    # approximation
## Two-sample test.
## Hollander & Wolfe (1973), 69f.
## Permeability constants of the human chorioamnion (a placental
## membrane) at term (x) and between 12 to 26 weeks gestational
## age (y). The alternative of interest is greater permeability
## of the human chorioamnion for the term pregnancy.
x <- c(0.80, 0.83, 1.89, 1.04, 1.45, 1.38, 1.91, 1.64, 0.73, 1.46)
y <- c(1.15, 0.88, 0.90, 0.74, 1.21)
wilcox.test(x, y, alternative = "g") # greater
wilcox.test(x, y, alternative = "greater",
    exact = FALSE, correct = FALSE) # H&W large sample
    # approximation
wilcox.test(rnorm(10), rnorm(10, 2), conf.int = TRUE)
## Formula interface.
data(airquality)
boxplot(Ozone ~ Month, data = airquality)
wilcox.test(Ozone ~ Month, data = airquality,
    subset = Month %in% c(5, 8))
```


## Chapter 3

## The eda package

```
line
Robust Line Fitting
```


## Description

Fit a line robustly as recommended in Exploratory Data Analysis.

## Usage

line(x, y)
coef(object, ...)
residuals(object, type, ...)
fitted(object, ...)
print(x, digits $=\max (3$, getOption("digits") - 3), ...)

## Arguments

| $\mathrm{x}, \mathrm{y}$ | the arguments can be any way of specifying x-y pairs. |
| :--- | :--- |
| object | a tukeyline object, typically the result of line (*). |
| digits | number of significant digits to use, see print. |
| type, ... | potentially further arguments, required by generic. |

## Value

An object of class "tukeyline".
Methods are available for the generic functions coef, residuals, fitted, and print.

## References

Tukey, J. W. (1977). Exploratory Data Analysis, Reading Massachusetts: Addison-Wesley.

See Also
lm.

## Examples

```
library(eda)
data(cars)
plot(cars)
(z <- line(cars))
abline(coef(z))
## Tukey-Anscombe Plot :
plot(residuals(z) ~ fitted(z), main = deparse(z$call))
```

```
medpolish Median Polish of a Matrix
```


## Description

Fits an additive model using Tukey's median polish procedure.

## Usage

medpolish(x, eps $=0.01$, maxiter $=10$, trace.iter $=$ TRUE)

## Arguments

x
a numeric matrix.
eps real number greater than 0 . A tolerance for convergence: see Details.
maxiter the maximum number of iterations
trace.iter logical. Should progress in convergence be reported?

## Details

The model fitted is additive (constant + rows + columns). The algorithm works by alternately removing the row and column medians, and continues until the proportional reduction in the sum of absolute residuals is less than eps or until there have been maxiter iterations. The sum of absolute residuals is printed at each iteration of the fitting process, if trace.iter is TRUE.
medpolish returns an object of class medpolish (see below). There are printing and plotting methods for this class, which are invoked via by the generics print and plot.

## Value

An object of class medpolish with the following named components:

| overall | the fitted constant term. |
| :--- | :--- |
| row | the fitted row effects. |
| col | the fitted column effects. |
| residuals | the residuals. |
| name | the name of the dataset. |

## References

Tukey, J. W. (1977). Exploratory Data Analysis, Reading Massachusetts: Addison-Wesley.

## See Also

 median; aov for a mean instead of median decomposition.
## Examples

```
## Deaths from sport parachuting; from ABC of EDA, p.224:
deaths <-
    rbind(c(14,15,14),
                c( 7, 4, 7),
                c( 8, 2,10),
                c(15, 9,10),
                c( 0, 2, 0))
dimnames(deaths) <- list(c("1-24", "25-74", "75-199", "200++", "NA"),
                    paste(1973:1975))
deaths
(med.d <- medpolish(deaths))
plot(med.d)
## Check decomposition:
all(deaths == med.d$overall + outer(med.d$row,med.d$col, "+") + med.d$resid)
```

smooth Tukey's (Running Median) Smoothing

## Description

Tukey's smoothers, 3RS3R, 3RSS, 3R, etc.

## Usage

```
smooth(x, kind = c("3RS3R", "3RSS", "3RSR", "3R", "3", "S"),
            twiceit = FALSE,
            endrule = "Tukey", do.ends = FALSE)
    print(x, ...)
summary(object, ...)
```


## Arguments

| x | a vector or time series |
| :---: | :---: |
| kind | a character string indicating the kind of smoother required; defaults to "3RS3R". |
| twiceit | logical, indicating if the result should be "twiced". Twicing a smoothe $S(y)$ means $S(y)+S(y-S(y))$, i.e., adding smoothed residuals to the smoothed values. This decreases bias (increasing variance). |
| endrule | a character string indicating the rule for smoothing at the boundary. Either "Tukey" (default) or "copy". |
| do.ends | logical, indicating if the 3 -splitting of ties should also happen at the boundaries ("ends"). This is only used for kind = "S". |
| object | (and $x$ in print(.)): object of class "tukeysmooth", typically the result of $\operatorname{smooth}($.$) .$ |
|  | potentially further arguments, required by generic. |

## Details

3 is Tukey's short notation for running medians of length 3, $3 R$ stands for Repeated 3 until convergence, and $S$ for Splitting of horizontal stretches of length 2 or 3.
Hence, $3 R S 3 R$ is a concatenation of $3 R$, $S$ and $3 R$, $3 R S S$ similarly, whereas $3 R S R$ means first 3R and then ( S and 3) Repeated until convergence - which can be bad.

## Value

An object of class "tukeysmooth" (which has print and summary methods) and is a vector or time series containing the smoothed values with additional attributes.

## Note

S and S-PLUS use a different (somewhat better) Tukey smoother in smooth(*). Note that there are other smoothing methods which provide rather better results. These were designed for hand calculations and may be used mainly for didactical purposes.
Since R version 1.2 , smooth does really implement Tukey's end-point rule correctly (see argument endrule).
kind $=$ "3RSR" has been the default till R-1.1, but it can have very bad properties, see the examples.
Note that repeated application of smooth(*) does smooth more, for the "3RS*" kinds.

## References

Tukey, J. W. (1977). Exploratory Data Analysis, Reading Massachusetts: Addison-Wesley.

## See Also

lowess; loess, supsmu and smooth.spline in package 'modreg'.

## Examples

```
## see also demo(smooth) !
x1 <- c(4, 1, 3, 6, 6, 4, 1, 6, 2, 4, 2) # very artificial
(x3R <- smooth(x1, "3R")) # 2 iterations of "3"
smooth(x3R, kind = "S")
sm.3RS <- function(x, ...)
    smooth(smooth(x, "3R", ...), "S", ...)
y<- c(1,1, 19:1)
plot(y, main = "misbehaviour of \"3RSR\"", col.main = 3)
lines(sm.3RS(y))
lines(smooth(y))
lines(smooth(y, "3RSR"), col = 3, lwd = 2)# the horror
x <- c(8:10,10, 0,0, 9,9)
plot(x, main = "breakdown of 3R and S and hence 3RSS")
matlines(cbind(smooth(x,"3R"),smooth(x,"S"), smooth(x,"3RSS"),smooth(x)))
data(presidents)
presidents[is.na(presidents)] <- 0 # silly
```

```
summary(sm3 <- smooth(presidents, "3R"))
summary(sm2 <- smooth(presidents,"3RSS"))
summary(sm <- smooth(presidents))
all.equal(c(sm2),c(smooth(smooth(sm3, "S"), "S"))) # 3RSS === 3R S S
all.equal(c(sm), c(smooth(smooth(sm3, "S"), "3R")))# 3RS3R === 3R S 3R
plot(presidents, main = "smooth(presidents0, *) : 3R and default 3RS3R")
lines(sm3,col = 3, lwd = 1.5)
lines(sm, col = 2, lwd = 1.25)
```


## Chapter 4

## The lqs package

## Description

Compute a multivariate location and scale estimate with a high breakdown point - this can be thought of as estimating the mean and covariance of the good part of the data. cov.mve and cov.mcd are compatibility wrappers.

## Usage

```
cov.rob(x, cor \(=\) FALSE, quantile.used \(=\) floor \(((n+p+1) / 2)\),
    method = c("mve", "mcd", "classical"), nsamp = "best", seed)
cov.mve(x, cor \(=\) FALSE, quantile.used \(=\) floor ( \((n+p+1) / 2)\),
    nsamp = "best", seed)
cov.mcd(x, cor = FALSE, quantile.used \(=\) floor( \((\mathrm{n}+\mathrm{p}+1) / 2)\),
    nsamp = "best", seed)
```


## Arguments

x
cor
quantile.used the minimum number of the data points regarded as good points.
method the method to be used - minimum volume ellipsoid, minimum covariance determinant or classical product-moment. Using cov.mve or cov.mcd forces mve or mcd respectively.
nsamp the number of samples or "best" or "exact" or "sample". If "sample" the number chosen is $\min (5 * \mathrm{p}, 3000)$, taken from Rousseeuw and Hubert (1997). If "best" exhaustive enumeration is done up to 5000 samples: if "exact" exhaustive enumeration will be attempted however many samples are needed.
seed the seed to be used for random sampling: see RNGkind. The current value of . Random.seed will be preserved if it is set.

## Details

For method "mve", an approximate search is made of a subset of size quantile.used with an enclosing ellipsoid of smallest volume; in method "mcd" it is the volume of the Gaussian confidence ellipsoid, equivalently the determinant of the classical covariance matrix, that is minimized. The mean of the subset provides a first estimate of the location, and the rescaled covariance matrix a first estimate of scatter. The Mahalanobis distances of all the points from the location estimate for this covariance matrix are calculated, and those points within the $97.5 \%$ point under Gaussian assumptions are declared to be good. The final estimates are the mean and rescaled covariance of the good points.
The rescaling is by the appropriate percentile under Gaussian data; in addition the first covariance matrix has an ad hoc finite-sample correction given by Marazzi.
For method "mve" the search is made over ellipsoids determined by the covariance matrix of $p$ of the data points. For method "mcd" an additional improvement step suggested by Rousseeuw and van Driessen (1999) is used, in which once a subset of size quantile.used is selected, an ellipsoid based on its covariance is tested (as this will have no larger a determinant, and may be smaller).

## Value

A list with components

| center | the final estimate of location. |
| :--- | :--- |
| cov | the final estimate of scatter. |
| cor | (only is cor = TRUE) the estimate of the correlation matrix. |
| sing | message giving number of singular samples out of total <br> crit |
| the value of the criterion on log scale. For MCD this is the determinant, |  |
| and for MVE it is proportional to the volume. |  |
| best | the subset used. For MVE the best sample, for MCD the best set of size <br> quantile.used. |
| n.obs | total number of observations. |

## Author(s)

B.D. Ripley

## References

P. J. Rousseeuw and A. M. Leroy (1987) Robust Regression and Outlier Detection. Wiley.
A. Marazzi (1993) Algorithms, Routines and S Functions for Robust Statistics. Wadsworth and Brooks/Cole.
P. J. Rousseeuw and B. C. van Zomeren (1990) Unmasking multivariate outliers and leverage points, Journal of the American Statistical Association, 85, 633-639.
P. J. Rousseeuw and K. van Driessen (1999) A fast algorithm for the minimum covariance determinant estimator. Technometrics 41, 212-223.
P. Rousseeuw and M. Hubert (1997) Recent developments in PROGRESS. In L1-Statistical Procedures and Related Topics ed Y. Dodge, IMS Lecture Notes volume 31, pp. 201-214.

## See Also

## Examples

```
data(stackloss)
set.seed(123)
cov.rob(stackloss)
cov.rob(stack.x, method = "mcd", nsamp = "exact")
```

lqs Resistant Regression

## Description

Fit a regression to the good points in the dataset, thereby achieving a regression estimator with a high breakdown point. lmsreg and ltsreg are compatibility wrappers.

## Usage

```
lqs \((x, \ldots)\)
lqs.formula(formula, data, ...,
    method = c("lts", "lqs", "lms", "S", "model.frame"),
    subset, na.action = na.fail, model = TRUE,
    \(\mathrm{x}=\mathrm{FALSE}, \mathrm{y}=\) FALSE, contrasts \(=\) NULL)
lqs.default(x, y, intercept = TRUE, method = c("lts", "lqs", "lms", "S"),
    quantile, control \(=\) lqs.control(...), \(k 0=1.548\), seed, ...)
lmsreg(...)
ltsreg(...)
```


## Arguments

| formula | a formula of the form $y \sim x 1+x 2+\ldots$ <br> data <br> data frame from which variables specified in formula are preferentially to |
| :--- | :--- |
| bubset | an index vector specifying the cases to be used in fitting. (NOTE: If given, <br> this argument must be named exactly.) <br> function to specify the action to be taken if NAs are found. The de- <br> fault action is for the procedure to fail. Alternatives include na.omit <br> and na.exclude, which lead to omission of cases with missing values on <br> any required variable. (NOTE: If given, this argument must be named <br> exactly.) |
| model | logical. If TRUE the model frame is returned. <br> an optional list. See the contrasts.arg of model.matrix.default. |
| contrasts | a matrix or data frame containing the explanatory variables. |
| y | the response: a vector of length the number of rows of x. |
| intercept | should the model include an intercept? |
| method | the method to be used. model.frame returns the model frame: for the <br> others see the Details section. Using lmsreg or ltsreg forces "lms" and |
| "lts" respectively. |  |

control additional control items: see Details.
k0 the cutoff / tuning constant used for $\chi()$ and $\psi()$ functions when method = "S", currently corresponding to Tukey's "biweight".
seed the seed to be used for random sampling: see .Random. seed. The current value of .Random.seed will be preserved if it is set..
... arguments to be passed to lqs.default or lqs.control, see control above and Details.

## Details

Suppose there are n data points and p regressors, including any intercept.
The first three methods minimize some function of the sorted squared residuals. For methods "lqs" and "lms" is the quantile squared residual, and for "lts" it is the sum of the quantile smallest squared residuals. "lqs" and "lms" differ in the defaults for quantile, which are floor $((n+p+1) / 2)$ and floor $((n+1) / 2)$ respectively. For "lts" the default is floor $(n / 2)+$ floor $((p+1) / 2)$.

The "S" estimation method solves for the scale s such that the average of a function chi of the residuals divided by $s$ is equal to a given constant.

The control argument is a list with components
psamp: the size of each sample. Defaults to p.
nsamp: the number of samples or "best" (the default) or "exact" or "sample". If "sample" the number chosen is min ( $5 * \mathrm{p}, 3000$ ), taken from Rousseeuw and Hubert (1997). If "best" exhaustive enumeration is done up to 5000 samples; if "exact" exhaustive enumeration will be attempted however many samples are needed.
adjust: should the intercept be optimized for each sample? Defaults to TRUE.

## Value

An object of class "lqs". This is a list with components
crit the value of the criterion for the best solution found, in the case of method == "S" before IWLS refinement.
sing character. A message about the number of samples which resulted in singular fits.
coefficients of the fitted linear model
bestone the indices of those points fitted by the best sample found (prior to adjustment of the intercept, if requested).
fitted.values the fitted values.
residuals the residuals.
scale estimate(s) of the scale of the error. The first is based on the fit criterion. The second (not present for method == "S") is based on the variance of those residuals whose absolute value is less than 2.5 times the initial estimate.

## Note

There seems no reason other than historical to use the lms and lqs options. LMS estimation is of low efficiency (converging at rate $n^{-1 / 3}$ ) whereas LTS has the same asymptotic efficiency as an M estimator with trimming at the quartiles (Marazzi, 1993, p.201). LQS and LTS have the same maximal breakdown value of ( $f$ loor $((n-p) / 2$ ) +1$) / n$ attained if floor $((n+p) / 2)$ <= quantile <= floor $((n+p+1) / 2)$. The only drawback mentioned of LTS is greater computation, as a sort was thought to be required (Marazzi, 1993, p.201) but this is not true as a partial sort can be used (and is used in this implementation).
Adjusting the intercept for each trial fit does need the residuals to be sorted, and may be significant extra computation if n is large and p small.

Opinions differ over the choice of psamp. Rousseeuw and Hubert (1997) only consider p; Marazzi (1993) recommends p+1 and suggests that more samples are better than adjustment for a given computational limit.

The computations are exact for a model with just an intercept and adjustment, and for LQS for a model with an intercept plus one regressor and exhaustive search with adjustment. For all other cases the minimization is only known to be approximate.

## Author(s)

B. D. Ripley

## References

P. J. Rousseeuw and A. M. Leroy (1987) Robust Regression and Outlier Detection. Wiley.
A. Marazzi (1993) Algorithms, Routines and S Functions for Robust Statistics. Wadsworth and Brooks/Cole.
P. Rousseeuw and M. Hubert (1997) Recent developments in PROGRESS. In L1-Statistical Procedures and Related Topics, ed Y. Dodge, IMS Lecture Notes volume 31, pp. 201-214.

## See Also

```
predict.lqs
```


## Examples

```
data(stackloss)
set.seed(123)
lqs(stack.loss ~ ., data = stackloss)
lqs(stack.loss ~ ., data = stackloss, method = "S", nsamp = "exact")
```

```
predict.lqs Predict from an lqs Fit
```


## Description

Predict from an resistant regression fitted by lqs.

## Usage

```
predict(object, newdata, ...)
```


## Arguments

| object | object inheriting from class "lqs" |
| :--- | :--- |
| newdata | matrix or data frame of cases to be predicted or, if object has a formula, <br> a data frame with columns of the same names as the variables used. A <br> vector will be interpreted as a row vector. If newdata is missing, an <br> attempt will be made to retrieve the data used to fit the lqs object. |
| $\ldots$ | arguments to be passed from or to other methods. |

## Details

This function is a method for the generic function predict() for class lqs. It can be invoked by calling predict( x ) for an object x of the appropriate class, or directly by calling predict.lqs(x) regardless of the class of the object.
Missing values in newdata are handled by returning NA if the linear discriminants cannot be evaluated. If newdata is omitted and the na.action of the fit omitted cases, these will be omitted on the prediction.

## Value

A vector of predictions.

## Author(s)

B.D. Ripley

## See Also

lqs

## Examples

```
data(stackloss)
set.seed(123)
fm <- lqs(stack.loss ~ ., data = stackloss, method = "S", nsamp = "exact")
predict(fm, stackloss)
```


## Chapter 5

## The methods package

. BasicFunsList List of Builtin and Special Functions

## Description

A named list providing instructions for turning builtin and special functions into generic functions.

Functions in R that are defined as .Primitive(<name>) are not suitable for formal methods, because they lack the basic reflectance property. You can't find the argument list for these functions by examining the function object itself.

Future versions of R may fix this by attaching a formal argument list to the corresponding function. While generally the names of arguments are not checked by the internal code implementing the function, the number of arguments frequently is.

In any case, some definition of a formal argument list is needed if users are to define methods for these functions. In particular, if methods are to be merged from multiple packages, the different sets of methods need to agree on the formal arguments.
In the absence of reflectance, this list provides the relevant information via a dummy function associated with each of the known specials for which methods are allowed.

At the same, the list flags those specials for which methods are meaningless (e.g., for) or just a very bad idea (e.g., . Primitive).

A generic function created via setMethod, for example, for one of these special functions will have the argument list from. BasicFunsList. If no entry exists, the argument list ( x , ...) is assumed.

## Description

These functions manage the relations that allow coercing an object to a given class.

```
Usage
    as(object, Class, strict=TRUE)
    as(object, Class) <- value
    setAs(from, to, def, replace, where = 1)
```


## Arguments

| object | Any object. |
| :--- | :--- |
| Class | The name of the class to which object should be coerced. |
| strict | A logical flag. If TRUE, the returned object must be strictly from the <br> target class (unless that class is a virtual class, in which case the object <br> will be from the closest actual class (often the original object, if that class <br> extends the virtual class directly). <br> If strict = FALSE, any simple extension of the target class will be re- <br> turned, without further change. A simple extension is, roughly, one that <br> just adds slots to an existing class. |
| value | The value to use to modify object (see the discussion below). You should <br> supply an object with class Class; some coercion is done, but you're <br> unwise to rely on it. |
| from, to | The classes between which def performs coercion. <br> (In the case of the coerce function these are objects from the classes, not |
| the names of the classes, but you're not expected to call coerce directly.) |  |
| def | A function of one argument. It will get an object from class from and <br> had better return an object of class to. (If you want to save setAs a little <br> work, make the name of the argument from, but don't worry about it, |
|  | setAs will do the conversion.) |
| replace | If supplied, the function to use as a replacement method. <br> The position or environment in which to store the resulting method for |
| where | Therce; by default, the global environment. <br> coercen |

## Summary of Functions

as: Returns the version of this object coerced to be the given Class.
If the corresponding is relation is true, it will be used. In particular, if the relation has a coerce method, the method will be invoked on object.
If the is relation is FALSE, and coerceFlag is TRUE, the coerce function will be called (which will throw an error if there is no valid way to coerce the two objects). Otherwise, NULL is returned.
Coerce methods are pre-defined for basic classes (including all the types of vectors, functions and a few others). The object asFunctions contains the list of such predefined relations: names (asFunctions) gives the names of all the classes.
Beyond these two sources of methods, further methods are defined by calls to the setAs function.
coerce: Coerce from to be of the same class as to.
Not a function you should usually call explicitly. The function setAs creates methods for coerce for the as function to use.
setAs: The function supplied as the third argument is to be called to implement as ( x , to) when x has class from. Need we add that the function should return a suitable object with class to.

## How Functions 'as' and 'setAs' Work

The function as contrives to turn object into an object with class Class. In doing so, it uses information about classes and methods, but in a somewhat special way. Keep in mind that objects from one class can turn into objects from another class either automatically or by an explicit call to the as function. Automatic conversion is special, and comes from the designer of one class of objects asserting that this class extends a another class (see setClass and setIs).
Because inheritance is a powerful assertion, it should be used sparingly (otherwise your computations may produce unexpected, and perhaps incorrect, results). But objects can also be converted explicitly, by calling as, and that conversion is designed to use any inheritance information, as well as explicit methods.

As a first step in conversion, the as function determines whether is(object, Class) is TRUE. This can be the case either because the class definition of object includes Class as a "super class" (directly or indirectly), or because a call to setIs established the relationship.
Either way, the inheritance relation defines a method to coerce object to Class. In the most common case, the method is just to extract from object the slots needed for Class, but it's also possible to specify a method explicitly in a setIs call.

So, if inheritance applies, the as function calls the appropriate method. If inheritance does not apply, and coerceFlag is FALSE, NULL is returned.
By default, coerceFlag is TRUE. In this case the as function goes on to look for a method for the function coerce for the signature $c$ (from = class (object), to = Class).
Method selection is used in the as function in two special ways. First, inheritance is applied for the argument from but not for the argument to (if you think about it, you'll probably agree that you wouldn't want the result to be from some class other than the Class specified). Second, the function tries to use inheritance information to convert the object indirectly, by first converting it to an inherited class. It does this by examining the classes that the from class extends, to see if any of them has an explicit conversion method. Suppose class "by" does: Then the as function implicitly computes as (as (object, "by"), Class).
With this explanation as background, the function setAs does a fairly obvious computation: It constructs and sets a method for the function coerce with signature c (from, to), using the def argument to define the body of the method. The function supplied as def can have one argument (interpreted as an object to be coerced) or two arguments (the from object and the to class). Either way, setAs constructs a function of two arguments, with the second defaulting to the name of the to class. The method will be called from as with the object as the only argument: The default for the second argument is provided so the method can know the intended to class.
The function coerce exists almost entirely as a repository for such methods, to be selected as desribed above by the as function. In fact, it would usually be a bad idea to call coerce directly, since then you would get inheritance on the to argument; as mentioned, this is not likely to be what you want.

## The Function 'as' Used in Replacements

When as appears on the left of an assignment, the intuitive meaning is "Replace the part of object that was inherited from Class by the value on the right of the assignment."

This usually has a straightforward interpretation, but you can control explicitly what happens, and sometimes you should to avoid possible corruption of objects.
When object inherits from Class in the usual way, by including the slots of Class, the default as method is to set the corresponding slots in object to those in value.

The default computation may be reasonable, but usually only if all other slots in object are unrelated to the slots being changed. Often, however, this is not the case. The class of object may have extended Class with a new slot whose value depends on the inherited slots. In this case, you may want to define a method for replacing the inherited information that recomputes all the dependent information. Or, you may just want to prohibit replacing the inherited information directly .
The way to control such replacements is through the replace argument to function setis. This argument is a method that function as calls when used for replacement. It can do whatever you like, including calling stop if you want to prohibit replacements. It should return a modified object with the same class as the object argument to as.
In R, you can also explicitly supply a replacement method, even in the case that inheritance does not apply, through the replace argument to setAs. It works essentially the same way, but in this case by constructing a method for "coerce<-". (Replace methods for coercion without inheritance are not in the original description and so may not be compatible with S-Plus, at least not yet.)
When inheritance does apply, coerce and replace methods can be specified through either setIs or setAs; the effect is essentially the same.

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## Examples

```
## using the definition of class "track" from Classes
setAs("track", "numeric", function(from)from@y)
t1 <- new("track", x=1:20, y=(1:20) ^2)
as(t1, "numeric")
## The next example shows:
## 1. A virtual class to define setAs for several classes at once.
## 2. as() using inherited information
setClass("ca", representation(a = "character", id = "numeric"))
setClass("cb", representation(b = "character", id = "numeric"))
```

```
setClass("id")
setIs("ca", "id")
setIs("cb", "id")
setAs("id", "numeric", function(from) from@id)
CA <- new("ca", a ="A", id = 1)
CB <- new("cb", b = "B", id = 2)
setAs("cb", "ca", function(from, to )new(to, a=from@b, id = from@id))
as(CB, "numeric")
```

BasicClasses Classes Corresponding to Basic Data Types

## Description

Formal classes exist corresponding to the basic R data types, allowing these types to be used in method signatures, as slots in class definitions, and to be extended by new classes.

## Usage

```
### The following are all basic vector classes.
### They can appear as class names in method signatures,
### in calls to as(), is(), and new().
"character"
"complex"
"double"
"expression"
"integer"
"list"
"logical"
"numeric"
"single"
### the class
"vector"
### is a virtual class, extended by all the above
### The following are additional basic classes
"NULL" # NULL objects
"function" # function objects, including primitives
"externalptr" # raw external pointers for use in C code
"ANY" # virtual classes used by the methods package itself
"VIRTUAL"
```

```
"missing"
```


## Objects from the Classes

Objects can be created by calls of the form new(Class, ...), where Class is the quoted class name, and the remaining arguments if any are objects to be interpreted as vectors of this class. Multiple arguments will be concatenated.

The class "expression" is slightly odd, in that the ... arguments will not be evaluated; therefore, don't enclose them in a call to quote().

## Extends

Class "vector", directly.

## Methods

coerce Methods are defined to coerce arbitrary objects to these classes, by calling the corresponding basic function, for example, as(x, "numeric") calls as.numeric(x).

## BasicFunctions Group Generic Functions

## Description

These are group generic functions. Methods defined for them will be used for any of the specific functions belonging to the particular group, provided no specific methods override. These functions should never be called directly (a suitable error message will result if they are).

## Usage

```
Arith (e1, e2)
Compare(e1, e2)
Ops(e1, e2)
Math (x)
Math2(x, digits)
Summary (x, ..., na.rm = FALSE)
Complex (z)
```


## Arguments

e1, e2
Arguments to the various binary operators.
x
The argument to the Math or Summary groups of functions.

## Details

The functions belonging to the various groups are as follows:

```
Arith "+", "-", "*", "^", "%%", "%/%", "/"
Compare "==", ">", "<", "!=", "<=", ">="
Ops "Arith", "Compare"
Math "log", "sqrt", "log10", "cumprod", "abs", "acos", "acosh", "asin",
    "asinh", "atan", "atanh", "ceiling", "cos", "cosh", "cumsum", "exp",
    "floor", "gamma", "lgamma", "sin", "sinh", "tan", "tanh", "trunc"
Math2 "round", "signif"
Summary "max", "min", "range", "prod", "sum", "any", "all"
Complex "Arg", "Conj", "Im", "Mod", "Re"
```

All the functions in these groups (other than the group generics themselves) are basic functions in R. They are not by default generic functions, and many of them are defined as primitives, meaning that they do not have formal arguments. However, you can still define methods for them. The effect of doing so is to create a generic function with the appropriate arguments, in the environment where the method definition is to be stored. It all works more or less as you might expect, admittedly via a bit of trickery in the background.

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.
The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## callNextMethod Call an Inherited Method

## Description

A call to callNextMethod can only appear inside a method definition. It then results in a call to the first inherited method after the current method, with the arguments to the current method passed down to the next method. The value of that method call is the value of callNextMethod.

## Usage

callNextMethod(...)

## Arguments

 note that the dispatch is as described below.) The recommendation for most applications is to use callNextMethod with no explicit arguments.
## Details

The definition of the first inherited method is that is the method which would have been called if the current method did not exist. This is more-or-less literally what happens: The current method is deleted from a copy of the methods for the current generic, and selectMethod is called to find the next method (the result is cached in a special object, so the search only typically happens once per session per combination of argument classes).
It is also legal, and often useful, for the method called by callNextMethod to itself have a call to callNextMethod. This works the same way, except that now two methods are deleted before selecting the next method, and so on for further nested calls to callNextMethod.

The statement that the method is called with the current arguments is more precisely as follows. Arguments that were missing in the current call are still missing (remember that "missing" is a valid class in a method signature). For a formal argument, say x, that appears in the original call, there is a corresponding argument in the next method call equivalent to " $x=x$ ". In effect, this means that the next method sees the same actual arguments, but arguments are evaluated only once.

## Value

The value returned by the selected method.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

Methods for the general behavior of method dispatch

## Examples

```
## some class definitions with simple inheritance
setClass("BO" , representation(b0 = "numeric"))
setClass("B1", "B0")
setClass("B2", representation("B1", b2 = "logical"))
## and a rather silly function to illustrate callNextMethod
f <- function(x) class(x)
setMethod("f", "BO", function(x) c(x@b0, callNextMethod()))
setMethod("f", "B2", function(x) c(x@b2, callNextMethod()))
```

```
    b2 <- new("B2", b2 = FALSE, b0 = 10)
```

    b1 <- new("B1", b0 = 2)
    f(b2)
    f (b1)
    class Class of an Object

## Description

Returns the name of the object's class as a character string.
In contrast to the version of this function in the base package, this version of class never returns NULL. For objects that do not have a formal class definition, and do not have the "class" attribute set, the value returned is effectively the same as data.class.
The replacement version of the function sets the class to the value provided. For classes that have a formal definition, directly replacing the class this way is strongly deprecated. The expression as (object, value) is the way to coerce an object to a particular class.

## Usage

class (object)
class(object) <- value

## Arguments

object Any R object (including basic objects for which no class is currently defined). When assigning the class, however, it must be possible to coerce the object to the specified class: the semantics of assigning a class to object are equivalent to object <- as (object, value).

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

## Classes <br> Class Definitions

## Description

Class definitions are objects that contain the formal definition of a class of R objects.

## Details

When a class is defined, an object is stored that contains the information about that class, including:
slots Each slot is a component object. Like elements of a list these may be extracted (by name) and set. However, they differ from list components in important ways.
All the objects from a particular class have the same set of slot names; specifically, the slot names that are contained in the class definition. Each slot in each object always has the same class; again, this is defined by the overall class definition.
Classes don't need to have any slots, and many useful classes do not. These objects usually extend other, simple objects, such as numeric or character vectors. Finally, classes can have no data at all- these are known as virtual classes and are in fact very important programming tools. They are used to group together ordinary classes that want to share some programming behavior, without necessarily restricting how the behavior is implemented.
extends The names of the classes that this class extends. A class Fancy, say, extends a class Simple if an object from the Fancy class has all the capabilities of the Simple class (and probably some more as well). In particular, and very usefully, any method defined to work for a Simple object can be applied to a Fancy object as well.
In other programming languages, this relationship is sometimes expressed by saying that Simple is a superclass of Fancy, or that Fancy is a subclass of Simple.
The actual class definition object contains the names of all the classes this class extends. But those classes can themselves extend other classes also, so the complete extension can only be known by obtaining all those class definitions.
Class extension is usually defined when the class itself is defined, by including the names of superclasses as unnamed elements in the representation argument to setClass.
An object from a given class will then have all the slots defined for its own class and all the slots defined for its superclasses as well.
Note that extends relations can be defined in other ways as well, by using the set Is function.
prototype Each class definition contains a prototype object from the class. This must have all the slots, if any, defined by the class definition.
The prototype most commonly just consists of the prototypes of all its slots. But that need not be the case: the definition of the class can specify any valid object for any of the slots.
There are a number of "basic" classes, corresponding to the ordinary kinds of data occurring in R. For example, "numeric" is a class corresponding to numeric vectors. These classes are predefined and can then be used as slots or as superclasses for any other class definitions. The prototypes for the vector classes are vectors of length 0 of the corresponding type.

There are also a few basic virtual classes, the most important being "vector", grouping together all the vector classes; and "language", grouping together all the types of objects making up the R language.

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

Methods, setClass, is, as, new, slot

```
classRepresentation-class
Class Objects
```


## Description

These are the objects that hold the definition of classes of objects. They are constructed and stored as meta-data by calls to the function setClass. Don't manipulate them directly, except perhaps to look at individual slots.

## Details

Class definitions are stored as metadata in various packages. Additional metadata supplies information on inheritance (the result of calls to setIs). Inheritance information implied by the class definition itself (because the class contains one or more other classes) is also constructed automatically.

When a class is to be used in an R session, this information is assembled to complete the class definition. The completion is a second object of class "classRepresentation", cached for the session or until something happens to change the information. A call to getClass returns the completed definition of a class; a call to getClassDef returns the stored definition (uncompleted).

In particular, completion fills in the upward- and downward-pointing inheritance information for the class, in slots contains and subclasses respectively. It's in principle important to note that this information can depend on which packages are installed, since these may define additional subclasses or superclasses.

## Slots

slots: A named list of the slots in this class; the elements of the list are the classes to which the slots must belong (or extend), and the names of the list gives the corresponding slot names.
contains: A named list of the classes this class "contains"; the elements of the list are objects of SClassExtension-class. The list may be only the direct extensions or all the currently known extensions (see the details).
virtual: Logical flag, set to TRUE if this is a virtual class.
prototype: The object that represents the standard prototype for this class; i.e., the data and slots returned by a call to new for this class with no special arguments. Don't mess with the prototype object directly.
validity: Optionally, a function to be used to test the validity of objects from this class. See validObject.
access: Access control information. Not currently used.
className: The character string name of the class.
package: The character string name of the package to which the class belongs. Nearly always the package on which the metadata for the class is stored, but in operations such as constructing inheritance information, the internal package name rules.
subclasses: A named list of the classes known to extend this class'; the elements of the list are objects of SClassExtension-class. The list is currently only filled in when completing the class definition. (see the details).
versionKey: Object of class "externalptr"; eventually will perhaps hold some versioning information, but not currently used.
sealed: Object of class "logical"; is this class sealed? If so, no modifications are allowed.

## See Also

See function setClass to supply the information in the class definition. See Classes for a more basic discussion of class information.

## Examples

\#\#---- Should be DIRECTLY executable !! ----

## EmptyMethodsList-class

Internal Class representing Empty Methods List

## Description

Objects from class "EmptyMethodsList" are generated during method selection to indicate failed search (forcing backtracking). Other classes described here are used internally in method dispatch. All these are for internal use.

## Usage

```
## class described below
"EmptyMethodsList"
### Other, virtual classes used in method dispatch
"OptionalMethods"
"PossibleMethod"
```

Slots
argument: Object of class "name" the argument names being selected on.
sublist: Object of class "list" (unused, and perhaps to be dropped in a later version.)

## Methods

No methods defined with class "EmptyMethodsList" in the signature.

## See Also

Function MethodsListSelect uses the objects; see MethodsList-class for the non-empty methods list objects.

```
environment-class Class "environment"
```


## Description

A formal class for R environments.

## Objects from the Class

Objects can be created by calls of the form new("environment", ...). The arguments in ..., if any, should be named and will be assigned to the newly created environment.

## Methods

coerce signature(from = "ANY", to = "environment"): calls as.environment.
initialize signature(object = "environment"): Implements the assignments in the new environment. Note that the object argument is ignored; a new environment is always created, since environments are not protected by copying.

See Also
new.env

```
genericFunction-class
```

Generic Function Objects

## Description

Generic functions (objects from or extending class genericFunction) are extended function objects, containing information used in creating and dispatching methods for this function. They also identify the package associated with the function and its methods.

## Objects from the Class

Generic functions are created and assigned by setGeneric or setGroupGeneric and, indirectly, by setMethod.

As you might expect setGeneric and setGroupGeneric create objects of class "genericFunction" and "groupGenericFunction" respectively.

## Slots

.Data: Object of class "function", the function definition of the generic, usually created automatically as a call to standardGeneric.
generic: Object of class "character", the name of the generic function.
package: Object of class "character", the name of the package to which the function definition belongs (and not necessarily where the generic function is stored). If the package is not specified explicitly in the call to setGeneric, it is usually the package on which the corresponding non-generic function exists.
group: Object of class "list", the group or groups to which this generic function belongs. Empty by default.
valueClass: Object of class "character"; if not an empty character vector, identifies one or more classes. It is asserted that all methods for this function return objects from these class (or from classes that extend them).
signature: Object of class "character", the vector of formal argument names that can appear in the signature of methods for this generic function. By default, it is all the formal arguments, except for .... Order matters for efficiency: the most commonly used arguments in specifying methods should come first.
default: Object of class "OptionalMethods", the default method for this function. Generated automatically and used to initialize method dispatch.
skeleton: Object of class "call", a slot used internally in method dispatch. Don't expect to use it directly.

## Extends

Class "function", from data part.
Class "OptionalMethods", by class "function".
Class "PossibleMethod", by class "function".

## Methods

Generic function objects are used in the creation and dispatch of formal methods; information from the object is used to create methods list objects and to merge or update the existing methods for this generic.

GenericFunctions Tools for Managing Generic Functions

## Description

The functions documented here manage collections of methods associated with a generic function, as well as providing information about the generic functions themselves.

## Usage

```
isGeneric(f, where, fdef, getName = FALSE)
isGroup(f, where, fdef)
removeGeneric(f, where)
standardGeneric(f)
dumpMethod(f, signature, file, where, def)
existsFunction(f, generic = TRUE, where)
findFunction(f, generic=TRUE)
dumpMethods(f, file, signature, methods, where)
signature(...)
removeMethods(f, where)
setReplaceMethod(f, ...)
getGenerics(where, searchForm = FALSE)
allGenerics(where, searchForm = FALSE)
callGeneric(...)
```


## Arguments

f
where

The character string naming the function.
Where on the search list of attached packages to look for functions or methods. By default, use the whole search list to find the relevant object(s).

```
signature The class signature of the relevant method. A signature is a named or
    unnamed vector of character strings. If named, the names must be formal
    argument names for the generic function. If signature is unnamed, the
    default is to use the first length(signature) formal arguments of the
    function.
file The file on which to dump method definitions.
def The function object defining the method; if omitted, the current method
    definition corresponding to the signature.
    ... Named or unnamed arguments to form a signature.
    generic In testing or finding functions, should generic functions be included. Sup-
    ply as FALSE to get only non-generic functions.
    fdef Optional, the generic function definition.
    Usually omitted in calls to isGeneric
    getName If TRUE, isGeneric returns the name of the generic. By default, it returns
        TRUE.
    methods The methods object containing the methods to be dumped. By default,
        the methods defined for this generic (optionally on the specified where
        location)
    searchForm In getGenerics, if TRUE, the package slot of the returned result is in
        the form used by search(), otherwise as the simple package name (e.g,
        "package:base" vs "base")
```


## Summary of Functions

isGeneric: Is there a function named $f$, and if so, is it a generic?
The getName argument allows a function to find the name from a function definition. If it is TRUE then the name of the generic is returned, or FALSE if this is not a generic function definition.
The behavior of isGeneric and getGeneric for primitive functions is slightly different. These functions don't exist as formal function objects (for efficiency and historical reasons), regardless of whether methods have been defined for them. A call to isGeneric tells you whether methods have been defined for this primitive function, anywhere in the current search list, or in the specified position where. In contrast, a call to getGeneric will return what the generic for that function would be, even if no methods have been currently defined for it.
removeGeneric, removeMethods: Remove the all the methods for the generic function of this name. In addition, removeGeneric removes the function itself; removeMethods restores the non-generic function which was the default method. If there was no default method, removeMethods leaves a generic function with no methods.
standardGeneric: Dispatches a method from the current function call for the generic function $f$.
getMethods: The list of methods for the specified generic.
dumpMethod: Dump the method for this generic function and signature.
existsFunction: Is there a function of this name. If generic is FALSE, generic functions are not counted.
findFunction: return all the elements of the search list on which a function definition for name exists.

NOTE: Use this rather than find with mode="function", which is not as meaningful, and has a few subtle bugs from its use of regular expressions.
selectMethod: Returns the method (a function) that $R$ would use to evaluate a call to this generic, with arguments corresponding to the specified signature.
$\mathrm{f}=$ the name of the generic function, signature is the signature of classes to match to the arguments of $f$.
dumpMethods: Dump all the methods for this generic.
signature: Returns a named list of classes to be matched to arguments of a generic function.
getGenerics: Returns the names of the generic functions that have methods defined on where; this argument can be an environment or an index into the search list. By default, the whole search list is used.
The methods definitions are stored with package qualifiers; for example, methods for function "initialize" might refer to two different functions of that name, on different packages. The package names corresponding to the method list object are contained in the slot package of the returned object. The form of the returned name can be plain (e.g., "base"), or in the form used in the search list ("package:base") according to the value of searchForm
callGeneric: In the body of a method, this function will make a call to the current generic function. If no arguments are passed to callGeneric, the arguments to the current call are passed down; otherwise, the arguments are interpreted as in a call to the generic function.

## Details

setGeneric: If there is already a non-generic function of this name, it will be used to define the generic unless def is supplied, and the current function will become the default method for the generic.
If def is supplied, this defines the generic function, and no default method will exist (often a good feature, if the function should only be available for a meaningful subset of all objects).
Arguments group and valueClass are retained for consistency with S-Plus, but are currently not used.
isGeneric: If the fdef argument is supplied, take this as the definition of the generic, and test whether it is really a generic, with $f$ as the name of the generic. (This argument is not available in S-Plus.)
removeGeneric: If where supplied, just remove the version on this element of the search list; otherwise, removes the first version encountered.
standardGeneric: Generic functions should usually have a call to standardGeneric as their entire body. They can, however, do any other computations as well.
The usual setGeneric (directly or through calling setMethod) creates a function with a call to standardGeneric.
getMethods: If the function is not a generic function, returns NULL. The $f$ argument can be either the character string name of the generic or the object itself.
The where argument optionally says where to look for the function, if $f$ is given as the name.
dumpMethod: The resulting source file will recreate the method
findFunction: If generic is FALSE, ignore generic functions.
selectMethod: The vector of strings for the classes can be named or not. If named, the names must match formal argument names of $f$. If not named, the signature is assumed to apply to the arguments of $f$ in order.

If mustFind is TRUE, an error results if there is no method (or no unique method) corresponding to this signature. Otherwise may return NULL or a MethodsList object.
dumpMethods: If signature is supplied only the methods matching this initial signature are dumped. (This feature is not found in S-Plus: don't use it if you want compatibility.) signature: The advantage of using signature is to provide a check on which arguments you meant, as well as clearer documentation in your method specification. In addition, signature checks that each of the elements is a single character string. removeMethods: Returns TRUE if $f$ was a generic function, FALSE (silently) otherwise.

If there is a default method, the function will be re-assigned as a simple function with this definition. Otherwise, the generic function remains but with no methods (so any call to it will generate an error). In either case, a following call to setMethod will consistently re-establish the same generic function as before.

## References

The $R$ package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections 1.6, 2.7, 2.8, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

setGeneric, setClass, showMethods

## getClass Get Class Definition

## Description

Get the definition of a class.

## Usage

getClass(Class,. Force $=$ FALSE)
getClassDef(Class, where=-1)

## Arguments

Class the character-string name of the class.
.Force if TRUE, return NULL if the class is undefined; otherwise, an undefined class results in an error.
where where to search for the definition; by default, anywhere on the current search list.

## Details

A call to getClass returns the complete definition of the class supplied as a string, including all slots, etc. in classes that this class extends. A call to getClassDef returns the definition of the class from the environment where, unadorned. It's usually getClass you want.

If you really want to know whether a class is formally defined, call isClass.

## Value

The object defining the class. This is an object of class "classRepEnvironment". However, do not deal with the contents of the object directly unless you are very sure you know what you're doing. Even then, it is nearly always better practice to use functions such as setClass and setIs. Messing up a class object will cause great confusion.

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

Classes, setClass, isClass.

## Examples

```
getClass("numeric") ## a built in class
```

```
getMethod Get or Test for the Definition of a Method
```


## Description

The functions getMethod and selectMethod get the definition of a particular method; the functions existsMethod and hasMethod test for the existence of a method. In both cases the first function only gets direct definitions and the second uses inheritance. The function findMethod returns the package(s) in the search list (or in the packages specified by the where argument) that contain a method for this function and signature.

The other functions are support functions: see the details below.

```
Usage
    getMethod(f, signature=character(), where, optional=FALSE)
    findMethod(f, signature, where)
    getMethods(f, where=-1)
    existsMethod(f, signature = character(), where)
    hasMethod(f, signature=character())
    selectMethod(f, signature, optional=FALSE, useInherited,
        mlist=getMethods(f), fdef = getGeneric(f))
    MethodsListSelect(f, env, mlist, fEnv, finalDefault, evalArgs,
        useInherited, fdef)
```


## Arguments

$\mathrm{f} \quad$ The character-string name of the generic function.
In getMethods only, this argument may be a function definition, in which case the special methods list object, if any, stored in the environment of the function is returned. (This usage is largely for internal purposes; you aren't likely to have such a function definition for direct use.)
signature The signature of classes to match to the arguments of f . The vector of strings for the classes should be named, and the names must match formal argument names of $f$. If not named, the signature is assumed to apply to the arguments of $f$ in order, but note below for selectMethod.
For selectMethod, the signature can optionally be an environment with classes assigned to the names of the corresponding arguments. Note: the names correspond to the names of the classes, not to the objects supplied in a call to the generic function.
where The position or environment in which to look for the method: by default, anywhere in the current search list.
optional If the selection does not produce a unique result, an error is generated, unless this argument is TRUE. In that case, the value returned is either a MethodsList object, if more than one method matches this signature, or NULL if no method matches.
mlist, fdef In selectMethod, the MethodsList object and/or the generic function object can be explicitly supplied. (Unlikely to be used, except in the recursive call that finds matches to more than one argument.)
env The environment in which argument evaluations are done in MethodsListSelect. Currently must be supplied, but should usually be sys.frame(sys.parent()) when calling the function explicitly for debugging purposes.
fEnv, finalDefault, evalArgs, useInherited
Internal-use arguments for the function's environment, the method to use as the overall default, whether to evaluate arguments, and which arguments should use inheritance.

## Details

A call to getMethod returns the method for a particular function and signature. As with other get functions, argument where controls where the function looks (by default anywhere in the search list) and argument optional controls whether the function returns NULL or generates an error if the method is not found. The search for the method makes no use of inheritance.

The function selectMethod also looks for a method given the function and signature, but makes full use of the method dispatch mechanism; i.e., inherited methods and group generics are taken into account just as they would be in dispatching a method for the corresponding signature, with the one exception that conditional inheritance is not used. Like getMethod, selectMethod returns NULL or generates an error if the method is not found, depending on the argument optional.

The functions existsMethod and hasMethod return TRUE or FALSE according to whether a method is found, the first corresponding to getMethod (no inheritance) and the second to selectMethod.

The function getMethods returns all the methods for a particular generic (in the form of a generic function with the methods information in its environment). The function is called from the evaluator to merge method information, and is not intended to be called directly.

The function MethodsListSelect performs a full search (including all inheritance and group generic information: see the Methods documentation page for details on how this works). The call returns a possibly revised methods list object, incorporating any method found as part of the allMethods slot.
Normally you won't call MethodsListSelect directly, but it is possible to use it for debugging purposes (only for distinctly advanced users!).
Note that the statement that MethodsListSelect corresponds to the selection done by the evaluator is a fact, not an assertion, in the sense that the evaluator code constructs and executes a call to MethodsListSelect when it does not already have a cached method for this generic function and signature. (The value returned is stored by the evaluator so that the search is not required next time.)

## Value

The call to selectMethod or getMethod returns a MethodDefinition-class object, the selected method, if a unique selection exists. (This class extends function, so you can use the result directly as a function if that is what you want.) Otherwise an error is thrown if optional is FALSE. If optional is TRUE, the value returned is NULL if no method matched, or a MethodsList object if multiple methods matched.

The call to getMethods returns the MethodsList object containing all the methods requested. If there are none, NULL is returned: getMethods does not generate an error in this case.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface
developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

```
getPackageName The name associated with a given package
```


## Description

The functions below produce the package associated with a particular environment or position on the search list, or of the package containing a particular function. They are primarily used to support computations that need to differentiate objects on multiple packages.

## Usage

getPackageName (where = 1)
functionPackageName (name)

## Arguments

where The environment or position on the search list associated with the desired package.
name The name of a function: functionPackageName finds the function and returns the package name corresponding. Note that if there are multiple instances of the function, all the corresponding names are returned.
pkg The name to be assigned to the package internally.

## Details

Package names are normally installed during loading of the package, by the INSTALL script or by the library function. (Currently, the name is stored as the object .packageName but don't trust this for the future.)

## Value

All the functions return the character-string name of the package (without the extraneous "package:" found in the search list).

## See Also

search

## Examples

\#\# both the following usually return "base" getPackageName(length(search()))
functionPackageName("rnorm")
hasArg Look for an Argument in the Call

## Description

Returns TRUE is name corresponds to an argument in the call, either a formal argument to the function, or a component of . . ., and FALSE otherwise.

## Usage

hasArg(name)

## Arguments

name The unquoted name of a potential argument.

## Details

The expression hasArg(x), for example, is similar to !missing $(x)$, with two exceptions. First, hasArg will look for an argument named x in the call if x is not a formal argument to the calling function, but ... is. Second, hasArg never generates an error if given a name as an argument, whereas missing( x ) generates an error if x is not a formal argument.

## Value

Always TRUE or FALSE as described above.

## See Also

```
missing
```


## Examples

```
ftest <- function(x1, ...) c(hasArg(x1), hasArg(y2))
ftest(1) ## c(TRUE, FALSE)
ftest(1, 2) ## c(TRUE, FALSE)
ftest(y2=2) ## c(FALSE, TRUE)
ftest(y=2) ## c(FALSE, FALSE) (no partial matching)
ftest(y2 = 2, x=1) ## c(TRUE, TRUE) partial match x1
```


## Description

is: With two arguments, tests whether object can be treated as from class2.
With one argument, returns all the super-classes of this object's class.
extends: Does the first class extend the second class? Returns maybe if the extension includes a test.
setIs: Defines class1 to be an extension of class2.

## Usage

```
is(object, class2)
extends(class1, class2, maybe=TRUE)
setIs(class1, class2, test=NULL, coerce=NULL, replace=NULL,
        by = NULL, where = 1)
```


## Arguments

| object | Any R object. |
| :---: | :---: |
| class1, class2 |  |
|  | The names of the classes between which is relations are to be defined. |
| maybe | What value to return if the relationship is conditional. |
| test, coerce, replace |  |
|  | Functions optionally supplied to test whether the relation is defined, to coerce the object to class2, and to alter the object so that is (object, class2) is identical to value. |
| by | The name of an intermediary class. Coercion will proceed by first coercing to this class and from there to the target class. (The intermediate coercions have to be valid.) |
| where | Where to store the metadata defining the relationship. Default is the global environment. |

## Details

```
setIs:
```

The relationship can be conditional, if a function is supplied as the test argument. If a function is supplied as the coerce argument, this function will be applied to any class1 object in order to turn it into a class2 object. If the relationship is to be defined indirectly through a third class, this class can be named in the by argument.
Extension may imply that a class1 object contains a class2 object. The default sense of containing is that all the slots of the simpler class are found in the more elaborate one. If the replace argument is supplied as an $S$ replacement function, this function will be used to implement as(obj, class2) <- value.

The coerce, replace, and by arguments behave as described for the setAs function. It's unlikely you would use the by argument directly, but it is used in defining cached information about classes. The value returned (invisibly) by setIs is the extension information, as a list.
Information about setIs relations can be stored in the metadata for either class1 (in the extends information) or in the metadata for class2 (in the subclasses information). For the information to be retained for a future session, one of these classes must be defined in the global environment, since only objects assigned there are saved by save.image. If neither class is defined in environment where, setIs generates an error.

Because only global environment information is saved, it rarely makes sense to give a value other than the default for argument where. One exception is where $=0$, which modifies the cached (i.e., session-scope) information about the class. Class completion computations use this version, but don't use it yourself unless you are quite sure you know what you're doing.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## Examples

```
## a class definition (see setClass for the example)
setClass("trackCurve",
    representation("track", smooth = "numeric"))
## A class similar to "trackCurve", but with different structure
## allowing matrices for the "y" and "smooth" slots
setClass("trackMultiCurve", representation(x="numeric", y="matrix", smooth="matrix"),
            prototype = structure(list(), x=numeric(), y=matrix(0,0,0), smooth= matrix(0,0,0)))
## Define a multi-curve to extend a single curve ONLY
## if the y data is one variable.
setIs("trackMultiCurve", "trackCurve", test = function(obj) {ncol(slot(obj, "y")) == 1},
    coerce = function(obj) { new("trackCurve", x = slot(obj, "x"),
            y = as.numeric(slot(obj,"y")), curve = as.numeric(slot(obj, "curve")))})
```

    language-class Classes to Represent Unevaluated Language Objects
    
## Description

The virtual class "language" and the specific classes that extend it represent unevaluated objects, as produced for example by the parser or by functions such as quote.

## Usage

\#\#\# each of these classes corresponds to an unevaluated object
\#\#\# in the $S$ language. The class name can appear in method signatures,
\#\#\# and in a few other contexts (such as some calls to as()).
" ("
"<-"
"call"
"for"
"if"
"repeat"
"while"
"name"
" $\{$ "
\#\#\# Each of the classes above extends the virtual class
"language"

## Objects from the Class

"language" is a virtual class; no objects may be created from it.
Objects from the other classes can be generated by a call to new (Class, ...), where Class is the quoted class name, and the ... arguments are either empty or a single object that is from this class (or an extension).

## Methods

coerce signature (from = "ANY", to = "call"). A method exists for as(object, "call"), calling as.call().
languageEl Elements of Language Objects

## Description

Internal routines to support some operations on language objects.

## Usage

languageEl(object, which)
isGrammarSymbol(symbol)

## Summary of Functions

languageEl: extract an element of a language object, consistently for different kinds of objects.
The 1st., etc. elements of a function are the corresponding formal arguments, with the default expression if any as value.
The first element of a call is the name or the function object being called.
The 2nd, 3rd, etc. elements are the 1st, 2nd, etc. arguments expressions. Note that the form of the extracted name is different for R and S-Plus. When the name (the first element) of a call is replaced, the languageEl replacement function coerces a character string to the internal form for each system.
The 1st, 2nd, 3rd elements of an if expression are the test, first, and second branch. The 1st element of a for object is the name (symbol) being used in the loop, the second is the expression for the range of the loop, the third is the body of the loop. The first element of a while object is the loop test, and the second the body of the loop.
isGrammarSymbol: Checks whether the symbol is part of the grammar. Don't use this function directly.

## LinearMethodsList-class

Class "LinearMethodsList"

## Description

A version of methods lists that has been "linearized" for producing summary information. The actual objects from class "MethodsList" used for method dispatch are defined recursively over the arguments involved.

## Objects from the Class

The function linearizeMlist converts an ordinary methods list object into the linearized form.

## Slots

methods: Object of class "list", the method definitions.
arguments: Object of class "list", the corresponding formal arguments.
classes: Object of class "list", the corresponding classes in the signatures.
fromClasses: Object of class "list"

## Future Note

The current version of linearizeMlist does not take advantage of the MethodDefinition class, and therefore does more work for less effect than it could. In particular, we may move to redefine both the function and the class to take advantage of the stored signatures. Don't write code depending precisely on the present form, although all the current information will be obtainable in the future.

## See Also

Function linearizeMlist for the computation, and MethodsList-class for the original, recursive form.

```
makeClassRepresentation
    Create a Class Definition
```


## Description

Constructs a classRepresentation-class object to describe a particular class. Mostly a utility function, but you can call it to create a class definition without assigning it, as setClass would do.

## Usage

makeClassRepresentation(name, slots=list(), superClasses=character(), prototype=NULL, package, validity, access, version, sealed, virtual=NA)

## Arguments

| name | character string name for the class <br> named list of slot classes as would be supplied to setClass, but without <br> the unnamed arguments for superClasses if any. |
| :--- | :--- |
| superClasses |  |
| prototype | what classes does this class extend <br> an object providing the default data for the class, e.g, the result of a call <br> to prototype. |
| package | The character string name for the package in which the class will be stored; <br> see getPackageName. <br> Optional validity method. See validObject, and the discussion of validity <br> methods in the reference. |
| validity | Access information. Not currently used. |
| access version <br> sealed Is this known to be a virtual class? <br> virtual  |  |

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

```
setClass
```

MethodDefinition-class

## Description

These classes extend the basic class "function" when functions are to be stored and used as method definitions.

## Details

Method definition objects are functions with additional information defining how the function is being used as a method. The target slot is the class signature for which the method will be dispatched, and the defined slot the signature for which the method was orignally specified (that is, the one that appeared in some call to setMethod).

## Objects from the Class

The action of setting a method creates an object of this class. It's unwise to create them directly.

## Slots

.Data: Object of class "function"; the data part of the definition.
target: Object of class "signature"; the signature for which the method was wanted.
defined: Object of class "signature"; the signature for which a method was found. If the method was inherited, this will not be identical to target.

## Extends

Class "function", from data part.
Class "PossibleMethod", directly.
Class "OptionalMethods", by class "function".

## See Also

class MethodsList-class for the objects defining sets of methods associated with a particular generic function. The individual method definitions stored in these objects are from class MethodDefinition, or an extension. MethodWithNext-class for an extension used by callNextMethod.

Methods General Information on Methods

## Description

This documentation section covers some general topics on how methods work and how the methods package interacts with the rest of R. The information is usually not needed to get started with methods and classes, but may be helpful for moderately ambitious projects, or when something doesn't work as expected.

The section How Methods Work describes the underlying mechanism; Class Inheritance and Method Selection provides more details on how class definitions determine which methods are used.

The section Changes with the Methods Package outlines possible effects on other computations when running with package methods.

## How Methods Work

A generic function is a function that has associated with it a collection of other functions (the methods), all of which agree in formal arguments with the generic. In R, the "collection" is an object of class "MethodsList", which contains a named list of methods (the methods slot), and the name of one of the formal arguments to the function (the argument slot). The names of the methods are the names of classes, and the corresponding element defines the method or methods to be used if the corresponding argument has that class. For example, suppose a function $f$ has formal arguments $x$ and $y$. The methods list object for that function has the object as.name("x") as its argument slot. An element of the methods named "track" is selected if the actual argument corresponding to x is an object of class "track". If there is such an element, it can generally be either a function or another methods list object.
In the first case, the function defines the method to use for any call in which x is of class "track". In the second case, the new methods list object defines the selection of methods depending on the remaining formal arguments, in this example, y. The same selection process takes place, recursively, using the new methods list. Eventually, the selection returns either a function or NULL, meaning that no method matched the actual arguments.
Each method selected corresponds conceptually to a signature; that is a named list of classes, with names corresponding to some or all of the formal arguments. In the previous example, if selecting class "track" for x , finding that the selection was another methods list and then selecting class "numeric" for y would produce a method associated with the signature $\mathrm{x}=$ "track", $\mathrm{y}=$ "numeric".
The actual selection is done recursively, but you can see the methods arranged by signature by calling the function showMethods, and objects with the methods arranged this way (in two different forms) are returned by the functions listFromMlist and linearizeMlist.
In an R session, each generic function has a single methods list object defining all the currently available methods. The session methods list object is created the first time the function is called by merging all the relevant method definitions currently visible. Whenever something happens that might change the definitions (such as attaching or detaching a package with methods for this function, or explicitly defining or removing methods), the merged methods list object is removed. The next call to the function will recompute the merged definitions.
When methods list are merged, they can come from two sources:

1. Methods list objects for the same function anywhere on the current search list. These are merged so that methods in an environment earlier in the search list override methods for the same function later in the search list. A method overrides only another method for the same signature. See the comments on class "ANY" in the section on Inheritance.
2. Methods list objects corresponding the group generic functions, if any, for this function. Any generic function can be defined to belong to a group generic. The methods for the group generic are available as methods for this function. The group generic can itself be defined as belong to a group; as a result there is a list of group generic functions. A method defined for a function and a particular signature overrides a method for the same signature for that function's group generic.

Merging is done first on all methods for a particular function, and then over the generic and its group generics.

The result is a single methods list object that contains all the methods directly defined for this function. As calls to the function occur, this information may be supplemented by inherited methods, which we consider next.

## Class Inheritance and Method Selection

If no method is found directly for the actual arguments in a call to a generic function, an attempt is made to match the available methods to the arguments by using inheritance.
Each class definition potentially includes the names of one or more classes that the new class extends. (These are sometimes called the superclasses of the new class.) These classes themselves may extend other classes. Putting all this information together produces the full list of superclasses for this class. (You can see this list for any class "A" from the expression extends("A").) In addition, any class implicitly extends class "ANY".
A method will be selected by inheritance if we can find a method in the methods list for a signature corresponding to any combination of superclasses for each of the relevant arguments. The search for such a method is performed by the function MethodsListSelect, working as follows.
For the first formal argument of the function, a list of classes is made up from the class itself, all its superclasses, and class "ANY". For each of these, the selection computation looks for an element of the methods with the corresponding name. Each time it finds one, it then calls the selection process recursively if necessary to select a method directly or by inheritance for the remaining arguments.

Each one of these recursive calls can fail or it can return a function (the method). As long as the calls fail, the selection process moves on to the next superclass and tries again. The last step corresponds to class "ANY", the default method defined at this level.
The effect of this definition of the selection process is to order all possible inherited methods, first by the superclasses for the first argument, then within this by the superclasses for the second argument, and so on. Superclasses are ordered by how direct they are: first, the direct superclasses, then the superclasses of these classes.

## Changes with the Methods Package

The methods package is designed to leave other computations in R unchanged. There are, however, a few areas where the default functions and behavior are overridden when running with the methods package attached. This section outlines those known to have some possible effect.
class: The methods package enforces the notion that every object has a class; in particular, class ( x ) is never NULL, as it would be for basic vectors, for example, when not using methods.
In addition, when assigning a class, the value is required to be a single string. (However, objects can have multiple class names if these were generated by old-style class computations. The methods package does not hide the "extra" class names.)
Computations using the notion of NULL class attributes or of class attributes with multiple class names are not really compatible with the ideas in the methods package. Formal classes and class inheritance are designed to give more flexible and reliable implementations of similar ideas.
If you do have to mix the two approaches, any operations that use class attributes in the old sense should be written in terms of attr ( $x$, "class"), not class(x). In particular, test for no class having been assigned with is.null(attr ( $x$, "class")).
Printing To provide appropriate printing automatically for objects with formal class definitions, the methods package overrides print.default, to look for methods for the generic function show, and to use a default method for objects with formal class definitions.
The revised version of print.default is intended to produce identical printing to the original version for any object that does not have a formally defined class, including honoring old-style print methods. So far, no exceptions are known.
plot A version of the plot function is included in the current methods package, differing from the one in the base package in that it has a y argument (necessary if methods for plot are to be defined for the y data; see the examples for setMethod). This version will move into base as soon as it is tested.

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

```
See Also
setGeneric, setClass
```

MethodsList MethodsList Objects

## Description

These functions create and manipulate MethodsList objects, the objects used in R to store methods for dispatch. You should not call any of these functions from code that you want to port to S-Plus. Instead, use the functions described in the references.

```
Usage
MethodsList(.ArgName, ...)
makeMethodsList(object, level=1)
SignatureMethod(names, signature, definition)
insertMethod(mlist, signature, args, def, cacheOnly)
inheritedSubMethodLists(object, thisClass, mlist, ev)
showMlist(mlist, includeDefs = TRUE, inherited = TRUE,
        classes, useArgNames, printTo = stdout() )
print.MethodsList(x, ...)
listFromMlist(mlist, prefix = list())
linearizeMlist(mlist, inherited = TRUE)
finalDefaultMethod(mlist, fname = "NULL")
mergeMethods(m1, m2)
loadMethod(method, fname, envir)
```


## Details

Note that MethodsList objects represent methods only in the R implementation. You can use them to find or manipulate information about methods, but avoid doing so if you want your code to port to S-Plus.

## Details

MethodsList: Create a MethodsList object out of the arguments.
Conceptually, this object is a named collection of methods to be dispatched when the (first) argument in a function call matches the class corresponding to one of the names. A final, unnamed element (i.e., with name " ") corresponds to the default method.
The elements can be either a function, or another MethodsList. In the second case, this list implies dispatching on the second argument to the function using that list, given a selection of this element on the first argument. Thus, method dispatching on an arbitrary number of arguments is defined.
MethodsList objects are used primarily to dispatch OOP-style methods and, in R, to emulate S4-style methods.
SignatureMethod: construct a MethodsList object containing (only) this method, corresponding to the signature; i.e., such that signature [[1]] is the match for the first argument, signature[[2]] for the second argument, and so on. The string "missing" means a match for a missing argument, and "ANY" means use this as the default setting at this level.

The first argument is the argument names to be used for dispatch corresponding to the signatures.
insertMethod: insert the definition def into the MethodsList object, mlist, corresponding to the signature. By default, insert it in the slot "methods", but cacheOnly=TRUE inserts it into the "allMethods" slot (used for dispatch but not saved).
inheritedSubMethodLists: Utility function to match the object or the class (if the object is NULL) to the elements of a methods list. Used in finding inherited methods, and not meant to be called directly.
showMlist: Prints the contents of the MethodsList. If includeDefs the signatures and the corresonding definitions will be printed; otherwise, only the signatures.
The function calls itself recursively: prev is the previously selected classes.
listFromMlistForPrint: Undo the recursive nature of the methods list, making a list of function defintions, with the names of the list being the corresponding signatures (designed for printing; for looping over the methods, use listFromMlist instead).
The function calls itself recursively: prev is the previously selected classes.
finalDefaultMethod: The true default method for the methods list object mlist (the method that matches class "ANY" for as many arguments as are used in methods matching for this generic function). If mlist is null, returns the function called fname, or NULL if there is no such function.
mergeMethods: Merges the methods in the second MethodsList object into the first, and returns the merged result. Called from getAllMethods.
loadMethod: Called, if necessary, just before a call to method is dispatched in the frame envir. The function exists so that methods can be defined for special classes of objects. Usually the point is to assign or modify information in the frame environment to be used evaluation. For example, the standard class MethodDefinition has a method that stores the target and defined signatures in the environment.
Any methods defined for loadMethod must return the function definition to be used for this call; typically, this is just the method argument.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

$$
\begin{array}{ll}
\text { MethodsList-class } & \begin{array}{l}
\text { Class MethodsList, Representation of Methods for a Generic } \\
\text { Function }
\end{array}
\end{array}
$$

## Description

Objects from this class are generated and revised by the definition of methods for a generic function.

## Slots

argument: Object of class "name". The name of the argument being used for dispatch at this level.
methods: A named list of the methods (and method lists) defined explicitly for this argument, with the names being the classes for which the methods have been defined.
allMethods: A named list, which may be empty if this object has not been used in dispatch yet. Otherwise, it contains all the directly defined methods from the methods slot, plus any inherited methods.

## Extends

Class "OptionalMethods", directly.

MethodSupport Additional (Support) Functions for Methods

## Description

These are support routines for computations on formal methods.

## Usage

getMethodsForDispatch(f, fdef)
cacheMethod(f, sig, def, args, fdef)
resetGeneric(f, fdef)

## Summary of Functions

resetGeneric: reset the currently defined methods for this generic by removing the corresponding definition from the methods metadata (to be remerged when the function is next called). Returns TRUE or FALSE according to whether information for the function was found in the metadata.
You must call this function when you change relevant inheritance information during a session, to guarantee that the new information is used if this generic has already been called.
cacheMethod: Store the definition for this function and signature in the method metadata for the function. Used to store extensions of coerce methods found through inheritance. No persistent effect, since the method metadata is session-scope only.
getMethodsForDispatch: Get the current methods list object representing the methods for function f , merged from the various packages and with any additional caching information stored in the allMethods slot.
If methods have not yet been merged, calling getMethodsForDispatch will cause the merge to take place.

```
methodUtilities Utility Functions for Methods and S-Plus Compatibility
```


## Description

These are utilities, currently in the methods package, that either provide some functionality needed by the package (e.g., element matching by name), or add compatibility with S-Plus, or both.

## Usage

functionBody(fun=sys.function(sys.parent()))
allNames(x)
getFunction(name, generic=TRUE, mustFind=TRUE, where)
el(object, where)
elNamed(x, name, mustFind=FALSE)
formalArgs (def)

Quote()
message(...)
showDefault(object, oldMethods = TRUE)
initMethodDispatch()

## Summary of Functions

allNames: the character vector of names (unlike names(), never returns NULL).
getFunction: find the object as a function.
elNamed: get the element of the vector corresponding to name. Unlike the [, [ [, and \$ operators, this function requires name to match the element name exactly (no partial matching).
formalArgs: Returns the names of the formal arguments of this function.
existsFunction: Is there a function of this name? If generic is FALSE, generic functions are not counted.
findFunction: return all the indices of the search list on which a function definition for name exists.
If generic is FALSE, ignore generic functions.
message: Output all the arguments, pasted together with no intervening spaces.
showDefault: Utility, used to enable show methods to be called by the automatic printing (via print.default).
initMethodDispatch: Turn on the internal method dispatch code. Called on attaching the package. Also, if dispatch has been turned off (by calling .isMethodsDispatchOn(FALSE) -a very gutsy thing to do), calling this function should turn dispatch back on again.

```
MethodWithNext-class Class MethodWithNext
```


## Description

Class of method definitions set up for callNextMethod

## Objects from the Class

Objects from this class are generated as a side-effect of calls to callNextMethod.

## Slots

.Data: Object of class "function"; the actual function definition.
nextMethod: Object of class "PossibleMethod" the method to use in response to a callNextMethod() call.
excluded: Object of class "list"; one or more signatures excluded in finding the next method.
target: Object of class "signature", from class "MethodDefinition"
defined: Object of class "signature", from class "MethodDefinition"

## Extends

Class "MethodDefinition", directly. Class "function", from data part. Class "PossibleMethod", by class "MethodDefinition". Class "OptionalMethods", by class "MethodDefinition".

## Methods

findNextMethod signature(method = "MethodWithNext"): used internally by method dispatch.
loadMethod signature(method $=$ "MethodWithNext"): used internally by method dispatch.
show signature(object = "MethodWithNext")

## See Also

callNextMethod, and MethodDefinition-class.

## Description

Given the the name or the definition of a class, plus optionally data to be included in the object, new returns an object from that class.

## Usage

new(Class, ...)
initialize(.Object, ...)

## Arguments

Class Either the name of a class (the usual case) or the object describing the class (e.g., the value returned by getClass).
... Data to include in the new object. Named arguments correspond to slots in the class definition. Unnamed arguments must be objects from classes that this class extends.
.Object An object: see the Details section.

## Details

The function new begins by copying the prototype object from the class definition. Then information is inserted according to the . . . arguments, if any.

The interpretation of the ... arguments can be specialized to particular classes, if an appropriate method has been defined for the generic function "initialize". The new function calls initialize with the object generated from the prototype as the .Object argument to initialize.

By default, unnamed arguments in the . . . are interpreted as objects from a superclass, and named arguments are interpreted as objects to be assigned into the correspondingly named slots. Thus, explicit slots override inherited information for the same slot, regardless of the order in which the arguments appear.

The initialize methods do not have to have ... as there second argument (see the examples), and generally it is better design not to have . . . as a formal argument, if only a fixed set of arguments make sense.

Note that the basic vector classes, "numeric", etc. are implicitly defined, so one can use new for these classes.

Author(s)
John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

Classes

## Examples

```
## using the definition of class "track" from Classes
## a new object with two slots specified
t1 <- new("track", x = seq(along=ydata), y = ydata)
# a new object including an object from a superclass, plus a slot
t2 <- new("trackCurve", t1, smooth = ysmooth)
### define a method for initialize, to ensure that new objects have
### equal-length x and y slots.
setMethod("initialize", "track", function(.Object, x = numeric(0), y = numeric(0)) {
    if(nargs() > 1) {
        if(length(x) != length(y))
            stop("specified x and y of different lengths")
            .Object@x <- x
            .Object@y <- y
        }
        .Object
    })
```

\#\#\# the next example will cause an error (x will be numeric(0)),
\#\#\# because we didn't build in defaults for $x$,
\#\#\# although we could with a more elaborate method for initialize
try(new("track", y = sort(rnorm(10))))
\#\# a better way to implement the previous initialize method.
\#\# Why? By using callNextMethod to call the default initialize method
\#\# we don't inhibit classes that extend "track" from using the general
\#\# form of the new() function. In the previous version, they could only
\#\# use x and y as arguments to new, unless they wrote their own
\#\# intialize method.
setMethod("initialize", "track", function(.Object, ...) \{
.Object <- callNextMethod()
if (length(.Object@x) != length(.Object@y))
stop("specified x and y of different lengths")
.Object
\})

```
ObjectsWithPackage-class
```

A Vector of Object Names, with associated Package Names

## Description

This class of objects is used to represent ordinary character string object names, extended with a package slot naming the package associated with each object.

## Objects from the Class

The function getGenerics returns an object of this class.

## Slots

.Data: Object of class "character": the object names.
package: Object of class "character" the package names.

## Extends

Class "character", from data part. Class "vector", by class "character".

## See Also

Methods for general background.

```
promptClass Generate a Shell for Documentation of a Formal Class
```


## Description

Assembles all relevant slot and method information for a class, with minimal markup for Rd processing; no QA facilities at present.

## Usage

```
promptClass(clName, filename, type = "class", keywords = "classes", where)
```


## Arguments

| clName | character string naming the class to be documented. |
| :--- | :--- |
| filename | Usually, the filename on which the documentation shell should be written. <br> By default it is the topic name for the class documentation, followed by <br> ".Rd". See the example below. The argument can also be any writable <br> connection. |
| type | The documentation type to be declared in the output file. |
| keywords | The keywords to include in the shell of the documentation. The keyword <br>  <br> "classes" should be one of them. <br> where |
|  | where to look for the definition of the class and of methods that use it: <br> by default, anywhere in the current search list. |

## Details

The class definition is found on the search list. Using that definition, information about classes extended and slots is determined.
In addition, the currently available generics with methods for this class are found (using getGenerics). Note that these methods need not be in the same environment as the class definition; in particular, this part of the output may depend on which packages are currently in the search list.
As with other prompt-style functions, the documentation shell is written to a file, which will need editing to give information about the meaning of the class. The output of promptClass can only contain information from the metadata about the formal definition and how it is used.

## Value

The name of the file to which the shell is written (the value is invisible). A message is also printed notifying the user about the file.

## Author(s)

VJ Carey 〈stvjc@channing.harvard.edu〉 and John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

prompt for documentation of functions, promptMethods for documentation of method definitions.

For processing of the edited documentation, either use $R$ CMD Rdconv, or include the edited file in the 'man' subdirectory of a package.

## Examples

```
> promptClass("track")
A shell of class documentation has been written to the
file "track-class.Rd".
```

promptMethods
Generate a Shell for Documentation of Formal Methods

## Description

Generates a shell of documentation for the methods of a generic function.

```
Usage
    promptMethods(f, filename, methods)
```


## Arguments

f
filename
methods

The name of the generic function whose methods are to be documented.
Optional file on which to write the documentation shell. If supplied, it can be the name of a file, a connection, or FALSE.
If FALSE, the text is returned, presumably to be inserted some other documentation file, such as the documentation of the generic function itself (see prompt).
The default file name is the same as the coded topic name for these methods (currently "f-methods.Rd".
Optional methods list object giving the methods to be documented. By default, the first methods object for this generic is used (for example, if the current global environment has some methods for $f$, these would be documented).
If this argument is supplied, it is likely to be getMethods(f, where), with where some package containing methods for $f$.

## Value

If filename is FALSE, the text generated; otherwise, the name of the file written.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

prompt and promptClass

RClassUtils
Utilities for Managing Class Definitions

## Description

These are various functions to support the definition and use of formal classes. Most of them are rarely suitable to be called directly.

## Usage

```
testVirtual(properties, extends, prototype)
makePrototypeFromClassDef(properties, prototype, extends)
newEmptyObject()
completeClassDefinition(Class, ClassDef)
getProperties(ClassDef)
getSlots(x, complete = TRUE)
getExtends(ClassDef)
getAccess(ClassDef)
getAllSuperClasses(ClassDef)
superClassDepth(ClassDef, soFar)
getPrototype(ClassDef)
getVirtual(ClassDef)
isVirtualClass(Class)
getSubclasses(ClassDef)
getClassName(ClassDef)
assignClassDef(Class, def, where=.GlobalEnv)
newBasic(Class, ...)
makeExtends(Class, to, coerce, test, replace, by, package, slots,
                                    classDef1, classDef2)
reconcilePropertiesAndPrototype(name, properties, prototype, superClasses)
tryNew(Class)
trySilent(expr)
empty.dump()
showClass(Class, complete=TRUE, propertiesAreCalled="Properties")
showExtends(ext, printTo = stdout())
getFromClassMetaData(name)
```

```
assignToClassMetaData(name, value)
removeFromClassMetaData(name)
possibleExtends(class1, class2)
completeExtends(ClassDef)
classMetaName(name)
methodsPackageMetaName(prefix, name)
metaNameUndo(strings, prefix = "M", searchForm = FALSE)
requireMethods(functions, signature, message)
checkSlotAssignment(obj, name, value)
defaultPrototype()
isClassDef(object)
validSlotNames(names)
getDataPart(object)
setDataPart(object, value)
```


## Summary of Functions

testVirtual: Test for a Virtual Class. Figures out, as well as possible, whether the class with these properties, extension, and prototype is a virtual class. Can be forced to be virtual by extending "VIRTUAL".
Otherwise, a class is virtual only if it has no slots, extends no non-virtual classes, and has a NULL Prototype.
makePrototypeFromClassDef: Makes the prototype implied by the class definition.
The following three rules are applied in this order.
If the class has slots, then the prototype for each slot is used by default, but a corresponding element in the explicitly supplied prototype, if there is one, is used instead (but it must be coercible to the class of the slot).
If there are no slots but a non-null prototype was specified, this is returned.
If there is a single non-virtual superclass (a class in the extends list), then its prototype is used.
If all three of the above fail, the prototype is NULL.
newEmptyObject: Utility function to create an empty object into which slots can be set.
Currently just creates an empty list with class "NULL".
Later version should create a special object reference that marks an object currently with no slots and no data.
completeClassDefinition: Completes the definition of Class, relative to the current session.

The completed definition is stored in the session's class metadata, to be retrieved the next time that getClass is called on this class, and is returned as the value of the call. If ClassDef is omitted, the initial definition is obtained from the first package having a meta-object for this class.
getFromClassDef: Extracts one of the intrinsically defined class definition properties (".Properties", etc.) Strictly a utility function.
getProperties: Extracts the class's Properties information from the class representation (only, not from the name of the class).
getSlots: Returns a named character vector. The names are the names of the slots, the values are the classes of the corresponding slots. If complete is TRUE, all slots from all superclasses will be included. The argument x can either be the name of a class or an object having that class.
getExtends: Extracts the class's Extends information from the class representation (only, not from the name of the class)
Contrast with the possibleExtends and is functions, both of which use indirect information as well.
getAllSuperClasses, superClassDepth: Get the names of all the classes that this class definition extends.
getAllSuperClasses is a utility function used to complete a class definition. It returns all the superclasses reachable from this class, in breadth-first order (which is the order used for matching methods); that is, the first direct superclass followed by all its superclasses, then the next, etc. (The order is relevant only in the case that some of the superclasses have multiple inheritance.)
superClassDepth, which is called from getAllSuperClasses, returns the same information, but as a list with components label and depth, the latter for the number of generations back each class is in the inheritance tree. The argument soFar is used to avoid loops in the network of class relationships.
getPrototype: extract the class's Prototype information from the class representation (only, not from the name of the class)
getAccess: extract the class's Access information from the class representation (only, not from the name of the class)
getVirtual: extract the class's Virtual information from the class representation (only, not from the name of the class)
isVirtualClass: Is the named class a virtual class?
A class is virtual if explicitly declared to be, and also if the class is not formally defined.
getSubclasses: extract the class's Subclasses information from the class representation (only, not from the name of the class)
getClassName: The internal property in the class definition for the class name.
assignClassDef: assign the definition of the class to the specially named object
newBasic: the implementation of the function new for basic classes that don't have a formal definition.
Any of these could have a formal definition, except for Class="NULL" (disallowed because NULL can't have attributes). For all cases except "NULL", the class of the result will be set to Class.
See new for the interpretation of the arguments.
makeExtends: convert the argument to a list defining the extension mechanism.
reconcilePropertiesAndPrototype: makes a list or a structure look like a prototype for the given class.
Specifically, returns a structure with attributes corresponding to the slot names in properties and values taken from prototype if they exist there, from new(classi) for the class, classi of the slot if that succeeds, and NULL otherwise.
The prototype may imply slots not in the properties list, since properties does not include inherited slots (these are left unresolved until the class is used in a session).
tryNew: Tries to generate a new element from this class, but if the attempt fails (as, e.g., when the class is undefined or virtual) just returns NULL.
This is inefficient and also not a good idea when actually generating objects, but is useful in the initial definition of classes.
showClass: Print the information about a class definition.
If complete is TRUE, include the indirect information about extensions.
showExtends: Print the elements of the list of extensions.
(Used also by promptClass to get the list of what and how for the extensions.)
possibleExtends: Find the information that says whether class1 extends class2, directly or indirectly.
This can be either a logical value or an object of class SClassExtension-class containing various functions to test and/or coerce the relationship.
completeExtends: complete the extends information in the class definition, by following transitive chains.
Elements in the immediate extends list may be added and current elements may be replaced, either by replacing a conditional relation with an unconditional one, or by adding indirect relations.
classMetaName: a name for the object storing this class's definition
methodsPackageMetaName: a name mangling device to hide metadata defining method and class information.
metaNameUndo As its name implies, this function undoes the name-mangling used to produce meta-data object names, and returns a object of class ObjectsWithPackage-class.
requireMethods: Require a subclass to implement methods for the generic functions, for this signature.
For each generic, setMethod will be called to define a method that throws an error, with the supplied message.
The requireMethods function allows virtual classes to require actual classes that extend them to implement methods for certain functions, in effect creating an API for the virtual class.
Otherwise, default methods for the corresponding function would be called, resulting in less helpful error messages or (worse still) silently incorrect results.
checkSlotAssignment: Check that the value provided is allowed for this slot, by consulting the definition of the class. Called from the C code that assigns slots.
For privileged slots (those that can only be set by accesor functions defined along with the class itself), the class designer may choose to improve efficiency by validating the value to be assigned in the accessor function and then calling slot<- with the argument check=FALSE, to prevent the call to checkSlotAssignment.
defaultPrototype: The prototype for a class which will have slots, is not a virtual class, and does not extend one of the basic classes. In future releases, this will likely be a non-vector R object type, but none of the current types (as of release 1.4) is suitable.

SessionClassMetaData: Contains the name of the special table in which class information is cached during the session.
.InitBasicClasses, .InitMethodsListClass, .setCoerceGeneric: These functions perform part of the initialization of classes and methods, and are called (only!) from .First.lib.
isClassDef: Is object a representation of a class?
validSlotNames: Returns names unless one of the names is reserved, in which case there is an error. (As of writing, "class" is the only reserved slot name.)
getDataPart, setDataPart: Utilities called from the base $C$ code to implement object@.Data.
representation Construct a Representation or a Prototype for a Class Definition

## Description

In calls to setClass, these two functions construct, respectively, the representation and prototype arguments. They do various checks and handle special cases. You're encouraged to use them when defining classes that, for example, extend other classes as a data part or have multiple superclasses, or that combine extending a class and slots.

## Usage

representation(...)
prototype(...)

## Arguments

... The call to representation takes arguments that are single character strings. Unnamed arguments are classes that a newly defined class extends; named arguments name the explicit slots in the new class, and specify what class each slot should have.
In the call to prototype, if an unnamed argument is supplied, it unconditionally forms the basis for the prototype object. Remaining arguments are taken to correspond to slots of this object. It is an error to supply more than one unnamed argument.

## Details

The representation function applies tests for the validity of the arguments. Each must specify the name of a class.
The classes named don't have to exist when representation is called, but if they do, then the function will check for any duplicate slot names introduced by each of the inherited classes.

The arguments to prototype are usually named initial values for slots, plus an optional first argument that gives the object itself. The unnamed argument is typically useful if there is a data part to the definition (see the examples below).

## Value

The value pf representation is just the list of arguments, after these have been checked for validity.

The value of prototype is the object to be used as the prototype. Slots will have been set consistently with the arguments, but the construction does not use the class definition to test validity of the contents (it hardly can, since the prototype object is usually supplied to create the definition).

## Author(s)

John Chambers

## References

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

## See Also

```
setClass
```


## Examples

```
## representation for a new class with a directly define slot "smooth"
## which should be a "numeric" object, and extending class "track"
representation("track", smooth ="numeric")
setClass("Character",representation("character"))
setClass("TypedCharacter",representation("Character",type="character"),
    prototype(character(0),type="plain"))
ttt <- new("TypedCharacter", "foo", type = "character")
```

RMethodUtils RMethodUtils

## Description

Utility functions to support the definition and use of formal methods. Most of these functions will not normally be called directly by the user.

## Usage

```
makeGeneric(f, fdef, fdefault, group=character(),
    valueClass=character(), package, signature, genericFunction)
makeStandardGeneric(f, fdef)
generic.skeleton(name, fdef, fdefault)
defaultDumpName(generic, signature)
getAllMethods(f, fdef, libs=search())
setAllMethodsSlot(mlist)
doPrimitiveMethod(name, def, call=sys.call(-1), ev=sys.frame(-2))
conformMethod(signature, mnames, fnames)
getGeneric(f, mustFind=FALSE)
getGroup(fdef, recursive=FALSE)
matchSignature(signature, fun)
## manage method metadata
getFromMethodMetaData(name)
assignToMethodMetaData(name, value)
removeFromMethodMetaData(name)
removeMethodsObject(f, where)
findUnique(what, doFind, message)
MethodAddCoerce(method, argName, thisClass, methodClass)
is.primitive(fdef)
cacheMetaData(where, attach = TRUE)
cacheGenericsMetaData(generics, attach = TRUE, where, package)
setPrimitiveMethods(f, fdef, code, generic, mlist)
printNoClass(x, digits, quote, na.print, print.gap, right, ...)
print.default(x, ...)
missingArg(symbol, envir = parent.frame(), eval)
balanceMethodsList(mlist, args, check = TRUE)
```

```
sigToEnv(signature)
rematchDefinition(definition, generic, mnames, fnames)
unRematchDefinition(definition)
asMethodDefinition(def, signature)
findNextMethod(method, f, mlist, optional = FALSE, envir)
```


## Summary of Functions

makeGeneric: Makes a generic function object corresponding to the given function name, optional definition and optional default method. Other arguments supply optional elements for the slots of genericFunction-class.
makeStandardGeneric: a utility function that makes a valid function calling standardGeneric for name f. Works (more or less) even if the actual definition, fdef, is not a proper function, that is, it's a primitive or internal.
conformMethod: If the formal arguments, mnames, are not identical to the formal arguments to the function, fnames, conformMethod determines whether the signature and the two sets of arguments conform, and returns the signature, possibly extended.
The method assignment conforms if either method and function have identical formal argument lists. It can also conform if the method omits some of the formal arguments of the function but: (1) the non-omitted arguments are a subset of the function arguments, appearing in the same order; (2) there are no arguments to the method that are not arguments to the function; and (3) the omitted formal arguments do not appear as explicit classes in the signature.
defaultDumpName: the default name to be used for dumping a method.
getAllMethods: a generic function (with methods) representing the merge of all the versions of $f$ on the specified packages (anything on the current search path by default). If the generic $f$ has a group generic, methods for this group generic (and further generations of group generics, if any) are also merged.
The merging rule is as follows: each generic is merged across packages, and the group generics are then merged, finally adding the directly defined methods of $f$.
The effect of the merging rule is that any method directly defined for $f$ on any included package overrides a method for the same signature defined for the group generic; similarly for the group generic and its group, if any, etc.
For $f$ or for a specific group generic, methods override in the order of the packages being searched. A method for a particular signature on a particular package overrides any methods for the same signature on packages later on in the list of packages being searched.
The slot "allMethods" of the merged methods list is set to a copy of the methods slot; this is the slot where inherited methods are stored.
doPrimitiveMethod: do a primitive call to builtin function name the definition and call provided, and carried out in the environment ev.
A call to doPrimitiveMethod is used when the actual method is a .Primitive. (Because primitives don't behave correctly as ordinary functions, not having either formal arguments nor a function body).
getGeneric: return the definition of the function named $f$ as a generic.
If there is no definition in the current search list, throws an error or returns NULL according to the value of mustFind.
Primitive functions are dealt with specially, since there is never a formal generic definition for them. The value returned is the formal definition used for assigning methods to this primitive. Not all primitives can have methods; if this one can't, then getGeneric returns NULL or throws an error.
getGroup: return the groups to which this generic belongs.
If recursive=TRUE, also all the group(s) of these groups.
matchSignature Matches the signature object (a partially or completely named subset of the signature arguments of the generic function object fun), and return a vector of all the classes in the order specified by fun@signature. The classes not specified by 'signature' will be "ANY" in the value, but extra trailing "ANY"'s are removed. When the input signature is empty, the returned signature is a single "ANY" matching the first formal argument (so the returned value is always non-empty).
Generates an error if any of the supplied signature names are not legal; that is, not in the signature slot of the generic function.
getMethodsMetaData, assignMethodsMetaData, mlistMetaName: utilities to manage methods list objects in a particular environment. Not to be called directly.
getFromMethodMetaData, assignToMethodMetaData, removeFromMethodMetaData Functions to manage the session metadata for methods. Don't call these directly.
MethodAddCoerce Possibly modify one or more methods to explicitly coerce this argument to methodClass, the class for which the method is explicitly defined. Only modifies the method if an explicit coerce is required to coerce from thisClass to methodClass.
is.primitive Is this object a primitive function (either a builtin or special)?
removeMethodsObject: remove the metadata object containing methods for $f$.
findUnique: Find the first position on the search list containing object what; if more than one is found, a warning message is generated, using message to identify what was being searched for.
If doFind is supplied, it's the version of find used to do the search (e.g., findFunction.
cacheMetaData, cacheGenericsMetaData, setPrimitiveMethods: Utilities for ensuring that the session-scope information about class and method definitions is up to date. Should normally be called automatically whenever needed (for example, when a method or class definition changes, or when a package is attached or detached.
The environment must be one of the environments on the current search list; note in particular that even on detaching (attach=FALSE), the environment will normally still be on the search list.
The setPrimitiveMethods function resets the caching information for a particular primitive function. Don't call it directly.
printNoClass,print.default: printNoClass is equivalent to the version of print.default in the base package. The methods package overrides the latter function to provide meaningful printing for formally defined classes, and printNoClass is used to get the original default printing.
missingArg: Returns TRUE if the symbol supplied is missing from the call corresponding to the environment supplied (by default, environment of the call to missingArg). If eval is true, the argument is evaluated to get the name of the symbol to test. Note that missingArg is closer to the "blue-book" sense of the missing function, not that of the current R base package implementation. But beware that it works reliably only
if no assignment has yet been made to the argument. (For method dispatch this is fine, because computations are done at the begining of the call.)
balanceMethodsList: Called from setMethod to ensure that all nodes in the list have the same depth (i.e., the same number of levels of arguments). Balance is needed to ensure that all necessary arguments are examined when inherited methods are being found and added to the allMethods slot. No actual recomputation is needed usually except when a new method uses a longer signature than has appeared before.
Balance requires that all methods be added to the generic via setMethod (how else could you do it?) or by the initial setGeneric call converting the ordinary function.
sigToEnv: Turn the signature (a named vector of classes) into an environment with the classes assigned to the names. The environment is then suitable for calling MethodsListSelect, with evalArgs=FALSE, to select a method corresponding to the signature. Usually not called directly: see selectMethod.
.saveImage: Flag, used in dynamically initializing the methods package from .First.lib

SClassExtension-class
Class to Represent Inheritance (Extension) Relations

## Description

An object from this class represents a single "is" relationship; lists of these objects are used to represent all the extensions (superclasses) and subclasses for a given class. The object contains information about how the relation is defined and methods to coerce, test, and replace correspondingly.

## Objects from the Class

Objects from this class are generated by setIs, both from direct calls

## Slots

superClass: The name of the class being extended.
package: The package to which that class belongs.
coerce: A function to carry out the as() computation implied by the relation. Note that these functions should not be used directly. They only deal with the strict=TRUE calls to the as function, with the full method constructed from this mechanically.
test: The function that would test whether the relation holds. Except for explicitly specified test arguments to setIs, this function is trivial.
replace: The method used to implement as(x, Class) <- value.
simple: A "logical" flag, TRUE if this is a simple relation, either because one class is contained in the definition of another, or because a class has been explicitly stated to extend a virtual class. For simple extensions, the three methods are generated automatically.
by: If this relation has been constructed transitively, the first intermediate class from the subclass.
dataPart: A "logical" flag, TRUE if the extended class is in fact the data part of the subclass. In this case the extended class is a basic class (i.e., a type).

## Methods

No methods defined with class "SClassExtension" in the signature.

## See Also

is, as and classRepresentation-class.

## Session Deprecated: Session Data and Debugging Tools

## Description

The functions traceOn and traceOff have been replaced by extended versions of the functions trace and untrace, and should not be used.

## Usage

sessionData()
traceOn(what, tracer=browseAll, exit=NULL)
traceOff(what)
browseAll()

## Details

sessionData: return the index of the session data in the search list, attaching it if it is not attached.
trace0n: initialize tracing on calls to function what. The function or expression tracer is called on entry, and the function or expression exit on exit.
traceOff: turn off tracing of this function.
browseAll: browse the current stack of function calls.
Uses the function debugger to set up browser calls on the frames. On exit from that function, computation continues after the call to browseAll. Computations done in the frames will have no effect.

## Author(s)

John Chambers

## References

See Programming with Data (John M. Chambers, Springer, 1998) for the equivalent functions.

```
setClass
Create a Class Definition
```


## Description

Create a formally defined class with specified slots and/or relationships to other classes. Also functions to remove a class definition, to test whether a class has been defined, to test whether an object is a class definition, and to reset the internal definition of a class.

## Usage

```
setClass(Class, representation, prototype, contains=character(), validity,
            access, where=1, version=FALSE, sealed, package)
removeClass(Class, where=-1, removeSubclassLinks = TRUE)
isClass(Class, formal=TRUE)
isClassDef(object)
getClasses(where)
findClass(Class)
resetClass(Class, resetSubclasses = TRUE)
```


## Arguments

Class character string name for the class
representation
the slots that the new class should have and/or other classes that this class extends. Usually a call to the representation function.
prototype an object (usually a list) providing the default data for the slots specified in the representation.
contains what classes does this class extend? (These are called superclasses in some languages.) When these classes have slots, all their slots will be contained in the new class as well.
where What environment to use to store or remove the definition (as metadata). By default, uses the global environment for setClass and searches for a definition to remove, for removeClass.
validity, access, version
Control arguments included for compatibility with the S-Plus API, but not currently used.
sealed If TRUE, the class definition will be sealed, so that another call to setClass will fail on this class name.
package An optional package name for the class. By default (and usually) the package where the class definition is assigned will be used.
$\mathrm{x} \quad$ an arbitrary object.
formal Should a formal definition be required?
object any R object.
removeSubclassLinks
When a class is removed, any links to that class from other classes will become invalid. If this argument is not supplied as FALSE, then removeClass will search for all such links and delete them. You can omit the argument, or supply it as the positions in the search list to look for these links (by default all attached object tables will be searched).
resetSubclasses
Should resetClass also reset all known subclasses. Usually TRUE, unless you know from the context these will be reset elsewhere.

## Details

These are the functions that create and manipulate formal class definitions. Brief documentation is provided below. See the references for an introduction and for more details.
setClass: Define Class to be an S-style class. The effect is to create an object, of class "classRepEnvironment", and store this (hidden) in the specified environment or database. Objects can be created from the class (e.g., by calling new), manipulated (e.g., by accessing the object's slots), and methods may be defined including the class name in the signature (see setMethod).
removeClass: Remove the definition of this class. Calling this always resets the version of the class cached for the session. If where $=0$, that's all it does. Otherwise, it removes the version from the specified environment or database (from the global environment by default).
isClass: Is this a the name of a formally defined class? (Argument formal is for compatibility and is ignored.)
isClassDef: Is this object a class definition (it will be, for example, if it is the value of a call to getClass, the complete definition of a class with its extensions, or to getClassDef, the local definition of the class).
getClasses: The names of all the classes formally defined on where. If called with no argument, all the classes currently known in the session (which does not include classes that may be defined on one of the attached packages, but have not yet been used in the session).
findClass: Where on the current search list the class named Class is defined. (If there is more than one definition, all corresponding elements of the search list are returned.)
unclass: Returns the object containing the values of all the slots in this object's class definition (specifically, if the returned object has attributes corresponding to each slot), in the case that the object's class is formally defined with slots. For classes that extend a single other class (e.g., a basic class such as "numeric") the result is an object of that class.
resetClass: Reset the internal definition of a class. The effect is that the next time the definition of this class is needed, it will be recomputed from the information in the currently attached packages.
This function is called when aspects of the class definition are changed. You would need to call it explicitly if you changed the definition of a class that this class extends (but doing that in the middle of a session is living dangerously, since it may invalidate existing objects).

## Inheritance and Prototypes

Defining new classes that inherit from ("extend") other classes is a powerful technique, but has to be used carefully and not over-used. Otherwise, you will often get unintended results when you start to compute with objects from the new class.
As shown in the examples below, the simplest and safest form of inheritance is to start with an explicit class, with some slots, that does not extend anything else. It only does what we say it does.
Then extensions will add some new slots and new behavior.
Another variety of extension starts with one of the basic classes, perhaps with the intension of modifying R's standard behavior for that class. Perfectly legal and sometimes quite helpful, but you may need to be more careful in this case: your new class will inherit much of the behavior of the basic (informally defined) class, and the results can be surprising. Just proceed with caution and plenty of testing.
As an example, the class "matrix" is included in the pre-defined classes, to behave essentially as matrices do without formal class definitions. Suppose we don't like all of this; in particular, we want the default matrix to have 0 rows and columns (not 1 by 1 as it is now).

```
setClass("myMatrix", "matrix", prototype = matrix(0,0,0))
```

The arguments above illustrate two short-cuts relevant to such examples. We abbreviated the representation argument to the single superclass, because the new class doesn't add anything to the representation of class "matrix". Also, we provided an object from the superclass as the prototype, not a list of slots.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

Methods, makeClassRepresentation

## Examples

```
## A simple class with two slots
setClass("track",
    representation(x="numeric", y="numeric"))
## A class extending the previous, adding one more slot
setClass("trackCurve",
    representation("track", smooth = "numeric"))
## A class similar to "trackCurve", but with different structure
## allowing matrices for the "y" and "smooth" slots
setClass("trackMultiCurve", representation(x="numeric", y="matrix", smooth="matrix"),
    prototype = list(x=numeric(), y=matrix(0,0,0), smooth= matrix(0,0,0)))
##
```

```
## Suppose we want trackMultiCurve to be like trackCurve when there's only
## one column
## First, the wrong way.
try(setIs("trackMultiCurve", "trackCurve",
    test = function(obj) {ncol(slot(obj, "y")) == 1}))
## why didn't that work? You can only override the slots "x", "y", and "smooth"
## if you provide an explicit coerce function to correct any inconsistencies:
setIs("trackMultiCurve", "trackCurve",
    test = function(obj) {ncol(slot(obj, "y")) == 1},
    coerce = function(obj) { new("trackCurve", x = slot(obj, "x"),
            y = as.numeric(slot(obj,"y")), smooth = as.numeric(slot(obj, "smooth")))})
```

```
setGeneric Define a New Generic Function
```


## Description

Create a new generic function of the given name, for which formal methods can then be defined. Typically, an existing non-generic function becomes the default method, but there is much optional control. See the details section.

## Usage

```
setGeneric(name, def, group=NULL, valueClass=NULL, where=1, package,
                        signature, useAsDefault, genericFunction)
setGroupGeneric(name, def, group=NULL, valueClass=NULL, knownMembers,
                package, where=1)
```


## Arguments

name The character string name of the generic function. In the simplest and most common case, a function of this name is already defined. The existing function may be non-generic or already a generic (see the details).
def An optional function object, defining the generic. This argument is usually only needed (and is then required) if there is no current function of this name. In that case, the formal arguments and default values for the generic are taken from def. You can also supply this argument if you want the generic function to do something other than just dispatch methods (an advanced topic best left alone unless you are sure you want it).
Note that def is not the default method; use argument useAsDefault if you want to specify the default separately.
group Optionally, a character string giving the group of generic functions to which this function belongs. Methods can be defined for the corresponding group generic, and these will then define methods for this specific generic function, if no method has been explicitly defined for the corresponding signature. See the references for more discussion.

| valueClass | An optional character vector or unevaluated expression. The value returned by the generic function must have (or extend) this class, or one of the classes; otherwise, an error is generated. See the details section for supplying an expression. |
| :---: | :---: |
| package | The name of the package with which this function is associated. Usually determined automatically (as the package containing the non-generic version if there is one, or else the package where this generic is to be saved). |
| where | Where to store the resulting initial methods definition, and possibly the generic function; by default, stored into the top-level environment. |
| signature | Optionally, the signature of arguments in the function that can be used in methods for this generic. By default, all arguments other than . . . can be used. The signature argument can prohibit methods from using some arguments. The argument, if provided, is a vector of formal argument names. |
| genericFunction |  |
|  | The object to be used as a (nonstandard) generic function definition. Supply this explicitly only if you know what you are doing! |
| useAsDefault | Override the usual choice of default argument (an existing non-generic function or no default if there is no such function). Argument useAsDefault can be supplied, either as a function to use for the default, or as a logical value. FALSE says not to have a default method at all, so that an error occurs if there is not an explicit or inherited method for a call. TRUE says to use the existing function as default, unconditionally (hardly ever needed as an explicit argument). See the section on details. |
| knownMembers | (For setGroupGeneric only) The names of functions that are known to be members of this group. This information is used to reset cached definitions of the member generics when information about the group generic is changed. |

## Details

The setGeneric function is called to initialize a generic function in an environment (usually the global environment), as preparation for defining some methods for that function.

The simplest and most common situation is that name is already an ordinary non-generic function, and you now want to turn this function into a generic. In this case you will most often supply only name. The existing function becomes the default method, and the special group and valueClass properties remain unspecified.
A second situation is that you want to create a new, generic function, unrelated to any existing function. In this case, you need to supply a skeleton of the function definition, to define the arguments for the function. The body of a generic function is usually a standard form, standardGeneric (name) where name is the quoted name of the generic function.
When calling setGeneric in this form, you would normally supply the def argument as a function of this form. If not told otherwise, setGeneric will try to find a non-generic version of the function to use as a default. If you don't want this to happen, supply the argument useAsDefault. That argument can be the function you want to be the default method. You can supply the argument as FALSE to force no default (i.e., to cause an error if there is not direct or inherited method on call to the function).

The same no-default situation occurs if there is no non-generic form of the function, and useAsDefault=FALSE. Remember, though, you can also just assign the default you want (even one that generates an error) rather than relying on the prior situation.

Usually, calling setGeneric if there is already a generic function of this name has no effect. If you want to force a new definition, supply doAssign $=$ TRUE (but it would be cleaner in most cases to remove the old generic before creating the new one; see removeGeneric). There is one absolute restriction: you cannot create an explicit generic for the primitive functions in the base library. These are dispatched from C code for efficiency and are not to be redefined in any case.
As mentioned, the body of a generic function usually does nothing except for dispatching methods by a call to standardGeneric. Under some circumstances you might just want to do some additional computation in the generic function itself. As long as your function eventually calls standardGeneric that is permissible (though perhaps not a good idea, in that it makes the behavior of your function different from the usual S model). If your explicit definition of the generic function does not call standardGeneric you are in trouble, because none of the methods for the function will ever be dispatched.

By default, the generic function can return any object. If valueClass is supplied, it should be a vector of class names; the value returned by a method is then required to satisfy is (object, Class) for one of the specified classes. An empty (i.e., zero length) vector of classes means anything is allowed. Note that more complicated requirements on the result can be specified explicitly, by defining a non-standard generic function.

The setGroupGeneric function behaves like setGeneric except that it constructs a group generic function, differing in two ways from an ordinary generic function. First, this function cannot be called directly, and the body of the function created will contain a stop call with this information. Second, the group generic function contains information about the known members of the group, used to keep the members up to date when the group definition changes, through changes in the search list or direct specification of methods, etc.

## Value

The setGeneric function exists for its side effect: saving the generic function to allow methods to be specified later. It returns name.

## Generic Functions and Primitive Functions

A number of the basic $R$ functions are specially implemented as primitive functions, to be evaluated directly in the underlying $C$ code rather than by evaluating an $S$ language definition. Primitive functions are eligible to have methods, but are handled differently by setGeneric and setGroupGeneric. A call to setGeneric for a primitive function does not create a new definition of the function, and the call is allowed only to "turn on" methods for that function.

A call to setGeneric for a primitive causes the evaluator to look for methods for that generic; a call to setGroupGeneric for any of the groups that include primitives ("Arith", "Logic", "Compare", "Ops", "Math", "Math2", "Summary", and "Complex") does the same for each of the functions in that group.

You usually only need to use either function if the methods are being defined only for the group generic. Defining a method for a primitive function, say "+", by a call to setMethod turns on method dispatch for that function. But in R defining a method for the corresponding group generic, "Arith", does not currently turn on method dispatch (for efficiency reasons). If there are no non-group methods for the functions, you have two choices.

You can turn on method dispatch for all the functions in the group by calling setGroupGeneric("Arith"), or you can turn on method dispatch for only some of the functions by calling setGeneric ("+"), etc. Note that in either case you should give the name of the generic function as the only argument.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections 1.6, 2.7, 2.8, and chapters 7 and 8.

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

Methods for a discussion of other functions to specify and manipulate the methods of generic functions.

## Examples

```
### A non-standard generic function. It insists that the methods
### return a non-empty character vector (a stronger requirement than
### valueClass = "character" in the call to setGeneric)
setGeneric("authorNames",
    function(text) {
        value <- standardGeneric("authorNames")
        if(!(is(value, "character") && any(nchar(value)>0)))
            stop("authorNames methods must return non-empty strings")
        value
        })
## An example of group generic methods, using the class
## "track"; see the documentation of setClass for its definition
#define a method for the Arith group
setMethod("Arith", c("track", "numeric"),
    function(e1, e2){
        e1@y <- callGeneric(e1@y , e2)
        e1
})
setMethod("Arith", c("numeric", "track"),
    function(e1, e2){
        e2@y <- callGeneric(e1, e2@y)
        e2
})
```

```
# now arithmetic operators will dispatch methods:
t1 <- new("track", x=1:10, y=sort(rnorm(10)))
t1 - 100
1/t1
```

setMethod

Create and Save a Method

## Description

Create and save a formal method for a given function and list of classes.

## Usage

```
setMethod(f, signature=character(), definition, where=1, valueClass)
removeMethod(f, signature, where)
```


## Arguments

```
f
signature
```

definition A function definition, which will become the method called when the ar- guments in a call to $f$ match the classes in signature, directly or through inheritance.
where The database in which to store the definition of the method; by default, the current global environment.
For removeMethod, the default is the location of the (first) instance of the method for this signature.
valueClass If supplied, this argument asserts that the method will return a value of this class. (At present this argument is stored but not explicitly used.)

## Details

R methods for a particular generic function are stored in an object of class MethodsList, which in turn is stored with the definition of the generic function. The effect of calling setMethod is to store definition in a MethodsList object in a definition of the generic function on database where. If no such function exists (on that database) one will be created, by copying the generic function from where it is found in the current search list. Finally, if $f$ doesn't exist as a generic function, but there is an ordinary function of the same name and the same formal arguments, a new generic function is created, and the previous non-generic version of $f$ becomes the default method.

Methods are stored in a hierarchical structure, by formal arguments to f: see MethodsList for details. The class names in the signature can be any formal class, plus predefined basic classes such as "numeric", "character", and "matrix". Two additional special class names can appear: "ANY", meaning that this argument can have any class at all; and "missing", meaning that this argument must not appear in the call in order to match this signature. Don't confuse these two: if an argument isn't mentioned in a signature, it corresponds implicitly to class "ANY", not to "missing". See the example below.

While $f$ can correspond to methods defined on several packages or environments, the underlying model is that these together make up the definition for a single generic function. When R proceeds to select and evaluate methods for $f$, the methods on the current search list are merged to form a single generic. In particular, all the versions of $f$ and all the methods must correspond to the same formal arguments (including, in the present definition, the same default expressions for the arguments). For compatibility with S-Plus, the current implementation enforces this partly with a warning and a reconstruction of a method that fails to match, but don't count on this for the future: Make the formal arguments of definition match those of the generic..

## Value

These functions exist for their side-effect, in setting or removing a method in the object defining methods for the specified generic.
The value returned by removeMethod is TRUE if a method was found to be removed.

## References

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections $1.6,2.7,2.8$, and chapters 7 and 8 .

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

## See Also

Methods, MethodsList for details of the implementation

## Examples

```
## methods for plotting track objects (see the example for setClass)
##
## First, with only one object as argument:
setMethod("plot", signature(x="track", y="missing"),
    function(x, y, ...) plot(slot(x, "x"), slot(x, "y"), ...)
)
## Second, plot the data from the track on the y-axis against anything
## as the x data.
setMethod("plot", signature(y = "track"),
    function(x, y, ...) plot(x, slot(y, "y"), ...)
)
```

```
## and similarly with the track on the x-axis (using the short form of
## specification for signatures)
setMethod("plot", "track",
    function(x, y, ...) plot(slot(x, "y"), y, ...)
)
t1 <- new("track", x=1:20, y=(1:20)^2)
tc1 <- new("trackCurve", t1)
slot(tc1, "smooth") <- smooth.spline(slot(tc1, "x"), slot(tc1, "y"))$y #$
plot(t1)
plot(qnorm(ppoints(20)), t1)
## An example of inherited methods, and of conforming method arguments
## (note the dotCurve argument in the method, which will be pulled out
## of ... in the generic.
setMethod("plot", c("trackCurve", "missing"),
function(x, y, dotCurve = FALSE, ...) {
    plot(as(x, "track"))
    if(length(slot(x, "smooth") > 0))
        lines(slot(x, "x"), slot(x, "smooth"),
            lty = if(dotCurve) 2 else 1)
    }
)
## the plot of tc1 alone has an added curve; other uses of tc1
## are treated as if it were a "track" object.
plot(tc1, dotCurve = TRUE)
plot(qnorm(ppoints(20)), tc1)
## defining methods for a special function.
## Although "[" and "length" are not ordinary functions
## methods can be defined for them.
setMethod("[", "track",
    function(x, i, j, ..., drop) {
        x@x <- x@x[i]; x@y <- x@y[i]
        x
        })
plot(t1[1:15])
setMethod("length", "track", function(x)length(x@y))
length(t1)
## methods can be defined for missing arguments as well
setGeneric("summary") ## make the function into a generic
## A method for summary()
## The method definition can include the arguments, but
## if they're omitted, class "missing" is assumed.
setMethod("summary", "missing", function() "<No Object>")
```


## Description

The Classes argument is an old-style class assignment. In particular, if there is more than one name, we expect old-style class inheritance. Calling setOldClass establishes similar inheritance for formal method dispatch, so that the class names can be used in a call to setMethod.

## Usage

```
setOldClass(Classes, where = 1)
```


## Arguments

Classes A character vector of names for old-style classes.
where Where to store the class definitions.

## Details

Each of the names will be defined as a virtual class, extending the remaining classes in Classes, and the class oldClass, which is the "root" of all old-style classes.
See the list .OldClassesList for the old-style classes that are defined by the methods package. Each element of the list is an old-style list, with multiple character strings if inheritance is included. Each element of the list was passed to setOldClass when creating the methods package.

## See Also

```
setClass, setMethod
```


## Examples

```
setOldClass(c("mlm", "lm"))
setGeneric("dfResidual", function(model)standardGeneric("dfResidual"))
setMethod("dfResidual", "lm", function(model)model$df.residual)
## dfResidual will work on mlm objects as well as lm objects
myData <- data.frame(time = 1:10, y = (1:10)^.5)
myLm <- lm(cbind(y, y^3) ~ time, myData)
```

```
rm(myData, myLm)
removeGeneric("dfResidual")
removeClass("mlm")
removeClass("lm")
```

```
show

\section*{Description}

Display the object, by printing, plotting or whatever suits its class.
The function show exists to be specialized by methods; the default method calls showDefault.

With library methods attached, methods for show will usually be invoked for automatic printing (see the details).
The function showDefault allows redirection of output and optional use of old-style print methods, but normally will not be called directly.

\section*{Usage}
```

show(object)
showDefault(object, oldMethods = TRUE)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & Any R object \\
oldMethods & \begin{tabular}{l} 
Should old-style print methods be used for this object? TRUE by default \\
if called directly, but FALSE when called from the methods package for \\
automatic printing (to avoid potential recursion; see the details below).
\end{tabular}
\end{tabular}

\section*{Details}

The methods package overrides the base definition of print.default to arrange for automatic printing to honor methods for the function show. This does not quite manage to override old-style printing methods, since the automatic printing in the evaluator will look first for the old-style method.

If you have a class myClass and want to define a method for show, all will be well unless there is already a function named print.myClass. In that case, to get your method dispatched for automatic printing, it will have to be a method for print. A slight cheat is to override the function print.myClass yourself, and then call that function also in the method for show with signature "myClass".

\section*{Value}
show returns an invisible NULL.
For showDefault, if printTo is FALSE, the value is a character vector containing the lines that would otherwise have been printed.

\section*{See Also}
showMethods prints all the methods for one or more functions; showMlist prints individual methods lists; showClass prints class definitions. Neither of the latter two normally needs to be called directly.

\section*{Examples}
```


## following the example shown in the setMethod documentation ...

setClass("track",
representation(x="numeric", y="numeric"))
setClass("trackCurve",

```
```

    representation("track", smooth = "numeric"))
    t1 <- new("track", x=1:20, y=(1:20)^2)
tc1 <- new("trackCurve", t1)
setMethod("show", "track",
function(object)print(rbind(x = object@x, y=object@y))
)

## The method will now be used for automatic printing of t1

```
t1
\begin{tabular}{lrrrrrrrrrrrr} 
& {\([, 1]\)} & {\([, 2]\)} & {\([, 3]\)} & {\([, 4]\)} & {\([, 5]\)} & {\([, 6]\)} & {\([, 7]\)} & {\([, 8]\)} & {\([, 9]\)} & {\([, 10]\)} & {\([, 11]\)} & {\([, 12]\)} \\
x & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 & 10 & 11 & 12 \\
y & 1 & 4 & 9 & 16 & 25 & 36 & 49 & 64 & 81 & 100 & 121 & 144 \\
& {\([, 13]\)} & {\([, 14]\)} & {\([, 15]\)} & {\([, 16]\)} & {\([, 17]\)} & {\([, 18]\)} & {\([, 19]\)} & {\([, 20]\)} & & \\
x & 13 & 14 & 15 & 16 & 17 & 18 & 19 & 20 & & \\
y & 169 & 196 & 225 & 256 & 289 & 324 & 361 & 400 & &
\end{tabular}
\#\# and also for tc1, an object of a class that extends "track"
tc1
\begin{tabular}{lrrrrrrrrrrrr} 
& {\([, 1]\)} & {\([, 2]\)} & {\([, 3]\)} & {\([, 4]\)} & {\([, 5]\)} & {\([, 6]\)} & {\([, 7]\)} & {\([, 8]\)} & {\([, 9]\)} & {\([, 10]\)} & {\([, 11]\)} & {\([, 12]\)} \\
x & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 & 10 & 11 & 12 \\
y & 1 & 4 & 9 & 16 & 25 & 36 & 49 & 64 & 81 & 100 & 121 & 144 \\
& {\([, 13]\)} & {\([, 14]\)} & {\([, 15]\)} & {\([, 16]\)} & {\([, 17]\)} & {\([, 18]\)} & {\([, 19]\)} & {\([, 20]\)} & & \\
x & 13 & 14 & 15 & 16 & 17 & 18 & 19 & 20 & & \\
y & 169 & 196 & 225 & 256 & 289 & 324 & 361 & 400 & &
\end{tabular}
```

showMethods Show all the methods for the specified function(s)

```

\section*{Description}

Show a summary of the methods for one or more generic functions, possibly restricted to those involving specified classes.

\section*{Usage}
```

showMethods(f=character(), where=-1, classes=NULL, includeDefs=FALSE,
inherited=TRUE, printTo = stdout())

```

\section*{Arguments}
\begin{tabular}{ll}
f & One or more function names. If omitted, all functions will be examined. \\
where & If where is supplied, the methods definition from that position will be \\
used; otherwise, the current definition is used (which will include inherited \\
methods that have arisen so far in the session). If f is omitted, where \\
controls where to look for generic functions. \\
classes & \begin{tabular}{l} 
If argument classes is supplied, it is a vector of class names that restricts \\
the displayed results to those methods whose signatures include one or \\
more of those classes.
\end{tabular}
\end{tabular}
includeDefs If includeDefs is TRUE, include the definitions of the individual methods in the printout.
inherited If inherits is TRUE, then methods that have been found by inheritance, so far in the session, will be included and marked as inherited.
printTo The connection on which the printed information will be written. If printTo is FALSE, the output will be collected as a character vector and returned as the value of the call to showMethod. See show.

\section*{Details}

The output style is different from S-Plus in that it does not show the database from which the definition comes, but can optionally include the method definitions.

\section*{Value}

If printTo is FALSE, the character vector that would have been printed is returned; otherwise the value is the connection or filename.

\section*{Author(s)}

John Chambers

\section*{References}

The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

The functions in this package emulate the facility for classes and methods described in Programming with Data (John M. Chambers, Springer, 1998). See this book for further details and examples.

\section*{See Also}
setMethod, and GenericFunctions for other tools involving methods; show

\section*{Examples}
```


## assuming the methods for plot

## are set up as in the documentation for setMethod,

## print (without definitions) the methods that involve

## class "track"

showMethods("plot", classes = "track")
Function "plot":
x = ANY, y = track
x = track, y = missing
x = track, y = ANY

```
signature-class Class "signature" For Method Definitions

\section*{Description}

This class represents the mapping of some of the formal arguments of a function onto the names of some classes. It is used as one of two slots in the MethodDefinition-class.

\section*{Objects from the Class}

Objects can be created by calls of the form new("signature", functionDef, ...). The functionDef argument, if it is supplied as a function object, defines the formal names. The other arguments define the classes.

\section*{Slots}
.Data: Object of class "character" the classes.
names: Object of class "character" the corresponding argument names.

\section*{Extends}

Class "character", from data part. Class "vector", by class "character".

\section*{Methods}
initialize signature(object = "signature"): see the discussion of objects from the class, above.

\section*{See Also}

MethodDefinition-class for the use of this class
```

slot The Slots in an Object from a Formal Class

```

\section*{Description}

These functions return or set information about the individual slots in an object.

\section*{Usage}
```

object@name
object@name <- value
slot(object, name)
slot(object, name) <- value
slotNames(x)

```

\section*{Arguments}
\[
\begin{array}{ll}
\text { object } & \text { An object from a formally defined class. } \\
\text { name } & \begin{array}{l}
\text { The character-string name of the slot. The name must be a valid slot } \\
\text { name: see Details below. } \\
\text { value }
\end{array} \\
\mathrm{x} & \begin{array}{l}
\text { Either the name of a class or an object from that class. Print } \\
\text { getClass (class) to see the full description of the slots. }
\end{array}
\end{array}
\]

\section*{Details}

The "@" operator and the slot function extract or replace the formally defined slots for the object. The operator takes a fixed name, which can be unquoted if it is syntactically a name in the language. A slot name can be any non-empty string, but if the name is not made up of letters, numbers, and ".", it needs to be quoted.

In the case of the slot function, the slot name can be any expression that evaluates to a valid slot in the class definition. Generally, the only reason to use the functional form rather than the simpler operator is because the slot name has to be computed.

The definition of the class contains the names of all slots diretly and indirectly defined. Each slot has a name and an associated class. Extracting a slot returns an object from that class. Setting a slot first coerces the value to the specified slot and then stores it.

Unlike attributes, slots are not partially matched, and asking for (or trying to set) a slot with an invalid name for that class generates an error.

\section*{Author(s)}

John Chambers

\section*{References}

Chambers, J. M. (1998) Programming with Data, Springer.
The web page http://www.omegahat.org/RSMethods/index.html is the primary documentation.

\section*{See Also}
```

@, Classes, Methods, getClass

```

\section*{Examples}
```

slot(myTrack, "x")
slot(myTrack, "y") <- log(slot(myTrack, "x"))
slotNames("track")

```

StructureClasses Classes Corresponding to Basic Structures

\section*{Description}

The virtual class structure and classes that extend it are formal classes analogous to S language structures such as arrays and time-series
```

Usage

## The folowing class names can appear in method signatures,

## as the class in as() and is() expressions, and, except for

## the classes commented as VIRTUAL, in calls to new()

"matrix"
"array"
"ts"
"structure" \#\# VIRTUAL

```

\section*{Objects from the Classes}

Objects can be created by calls of the form new (Class, ...), where Class is the quoted name of the specific class (e.g., "matrix"), and the other arguments, if any, are interpreted as arguments to the corresponding function, e.g., to function matrix(). There is no particular advantage over calling those functions directly, unless you are writing software designed to work for multiple classes, perhaps with the class name and the arguments passed in.

\section*{Extends}

The specific classes all extend class "structure", directly, and class "vector", by class "structure".

\section*{Methods}
coerce Methods are defined to coerce arbitrary objects to these classes, by calling the corresponding basic function, for example, as(x, "matrix") calls as.matrix(x).
```

substituteDirect SubstituteDirect

```

\section*{Description}

Substitute for the variables named in the second argument the corresponding objects, substituting into object. The argument frame is a named list; if omitted, the environment of the caller is used.

This function differs from the ordinary substitute in that it treats its first argument in the standard \(S\) way, by evaluating it. In contrast, substitute does not evaluate its first argument.
The goal is to replace this with an eval= argument to substitute.

\section*{Usage}
```

    substituteDirect(object, frame, cleanFunction=TRUE)
    ```

TraceClasses Classes Used Internally to Control Tracing

\section*{Description}

The classes described here are used by the R function trace to create versions of functions and methods including browser calls, etc., and also to untrace the same objects.
```

Usage
\#\#\# Objects from the following classes are generated

### by calling trace() on an object from the corresponding

### class without the "WithTrace" in the name.

"functionWithTrace"
"MethodDefinitionWithTrace"
"MethodWithNextWithTrace"
"genericFunctionWithTrace"
"groupGenericFunctionWithTrace"

### the following is a virtual class extended by each of the

### classes above

"traceable"

```

\section*{Objects from the Class}

Objects will be created from these classes by calls to trace. (There is an initialize method for class "traceable", but you are unlikely to need it directly.)

\section*{Slots}
.Data: The data part, which will be "function" for class "functionWithTrace", and similarly for the other classes.
original: Object of the original class; e.g., "function" for class "functionWithTrace".

\section*{Extends}

Each of the classes extends the corresponding untraced class, from the data part; e.g., "functionWithTrace" extends "function". Each of the specific classes extends "traceable", directly, and class "VIRTUAL", by class "traceable".

\section*{Methods}

The point of the specific classes is that objects generated from them, by function trace(), remain callable or dispatchable, in addition to their new trace information.

\section*{See Also}
function trace
```

validObject Test the Validity of an Object

```

\section*{Description}

The validity of object related to its class definition is tested. If the object is valid, TRUE is returned; otherwise, either a vector of strings describing validity failures is returned, or an error is generated (according to whether test is TRUE).
The functions getValidity and setValidity get and set the validity method of a class. This method is a function of one object that returns TRUE or a description of the non-validity.

\section*{Usage}
```

validObject(object, test)
getValidity(ClassDef)
setValidity(Class, method, where = 1 )

```

\section*{Arguments}
object Any object, but not much will happen unless the object's class has a formal definition.
test If test is TRUE, and validity fails the function returns a vector of strings describing the problems. If test is FALSE (the default) validity failure generates an error.
Class The name or class definition of the class whose validity method is to be set.

ClassDef The class definition of the class whose validity method is to be retrieved.
method A validity method; that is, either NULL or a function of one argument (the object). Like validObject, the function should return TRUE if the object is valid, and one or more descriptive strings if any problems are found. Unlike validObject, it should never generate an error.
where The modified class definition will be stored in this position on the search list.

Note that validity methods do not have to check validity of any slots or superclasses: the logic of validObject ensures these tests are done once only. As a consequence, if one validity method wants to use another, it should extract and call the method from the other definition of the other class by calling getValidity: it should not call validObject.

\section*{Details}

Validity testing takes place "bottom up": first the validity of the object's slots, if any, is tested. Then for each of the classes that this class extends (the "superclasses"), the explicit validity method of that class is called, if one exists. Finally, the validity method of object's class is called, if there is one.
Testing generally stops at the first stage of finding an error, except that all the slots will be examined even if a slot has failed its validity test.

\section*{Value}
validObject returns TRUE if the object is valid. Otherwise a vector of strings describing problems found, except that if test is FALSE, validity failure generates an error, with the corresponding strings in the error message.

\section*{References}

The R package methods implements, with a few exceptions, the programming interface for classes and methods in the book Programming with Data (John M. Chambers, Springer, 1998), in particular sections \(1.6,2.7,2.8\), and chapters 7 and 8 .

While the programming interface for the methods package follows the reference, the R software is an original implementation, so details in the reference that reflect the S 4 implementation may appear differently in R. Also, there are extensions to the programming interface developed more recently than the reference. For a discussion of details and ongoing development, see the web page http://developer.r-project.org/methodsPackage.html and the pointers from that page.

\section*{See Also}

\section*{Examples}
```

setClass("track",
representation(x="numeric", y = "numeric"))
t1 <- new("track", x=1:10, y=sort(rnorm(10)))

## A valid "track" object has the same number of x, y values

validTrackObject <- function(x){
if(length(x@x) == length(x@y)) TRUE
else paste("Unequal x,y lengths: ", length(x@x), ", ", length(x@y),
sep="")
}

## assign the function as the validity method for the class

setValidity("track", validTrackObject)

## t1 should be a valid "track" object

validObject(t1)

## Now we do something bad

t1@x <- 1:20

## This should generate an error

try(validObject(t1))

```

\section*{Chapter 6}

\section*{The modreg package}

\section*{ksmooth \\ Kernel Regression Smoother}

\section*{Description}

The Nadaraya-Watson kernel regression estimate.

\section*{Usage}
ksmooth(x, y, kernel = c("box", "normal"), bandwidth = 0.5, range. \(x=\) range \((x)\), n.points \(=\max (100\), length \((x))\), \(x . p o i n t s)\)

\section*{Arguments}
\(\mathrm{x} \quad\) input x values
\(\mathrm{y} \quad\) input y values
kernel the kernel to be used.
bandwidth the bandwidth. The kernels are scaled so that their quartiles (viewed as probability densities) are at \(\pm 0.25 *\) bandwidth.
range.x the range of points to be covered in the output.
n.points the number of points at which to evaluate the fit.
x .points points at which to evaluate the smoothed fit. If missing, n.points are chosen uniformly to cover range. x .

\section*{Value}

A list with components
x values at which the smoothed fit is evaluated. Guaranteed to be in increasing order.
\(\mathrm{y} \quad\) fitted values corresponding to x .

\section*{Note}

This function is implemented purely for compatibility with \(S\), although it is nowhere near as slow as the S function. Better kernel smoothers are available in other packages.

\section*{Author(s)}
B. D. Ripley

\section*{Examples}
```

data(cars)
attach(cars)
plot(speed, dist)
lines(ksmooth(speed, dist, "normal", bandwidth=2), col=2)
lines(ksmooth(speed, dist, "normal", bandwidth=5), col=3)
lines(ksmooth(speed, dist, "normal", bandwidth=10), col=4)
detach()

```

\section*{loess Local Polynomial Regression Fitting}

\section*{Description}

Fit a polynomial surface determined by one or more numerical predictors, using local fitting.

\section*{Usage}
```

loess(formula, data, weights, subset, na.action, model = FALSE,
span = 0.75, enp.target, degree = 2,
parametric = FALSE, drop.square = FALSE, normalize = TRUE,
family = c("gaussian", "symmetric"),
method = c("loess", "model.frame"),
control = loess.control(...), ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
formula & \begin{tabular}{l} 
a formula specifying the response and one to four numeric predictors (best \\
specified via an interaction, but can also be specified additively). \\
an optional data frame within which to look first for the response, pre- \\
dictors and weights.
\end{tabular} \\
data & \begin{tabular}{l} 
optional weights for each case.
\end{tabular} \\
weights & an optional specification of a subset of the data to be used. \\
subset & the action to be taken with missing values in the response or predictors. \\
na.action & \begin{tabular}{l} 
The default is to stop. \\
should the model frame be returned?
\end{tabular} \\
model & \begin{tabular}{l} 
the parameter \(\alpha\) which controls the degree of smoothing.
\end{tabular} \\
span & an alternative way to specify span, as the approximate equivalent number \\
enp.target & \begin{tabular}{l} 
of parameters to be used.
\end{tabular} \\
degree & \begin{tabular}{l} 
the degree of the polynomials to be used, up to 2.
\end{tabular} \\
parametric & \begin{tabular}{l} 
should any terms be fitted globally rather than locally? Terms can be \\
specified by name, number or as a logical vector of the same length as the
\end{tabular} \\
number of predictors.
\end{tabular}
\begin{tabular}{ll} 
drop.square & \begin{tabular}{l} 
for fits with more than one predictor and degree=2, should the quadratic \\
term (and cross-terms) be dropped for particular predictors? Terms are \\
specified in the same way as for parametric.
\end{tabular} \\
normalize & \begin{tabular}{l} 
should the predictors be normalized to a common scale if there is more \\
than one? The normalization used is to set the \(10 \%\) trimmed standard \\
deviation to one. Set to false for spatial coordinate predictors and others \\
know to be a common scale.
\end{tabular} \\
family & \begin{tabular}{l} 
if "gaussian" fitting is by least-squares, and if "symmetric" a re- \\
descending M estimator is used with Tukey's biweight function.
\end{tabular} \\
method & \begin{tabular}{l} 
fit the model or just extract the model frame.
\end{tabular} \\
control & \begin{tabular}{l} 
control parameters: see loess.control.
\end{tabular} \\
\(\ldots\) & control parameters can also be supplied directly.
\end{tabular}

\section*{Details}

Fitting is done locally. That is, for the fit at point \(x\), the fit is made using points in a neighbourhood of \(x\), weighted by their distance from \(x\) (with differences in 'parametric' variables being ignored when computing the distance). The size of the neighbourhood is controlled by \(\alpha\) (set by span or enp.target). For \(\alpha<1\), the neighbourhood includes proportion \(\alpha\) of the points, and these have tricubic weighting (proportional to \(\left(1-(\text { dist } / \text { maxdist })^{3}\right)^{3}\). For \(\alpha>1\), all points are used, with the 'maximum distance' assumed to be \(\alpha^{1 / p}\) times the actual maximum distance for \(p\) explanatory variables.

For the default family, fitting is by (weighted) least squares. For family="symmetric" a few iterations of an M-estimation procedure with Tukey's biweight are used. Be aware that as the initial value is the least-squares fit, this need not be a very resistant fit.

It can be important to tune the control list to achieve acceptable speed. See loess.control for details.

\section*{Value}

An object of class "loess".

\section*{Note}

As this is based on the cloess package available at netlib, it is similar to but not identical to the loess function of S . In particular, conditioning is not implemented.
The memory usage of this implementation of loess is roughly quadratic in the number of points, with 1000 points taking about 10 Mb .

\section*{Author(s)}
B.D. Ripley, based on the cloess package of Cleveland, Grosse and Shyu.

\section*{References}
W.S. Cleveland, E. Grosse and W.M. Shyu (1992) Local regression models. Chapter 8 of Statistical Models in \(S\) eds J.M. Chambers and T.J. Hastie, Wadsworth \& Brooks/Cole.

\section*{See Also}
loess.control, predict.loess.
lowess, the ancestor of loess (with different defaults!).

\section*{Examples}
```

data(cars)
cars.lo <- loess(dist ~ speed, cars)
predict(cars.lo, data.frame(speed = seq(5, 30, 1)), se = TRUE)

# to allow extrapolation

cars.lo2 <- loess(dist ~ speed, cars,
control = loess.control(surface = "direct"))
predict(cars.lo2, data.frame(speed = seq(5, 30, 1)), se = TRUE)

```
```

loess.control Set Parameters for Loess

```

\section*{Description}

Set control parameters for loess fits.

\section*{Usage}
```

loess.control(surface = c("interpolate", "direct"),
statistics = c("approximate", "exact"),
trace.hat = c("exact", "approximate"),
cell = 0.2, iterations = 4, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
surface & \begin{tabular}{l} 
should be fitted surface be computed exactly or via interpolation from a \\
kd tree?
\end{tabular} \\
statistics & \begin{tabular}{l} 
should the statistics be computed exactly or approximately? Exact com- \\
putation can be very slow.
\end{tabular} \\
trace.hat & \begin{tabular}{l} 
should the trace of the smoother matrix be computed exactly or approxi- \\
mately? It is recommended to use the approximation for more than about \\
1000 data points. \\
if interpolation is used this controls the accuracy of the approximation \\
via the maximum number of points in a cell in the kd tree. Cells with \\
more than floor (n*span*cell) points are subdivided.
\end{tabular} \\
cell & \begin{tabular}{l} 
the number of iterations used in robust fitting.
\end{tabular} \\
iterations \\
... & \begin{tabular}{l} 
further arguments which are ignored.
\end{tabular}
\end{tabular}

\section*{Value}

A list with components
surface
statistics
trace.hat
cell
iterations
with meanings as explained under 'Arguments'.

Author(s)
B.D. Ripley

See Also
loess
modreg-internal Internal modreg functions

\section*{Description}

Internal modreg functions.

\section*{Usage}
```

predLoess(y, x, newx, s, weights, robust, span, degree, normalize,
parametric, drop.square, surface, cell, family, kd, divisor,
se = FALSE)
simpleLoess(y, x, weights, span = 0.75, degree = 2, parametric = FALSE,
drop.square = FALSE, normalize = TRUE, statistics = "approximate",
surface = "interpolate", cell = 0.2, iterations = 1,
trace.hat = "exact")
pointwise(results, coverage)

```

\section*{Details}

These are not to be called by the user.
plot.ppr
Plot Ridge Functions for Projection Pursuit Regression Fit

\section*{Description}

Plot ridge functions for projection pursuit regression fit.

\section*{Usage}
```

plot(x, ask, type = "o", ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & A fit of class "ppr" as produced by a call to ppr. \\
ask & the graphics parameter ask: see par for details. If set to TRUE will ask \\
& between the plot of each cross-section. \\
type & the type of line to draw \\
\(\ldots\) & further graphical parameters
\end{tabular}

\section*{Value}

None

\section*{Side Effects}

A series of plots are drawn on the current graphical device, one for each term in the fit.

\section*{See Also}
```

ppr, par

```

\section*{Examples}
```

data(rock)
attach(rock)
area1 <- area/10000; peri1 <- peri/10000
par(mfrow=c(3,2))\# maybe: , pty="s")
rock.ppr <- ppr(log(perm) ~ area1 + peri1 + shape,
data = rock, nterms = 2, max.terms = 5)
plot(rock.ppr, main="ppr(log(perm) ~ ., nterms=2, max.terms=5)")
plot(update(rock.ppr, bass=5), main = "update(..., bass = 5)")
plot(update(rock.ppr, sm.method="gcv", gcvpen=2),
main = "update(..., sm.method=\"gcv\", gcvpen=2)")

```
ppr Projection Pursuit Regression

\section*{Description}

Fit a projection pursuit regression model.

\section*{Usage}
```

ppr(formula, data = sys.parent(), weights,
subset, na.action, contrasts = NULL,
ww = rep(1,q), nterms, max.terms = nterms, optlevel = 2,
sm.method = c("supsmu", "spline", "gcvspline"),
bass = 0, span = 0, df = 5, gcvpen = 1)
ppr(x, y, weights = rep(1,n),
ww = rep(1,q), nterms, max.terms = nterms, optlevel = 2,
sm.method = c("supsmu", "spline", "gcvspline"),
bass = 0, span = 0, df = 5, gcvpen = 1)

```

\section*{Arguments}
formula a formula specifying one or more response variables and the explanatory variables.
x
matrix of explanatory variables. Rows represent observations, and columns represent variables. Missing values are not accepted.
y matrix of response variables. Rows represent observations, and columns represent variables. Missing values are not accepted.
\begin{tabular}{|c|c|}
\hline nterms & number of terms to include in the final model. \\
\hline data & data frame from which variables specified in formula are preferentially to be taken. \\
\hline weights & a vector of weights w_i for each case. \\
\hline ww & a vector of weights for each response, so the fit criterion is the sum ove case \(i\) and responses \(j\) of \(w_{-} i w_{-} j\left(y \_i j-f i t \_i j\right) \wedge 2\) divided by the sum of w_i. \\
\hline subset & an index vector specifying the cases to be used in the training sample. (NOTE: If given, this argument must be named.) \\
\hline na.action & a function to specify the action to be taken if NAs are found. The default action is for the procedure to fail. An alternative is na.omit, which leads to rejection of cases with missing values on any required variable. (NOTE If given, this argument must be named.) \\
\hline contrasts & the contrasts to be used when any factor explanatory variables are coded. \\
\hline max.terms & maximum number of terms to choose from when building the model \\
\hline optlevel & integer from 0 to 3 which determines the thoroughness of an optimization routine in the SMART program. See the Details section. \\
\hline sm.method & the method used for smoothing the ridge functions. The default is to use Friedman's super smoother supsmu. The alternatives are to use the smoothing spline code underlying smooth.spline, either with a specified (equivalent) degrees of freedom for each ridge functions, or to allow the smoothness to be chosen by GCV. \\
\hline bass & super smoother bass tone control used with automatic span selection (see supsmu); the range of values is 0 to 10 , with larger values resulting in increased smoothing. \\
\hline span & super smoother span control (see supsmu). The default, 0 , results in automatic span selection by local cross validation. span can also take a value in ( 0,1 ]. \\
\hline df & if sm.method is "spline" specifies the smoothness of each ridge term via the requested equivalent degrees of freedom. \\
\hline gcvpen & if sm.method is "gcvspline" this is the penalty used in the GCV selection for each degree of freedom used. \\
\hline
\end{tabular}

\section*{Details}

The basic method is given by Friedman (1984), and is essentially the same code used by S-PLUS's ppreg. This code is extremely sensitive to the compiler used.

The algorithm first adds up to max.terms ridge terms one at a time; it will use less if it is unable to find a term to add that makes sufficient difference. It then removes the least "important" term at each step until nterm terms are left.

The levels of optimization (argument optlevel) differ in how thoroughly the models are refitted during this process. At level 0 the existing ridge terms are not refitted. At level 1 the projection directions are not refitted, but the ridge functions and the regression coefficients are. Levels 2 and 3 refit all the terms and are equivalent for one response; level 3 is more careful to re-balance the contributions from each regressor at each step and so is a little less likely to converge to a saddle point of the sum of squares criterion.

\section*{Value}

A list with the following components, many of which are for use by the method functions.
call the matched call
\(\mathrm{p} \quad\) the number of explanatory variables (after any coding)
q the number of response variables
mu the argument nterms
ml the argument max.terms
gof the overall residual (weighted) sum of squares for the selected model
gofn the overall residual (weighted) sum of squares against the number of terms, up to max.terms. Will be invalid (and zero) for less than nterms.
\(\mathrm{df} \quad\) the argument df
edf if sm.method is "spline" or "gcvspline" the equivalent number of degrees of freedom for each ridge term used.
xnames the names of the explanatory variables
ynames the names of the response variables
alpha a matrix of the projection directions, with a column for each ridge term
beta a matrix of the coefficients applied for each response to the ridge terms: the rows are the responses and the columns the ridge terms
\(\mathrm{yb} \quad\) the weighted means of each response
ys the overall scale factor used: internally the responses are divided by ys to have unit total weighted sum of squares.
fitted.values the fitted values, as a matrix if \(q>1\).
residuals the residuals, as a matrix if \(q>1\).
smod internal work array, which includes the ridge functions evaluated at the training set points.

\section*{References}

Friedman, J. H. and Stuetzle, W. (1981) Projection pursuit regression. Journal of the American Statistical Association, 76, 817-823.
Friedman, J. H. (1984) SMART User's Guide. Laboratory for Computational Statistics, Stanford University Technical Report No. 1.
Venables, W. N. \& Ripley, B. D. (1999) Modern Applied Statistics with S-PLUS. Springer.

\section*{See Also}
```

plot.ppr, supsmu, smooth.spline

```

\section*{Examples}
```


# Note: your numerical values may differ

data(rock)
attach(rock)
area1 <- area/10000; peri1 <- peri/10000
rock.ppr <- ppr(log(perm) ~ area1 + peri1 + shape,
data = rock, nterms = 2, max.terms = 5)
rock.ppr

```
```


# Call:

# ppr.formula(formula = log(perm) ~ area1 + peri1 + shape, data = rock,

# nterms = 2, max.terms = 5)

# 

# Goodness of fit:

# 2 terms 3 terms 4 terms 5 terms

# 8.737806 5.289517 4.7457994.490378

summary(rock.ppr)

# ..... (same as above)

# .....

# 

# Projection direction vectors:

# term 1 term 2

# area1 0.34357179 0.37071027

# peri1 -0.93781471 -0.61923542

# shape 0.04961846 0.69218595

# 

# Coefficients of ridge terms:

# term 1 term 2

# 1.6079271 0.5460971

par(mfrow=c(3,2))\# maybe: , pty="s")
plot(rock.ppr, main="ppr(log(perm)~ ., nterms=2, max.terms=5)")
plot(update(rock.ppr, bass=5), main = "update(..., bass = 5)")
plot(update(rock.ppr, sm.method="gcv", gcvpen=2),
main = "update(..., sm.method=\"gcv\", gcvpen=2)")
detach()

```
```

predict.loess Predict Loess Curve or Surface

```

\section*{Description}

Predictions from a loess fit, optionally with standard errors.

\section*{Usage}
```

predict(object, newdata = NULL, se = FALSE, ...)

```

\section*{Arguments}
object an object fitted by loess.
newdata an optional data frame specifying points at which to do the predictions. If missing, the original data points are used.
se should standard errors be computed?
... arguments passed to or from other methods.

\section*{Details}

The standard errors calculation is slower than prediction.
When the fit was made using surface="interpolate" (the default), predict.loess will not extrapolate - so points outside an axis-aligned hypercube enclosing the original data will have missing (NA) predictions and standard errors.

\section*{Value}

If se = FALSE, a vector giving the prediction for each row of newdata (or the original data).
If se \(=\) TRUE, a list containing components
\begin{tabular}{ll} 
fit & the predicted values. \\
se & an estimated standard error for each predicted value. \\
residual.scale
\end{tabular}\(\quad\)\begin{tabular}{l} 
the estimated scale of the residuals used in computing the standard errors. \\
df
\end{tabular}\(\quad\)\begin{tabular}{l} 
an estimate of the effective degrees of freedom used in estimating the \\
residual scale, intended for use with t-based confidence intervals.
\end{tabular}

\section*{Author(s)}
B.D. Ripley, based on the cloess package of Cleveland, Grosse and Shyu.

\section*{See Also}
loess

\section*{Examples}
```

data(cars)
cars.lo <- loess(dist ~ speed, cars)
predict(cars.lo, data.frame(speed=seq(5, 30, 1)), se=TRUE)

# to get extrapolation

cars.lo2 <- loess(dist ~ speed, cars,
control=loess.control(surface="direct"))
predict(cars.lo2, data.frame(speed=seq(5, 30, 1)), se=TRUE)

```
```

predict.smooth.spline

```

Predict from Smoothing Spline Fit

\section*{Description}

Predict a smoothing spline fit at new points, return the derivative if desired. The predicted fit is linear beyond the original data.

\section*{Usage}
```

predict(object, x, deriv = 0, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & a fit from smooth.spline. \\
x & the new values of x. \\
deriv & integer; the order of the derivative required. \\
\(\ldots\) & further arguments passed to or from other methods.
\end{tabular}

\section*{Value}

A list with components
x
The input x .
y
The fitted values or derivatives at x .

\section*{Author(s)}
B.D. Ripley

\section*{See Also}
```

smooth.spline

```

\section*{Examples}
```

data(cars)
attach(cars)
cars.spl <- smooth.spline(speed, dist, df=6.4)

## "Proof" that the derivatives are okay, by comparing with approximation

diff.quot <- function(x,y) {
\#\# Difference quotient (central differences where available)
n <- length(x); i1 <- 1:2; i2 <- (n-1):n
c(diff(y[i1]) / diff(x[i1]), (y[-i1] - y[-i2]) / (x[-i1] - x[-i2]),
diff(y[i2]) / diff(x[i2]))
}
xx <- unique(sort(c(seq(0,30, by = . 2), kn <- unique(speed))))
i.kn <- match(kn, xx)\# indices of knots within xx
op <- par(mfrow = c(2,2))
plot(speed, dist, xlim = range(xx), main = "Smooth.spline \& derivatives")
lines(pp <- predict(cars.spl, xx), col = "red")
points(kn, pp$y[i.kn], pch = 3, col="dark red")
mtext("s(x)", col = "red")
for(d in 1:3){
    n <- length(pp$x)
plot(pp$x, diff.quot(pp$x,pp$y), type = 'l', xlab="x", ylab="",
                col = "blue", col.main = "red",
            main= paste("s",paste(rep(")",d), collapse=""),"(x)", sep=""))
    mtext("Difference quotient approx.(last)", col = "blue")
    lines(pp <- predict(cars.spl, xx, deriv = d), col = "red")
    points(kn, pp$y[i.kn], pch = 3, col="dark red")
abline(h=0, lty = 3, col = "gray")
}
detach(); par(op)

```

\section*{rock Measurements on Petroleum Rock Samples}

\section*{Description}

Measurements on 48 rock samples from a petroleum reservoir.

\section*{Usage}
data(rock)

\section*{Format}

A data frame with 48 rows and 4 numeric columns.
\begin{tabular}{lll}
{\([, 1]\)} & area & area of pores space, in pixels out of 256 by 256 \\
{\([, 2]\)} & peri & perimeter in pixels \\
{\([, 3]\)} & shape & perimeter/sqrt(area) \\
{\([, 4]\)} & perm & permeability in milli-Darcies
\end{tabular}

\section*{Details}

Twelve core samples from petroleum reservoirs were sampled by 4 cross-sections. Each core sample was measured for permeability, and each cross-section has total area of pores, total perimeter of pores, and shape.

\section*{Source}

Data from BP Research, image analysis by Ronit Katz, U. Oxford.
```

scatter.smooth Scatter Plot with Smooth Curve Fitted by Loess

```

\section*{Description}

Plot and add a smooth curve computed by loess to a scatter plot.

\section*{Usage}
```

scatter.smooth(x, y, span = 2/3, degree = 1,
family = c("symmetric", "gaussian"),
xlab = deparse(substitute(x)), ylab = deparse(substitute(y)),
ylim = range(y, prediction\$y), evaluation = 50, ...)
loess.smooth(x, y, span = 2/3, degree = 1,
family = c("symmetric", "gaussian"), evaluation=50, ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & x coordinates for scatter plot. \\
y & y coordinates for scatter plot. \\
span & smoothness parameter for loess. \\
degree & degree of local polynomial used. \\
family & \begin{tabular}{l} 
if "gaussian" fitting is by least-squares, and if family="symmetric" a \\
\\
re-descending M estimator is used.
\end{tabular} \\
xlab & label for x axis. \\
ylab & label for y axis. \\
ylim & the y limits of the plot. \\
evaluation & number of points at which to evaluate the smooth curve. \\
\(\ldots\) & graphical parameters.
\end{tabular}

\section*{Details}
loess.smooth is an auxiliary function.

\section*{Value}

None.

\section*{Author(s)}
B.D. Ripley

\section*{See Also}
loess

\section*{Examples}
```

data(cars)
attach(cars)
scatter.smooth(speed, dist)
detach()

```
smooth.spline Fit a Smoothing Spline

\section*{Description}

Fits a cubic smoothing spline to the supplied data.

\section*{Usage}
```

smooth.spline(x, y = NULL, w = NULL, df, spar = NULL,
cv = FALSE, all.knots = FALSE, nknots = NULL,
df.offset = 0, penalty = 1, control.spar = list())

```

\section*{Arguments}

X
y
W
df
spar smoothing parameter, typically (but not necessarily) in ( 0,1\(]\). The coefficient \(\lambda\) of the integral of the squared second derivative in the fit (penalized \(\log\) likelihood) criterion is a monotone function of spar, see the details below.
cv ordinary (TRUE) or "generalized" cross-validation (GCV) when FALSE.
all.knots if TRUE, all distinct points in \(x\) are used as knots. If FALSE (default), a subset of x[] is used, specifically \(\mathrm{x}[\mathrm{j}]\) where the nknots indices are evenly spaced in \(1: n\), see also the next argument nknots.
nknots integer giving the number of knots to use when all.knots=FALSE. Per default, this is less than \(n\), the number of unique x values for \(n>49\).
df.offset allows the degrees of freedom to be increased by df.offset in the GCV criterion.
penalty the coefficient of the penalty for degrees of freedom in the GCV criterion.
control.spar optional list with named components controlling the root finding when the smoothing parameter spar is computed, i.e., missing or NULL, see below.
Note that this is partly experimental and may change with general spar computation improvements!
low: lower bound for spar; defaults to -1.5 (used to implicitly default to 0 in R versions earlier than 1.4).
high: upper bound for spar; defaults to +1.5 .
tol: the absolute precision (tolerance) used; defaults to 1e-4 (formerly \(1 \mathrm{e}-3)\).
eps: the relative precision used; defaults to \(2 \mathrm{e}-8\) (formerly 0.00244 ).
trace: logical indicating if iterations should be traced.
maxit: integer giving the maximal number of iterations; defaults to 500 .
Note that spar is only searched for in the interval [low, high].

\section*{Details}

The x vector should contain at least four distinct values. Distinct here means "distinct after rounding to 6 significant digits", i.e., \(x\) will be transformed to unique(sort(signif (x, 6 )) ), and y and w are pooled accordingly.
The computational \(\lambda\) used (as a function of \(s=s p a r\) ) is \(\lambda=r * 256^{3 s-1}\) where \(r=\) \(\operatorname{tr}\left(X^{\prime} W X\right) / \operatorname{tr}(\Sigma), \Sigma\) is the matrix given by \(\Sigma_{i j}=\int B_{i}^{\prime \prime}(t) B_{j}^{\prime \prime}(t) d t, X\) is given by \(X_{i j}=\) \(B_{j}\left(x_{i}\right), W\) is the diagonal matrix of weights (scaled such that its trace is \(n\), the original number of observations) and \(B_{k}(\).\() is the k\)-th B -spline.

Note that with these definitions, \(f_{i}=f\left(x_{i}\right)\), and the B-spline basis representation \(f=X c\) (i.e. \(c\) is the vector of spline coefficients), the penalized \(\log\) likelihood is \(L=(y-f)^{\prime} W(y-\) \(f)+\lambda c^{\prime} \Sigma c\), and hence \(c\) is the solution of the (ridge regression) \(\left(X^{\prime} W X+\lambda \Sigma\right) c=X^{\prime} W y\).

If spar is missing or NULL, the value of df is used to determine the degree of smoothing. If both are missing, leave-one-out cross-validation (ordinary or "generalized" as determined by cv ) is used to determine \(\lambda\). Note that from the above relation, spar is \(s=s 0+0.0601 * \log \lambda\), which is intentionally different from the S-plus implementation of smooth.spline (where spar is proportional to \(\lambda\) ). In R's \((\log \lambda)\) scale, it makes more sense to vary spar linearly.
Note however that currently the results may become very unreliable for spar values smaller than about -1 or -2 . The same may happen for values larger than 2 or so. Don't think of setting spar or the controls low and high outside such a safe range, unless you know what you are doing!

The "generalized" cross-validation method will work correctly when there are duplicated points in x . However, it is ambiguous what leave-one-out cross-validation means with duplicated points, and the internal code uses an approximation that involves leaving out groups of duplicated points. \(\mathrm{cv}=\) TRUE is best avoided in that case.

\section*{Value}

An object of class "smooth.spline" with components
\begin{tabular}{|c|c|}
\hline x & the distinct x values in increasing order, see the Details above. \\
\hline y & the fitted values corresponding to x . \\
\hline w & the weights used at the unique values of x . \\
\hline yin & the y values used at the unique y values. \\
\hline lev & leverages, the diagonal values of the smoother matrix. \\
\hline cv.crit & (generalized) cross-validation score. \\
\hline pen.crit & penalized criterion \\
\hline crit & the criterion value minimized in the underlying .Fortran routine 'sslvrg'. \\
\hline df & equivalent degrees of freedom used. Note that (currently) this value may become quite unprecise when the true df is between and 1 and 2 . \\
\hline spar & the value of spar computed or given. \\
\hline lambda & the value of \(\lambda\) corresponding to spar, see the details above. \\
\hline iparms & named integer(3) vector where . . \$ipars ["iter"] gives number of spar computing iterations used. \\
\hline fit & list for use by predict.smooth.spline, with components \\
\hline & \begin{tabular}{l}
knot: the knot sequence (including the repeated boundary knots). nk: number of coefficients or number of "proper" knots plus 2. \\
coef: coefficients for the spline basis used. \\
min, range: numbers giving the corresponding quantities of x .
\end{tabular} \\
\hline call & the matched call. \\
\hline
\end{tabular}

\section*{Note}

The default all.knots = FALSE and nknots = NULL entails using only \(O\left(n^{0.2}\right)\) knots instead of \(n\) for \(n>49\). This cuts speed and memory requirements, but not drastically anymore since R version 1.5 .1 where it is only \(O\left(n_{k}\right)+O(n)\) where \(n_{k}\) is the number of knots. In this case where not all unique x values are used as knots, the result is not a smoothing spline in the strict sense, but very close unless a small smoothing parameter (or large df) is used.

\section*{Author(s)}
B.D. Ripley and Martin Maechler (spar/lambda, etc).

\section*{References}

Green, P. J. and Silverman, B. W. (1994) Nonparametric Regression and Generalized Linear Models: A Roughness Penalty Approach; Chapman and Hall.

\section*{See Also}
predict.smooth.spline for evaluating the spline and its derivatives.

\section*{Examples}
```

data(cars)
attach(cars)
plot(speed, dist, main = "data(cars) \& smoothing splines")
cars.spl <- smooth.spline(speed, dist)
(cars.spl)

## This example has duplicate points, so avoid cv=TRUE

lines(cars.spl, col = "blue")
lines(smooth.spline(speed, dist, df=10), lty=2, col = "red")
legend(5,120,c(paste("default [C.V.] => df =",round(cars.spl$df,1)),
    "s( * , df = 10)"), col = c("blue","red"), lty = 1:2,
    bg='bisque')
detach()
##-- artificial example
y18 <- c(1:3,5,4,7:3,2*(2:5),rep(10,4))
xx <- seq(1,length(y18), len=201)
(s2 <- smooth.spline(y18)) # GCV
(s02 <- smooth.spline(y18, spar = 0.2))
plot(y18, main=deparse(s2$call), col.main=2)
lines(s2, col = "gray"); lines(predict(s2, xx), col = 2)
lines(predict(s02, xx), col = 3); mtext(deparse(s02\$call), col = 3)

## The following shows the problematic behavior of 'spar' searching:

(s2 <- smooth.spline(y18, con=list(trace=TRUE,tol=1e-6, low= -1.5)))
(s2m <- smooth.spline(y18, cv=TRUE, con=list(trace=TRUE,tol=1e-6, low= -1.5)))

## both above do quite similarly (Df = 8.5 +- 0.2)

```
supsmu Friedman's SuperSmoother

\section*{Description}

Smooth the ( \(\mathrm{x}, \mathrm{y}\) ) values by Friedman's "super smoother".

\section*{Usage}
```

supsmu(x, y, wt = rep(1, length(y)), span = "cv", periodic = FALSE,
bass = 0)

```

\section*{Arguments}
```

x x values for smoothing
y y values for smoothing
wt case weights
span the fraction of the observations in the span of the running lines smoother,
or "cv" to choose this by leave-one-out cross-validation.
periodic if TRUE, the x values are assumed to be in [0, 1] and of period 1.
bass controls the smoothness of the fitted curve. Values of up to 10 indicate
increasing smoothness.

```

\section*{Details}
supsmu is a running lines smoother which chooses between three spans for the lines. The running lines smoothers are symmetric, with \(\mathrm{k} / 2\) data points each side of the predicted point, and values of k as \(0.5 * \mathrm{n}, 0.2 * \mathrm{n}\) and \(0.05 * \mathrm{n}\), where n is the number of data points. If span is specified, a single smoother with span span \(* \mathrm{n}\) is used.
The best of the three smoothers is chosen by cross-validation for each prediction. The best spans are then smoothed by a running lines smoother and the final prediction chosen by linear interpolation.
The FORTRAN code says: "For small samples ( \(\mathrm{n}<40\) ) or if there are substantial serial correlations between observations close in x - value, then a prespecified fixed span smoother ( span \(>0\) ) should be used. Reasonable span values are 0.2 to 0.4."

\section*{Value}

A list with components
\(\mathrm{x} \quad\) the input values in increasing order with duplicates removed.
\(y \quad\) the corresponding \(y\) values on the fitted curve.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Friedman, J. H. (1984) SMART User's Guide. Laboratory for Computational Statistics, Stanford University Technical Report No. 1.
Friedman, J. H. (1984) A variable span scatterplot smoother. Laboratory for Computational Statistics, Stanford University Technical Report No. 5.

\section*{See Also}
ppr

\section*{Examples}
```

data(cars)
attach(cars)
plot(speed, dist)
lines(supsmu(speed, dist))
lines(supsmu(speed, dist, bass=7), lty=2)
detach()

```

\section*{Chapter 7}

\section*{The mva package}

\section*{ability.cov Ability and Intelligence Tests}

\section*{Description}

Six tests were given to 112 individuals. The covariance matrix is given in this object.

\section*{Usage}
```

data(ability.cov)

```

\section*{Details}

The tests are described as
general: a non-verbal measure of general intelligence using Cattell's culture-fair test.
picture: a picture-completion test
blocks: block design
maze: mazes
reading: reading comprehension
vocab: vocabulary
Bartholomew gives both covariance and correlation matrices, but these are inconsistent. Neither are in the original paper.

\section*{Source}

Barthlomew, D. J. (1987) Latent Variable Analysis and Factor Analysis. Griffin.
Barthlomew, D. J. and Knott, M. (1990) Latent Variable Analysis and Factor Analysis. Second Edition, Arnold.

\section*{References}

Smith, G. A. and Stanley G. (1983) Clocking \(g\) : relating intelligence and measures of timed performance. Intelligence, 7, 353-368.

\section*{Examples}
```

data(ability.cov)
(ability.FA <- factanal(factors = 1, covmat=ability.cov))
update(ability.FA, factors=2)
update(ability.FA, factors=2, rotation="promax")

```
```

as.hclust Convert Objects to Class hclust

```

\section*{Description}

Converts objects from other hierarchical clustering functions to class "hclust".

\section*{Usage}
```

as.hclust(x, ...)
as.hclust.default(x, ...)
as.hclust.twins(x, ...)

```

\section*{Arguments}
x Hierarchical clustering object
... further arguments passed to or from other methods.

\section*{Details}

Currently there is only support for converting objects of class "twins" as produced by the functions diana and agnes from the package 'cluster'.

\section*{Value}

An object of class "hclust".

\section*{See Also}
hclust, diana, agnes

\section*{Examples}
```

x <- matrix(rnorm(30), ncol=3)
hc <- hclust(dist(x), method="complete")
if(require(cluster)) {\# is a required package
ag <- agnes(x, method="complete")
hcag <- as.hclust(ag)
\#\# The dendrograms order slightly differently:
op <- par(mfrow=c(1,2))
plot(hc) ; mtext("hclust", side=1)
plot(hcag); mtext("agnes", side=1)
}
stopifnot(identical(hc, hhc <- as.hclust(hc)),
identical(hhc, as.hclust(hhc)))

```
biplot Biplot of Multivariate Data

\section*{Description}

Plot a biplot on the current graphics device.

\section*{Usage}
biplot(x, ...)
biplot.default(x, y, var.axes = TRUE, col, cex = rep(par("cex"), 2), xlabs = NULL, ylabs = NULL, expand = 1, xlim \(=\) NULL, ylim \(=\) NULL, arrow.len \(=0.1, \ldots\) )

\section*{Arguments}
x
\(\mathrm{y} \quad\) The second set of points (a two-column matrix), usually associated with variables.
var.axes If TRUE the second set of points have arrows representing them as (unscaled) axes.
col A vector of length 2 giving the colours for the first and second set of points respectively (and the corresponding axes). If a single colour is specified it will be used for both sets.
cex The character expansion factor used for labelling the points. The labels can be of different sizes for the two sets by supplying a vector of length two.
xlabs A vector of character strings to label the first set of points: the default is to use the row dimname of x , or \(1: \mathrm{n}\) is the dimname is NULL.
ylabs A vector of character strings to label the second set of points: the default is to use the row dimname of y , or \(1: \mathrm{n}\) is the dimname is NULL.
expand An expansion factor to apply when plotting the second set of points relative to the first. This can be used to get the two sets on to a physically comparable scale.
arrow.len The length of the arrow heads on the axes plotted in var.axes is true. The arrow head can be suppressed by arrow.len \(=0\).
xlim, ylim, ... graphical parameters.

\section*{Details}

A biplot is plot which aims to represent both the observations and variables of a matrix of multivariate data on the same plot. There are many variations on biplots (see the references) and perhaps the most widely used one is implemented by biplot.princomp. The function biplot.default merely provides the underlying code to plot two sets of variables on the same figure.
Graphical parameters can also be given to biplot.

\section*{Side Effects}
a plot is produced on the current graphics device.

\section*{Author(s)}
B.D. Ripley

\section*{References}
K. R. Gabriel (1971). The biplot graphical display of matrices with application to principal component analysis. Biometrika 58, 453-467.
J.C. Gower and D. J. Hand (1996). Biplots. Chapman \& Hall.

\section*{See Also}
biplot.princomp, also for examples.
```

biplot.princomp Biplot for Principal Components

```

\section*{Description}

Produces a biplot (in the strict sense) from the output of princomp.
```

Usage
biplot(x, choices $=1: 2$, scale $=1$, pc.biplot = FALSE, ...)

```

\section*{Arguments}
\[
\begin{array}{ll}
\mathrm{x} & \text { an object of class "princomp". } \\
\text { choices } & \begin{array}{l}
\text { length } 2 \text { vector specifying the components to plot. Only the default is a } \\
\text { biplot in the strict sense. }
\end{array} \\
\text { scale } & \begin{array}{l}
\text { The variables are scaled by lambda }- \text { scale and the observations are } \\
\text { scaled by lambda }-(1-\text { scale }) \text { where lambda are the singular values as } \\
\text { computed by princomp. Normally } 0<=\text { scale }<=1 \text {, and a warning will } \\
\text { be issued if the specified scale is outside this range. }
\end{array} \\
\text { pc.biplot } & \begin{array}{l}
\text { If true, use what Gabriel (1971) refers to as a "principal component bi- } \\
\text { plot", with lambda }=1 \text { and observations scaled up by sqrt(n) and vari- } \\
\text { ables scaled down by sqrt(n). Then inner products between variables ap- } \\
\text { proximate covariances and distances between observations approximate }
\end{array} \\
\text { Mahalanobis distance. }
\end{array}
\]

\section*{Details}

This is a method for the generic function biplot. There is considerable confusion over the precise definitions: those of the original paper, Gabriel (1971), are followed here. Gabriel and Odoroff (1990) use the same definitions, but their plots actually correspond to pc.biplot \(=\) TRUE.

\section*{Side Effects}
a plot is produced on the current graphics device.

\section*{References}

Gabriel, K. R. (1971). The biplot graphical display of matrices with applications to principal component analysis. Biometrika, 58, 453-467.
Gabriel, K. R. and Odoroff, C. L. (1990). Biplots in biomedical research. Statistics in Medicine, 9, 469-485.

\section*{See Also}
biplot, princomp.

\section*{Examples}
```

data(USArrests)
biplot(princomp(USArrests))

```
cancor Canonical Correlations

\section*{Description}

Compute the canonical correlations between two data matrices.

\section*{Usage}
cancor ( \(\mathrm{x}, \mathrm{y}, \mathrm{xcenter}=\) TRUE, ycenter \(=\) TRUE)

\section*{Arguments}

\section*{x}
numeric matrix \(\left(n \times p_{1}\right)\), containing the x coordinates.
\(\mathrm{y} \quad\) numeric matrix \(\left(n \times p_{2}\right)\), containing the y coordinates.
xcenter logical or numeric vector of length \(p_{1}\), describing any centering to be done on the x values before the analysis. If TRUE (default), subtract the column means. If FALSE, do not adjust the columns. Otherwise, a vector of values to be subtracted from the columns.
ycenter analogous to xcenter, but for the \(y\) values.

\section*{Details}

The canonical correlation analysis seeks linear combinations of the y variables which are well explained by linear combinations of the x variables. The relationship is symmetric as 'well explained' is measured by correlations.

\section*{Value}

A list containing the following components:
cor correlations.
\(\mathrm{xcoef} \quad\) estimated coefficients for the x variables.
ycoef estimated coefficients for the y variables.
xcenter the values used to adjust the x variables.
ycenter the values used to adjust the x variables.

\section*{References}

Hotelling H. (1936). Relations between two sets of variables. Biometrika, 28, 321-327.
Seber, G. A. F. (1984). Multivariate Observations. New York: Wiley, p. 506f.

\section*{See Also}
qr, svd.

\section*{Examples}
```

data(LifeCycleSavings)
pop <- LifeCycleSavings[, 2:3]
oec <- LifeCycleSavings[, -(2:3)]
cancor(pop, oec)
x <- matrix(rnorm(150), 50, 3)
y <- matrix(rnorm(250), 50, 5)
(cxy <- cancor(x, y))
all(abs(cor(x %*% cxy$xcoef,
    y %*% cxy$ycoef)[,1:3] - diag(cxy \$ cor)) < 1e-15)
all(abs(cor(x %*% cxy$xcoef) - diag(3)) < 1e-15)
all(abs(cor(y %*% cxy$ycoef) - diag(5)) < 1e-15)

```
cmdscale Classical (Metric) Multidimensional Scaling

\section*{Description}

Classical multidimensional scaling of a data matrix.

\section*{Usage}
```

cmdscale(d, k = 2, eig = FALSE, add = FALSE, x.ret = FALSE)

```

\section*{Arguments}
add logical indicating if an additive constant \(c *\) should be computed, and
d
k
eig
x.ret
a distance structure such as that returned by dist or a full symmetric matrix containing the dissimilarities.
the dimension of the space which the data are to be represented in; must be in \(\{1,2, \ldots, n-1\}\). added to the non-diagonal dissimilarites such that all \(n-1\) eigenvalues are non-negative.
indicates whether the doubly centered symmetric distance matrix should be returned.

\section*{Details}

Multidimensional scaling takes a set of dissimilarities and returns a set of points such that the distances between the points are approximately equal to the dissimilarities.
The functions isoMDS and sammon in package MASS provide alternative ordination techniques.
When add \(=\) TRUE, an additive constant \(c *\) is computed, and the dissimilarities \(d_{i j}+c *\) are used instead of the original \(d_{i j}\) 's.

Whereas S-PLUS computes this constant using an approximation suggested by Torgerson, R uses the exact analytical solution of Cailliez (1983), see also Cox and Cox (1994).

\section*{Value}

If eig = FALSE and x. ret \(=\) FALSE (default), a matrix with k columns whose rows give the coordinates of the points chosen to represent the dissimilarities.
Otherwise, a list containing the following components.
points a matrix with k columns whose rows give the coordinates of the points chosen to represent the dissimilarities.
eig the \(n-1\) eigenvalues computed during the scaling process if eig is true. \(\mathrm{x} \quad\) the doubly centered distance matrix if x. ret is true.
GOF a numeric vector of length 2 , equal to say \(\left(g_{1}, g_{2}\right)\), where \(g_{i}=\) \(\left(\sum_{j=1}^{k} \lambda_{j}\right) /\left(\sum_{j=1}^{n} T_{i}\left(\lambda_{j}\right)\right)\), where \(\lambda_{j}\) are the eigenvalues (sorted decreasingly), \(T_{1}(v)=|v|\), and \(T_{2}(v)=\max (v, 0)\).

\section*{References}

Cox, T. F. and Cox, M. A. A. (1994) Multidimensional Scaling. Chapman and Hall.
Mardia, K. V., Kent, J. T. and Bibby, J. M. (1979). Chapter 14 of Multivariate Analysis, London: Academic Press.

Seber, G. A. F. (1984). Multivariate Observations. New York: Wiley.
Torgerson, W. S. (1958). Theory and Methods of Scaling. New York: Wiley.
Cailliez, F. (1983) The analytical solution of the additive constant problem. Psychometrika 48, 343-349.

\section*{See Also}
dist. Also isoMDS and sammon in package MASS.

\section*{Examples}
```

data(eurodist)
loc <- cmdscale(eurodist)
x <- loc[,1]
y <- -loc[,2]
plot(x, y, type="n", xlab="", ylab="", main="cmdscale(eurodist)")
text(x, y, names(eurodist), cex=0.8)
cmdsE <- cmdscale(eurodist, k=20, add = TRUE, eig = TRUE, x.ret = TRUE)
str(cmdsE)

```
cophenetic Cophenetic Distances for a Hierarchical Clustering

\section*{Description}

Computes the cophenetic distances for a hierarchical clustering.

\section*{Usage}
cophenetic(x)

\section*{Arguments}
x
an object of class hclust or with a method for as.hclust() such as agnes.

\section*{Details}

The cophenetic distance between two observations that have been clustered is defined to be the intergroup dissimilarity at which the two observations are first combined into a single cluster. Note that this distance has many ties and restrictions.

It can be argued that a dendrogram is an appropriate summary of some data if the correlation between the original distances and the cophenetic distances is high. Otherwise, it should simply be viewed as the description of the output of the clustering algorithm.

\section*{Value}

An object of class dist.

\section*{Author(s)}

Robert Gentleman

\section*{References}

Sneath, P.H.A. and Sokal, R.R (1973) Numerical Taxonomy: The Principles and Practice of Numerical Classification, p. 278 ff; Freeman, San Francisco.

\author{
See Also
}
```

dist, hclust

```

\section*{Examples}
```

data(USArrests)
library(mva)
d1 <- dist(USArrests)
hc <- hclust(d1, "ave")
d2 <- cophenetic(hc)
cor(d1,d2) \# 0.7659

## Example from Sneath \& Sokal, Fig. 5-29, p. }27

d0 <- c(1,3.8,4.4,5.1, 4,4.2,5, 2.6,5.3, 5.4)
attributes(d0) <- list(Size = 5, diag=TRUE)
class(d0) <- "dist"
names(d0) <- letters[1:5]
d0
str(upgma <- hclust(do, method = "average"))
plot(upgma, hang = -1)

# 

(d.coph <- cophenetic(upgma))
cor(dO, d.coph) \# 0.9911

```
```

cutree Cut a tree into groups of data

```

\section*{Description}

Cuts a tree, e.g., as resulting from hclust, into several groups either by specifying the desired number(s) of groups or the cut height(s).

\section*{Usage}
cutree(tree, k=NULL, h=NULL)

\section*{Arguments}
tree a tree as produced by hclust. cutree() only expects a list with components merge, height, and labels, of appropriate content each.
\(k \quad\) an integer scalar or vector with the desired number of groups
\(\mathrm{h} \quad\) numeric scalar or vector with heights where the tree should be cut.
At least one of \(k\) or \(h\) must be specified, \(k\) overrides \(h\) if both are given.

\section*{Value}
cutree returns a vector with group memberships if k or h are scalar, otherwise a matrix with group meberships is returned where each column corresponds to the elements of k or h , respectively (which are also used as column names).

\section*{See Also}
hclust

\section*{Examples}
```

require(mva)
data(USArrests)
hc <- hclust(dist(USArrests))
cutree(hc, k=1:5)\#k = 1 is trivial
cutree(hc, h=250)

## Compare the 2 and 3 grouping:

g24 <- cutree(hc, k = c(2,4))
table(g24[,"2"], g24[,"4"])

```
dendrogram General Tree Structures

\section*{Description}

Class "dendrogram" provides general functions for handling tree-like structures. It is intended as a replacement for similar functions in hierarchical clustering and classification/regression trees, such that all of these can use the same engine for plotting or cutting trees.

The code is still in testing stage and the API may change in the future.

\section*{Usage}
```

as.dendrogram(object, ...)
as.dendrogram.hclust(object, ...)
plot(x, type=c("rectangle", "triangle"),
center=FALSE, edge.root= !is.null(attr(x,"edgetext")),
nodePar = NULL, edgePar = list(), xlab="", ylab="",
horiz = FALSE, ...)
cut(x, h, ...)
print(x, digits, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & any R object that can be made into a class "dendrogram" one. \\
x & object of class "dendrogram". \\
type & \begin{tabular}{l} 
type of plot.
\end{tabular} \\
center & \begin{tabular}{l} 
logical; if TRUE, nodes are plotted centered with respect to the leaves in \\
the branch. Otherwise (default), plot them in the middle of all direct \\
child nodes.
\end{tabular} \\
edge.root & \begin{tabular}{l} 
logical; if true, draw an edge to the root node.
\end{tabular} \\
nodePar & \begin{tabular}{l} 
a list of plotting parameters to use for the nodes (see points) or NULL by \\
default which does not draw symbols at the nodes. The list may contain
\end{tabular} \\
& \begin{tabular}{l} 
components named pch, cex, col, and/or bg each of which can have \\
length two for specifying separate attributes for inner nodes and leaves.
\end{tabular}
\end{tabular}
```

edgePar a list of plotting parameters to use for the edge (see lines). The list
may contain components named col, lty and/or lwd.
horiz logical indicating if the dendrogram should be draw horizontally or not.
h height at which the tree is cut.

```
..., xlab, ylab
                                    graphical parameters, or arguments for other methods.
digits integer specifiying the precision for printing, see print.default.

\section*{Details}

Warning: This documentation is preliminary
The dendrogram is directly represented as a nested list where each component corresponds to a branch of the tree. Hence, the first branch of tree \(z\) is \(z[[1]]\), the second branch of the corresponding subtree is \(\mathbf{z}[[1]][[2]]\) etc.. Each node of the tree carries some information needed for efficient plotting or cutting as attributes:
members total number of leaves in the branch
height numeric non-negative height at which the node is plotted.
midpoint numeric horizontal distance of the node from the left border of the branch. This is needed at least for plot (*, center=FALSE).
text character; the label of the node
edgetext character; the label for the edge leading to the node
nodePar a named list of length one vectors specifying node-specific attributes for points plotting, see the nodePar argument above.
edgePar a named list of length one vectors specifying attributes for segments plotting of the edge leading to the node, see the edgePar argument above.
leaf logical, if TRUE, the node is a leaf of the tree.
cut.dendrogram() returns a list with components \$upper and \$lower, the first is a truncated version of the original tree, also of class dendrogram, the latter a list with the branches obtained from cutting the tree, each a dendrogram.

There are [ [, print, and str methods for "dendrogram" objects where the first one (extraction) ensures that selecting sub-branches keeps the class.

Objects of class "hclust" can be converted to class "dendrogram" using method as.dendrogram.
isLeaf(), plotNode() and plotNodeLimit() are helper functions.

\section*{Note}

When using type = "triangle", center = TRUE often looks better.

\section*{Examples}
```

library(mva)
data(USArrests)
hc <- hclust(dist(USArrests), "ave")
(dend1 <- as.dendrogram(hc)) \# "print()" method
str(dend1) \# "str()" method
op <- par(mfrow= c(2,2), mar = c(3,3,1,1))

```
```

plot(dend1)

## "triangle" type and show inner nodes:

plot(dend1, nodePar=list(pch = c(1,NA),cex=0.8), type = "t", center=TRUE)
plot(dend1, edgePar=list(col = 1:2, lty = 2:3), edge.root = TRUE)
plot(dend1, nodePar=list(pch = 2:1, cex=.4*2:1, col = 2:3), horiz = TRUE)
dend2 <- cut(dend1, h=70)
plot(dend2\$upper)

## leafs are wrong horizontally:

plot(dend2\$upper, nodePar=list(pch = c(1,7), col = 2:1))

## dend2\$lower is *NOT* a dendrogram, but a list of .. :

plot(dend2\$lower[[3]], nodePar=list(col=4), horiz = TRUE, type = "tr")

## "inner" and "leaf" edges in different type \& color :

plot(dend2\$lower[[2]], nodePar=list(col=1),\# non empty list
edgePar = list(lty=1:2, col=2:1), edge.root=TRUE)
par(op)

```
dist Distance Matrix Computation

\section*{Description}

This function computes and returns the distance matrix computed by using the specified distance measure to compute the distances between the rows of a data matrix.

\section*{Usage}
```

dist(x, method = "euclidean", diag = FALSE, upper = FALSE)
print.dist(x, diag = NULL, upper = NULL, ...)
as.matrix.dist(x)
as.dist(m, diag = FALSE, upper = FALSE)

```

\section*{Arguments}
\(\mathrm{x} \quad\) numeric matrix or (data frame). Distances between the rows of x will be computed.
method the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra" or "binary". Any unambiguous substring can be given.
diag logical value indicating whether the diagonal of the distance matrix should be printed by print.dist.
upper logical value indicating whether the upper triangle of the distance matrix should be printed by print.dist.
m A matrix of distances to be converted to a "dist" object (only the lower triangle is used, the rest is ignored).
... further arguments, passed to the (next) print method.

\section*{Details}

Available distance measures are (written for two vectors \(x\) and \(y\) ):
euclidean: Usual square distance between the two vectors (2 norm).
maximum: Maximum distance between two components of \(x\) and \(y\) (supremum norm)
manhattan: Absolute distance between the two vectors (1 norm).
canberra: \(\sum_{i}\left|x_{i}-y_{i}\right| /\left|x_{i}+y_{i}\right|\). Terms with zero numerator and denominator are omitted from the sum and treated as if the values were missing.
binary: (aka asymmetric binary): The vectors are regarded as binary bits, so non-zero elements are 'on' and zero elements are 'off'. The distance is the proportion of bits in which only one is on amongst those in which at least one is on.

Missing values are allowed, and are excluded from all computations involving the rows within which they occur. Further, when Inf values are involved, all pairs of values are excluded when their contribution to the distance gave NaN or NA.
If some columns are excluded in calculating a Euclidean, Manhattan or Canberra distance, the sum is scaled up proportionally to the number of columns used. If all pairs are excluded when calculating a particular distance, the value is NA.

The functions as.matrix.dist() and as.dist() can be used for conversion between objects of class "dist" and conventional distance matrices and vice versa.

\section*{Value}

An object of class "dist".
The lower triangle of the distance matrix stored by columns in a vector, say do. If \(n\) is the number of observations, i.e., \(\mathrm{n}<-\operatorname{attr}(\mathrm{do}\), "Size"), then for \(i<j<=n\), the dissimilarity between (row) \(i\) and \(j\) is do[n*(i-1) - \(i *(i-1) / 2+j-i]\). The length of the vector is \(n *(n-1) / 2\), i.e., of order \(n^{2}\).

The object has the following attributes (besides "class" equal to "dist"):
Size integer, the number of observations in the dataset.
Labels optionally, contains the labels, if any, of the observations of the dataset.
Diag, Upper logicals corresponding to the arguments diag and upper above, specifying how the object should be printed.
call optionally, the call used to create the object.
methods optionally, the distance method used; resulting form dist(), the (match.arg()ed) method argument.

\section*{References}

Mardia, K. V., Kent, J. T. and Bibby, J. M. (1979) Multivariate Analysis. London: Academic Press.

\section*{See Also}
daisy in the 'cluster' package with more possibilities in the case of mixed (contiuous / categorical) variables. hclust.

\section*{Examples}
```

x <- matrix(rnorm(100), nrow=5)
dist(x)
dist(x, diag = TRUE)
dist(x, upper = TRUE)
m <- as.matrix(dist(x))
d <- as.dist(m)
stopifnot(d == dist(x))
names(d) <- LETTERS[1:5]
print(d, digits = 3)

## example of binary and canberra distances.

x <- c(0, 0, 1, 1, 1, 1)
y <- c(1, 0, 1, 1, 0, 1)
dist(rbind(x,y), method="binary")

## answer 0.4 = 2/5

dist(rbind(x,y), method="canberra")

## answer 2 * (6/5)

## Examples involving "Inf" :

## 1)

x[6] <- Inf
(m2 <- rbind(x,y))
dist(m2, method="binary")\# warning, answer 0.5 = 2/4

## These all give "Inf":

stopifnot(Inf == dist(m2, method= "euclidean"),
Inf == dist(m2, method= "maximum"),
Inf == dist(m2, method= "manhattan"))

## "Inf" is same as very large number:

x1 <- x; x1[6] <- 1e100
stopifnot(dist(cbind(x ,y), method="canberra") ==
print(dist(cbind(x1,y), method="canberra")))

## 2)

y[6] <- Inf \#-> 6-th pair is excluded
dist(rbind(x,y), method="binary")\# warning; 0.5
dist(rbind(x,y), method="canberra")\# 3
dist(rbind(x,y), method="maximum") \# 1
dist(rbind(x,y), method="manhattan")\# 2.4

```
```

factanal Factor Analysis

```

\section*{Description}

Perform maximum-likelihood factor analysis on a covariance matrix or data matrix.

\section*{Usage}
```

factanal(x, factors, data = NULL, covmat = NULL, n.obs = NA,
subset, na.action,
start = NULL, scores = c("none", "regression", "Bartlett"),
rotation = "varimax", control = NULL, ...)

```

\section*{Arguments}
n.obs The number of observations, used if covmat is a covariance matrix.
x
factors The number of factors to be fitted.
data
covmat
subset
na.action
start
scores
rotation a numeric matrix.

A data frame. correlation matrices are covariance matrices. The na.action to be used if x is used as a formula. uniquenesses.

Either a formula or a numeric matrix or an object that can be coerced to

A covariance matrix, or a covariance list as returned by cov.wt. Of course,

A specification of the cases to be used, if x is used as a matrix or formula.

NULL or a matrix of starting values, each column giving an initial set of

Type of scores to produce, if any. The default is none, "regression" gives Thompson's scores, "Bartlett" given Bartlett's weighted least-squares scores. Partial matching allows these names to be abbreviated.
character. "none" or the name of a function to be used to rotate the factors: it will be called with first argument the loadings matrix, and should return a list with component loadings giving the rotated loadings, or just the rotated loadings.
control A list of control values,
nstart The number of starting values to be tried if start \(=\) NULL. Default 1.
trace logical. Output tracing information? Default FALSE.
lower The lower bound for uniquenesses during optimization. Should be \(>0\). Default 0.005 .
opt A list of control values to be passed to optim's control argument. rotate a list of additional arguments for the rotation function.
... Components of control can also be supplied as named arguments to factanal.

\section*{Details}

The factor analysis model is
\[
x=\Lambda f+e
\]
for a \(p\)-element row-vector \(x\), a \(p \times k\) matrix of loadings, a \(k\)-element vector of scores and a \(p\)-element vector of errors. None of the components other than \(x\) is observed, but the major restriction is that the scores be uncorrelated and of unit variance, and that the errors be independent with variances \(\Phi\), the uniquenesses. Thus factor analysis is in essence a model for the covariance matrix of \(x\),
\[
\Sigma=\Lambda^{\prime} \Lambda+\Psi
\]

There is still some indeterminacy in the model for it is unchanged if \(\Lambda\) is replaced by \(G \Lambda\) for any orthogonal matrix \(G\). Such matrices \(G\) are known as rotations (although the term is applied also to non-orthogonal invertible matrices).

If covmat is supplied it is used. Otherwise x is used if it is a matrix, or a formula x is used with data to construct a model matrix, and that is used to construct a covariance matrix. (It makes no sense for the formula to have a response.) Once a covariance matrix is found
or calculated from x , it is converted to a correlation matrix for analysis. The correlation matrix is returned as component correlation of the result.
The fit is done by optimizing the log likelihood assuming multivariate normality over the uniquenesses. (The maximizing loadings for given uniquenesses can be found analytically: Lawley \& Maxwell (1971, p. 27).) All the starting values supplied in start are tried in turn and the best fit obtained is used. If start = NULL then the first fit is started at the value suggested by Jöreskog (1963) and given by Lawley \& Maxwell (1971, p. 31), and then control\$nstart - 1 other values are tried, randomly selected as equal values of the uniquenesses.

The uniquenesses are technically constrained to lie in [0, 1], but near-zero values are problematical, and the optimization is done with a lower bound of control\$lower, default 0.005 (Lawley \& Maxwell, 1971, p. 32).
Scores can only be produced if a data matrix is supplied and used. The first method is the regression method of Thomson (1951), the second the weighted least squares method of Bartlett (1937, 8). Both are estimates of the unobserved scores \(f\). Thomson's method regresses (in the population) the unknown \(f\) on \(x\) to yield
\[
\hat{f}=\Lambda^{\prime} \Sigma^{-1} x
\]
and then substitutes the sample estimates of the quantities on the right-hand side. Bartlett's method minimizes the sum of squares of standardized errors over the choice of \(f\), given (the fitted) \(\Lambda\).
If x is a formula then the standard NA-handling is applied to the scores (if requested): see napredict.

\section*{Value}

An object of class "factanal" with components
\begin{tabular}{ll} 
loadings & \begin{tabular}{l} 
A matrix of loadings, one column for each factor. The factors are ordered \\
in decreasing order of sums of squares of loadings, and given the sign that \\
will make the sum of the loadings positive.
\end{tabular} \\
uniquenesses & \begin{tabular}{l} 
The uniquenesses computed.
\end{tabular} \\
\begin{tabular}{l} 
correlation \\
criteria
\end{tabular} & \begin{tabular}{l} 
The correlation matrix used. \\
The results of the optimization: the value of the negative log-likelihood \\
and information on the iterations used.
\end{tabular} \\
factors & \begin{tabular}{l} 
The argument factors. \\
dof
\end{tabular} \\
method & The number of degrees of freedom of the factor analysis model. \\
scores & The method: always "mle". \\
n.obs & If requested, a matrix of scores. \\
call & The number of observations if available, or NA. \\
na.action & The matched call.
\end{tabular}

\section*{Note}

There are so many variations on factor analysis that it is hard to compare output from different programs. Further, the optimization in maximum likelihood factor analysis is hard, and many other examples we compared had less good fits than produced by this function. In particular, solutions which are Heywood cases (with one or more uniquenesses essentially zero) are much often common than most texts and some other programs would lead one to believe.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Bartlett, M. S. (1937) The statistical conception of mental factors. British Journal of Psychology, 28, 97-104.
Bartlett, M. S. (1938) Methods of estimating mental factors. Nature, 141, 609-610.
Jöreskog, K. G. (1963) Statistical Estimation in Factor Analysis. Almqvist and Wicksell.
Lawley, D. N. and Maxwell, A. E. (1971) Factor Analysis as a Statistical Method. Second edition. Butterworths.

Thomson, G. H. (1951) The Factorial Analysis of Human Ability. London University Press.

\section*{See Also}
print.loadings, varimax, princomp, ability.cov, Harman23.cor, Harman74.cor

\section*{Examples}
```


# A little demonstration, v2 is just v1 with noise,

# and same for v4 vs. v3 and v6 vs. v5

# Last four cases are there to add noise

# and introduce a positive manifold (g factor)

v1 <- c(1,1,1,1,1,1,1,1,1,1,3,3,3,3,3,4,5,6)
v2 <- c(1,2,1,1,1,1,2,1,2,1,3,4,3,3,3,4,6,5)
v3 <- c(3,3,3,3,3,1,1,1,1,1,1,1,1,1,1,5,4,6)
v4 <- c(3,3,4,3,3,1,1,2,1,1,1,1,2,1,1,5,6,4)
v5 <- c(1,1,1,1,1,3,3,3,3,3,1,1,1,1,1,6,4,5)
v6 <- c(1,1,1,2,1,3,3,3,4,3,1,1,1,2,1,6,5,4)
m1 <- cbind(v1,v2,v3,v4,v5,v6)
cor(m1)
factanal(m1, factors=3) \# varimax is the default
factanal(m1, factors=3, rotation="promax")

# The following shows the g factor as PC1

prcomp(m1)

## formula interface

factanal(~v1+v2+v3+v4+v5+v6, factors = 3,
scores = "Bartlett")\$scores

## a realistic example from Barthlomew (1987, pp. 61-65)

example(ability.cov)

```

\section*{Harman23.cor Harman Example 2.3}

\section*{Description}

A correlation matrix of eight physical measurements on 305 girls between ages seven and seventeen.

\section*{Usage}
data(Harman23.cor)

\section*{Source}

Harman, H. H. (1976) Modern Factor Analysis, Third Edition Revised, University of Chicago Press, Table 2.3.

\section*{Examples}
```

data(Harman23.cor)
(Harman23.FA <- factanal(factors = 1, covmat = Harman23.cor))
for(factors in 2:4) print(update(Harman23.FA, factors = factors))

```
```

Harman74.cor Harman Example 7.4

```

\section*{Description}

A correlation matrix of 24 psychological tests given to 145 seventh and eight-grade children in a Chicago suburb by Holzinger and Swineford.

\section*{Usage}
data(Harman74.cor)

\section*{Source}

Harman, H. H. (1976) Modern Factor Analysis, Third Edition Revised, University of Chicago Press, Table 7.4.

\section*{Examples}
```

data(Harman74.cor)
(Harman74.FA <- factanal(factors = 1, covmat = Harman74.cor))
for(factors in 2:5) print(update(Harman74.FA, factors = factors))
Harman74.FA <- factanal(factors = 5, covmat = Harman74.cor,
rotation="promax")
print(Harman74.FA\$loadings, sort = TRUE)

```

\section*{hclust Hierarchical Clustering}

\section*{Description}

Hierarchical cluster analysis on a set of dissimilarities and methods for analyzing it.
```

Usage
hclust(d, method = "complete", members=NULL)
plot(x, labels = NULL, hang = 0.1,
axes = TRUE, frame.plot = FALSE, ann = TRUE,
main = "Cluster Dendrogram",
sub = NULL, xlab = NULL, ylab = "Height", ...)
plclust(tree, hang = 0.1, unit = FALSE, level = FALSE, hmin = 0,
square = TRUE, labels = NULL, plot. = TRUE,
axes = TRUE, frame.plot = FALSE, ann = TRUE,
main = "", sub = NULL, xlab = NULL, ylab = "Height")

```

\section*{Arguments}
```

d a dissimilarity structure as produced by dist.
method the agglomeration method to be used. This should be (an unambigu-
ous abbreviation of) one of "ward", "single", "complete", "average",
"mcquitty", "median" or "centroid".
members NULL or a vector with length size of d.
x,tree an object of the type produced by hclust.
hang The fraction of the plot height by which labels should hang below the rest
of the plot. A negative value will cause the labels to hang down from 0.
labels A character vector of labels for the leaves of the tree. By default the row
names or row numbers of the original data are used. If labels=FALSE no
labels at all are plotted.
axes, frame.plot, ann
logical flags as in plot.default.
main, sub, xlab, ylab
character strings for title. sub and xlab have a non-NULL default when
there's a tree\$call.
... Further graphical arguments.
unit, level, hmin, square, plot.
as yet unimplemented arguments of plclust for S-plus compatibility.

```

\section*{Details}

This function performs a hierarchical cluster analysis using a set of dissimilarities for the \(n\) objects being clustered. Initially, each object is assigned to its own cluster and then the algorithm proceeds iteratively, at each stage joining the two most similar clusters, continuing until there is just a single cluster. At each stage distances between clusters are recomputed
by the Lance-Williams dissimilarity update formula according to the particular clustering method being used.

A number of different clustering methods are provided. Ward's minimum variance method aims at finding compact, spherical clusters. The complete linkage method finds similar clusters. The single linkage method (which is closely related to the minimal spanning tree) adopts a 'friends of friends' clustering strategy. The other methods can be regarded as aiming for clusters with characteristics somewhere between the single and complete link methods.

If members!=NULL, then d is taken to be a dissimilarity matrix between clusters instead of dissimilarities between singletons and members gives the number of observations per cluster. This way the hierarchical cluster algorithm can be "started in the middle of the dendrogram", e.g., in order to reconstruct the part of the tree above a cut (see examples). Dissimilarities between clusters can be efficiently computed (i.e., without hclust itself) only for a limited number of distance/linkage combinations, the simplest one being squared Euclidean distance and centroid linkage. In this case the dissimilarities between the clusters are the squared Euclidean distances between cluster means.
In hierarchical cluster displays, a decision is needed at each merge to specify which subtree should go on the left and which on the right. Since, for \(n\) observations there are \(n-1\) merges, there are \(2^{(n-1)}\) possible orderings for the leaves in a cluster tree, or dendrogram. The algorithm used in hclust is to order the subtree so that the tighter cluster is on the left (the last, i.e. most recent, merge of the left subtree is at a lower value than the last merge of the right subtree). Single observations are the tightest clusters possible, and merges involving two observations place them in order by their observation sequence number.

\section*{Value}

An object of class hclust which describes the tree produced by the clustering process. The object is a list with components:
\begin{tabular}{ll} 
merge & \begin{tabular}{l} 
an \(n-1\) by 2 matrix. Row \(i\) of merge describes the merging of clusters \\
at step \(i\) of the clustering. If an element \(j\) in the row is negative, then \\
observation \(-j\) was merged at this stage. If \(j\) is positive then the merge \\
was with the cluster formed at the (earlier) stage \(j\) of the algorithm. \\
Thus negative entries in merge indicate agglomerations of singletons, and \\
positive entries indicate agglomerations of non-singletons. \\
a set of \(n-1\) non-decreasing real values. The clustering height: that is, \\
the value of the criterion associated with the clustering method for the \\
particular agglomeration. \\
height \\
a vector giving the permutation of the original observations suitable for \\
plotting, in the sense that a cluster plot using this ordering and matrix \\
merge will not have crossings of the branches.
\end{tabular} \\
labels & \begin{tabular}{l} 
labels for each of the objects being clustered. \\
call \\
the call which produced the result.
\end{tabular} \\
method & \begin{tabular}{l} 
dhe cluster method that has been used.
\end{tabular} \\
& \begin{tabular}{l} 
the distance that has been used to create d (only returned if the distance \\
object has a "method" attribute).
\end{tabular}
\end{tabular}

There is a print and a plot method for hclust objects. The plclust() function is basically the same as the plot method, plot.hclust, primarily for back compatibility with S-plus. Its extra arguments are not yet implemented.

\section*{Author(s)}

The hclust function is based on Fortran code contributed to STATLIB by F. Murtagh.

\section*{References}

Everitt, B. (1974). Cluster Analysis. London: Heinemann Educ. Books.
Hartigan, J. A. (1975). Clustering Algorithms. New York: Wiley
Sneath, P. H. A. and R. R. Sokal (1973). Numerical Taxonomy. San Francisco: Freeman.
Anderberg, M. R. (1973). Cluster Analysis for Applications. Academic Press: New York.
Gordon, A. D. (1999). Classification. Second Edition. London: Chapman and Hall / CRC
Murtagh, F. (1985). "Multidimensional Clustering Algorithms", in COMPSTAT Lectures 4. Wuerzburg: Physica-Verlag (for algorithmic details of algorithms used).

\section*{See Also}
kmeans.

\section*{Examples}
```

library(mva)
data(USArrests)
hc <- hclust(dist(USArrests), "ave")
plot(hc)
plot(hc, hang = -1)

## Do the same with centroid clustering and squared Euclidean distance,

## cut the tree into ten clusters and reconstruct the upper part of the

## tree from the cluster centers.

hc <- hclust(dist(USArrests)^2, "cen")
memb <- cutree(hc, k = 10)
cent <- NULL
for(k in 1:10){
cent <- rbind(cent, colMeans(USArrests[memb == k, , drop = FALSE]))
}
hc1 <- hclust(dist(cent)^2, method = "cen", members = table(memb))
opar <- par(mfrow = c(1, 2))
plot(hc, labels = FALSE, hang = -1, main = "Original Tree")
plot(hc1, labels = FALSE, hang = -1, main = "Re-start from 10 clusters")
par(opar)

```
identify.hclust Identify Clusters in a Dendrogram

\section*{Description}
identify.hclust reads the position of the graphics pointer when the (first) mouse button is pressed. It then cuts the tree at the vertical position of the pointer and highlights the cluster containing the horizontal position of the pointer. Optionally a function is applied to the index of data points contained in the cluster.

\section*{Usage}
```

identify(x, FUN $=$ NULL, $N=20, \operatorname{MAXCLUSTER~}=20$,

```
    DEV.FUN = NULL, ...)

\section*{Arguments}
x
an object of the type produced by hclust.
FUN (optional) function to be applied to the index numbers of the data points in a cluster (see Details below).
N
the maximum number of clusters to be identified.
MAXCLUSTER The maximum number of clusters that can be produced by a cut (limits the effective vertical range of the pointer).
DEV.FUN (optional) integer scalar. If specified, the corresponding graphics device is made active before FUN is applied.
... further arguments to FUN.

\section*{Details}

By default clusters can be identified using the mouse and an invisible list of indices of the respective data points is returned.
If FUN is not NULL, then the index vector of data points is passed to this function as first argument, see the examples below. The active graphics device for FUN can be specified using DEV.FUN.

The identification process is terminated by pressing any mouse button other than the first, or by clicking outside the graphics window.

\section*{Value}

Either a list of data point index vectors or a list of return values of FUN.

\section*{See Also}
hclust, rect.hclust

\section*{Examples}
```

library(mva)
data(USArrests)
hca <- hclust(dist(USArrests))
plot(hca)
(x <- identify.hclust(hca))
data(iris)
hci <- hclust(dist(iris[,1:4]))
plot(hci)
identify.hclust(hci, function(k) print(table(iris[k,5])))

# open a new device

dev.set(2)
plot(hci)
identify.hclust(hci, function(k) barplot(table(iris[k,5])), DEV.FUN=3)

```
\begin{tabular}{l} 
kmeans \(\quad\) K-Means Clustering \\
\hline
\end{tabular}

\section*{Description}

Perform k-means clustering on a data matrix.

\section*{Usage}
kmeans( x , centers, iter.max \(=10\) )

\section*{Arguments}
\begin{tabular}{ll}
x & A numeric matrix of data, or an object that can be coerced to such a ma- \\
trix (such as a numeric vector or a data frame with all numeric columns). \\
centers & Either the number of clusters or a set of initial cluster centers. If the first, \\
a random set of rows in \(x\) are chosen as the initial centers. \\
iter.max & The maximum number of iterations allowed.
\end{tabular}

\section*{Details}

The data given by x is clustered by the k-means algorithm. When this terminates, all cluster centres are at the mean of their Voronoi sets (the set of data points which are nearest to the cluster centre).
The algorithm of Hartigan and Wong (1979) is used.

\section*{Value}

A list with components:
\begin{tabular}{ll} 
cluster & A vector of integers indicating the cluster to which each point is allocated. \\
centers & A matrix of cluster centres. \\
withinss & The within-cluster sum of squares for each cluster. \\
size & The number of points in each cluster.
\end{tabular}

\section*{References}

Hartigan, J.A. and Wong, M.A. (1979). A K-means clustering algorithm. Applied Statistics 28, 100-108.

\section*{Examples}
```


# a 2-dimensional example

x <- rbind(matrix(rnorm(100, sd = 0.3), ncol = 2),
matrix(rnorm(100, mean = 1, sd = 0.3), ncol = 2))
cl <- kmeans(x, 2, 20)
plot(x, col = cl$cluster)
points(cl$centers, col = 1:2, pch = 8)

```
```

loadings Print Loadings in Factor Analysis

```

\section*{Description}

Extract or print loadings in factor analysis (or principal components analysis).

\section*{Usage}
```

loadings(x)
print.loadings(x, digits = 3, cutoff = 0.1, sort = FALSE, ...)
print.factanal(x, digits = 3, ...)

```

\section*{Arguments}
\(\mathrm{x} \quad\) an object of class "factanal" or "princomp" or the loadings component of such an object.
digits number of decimal places to use in printing uniquenesses and loadings.
cutoff loadings smaller than this (in absolute value) are suppressed.
sort logical. If true, the variables are sorted by their importance on each factor. Each variable with any loading larger than 0.5 (in modulus) is assigned to the factor with the largest loading, and the variables are printed in the order of the factor they are assigned to, then those unassigned.
... further arguments for other methods, such as cutoff and sort for print.factanal.

Author(s)
B. D. Ripley

\section*{See Also}
factanal, princomp

\section*{prcomp Principal Components Analysis}

\section*{Description}

Performs a principal components analysis on the given data matrix and returns the results as an object of class prcomp.

\section*{Usage}
```

prcomp(x, retx = TRUE, center = TRUE, scale. = FALSE, tol = NULL)

```

\section*{Arguments}
x
retx
center
scale. a logical value indicating whether the variables should be scaled to have unit variance before the analysis takes place. The default is FALSE for consistency with S, but in general scaling is advisable. Alternately, a vector of length equal the number of columns of \(x\) can be supplied. The value is passed to scale.
tol
a value indicating the magnitude below which components should be omitted. (Components are omitted if their standard deviations are less than or equal to tol times the standard deviation of the first component.) With the default null setting, no components are omitted. Other settings for tol could be tol \(=0\) or tol \(=\) sqrt(.Machine\$double.eps), which would omit essentially constant components.

\section*{Details}

The calculation is done by a singular value decomposition of the (centered and scaled) data matrix, not by using eigen on the covariance matrix. This is generally the preferred method for numerical accuracy. The print method for the these objects prints the results in a nice format and the plot method produces a scree plot.

\section*{Value}
prcomp returns an list with class "prcomp" containing the following components:
sdev the standard deviations of the principal components (i.e., the square roots of the eigenvalues of the covariance/correlation matrix, though the calculation is actually done with the singular values of the data matrix).
rotation the matrix of variable loadings (i.e., a matrix whose columns contain the eigenvectors). The function princomp returns this in the element loadings.

X
if retx is true the value of the rotated data (the data multiplied by the rotation matrix) is returned.

\section*{References}

Mardia, K. V., J. T. Kent, and J. M. Bibby (1979) Multivariate Analysis, London: Academic Press.

Venables, W. N. and B. D. Ripley (1997, 9) Modern Applied Statistics with S-PLUS, Springer-Verlag.

\section*{See Also}
princomp, cor, cov, svd, eigen.

\section*{Examples}
```


## the variances of the variables in the

## USArrests data vary by orders of magnitude, so scaling is appropriate

data(USArrests)
prcomp(USArrests) \# inappropriate
prcomp(USArrests, scale = TRUE)
plot(prcomp(USArrests))
summary(prcomp(USArrests, scale = TRUE))

```
princomp Principal Components Analysis

\section*{Description}
princomp performs a principal components analysis on the given data matrix and returns the results as an object of class princomp.

\section*{Usage}
```

princomp(x, data = NULL, subset, na.action, ...)
princomp(x, cor = FALSE, scores = TRUE, covmat = NULL,
subset = rep(TRUE, nrow(as.matrix(x))), ...)

```

\section*{Arguments}
x
data an optional data frame containing the variables in the formula x . By default the variables are taken from environment ( x ).
subset an optional vector used to select rows (observations) of the data matrix x .
na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The "factory-fresh" default is na.omit.
cor a logical value indicating whether the calculation should use the correlation matrix or the covariance matrix.
scores a logical value indicating whether the score on each principal component should be calculated.
covmat a covariance matrix, or a covariance list as returned by cov.wt, cov.mve or cov.mcd. If supplied, this is used rather than the covariance matrix of x .
... arguments passed to or from other methods. If x is a formula one might specify cor or scores.

\section*{Details}
princomp is a generic function with "formula" and "default" methods.
The calculation is done using eigen on the correlation or covariance matrix, as determined by cor. This is done for compatibility with the S-PLUS result. A preferred method of calculation is to use svd on x , as is done in prcomp.
Note that the default calculation uses divisor N for the covariance matrix.
The print method for the these objects prints the results in a nice format and the plot method produces a scree plot (screeplot). There is also a biplot method.
If x is a formula then the standard NA-handling is applied to the scores (if requested): see napredict.

\section*{Value}
princomp returns a list with class "princomp" containing the following components:
sdev the standard deviations of the principal components.
loadings the matrix of variable loadings (i.e., a matrix whose columns contain the eigenvectors). This is of class "loadings": see loadings for its print method.
center the means that were subtracted.
scale the scalings applied to each variable.
n.obs the number of observations.
scores if scores = TRUE, the scores of the supplied data on the principal components.
call the matched call.
na.action If relevant.

\section*{References}

Mardia, K. V., J. T. Kent and J. M. Bibby (1979). Multivariate Analysis, London: Academic Press.

Venables, W. N. and B. D. Ripley (1997, 9). Modern Applied Statistics with S-PLUS, Springer-Verlag.

\section*{See Also}
```

summary.princomp, screeplot, biplot.princomp, prcomp, cor, cov, eigen.

```

\section*{Examples}
```


## The variances of the variables in the

## USArrests data vary by orders of magnitude, so scaling is appropriate

data(USArrests)
(pc.cr <- princomp(USArrests)) \# inappropriate
princomp(USArrests, cor = TRUE) \# =^= prcomp(USArrests, scale=TRUE)

## Similar, but different:

## The standard deviations differ by a factor of sqrt(49/50)

summary(pc.cr <- princomp(USArrests, cor = TRUE))
loadings(pc.cr) \#\# note that blank entries are small but not zero
plot(pc.cr) \# shows a screeplot.

```
```

biplot(pc.cr)

## Formula interface

princomp(~ ., data = USArrests, cor = TRUE)

# NA-handling

USArrests[1, 2] <- NA
pc.cr <- princomp(~ ., data = USArrests, na.action=na.exclude, cor = TRUE)
pc.cr\$scores

```
rect.hclust Draw Rectangles Around Hierarchical Clusters

\section*{Description}

Draws rectangles around the branches of a dendrogram highlighting the corresponding clusters. First the dendrogram is cut at a certain level, then a rectangle is drawn around selected branches.

\section*{Usage}
```

rect.hclust(tree, k = NULL, which = NULL, x = NULL, h = NULL,
border = 2, cluster = NULL)

```

\section*{Arguments}
tree an object of the type produced by hclust.
\(\mathrm{k}, \mathrm{h} \quad\) Scalar. Cut the dendrogram such that either exactly k clusters are produced or by cutting at height \(h\).
which, x A vector selecting the clusters around which a rectangle should be drawn. which seleccts clusters by number (from left to right in the tree), \(x\) selects clusters containing the respective horizontal coordinates. Default is which \(=1 \mathrm{k}\).
border Vector with border colors for the rectangles.
cluster Optional vector with cluster memberships as returned by cutree(hclust.obj, \(k=k\) ), can be specified for efficiency if already computed.

\section*{Value}
(Invisibly) returns a list where each element contains a vector of data points contained in the respective cluster.

\section*{See Also}
hclust, identify.hclust.

\section*{Examples}
```

library(mva)
data(USArrests)
hca <- hclust(dist(USArrests))
plot(hca)
rect.hclust(hca, k=3, border="red")
x <- rect.hclust(hca, h=50, which=c(2,7), border=3:4)
x

```
screeplot Screeplot of PCA Results

\section*{Description}
screeplot plots the variances against the number of the principal component. This is also the plot method for class "princomp".

\section*{Usage}
```

screeplot(x, npcs = min(10, length(x\$sdev)),
type = c("barplot", "lines"), main = deparse(substitute(x)), ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & an object of class "princomp", as from princomp(). \\
npcs & the number of principal components to be plotted. \\
type & the type of plot. \\
main,\(\ldots\) & graphics parameters.
\end{tabular}

\section*{References}

Mardia, K. V., J. T. Kent and J. M. Bibby (1979). Multivariate Analysis, London: Academic Press.

Venables, W. N. and B. D. Ripley (1997, 9). Modern Applied Statistics with S-PLUS, Springer-Verlag.

\section*{See Also}
princomp.

\section*{Examples}
```


## The variances of the variables in the

## USArrests data vary by orders of magnitude, so scaling is appropriate

data(USArrests)
(pc.cr <- princomp(USArrests, cor = TRUE)) \# inappropriate
screeplot(pc.cr)
data(Harman74.cor)
fit <- princomp(covmat=Harman74.cor)
screeplot(fit)
screeplot(fit, npcs=24, type="lines")

```
summary.princomp Summary method for Principal Components Analysis

\section*{Description}

The summary method for class "princomp".

\section*{Usage}
```

summary(object, loadings = FALSE, cutoff = 0.1, ...)
print(x, digits $=3$, loadings $=x \$ p r i n t . l o a d i n g s$,
cutoff = x\$cutoff, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & an object of class "princomp", as from princomp(). \\
loadings & \begin{tabular}{l} 
logical. Should loadings be included? \\
cutoff
\end{tabular} \\
\begin{tabular}{l} 
numeric. Loadings below this cutoff in absolute value are shown as blank \\
in the output.
\end{tabular} \\
x & \begin{tabular}{l} 
an object of class "summary.princomp".
\end{tabular} \\
digits & the number of significant digits to be used in listing loadings. \\
\(\ldots\) & arguments to be passed to or from other methods.
\end{tabular}

Value
object with additional components cutoff and print.loadings.

\section*{See Also}

\section*{Examples}
```

data(USArrests)
summary(pc.cr <- princomp(USArrests, cor=TRUE))
print(summary(princomp(USArrests, cor=TRUE),
loadings = TRUE, cutoff = 0.2), digits = 2)

```
varimax Rotation Methods for Factor Analysis

\section*{Description}

These functions 'rotate' loading matrices in factor analysis.

\section*{Usage}
varimax (x, normalize \(=\) TRUE, eps \(=1 \mathrm{e}-5\) )
promax (x, m = 4)

\section*{Arguments}
\(\mathrm{x} \quad\) A loadings matrix, with \(p\) rows and \(k<p\) columns
\(\mathrm{m} \quad\) The power used the target for promax. Values of 2 to 4 are recommended.
normalize logical. Should Kaiser normalization be performed? If so the rows of x are re-scaled to unit length before rotation, and scaled back afterwards.
eps The tolerance for stopping: the relative change in the sum of singular values.

\section*{Details}

These seek a 'rotation' of the factors \(\times \% * \% \mathrm{~T}\) that aims to clarify the structure of the loadings matrix. The matrix T is a rotation (possibly with reflection) for varimax, but a general linear transformation for promax, with the variance of the factors being preserved.

\section*{Value}

A list with components
loadings The 'rotated' loadings matrix, x \% \% \% rotmat.
rotmat The 'rotation matrix.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Hendrickson, A. E. and White, P. O. (1964) Promax: a quick method for rotation to orthogonal oblique structure. British Journal of Statistical Psychology, 17, 65-70.
Horst, P. (1965) Factor Analysis of Data Matrices. Holt, Rinehart and Winston. Chapter 10.

Kaiser, H. F. (1958) The varimax criterion for analytic rotation in factor analysis. Psychometrika 23, 187-200.

Lawley, D. N. and Maxwell, A. E. (1971) Factor Analysis as a Statistical Method. Second edition. Butterworths.

\section*{See Also}
```

factanal, Harman74.cor.

```

\section*{Examples}
```

data(swiss)

## varimax with normalize = T is the default

fa <- factanal( ~., 2, data = swiss)
varimax(fa$loadings, normalize = FALSE)
promax(fa$loadings)

```

\section*{Chapter 8}

\section*{The nls package}
asOneSidedFormula Convert to One-Sided Formula

\section*{Description}

Names, expressions, numeric values, and character strings are converted to one-sided formulas. If object is a formula, it must be one-sided, in which case it is returned unaltered.

\section*{Usage}
asOneSidedFormula(object)

\section*{Arguments}
object a one-sided formula, an expression, a numeric value, or a character string.

Value
a one-sided formula representing object

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
formula

\section*{Examples}
```

asOneSidedFormula("age")

```
asOneSidedFormula(~ age)

\section*{Description}

The BOD data frame has 6 rows and 2 columns giving the biochemical oxygen demand versus time in an evaluation of water quality.

\section*{Usage}
data(BOD)

\section*{Format}

This data frame contains the following columns:
Time A numeric vector giving the time of the measurement (days).
demand A numeric vector giving the biochemical oxygen demand ( \(\mathrm{mg} / \mathrm{l}\) ).

\section*{Source}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley, Appendix A1.4.

Originally from Marske (1967), Biochemical Oxygen Demand Data Interpretation Using Sum of Squares Surface M.Sc. Thesis, University of Wisconsin - Madison.

\section*{Examples}
```

data(BOD)

# simplest form of fitting a first-order model to these data

fm1 <- nls(demand ~ A*(1-exp(-exp(lrc)*Time)), data = BOD,
start = c(A = 20, lrc = log(.35)))
coef(fm1)
print(fm1)

# using the plinear algorithm

fm2 <- nls(demand ~ (1-exp(-exp(lrc)*Time)), data = BOD,
start = c(lrc = log(.35)), algorithm = "plinear", trace = TRUE)

# using a self-starting model

fm3 <- nls(demand ~ SSasympOrig(Time, A, lrc), data = BOD)
summary( fm3 )

```
ChickWeight Weight versus age of chicks on different diets

\section*{Description}

The ChickWeight data frame has 578 rows and 4 columns from an experiment on the effect of diet on early growth of chicks.

\section*{Usage}
```

data(ChickWeight)

```

\section*{Format}

This data frame contains the following columns:
weight a numeric vector giving the body weight of the chick (gm).
Time a numeric vector giving the number of days since birth when the measurement was made.

Chick an ordered factor with levels \(18<\ldots<48\) giving a unique identifier for the chick. The ordering of the levels groups chicks on the same diet together and orders them according to their final weight (lightest to heaviest) within diet.
Diet a factor with levels \(1, \ldots, 4\) indicating which experimental diet the chick received.

\section*{Details}

The body weights of the chicks were measured at birth and every second day thereafter until day 20. They were also measured on day 21. There were four groups on chicks on different protein diets.

\section*{Source}

Crowder, M. and Hand, D. (1990), Analysis of Repeated Measures, Chapman and Hall (example 5.3)
Hand, D. and Crowder, M. (1996), Practical Longitudinal Data Analysis, Chapman and Hall (table A.2)

Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in S and S-PLUS, Springer.

\section*{Examples}
```

data(ChickWeight)
coplot(weight ~ Time | Chick, data = ChickWeight,
type = "b", show = FALSE)

## fit a representative chick

fm1 <- nls(weight ~ SSlogis( Time, Asym, xmid, scal ),
data = ChickWeight, subset = Chick == 1)
summary( fm1 )

```
```

clearNames Remove the Names from an Object

```

\section*{Description}

This function sets the names attribute of object to NULL and returns the object.

\section*{Usage}
clearNames (object)

\section*{Arguments}
object an object that may have a names attribute

\section*{Value}

An object similar to object but without names.

\section*{Author(s)}

Douglas Bates and Saikat DebRoy

\section*{See Also}
setNames

\section*{Examples}
```

data( women )
lapply( women, mean ) \# has a names attribute
clearNames( lapply( women, mean ) ) \# removes the names

```

CO2
Carbon Dioxide uptake in grass plants

\section*{Description}

The CO2 data frame has 84 rows and 5 columns of data from an experiment on the cold tolerance of the grass species Echinochloa crus-galli.

\section*{Usage}
data(CO2)

\section*{Format}

This data frame contains the following columns:
Plant an ordered factor with levels \(\mathrm{Qn} 1<\mathrm{Qn} 2<\mathrm{Qn} 3<\ldots<\) Mc1 giving a unique identifier for each plant.
Type a factor with levels Quebec Mississippi giving the origin of the plant
Treatment a factor with levels nonchilled chilled
conc a numeric vector of ambient carbon dioxide concentrations ( \(\mathrm{mL} / \mathrm{L}\) ).
uptake a numeric vector of carbon dioxide uptake rates ( \(\mu \mathrm{mol} / m^{2} \mathrm{sec}\) ).

\section*{Details}

The \(\mathrm{CO}_{2}\) uptake of six plants from Quebec and six plants from Mississippi was measured at several levels of ambient \(\mathrm{CO}_{2}\) concentration. Half the plants of each type were chilled overnight before the experiment was conducted.

\section*{Source}

Potvin, C., Lechowicz, M. J. and Tardif, S. (1990) "The statistical analysis of ecophysiological response curves obtained from experiments involving repeated measures", Ecology, 71, 1389-1400.

Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in S and S-PLUS, Springer.

\section*{Examples}
```

data(CO2)
coplot(uptake ~ conc | Plant, data = CO2, show = FALSE, type = "b")

## fit the data for the first plant

fm1 <- nls(uptake ~ SSasymp(conc, Asym, lrc, c0),
data = CO2, subset = Plant == 'Qn1')
summary(fm1)

## fit each plant separately

fmlist <- list()
for (pp in levels(CO2\$Plant)) {
fmlist[[pp]] <- nls(uptake ~ SSasymp(conc, Asym, lrc, c0),
data = CO2, subset = Plant == pp)
}

## check the coefficients by plant

sapply(fmlist, coef)

```

\section*{Description}

The DNase data frame has 176 rows and 3 columns of data obtained during development of an ELISA assay for the recombinant protein DNase in rat serum.

\section*{Usage}
data(DNase)

\section*{Format}

This data frame contains the following columns:
Run an ordered factor with levels \(10<\ldots<3\) indicating the assay run.
conc a numeric vector giving the known concentration of the protein.
density a numeric vector giving the measured optical density (dimensionless) in the assay. Duplicate optical density measurements were obtained.

\section*{Source}

Davidian, M. and Giltinan, D. M. (1995) Nonlinear Models for Repeated Measurement Data, Chapman \& Hall (section 5.2.4, p. 134)
Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in S and S-PLUS, Springer.

\section*{Examples}
```

data(DNase)
coplot(density ~ conc | Run, data = DNase,
show = FALSE, type = "b")
coplot(density ~ log(conc) | Run, data = DNase,
show = FALSE, type = "b")

## fit a representative run

fm1 <- nls(density ~ SSlogis( log(conc), Asym, xmid, scal ),
data = DNase, subset = Run == 1)

```
```


## compare with a four-parameter logistic

fm2 <- nls(density ~ SSfpl( log(conc), A, B, xmid, scal ),
data = DNase, subset = Run == 1)
summary(fm2)
anova(fm1, fm2)

```
```

formula.nls Extract Model Formula from nls Object

```

\section*{Description}

Returns the model used to fit object.

\section*{Usage}
formula(x, ...)

\section*{Arguments}
x
an object inheriting from class nls, representing a nonlinear least squares fit.
... further arguments passed to or from other methods.

\section*{Value}
a formula representing the model used to obtain object.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, formula

\section*{Examples}
```

data(Orange)
fm1 <- nls(circumference ~ A/(1+exp((B-age)/C)), Orange,
start = list(A=160, B=700, C = 350))
formula(fm1)

```
```

getInitial Get Initial Parameter Estimates

```

\section*{Description}

This function evaluates initial parameter estimates for a nonlinear regression model. If data is a parameterized data frame or pframe object, its parameters attribute is returned. Otherwise the object is examined to see if it contains a call to a selfStart object whose initial attribute can be evaluated.

\section*{Usage}
getInitial(object, data, ...)

\section*{Arguments}
object a formula or a selfStart model that defines a nonlinear regression model
data a data frame in which the expressions in the formula or arguments to the selfStart model can be evaluated
... optional additional arguments

\section*{Value}

A named numeric vector or list of starting estimates for the parameters. The construction of many selfStart models is such that these "starting" estimates are, in fact, the converged parameter estimates.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

nls, selfStart, selfStart.default, selfStart.formula

```

\section*{Examples}
```

data(Puromycin)
PurTrt <- Puromycin[ Puromycin\$state == "treated", ]
getInitial( rate ~ SSmicmen( conc, Vm, K ), PurTrt )

```

Indometh Pharmacokinetics of Indomethicin

\section*{Description}

The Indometh data frame has 66 rows and 3 columns of data on the pharmacokinetics of indomethicin.

\section*{Usage}
data(Indometh)

\section*{Format}

This data frame contains the following columns:

Subject an ordered factor with containing the subject codes. The ordering is according to increasing maximum response.
time a numeric vector of times at which blood samples were drawn (hr).
conc a numeric vector of plasma concentrations of indomethicin ( \(\mathrm{mcg} / \mathrm{ml}\) ).

\section*{Details}

Each of the six subjects were given an intravenous injection of indomethicin.

\section*{Source}

Kwan, Breault, Umbenhauer, McMahon and Duggan (1976), "Kinetics of Indomethicin absorption, elimination, and enterohepatic circulation in man", Journal of Pharmacokinetics and Biopharmaceutics, 4, 255-280.

Davidian, M. and Giltinan, D. M. (1995) Nonlinear Models for Repeated Measurement Data, Chapman \& Hall (section 5.2.4, p. 134)

Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in \(S\) and S-PLUS, Springer.

\section*{Examples}
```

data(Indometh)
fm1 <- nls(conc ~ SSbiexp(time, A1, lrc1, A2, lrc2),
data = Indometh, subset = Subject == 1)
summary(fm1)

```
```

Loblolly Growth of Loblolly pine trees

```

\section*{Description}

The Loblolly data frame has 84 rows and 3 columns of records of the growth of Loblolly pine trees.

\section*{Usage}
```

data(Loblolly)

```

\section*{Format}

This data frame contains the following columns:
height a numeric vector of tree heights ( ft ).
age a numeric vector of tree ages (yr).
Seed an ordered factor indicating the seed source for the tree. The ordering is according to increasing maximum height.

\section*{Source}

Kung, F. H. (1986), "Fitting logistic growth curve with predetermined carrying capacity", Proceedings of the Statistical Computing Section, American Statistical Association, 340-343.
Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in S and S-PLUS, Springer.

\section*{Examples}
```

data(Loblolly)
plot(height ~ age, data = Loblolly, subset = Seed == 329,
xlab = "Tree age (yr)", las = 1,
ylab = "Tree height (ft)",
main = "Loblolly data and fitted curve (Seed 329 only)")
fm1 <- nls(height ~ SSasymp(age, Asym, R0, lrc),
data = Loblolly, subset = Seed == 329)
summary(fm1)
age <- seq(0, 30, len = 101)
lines(age, predict(fm1, list(age = age)))

```
nls \(\quad\) Nonlinear Least Squares

\section*{Description}

Determine the nonlinear least squares estimates of the nonlinear model parameters and return a class nls object.

\section*{Usage}
```

nls(formula, data = parent.frame(), start, control = nls.control(),
algorithm = "default", trace = FALSE, subset,
weights, na.action)

```

\section*{Arguments}
formula a nonlinear model formula including variables and parameters data an optional data frame in which to evaluate the variables in formula start a named list or named numeric vector of starting estimates control an optional list of control settings. See nls.control for the names of the settable control values and their effect.
algorithm character string specifying the algorithm to use. The default algorithm is a Gauss-Newton algorithm. The other alternative is "plinear", the GolubPereyra algorithm for partially linear least-squares models.
trace logical value indicating if a trace of the iteration progress should be printed. Default is FALSE. If TRUE the residual sum-of-squares and the parameter values are printed at the conclusion of each iteration. When the "plinear" algorithm is used, the conditional estimates of the linear parameters are printed after the nonlinear parameters.
subset an optional vector specifying a subset of observations to be used in the fitting process.
weights an optional numeric vector of (fixed) weights. When present, the objective function is weighted least squares. not yet implemented
na.action a function which indicates what should happen when the data contain NAs.

\section*{Details}

\section*{Do not use nls on artificial "zero-residual" data.}

The nls function uses a relative-offset convergence criterion that compares the numerical imprecision at the current parameter estimates to the residual sum-of-squares. This performs well on data of the form
\[
y=f(x, \theta)+\epsilon
\]
(with \(\operatorname{var}(\mathrm{eps})>0\) ). It fails to indicate convergence on data of the form
\[
y=f(x, \theta)
\]
because the criterion amounts to comparing two components of the round-off error. If you wish to test nls on artificial data please add a noise component, as shown in the example below.

An nls object is a type of fitted model object. It has methods for the generic functions coef, formula, resid, print, summary, AIC, fitted and vcov.

\section*{Value}

A list of
\[
\begin{array}{ll}
\mathrm{m} & \text { an nlsModel object incorporating the model } \\
\text { data } & \text { the expression that was passed to } n l s \text { as the data argument. The actual } \\
\text { data values are present in the environment of the } m \text { component. }
\end{array}
\]

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{References}

Bates, D.M. and Watts, D.G. (1988) Nonlinear Regression Analysis and Its Applications, Wiley

\section*{See Also}
nlsModel

\section*{Examples}
```

data( DNase )
DNase1 <- DNase[ DNase\$Run == 1, ]

## using a selfStart model

fm1DNase1 <- nls( density ~ SSlogis( log(conc), Asym, xmid, scal ), DNase1 )
summary( fm1DNase1 )

## using conditional linearity

fm2DNase1 <- nls( density ~ 1/(1 + exp(( xmid - log(conc) )/scal ) ),
data = DNase1,
start = list( xmid = 0, scal = 1 ),
alg = "plinear", trace = TRUE )
summary( fm2DNase1 )

## without conditional linearity

fm3DNase1 <- nls( density ~ Asym/(1 + exp(( xmid - log(conc) )/scal ) ),
data = DNase1,
start = list( Asym = 3, xmid = 0, scal = 1 ),
trace = TRUE )
summary( fm3DNase1 )

## weighted nonlinear regression

data(Puromycin)
Treated <- Puromycin[Puromycin\$state == "treated", ]
weighted.MM <- function(resp, conc, Vm, K)
{
\#\# Purpose: exactly as white book p.451 -- RHS for nls()
\#\# Weighted version of Michaelis-Menten model
\#\# -------------------------------------------------------------------------
\#\# Arguments: ' }\textrm{y}\mathrm{ ', ' ' }\textrm{x}\mathrm{ ' and the two parameters (see book)
\#\# ---------------------------------------------------------------------------
\#\# Author: Martin Maechler, Date: 23 Mar 2001, 18:48
pred <- (Vm * conc)/(K + conc)
(resp - pred) / sqrt(pred)
}
Pur.wt <- nls( ~ weighted.MM(rate, conc, Vm, K), data = Treated,
start = list(Vm = 200, K = 0.1),
trace = TRUE)

```

\section*{Description}

Allow the user to set some characteristics of the nls nonlinear least squares algorithm.

\section*{Usage}
```

nls.control(maxiter=50, tol=1e-05, minFactor=1/1024)

```

\section*{Arguments}
maxiter A positive integer specifying the maximum number of iterations allowed.
tol A positive numeric value specifying the tolerance level for the relative offset convergence criterion.
minFactor A positive numeric value specifying the minimum step-size factor allowed on any step in the iteration. The increment is calculated with a Gauss-Newton algorithm and successively halved until the residual sum of squares has been decreased or until the step-size factor has been reduced below this limit.

\section*{Value}

A list with exactly three components:
maxiter
tol
minFactor

\section*{Author(s)}

Douglas Bates and Saikat DebRoy

\section*{References}

Bates and Watts (1988), Nonlinear Regression Analysis and Its Applications, Wiley.

\section*{See Also}
nls

\section*{Examples}
```

nlsModel Create an nlsModel Object

```

\section*{Description}

This is the constructor for nlsModel objects, which are function closures for several functions in a list. The closure includes a nonlinear model formula, data values for the formula, as well as parameters and their values.

\section*{Usage}
nlsModel(form, data, start)

\section*{Arguments}
\begin{tabular}{ll} 
form & a nonlinear model formula \\
data & \begin{tabular}{l} 
a data frame or a list in which to evaluate the variables from the model \\
formula
\end{tabular} \\
start & \begin{tabular}{l} 
a named list or named numeric vector of starting estimates for the pa- \\
rameters in the model
\end{tabular}
\end{tabular}

\section*{Details}

An nlsModel object is primarily used within the nls function. It encapsulates the model, the data, and the parameters in an environment and provides several methods to access characteristics of the model. It forms an important component of the object returned by the nls function.

\section*{Value}

The value is a list of functions that share a common environment.
\begin{tabular}{ll}
\begin{tabular}{l} 
resid \\
fitted
\end{tabular} & \begin{tabular}{l} 
returns the residual vector evaluated at the current parameter values \\
returns the fitted responses and their gradient at the current parameter \\
values
\end{tabular} \\
formula & \begin{tabular}{l} 
returns the model formula
\end{tabular} \\
deviance & \begin{tabular}{l} 
returns the residual sum-of-squares at the current parameter values
\end{tabular} \\
gradient & \begin{tabular}{l} 
returns the gradient of the model function at the current parameter values \\
conv
\end{tabular} \\
returns the relative-offset convergence criterion evaluated at the current \\
parmeter values \\
incr & \begin{tabular}{l} 
returns the parameter increment calculated according to the Gauss- \\
Newton formula
\end{tabular} \\
setPars & \begin{tabular}{l} 
a function with one argument, pars. It sets the parameter values for the \\
nlsModel object and returns a logical value denoting a singular gradient \\
array.
\end{tabular} \\
getPars & \begin{tabular}{l} 
returns the current value of the model parameters as a numeric vector
\end{tabular} \\
getAllPars & \begin{tabular}{l} 
returns the current value of the model parameters as a numeric vector
\end{tabular} \\
getEnv & returns the environment shared by these functions
\end{tabular}
trace the function that is called at each iteration if tracing is enabled
Rmat the upper triangular factor of the gradient array at the current parameter values
predict takes as argument newdata, a data.frame and returns the predicted response for newdata.

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{References}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley

\section*{See Also}
nls

\section*{Examples}
```

data( DNase )
DNase1 <- DNase[ DNase$Run == 1, ]
mod <-
    nlsModel(density ~ SSlogis( log(conc), Asym, xmid, scal ),
        DNase1, list( Asym = 3, xmid = 0, scal = 1 ))
mod$getPars() \# returns the parameters as a list
mod$deviance() # returns the residual sum-of-squares
mod$resid() \# returns the residual vector and the gradient
mod$incr() # returns the suggested increment
mod$setPars( unlist(mod$getPars()) + mod$incr() ) \# set new parameter values
mod$getPars() # check the parameters have changed
mod$deviance() \# see if the parameter increment was successful
mod$trace() # check the tracing
mod$Rmat() \# R matrix from the QR decomposition of the gradient

```

NLSstAsymptotic Fit the Asymptotic Regression Model

\section*{Description}

Fits the asymptotic regression model, in the form b0 \(+\mathrm{b} 1 *(1-\exp (-\exp (\operatorname{lrc}) * \mathrm{x})\) to the xy data. This can be used as a building block in determining starting estimates for more complicated models.

\section*{Usage}

NLSstAsymptotic (xy)

\section*{Arguments}

\section*{Value}

A numeric value of length 3 with components labelled b0, b1, and lrc. b0 is the estimated intercept on the y -axis, b 1 is the estimated difference between the asymptote and the y intercept, and lrc is the estimated logarithm of the rate constant.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}

SSasymp

\section*{Examples}
```

data( Loblolly )
Lob.329 <- Loblolly[ Loblolly\$Seed == "329", ]
NLSstAsymptotic(sortedXyData(expression(age), expression(height), Lob.329 ))

```
```

NLSstClosestX Inverse Interpolation

```

\section*{Description}

Use inverse linear interpolation to approximate the x value at which the function represented by xy is equal to yval.

\section*{Usage}

NLSstClosestX(xy, yval)

\section*{Arguments}
xy a sortedXyData object
yval a numeric value on the y scale

\section*{Value}

A single numeric value on the x scale.
Author(s)
Jose Pinheiro and Douglas Bates

\section*{See Also}
sortedXyData, NLSstLfAsymptote, NLSstRtAsymptote, selfStart

\section*{Examples}
```

data( DNase )
DNase.2 <- DNase[ DNase\$Run == "2", ]
DN.srt <- sortedXyData( expression(log(conc)), expression(density), DNase.2 )
NLSstClosestX( DN.srt, 1.0 )

```

\section*{Description}

Provide an initial guess at the horizontal asymptote on the left side (i.e. small values of x ) of the graph of y versus x from the xy object. Primarily used within initial functions for self-starting nonlinear regression models.

\section*{Usage}
```

NLSstLfAsymptote(xy)

```

\section*{Arguments}
xy a sortedXyData object

\section*{Value}

A single numeric value estimating the horizontal asymptote for small x .

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
sortedXyData, NLSstClosestX, NLSstRtAsymptote, selfStart

\section*{Examples}
```

data( DNase )
DNase.2 <- DNase[ DNase\$Run == "2", ]
DN.srt <- sortedXyData( expression(log(conc)), expression(density), DNase.2 )
NLSstLfAsymptote( DN.srt )

```

NLSstRtAsymptote Horizontal Asymptote on the Right Side

\section*{Description}

Provide an initial guess at the horizontal asymptote on the right side (i.e. large values of x ) of the graph of y versus x from the xy object. Primarily used within initial functions for self-starting nonlinear regression models.

\section*{Usage}

NLSstRtAsymptote(xy)

\section*{Arguments}

\section*{Value}

A single numeric value estimating the horizontal asymptote for large x .

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

sortedXyData, NLSstClosestX, NLSstRtAsymptote, selfStart

```

\section*{Examples}
```

data( DNase )
DNase.2 <- DNase[ DNase\$Run == "2", ]
DN.srt <- sortedXyData( expression(log(conc)), expression(density), DNase.2 )
NLSstRtAsymptote( DN.srt )

```

\section*{numericDeriv Evaluate derivatives numerically}

\section*{Description}
numericDeriv numerically evaluates the gradient of an expression.

\section*{Usage}
numericDeriv(expr, theta, rho=parent.frame())

\section*{Arguments}
expr The expression to be differentiated. The value of this expression should be a numeric vector.
theta A character vector of names of variables used in expr
rho An environment containing all the variables needed to evaluate expr

\section*{Details}

This is a front end to the C function numeric_deriv, which is described in Writing \(R\) Extensions.

\section*{Value}

The value of eval (expr, env \(=\) rho) plus a matrix attribute called gradient. The columns of this matrix are the derivatives of the value with respect to the variables listed in theta.

Author (s)
Saikat DebRoy 〈saikat@stat.wisc.edu〉

\section*{Examples}
```

myenv <- new.env()
assign("mean", 0., env = myenv)
assign("sd", 1., env = myenv)
assign("x", seq(-3., 3., len = 31), env = myenv)
numericDeriv(quote(pnorm(x, mean, sd)), c("mean", "sd"), myenv)

```
Orange Growth of orange trees

\section*{Description}

The Orange data frame has 35 rows and 3 columns of records of the growth of orange trees.

\section*{Usage}
data(Orange)

\section*{Format}

This data frame contains the following columns:
Tree an ordered factor indicating the tree on which the measurement is made. The ordering is according to increasing maximum diameter.
age a numeric vector giving the age of the tree (days since 1968/12/31)
circumference a numeric vector of trunk circumferences (mm). This is probably "circumference at breast height", a standard measurement in forestry.

\section*{Source}

Draper, N. R. and Smith, H. (1998), Applied Regression Analysis (3rd ed), Wiley (exercise 24.N).

Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in \(S\) and S-PLUS, Springer.

\section*{Examples}
```

data(Orange)
coplot(circumference ~ age | Tree, data = Orange, show = FALSE)
fm1 <- nls(circumference ~ SSlogis(age, Asym, xmid, scal),
data = Orange, subset = Tree == 3)
plot(circumference ~ age, data = Orange, subset = Tree == 3,
xlab = "Tree age (days since 1968/12/31)",
ylab = "Tree circumference (mm)", las = 1,
main = "Orange tree data and fitted model (Tree 3 only)")
age <- seq(0, 1600, len = 101)
lines(age, predict(fm1, list(age = age)))

```
```

plot.profile.nls Plot a profile.nls Object

```

\section*{Description}

Displays a series of plots of the profile \(t\) function and interpolated confidence intervals for the parameters in a nonlinear regression model that has been fit with nls and profiled with profile.nls.

\section*{Usage}
plot(x, levels, conf= c(99, 95, 90, 80, 50)/100, nseg \(=50\), absVal =TRUE, ...)

\section*{Arguments}
\begin{tabular}{ll}
x & an object of class "profile.nls" \\
levels & \begin{tabular}{l} 
levels, on the scale of the absolute value of a t statistic, at which to inter- \\
polate intervals. Usually conf is used instead of giving levels explicitly.
\end{tabular} \\
conf & \begin{tabular}{l} 
a numeric vector of confidence levels for profile-based confidence intervals \\
on the parameters. Defaults to \(c(0.99,0.95,0.90,0.80,0.50)\).
\end{tabular} \\
nseg & \begin{tabular}{l} 
an integer value giving the number of segments to use in the spline inter- \\
polation of the profile t curves. Defaults to 50.
\end{tabular} \\
absVal & \begin{tabular}{l} 
a logical value indicating whether or not the plots should be on the scale \\
of the absolute value of the profile \(t . ~ D e f a u l t s ~ t o ~ T R U E . ~\)
\end{tabular} \\
\(\ldots\) & \begin{tabular}{l} 
other arguments to the plot function can be passed here.
\end{tabular}
\end{tabular}

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{References}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley (chapter 6)

\section*{See Also}
nls, profile, profile.nls

\section*{Examples}
```

data( BOD )

# obtain the fitted object

fm1 <- nls(demand ~ SSasympOrig( Time, A, lrc ), data = BOD)

# get the profile for the fitted model

pr1 <- profile( fm1 )
opar <- par(mfrow = c(2,2), oma = c(1.1, 0, 1.1, 0), las = 1)
plot(pr1, conf = c(95, 90, 80, 50)/100)
plot(pr1, conf = c(95, 90, 80, 50)/100, absVal = FALSE)
mtext("Confidence intervals based on the profile sum of squares",
side = 3, outer = TRUE)

```
```

mtext("BOD data - confidence levels of 50%, 80%, 90% and 95%",
side = 1, outer = TRUE)
par(opar)

```
predict.nls Predicting from Nonlinear Least Squares Fits

\section*{Description}
predict.nls produces predicted values, obtained by evaluating the regression function in the frame newdata. If the logical se.fit is TRUE, standard errors of the predictions are calculated. If the numeric argument scale is set (with optional df), it is used as the residual standard deviation in the computation of the standard errors, otherwise this is extracted from the model fit. Setting intervals specifies computation of confidence or prediction (tolerance) intervals at the specified level.

At present se.fit and interval are ignored.
```

Usage
predict(object, newdata, se.fit = FALSE, scale = NULL, df = Inf,
interval = c("none", "confidence", "prediction"),
level = 0.95, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & An object that inherits from class nls. \\
newdata & \begin{tabular}{l} 
A named list or data frame with values of the input variables for the \\
model in object. If newdata is missing the fitted values at the original \\
data points are returned.
\end{tabular} \\
se.fit & \begin{tabular}{l} 
A logical value indicating if the standard errors of the predictions should \\
be calculated. Defaults to FALSE. At present this argument is ignored.
\end{tabular} \\
scale & \begin{tabular}{l} 
A numeric scalar. If it is set (with optional df), it is used as the residual \\
standard deviation in the computation of the standard errors, otherwise \\
this information is extracted from the model fit. At present this argument \\
is ignored.
\end{tabular} \\
df & \begin{tabular}{l} 
A positive numeric scalar giving the number of degrees of freedom for the \\
scale estimate. At present this argument is ignored.
\end{tabular} \\
interval & \begin{tabular}{l} 
A character string indicating if prediction intervals or a confidence interval \\
on the mean responses are to be calculated. At present this argument is \\
ignored.
\end{tabular} \\
level & \begin{tabular}{l} 
A numeric scalar between 0 and 1 giving the confidence level for the \\
intervals (if any) to be calculated. At present this argument is ignored.
\end{tabular} \\
\(\ldots\) & \begin{tabular}{l} 
Additional optional arguments. At present no optional arguments are \\
used.
\end{tabular}
\end{tabular}

\section*{Value}
predict.nls produces a vector of predictions or a matrix of predictions and bounds with column names fit, lwr, and upr if interval is set. If se.fit is TRUE, a list with the following components is returned:
\begin{tabular}{ll} 
fit & vector or matrix as above \\
se.fit & standard error of predictions \\
residual.scale
\end{tabular}

\section*{See Also}

The model fitting function nls, predict.

\section*{Examples}
```

data( BOD )
fm <- nls(demand ~ SSasympOrig(Time, A, lrc), data = BOD)
predict(fm) \# fitted values at observed times

## Form data plot and smooth line for the predictions

opar <- par(las = 1)
plot(demand ~ Time, data = BOD, col = 4,
main = "BOD data and fitted first-order curve",
xlim = c(0,7), ylim = c(0, 20) )
tt <- seq(0, 8, length = 101)
lines(tt, predict(fm, list(Time = tt)))
par(opar)

```
profile.nls Method for Profiling nls Objects

\section*{Description}

Investigates behavior of the log-likelihood function near the solution represented by fitted.

\section*{Usage}
```

profile(fitted, which=1:npar, maxpts=100, alphamax=0.01,
delta.t=cutoff/5, ...)

```

\section*{Arguments}
fitted the original fitted model object.
which the original model parameters which should be profiled. By default, all parameters are profiled.
maxpts maximum number of points to be used for profiling each parameter.
alphamax maximum significance level allowed for the profile t-statistics.
delta.t suggested change on the scale of the profile \(t\)-statistics. Default value chosen to allow profiling at about 10 parameter values.
... further arguments passed to or from other methods.

\section*{Details}

The profile t-statistics is defined as the square root of change in sum-of-squares divided by residual standard error with an appropriate sign.

\section*{Value}

A list with an element for each parameter being profiled. The elements are data-frames with two variables par.vals a matrix of parameter values for each fitted model.
tau The profile t-statistics.

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{References}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley (chapter 6)

\section*{See Also}
nls, profile, profiler.nls, plot.profile.nls

\section*{Examples}
```

data( BOD )

# obtain the fitted object

fm1 <- nls(demand ~ SSasympOrig( Time, A, lrc ), data = BOD)

# get the profile for the fitted model

pr1 <- profile( fm1 )

# profiled values for the two parameters

pr1$A
pr1$lrc

```
```

profiler Constructor for Profiler Objects for Nonlinear Models

```

\section*{Description}

Create a profiler object for the model object fitted.

\section*{Usage}
profiler(fitted, ...)

\section*{Arguments}
\begin{tabular}{ll} 
fitted & the original fitted model object. \\
\(\ldots\) & Additional parameters. See documentation on individual methods.
\end{tabular}

\section*{Value}

An object of class "profiler" which is a list with function elements
getFittedPars()
the parameters in fitted
setDefault(varying, params)
this is used for changing the default settings for profiling. In absence of both parameters, the default is set to the original fitted parameters with all parameters varying. The arguments are
varying: a logical, integer or character vector giving parameters to be varied. params: the default value at which profiling is to take place.
getProfile(varying, params)
this can be used in conjunction with setDefault without any arguments.
Alternatively, the parameters to be varied and the values for fixed parameters can be specified using the arguments. The arguments are
varying: a logical vector giving parameters to be varied. This can be omitted if params is a named list or numeric vector.
params: values for parameters to be held fixed.
It returns a list with elements
parameters: the parameter values for the profiled optimum.
fstat: a profile statistics. See individual methods for details.
varying: a logical vector indicating parameters which were varied.

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{See Also}
profiler.nls, profile

\section*{Examples}
\# see documentation on individual methods
profiler.nls Constructor for Profiler Objects from nls Objects

\section*{Description}

Create a profiler object for the model object fitted of class nls.
```

Usage
profiler(fitted, ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
fitted & the original fitted model object of class nls. \\
\(\ldots\) & Additional parameters. None are used.
\end{tabular}

\section*{Value}

An object of class profiler.nls which is a list with function elements
getFittedModel()
the nlsModel object corresponding to fitted
getFittedPars()
See documentation for profiler
setDefault(varying, params)
See documentation for profiler
getProfile(varying, params)
In the returned list, fstat is the ratio of change in sum-of-squares and the residual standard error.
For other details, see documentation for profiler

\section*{WARNING}

When using setDefault and getProfile together, the internal state of the fitted model may get changed. So after completing the profiling for a parameter, the internal states should be restored by a call to setDefault without any arguments. For example see below or the source for profile.nls.

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{References}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley

\section*{See Also}
```

nls, nlsModel, profiler, profile.nls

```

\section*{Examples}
```

data( BOD )

## obtain the fitted object

fm1 <- nls(demand ~ SSasympOrig( Time, A, lrc ), data = BOD)

## get the profile for the fitted model

prof1 <- profiler( fm1 )

## profile with A fixed at 16.0

prof1\$getProfile(c(FALSE, TRUE), 16.0)

## vary lrc

prof1\$setDefault(varying = c(FALSE, TRUE))

## fix A at 14.0 and starting estimate of lrc at -0.2

prof1\$setDefault(params = c(14.0, -0.2))

## and get the profile

prof1\$getProfile()

## finally, set defaults back to original estimates

prof1\$setDefault()

```

\section*{Puromycin Reaction velocity of an enzymatic reaction}

\section*{Description}

The Puromycin data frame has 23 rows and 3 columns of the reaction velocity versus substrate concentration in an enzymatic reaction involving untreated cells or cells treated with Puromycin.

\section*{Usage}
data(Puromycin)

\section*{Format}

This data frame contains the following columns:
conc a numeric vector of substrate concentrations (ppm)
rate a numeric vector of instantaneous reaction rates (counts \(/ \mathrm{min} / \mathrm{min}\) )
state a factor with levels treated untreated

\section*{Details}

Data on the "velocity" of an enzymatic reaction were obtained by Treloar (1974). The number of counts per minute of radioactive product from the reaction was measured as a function of substrate concentration in parts per million ( ppm ) and from these counts the initial rate, or "velocity," of the reaction was calculated (counts \(/ \mathrm{min} / \mathrm{min}\) ). The experiment was conducted once with the enzyme treated with Puromycin, and once with the enzyme untreated.

\section*{Source}

Bates, D.M. and Watts, D.G. (1988), Nonlinear Regression Analysis and Its Applications, Wiley, Appendix A1.3.
Treloar, M. A. (1974), Effects of Puromycin on Galactosyltransferase in Golgi Membranes, M.Sc. Thesis, U. of Toronto.

\section*{Examples}
```

data(Puromycin)
plot(rate ~ conc, data = Puromycin, las = 1,
xlab = "Substrate concentration (ppm)",
ylab = "Reaction velocity (counts/min/min)",
pch = as.integer(Puromycin$state),
    col = as.integer(Puromycin$state),
main = "Puromycin data and fitted Michaelis-Menten curves")

## simplest form of fitting the Michaelis-Menten model to these data

fm1 <- nls(rate ~ Vm * conc/(K + conc), data = Puromycin,
subset = state == "treated",
start = c(Vm = 200, K = 0.05), trace = TRUE)
fm2 <- nls(rate ~ Vm * conc/(K + conc), data = Puromycin,
subset = state == "untreated",
start = c(Vm = 160, K = 0.05), trace = TRUE)

```
```

summary(fm1)
summary(fm2)

## using partial linearity

fm3 <- nls(rate ~ conc/(K + conc), data = Puromycin,
subset = state == "treated", start = c(K = 0.05),
algorithm = "plinear", trace = TRUE)

## using a self-starting model

fm4 <- nls(rate ~ SSmicmen(conc, Vm, K), data = Puromycin,
subset = state == "treated")
summary(fm4)

## add fitted lines to the plot

conc <- seq(0, 1.2, len = 101)
lines(conc, predict(fm1, list(conc = conc)), lty = 1, col = 1)
lines(conc, predict(fm2, list(conc = conc)), lty = 2, col = 2)
legend(0.8, 120, levels(Puromycin\$state),
col = 1:2, lty = 1:2, pch = 1:2)

```
selfStart Construct Self-starting Nonlinear Models

\section*{Description}

This function is generic; methods functions can be written to handle specific classes of objects. Available methods include selfStart.default and selfStart.formula. See the documentation on the appropriate method function.

\section*{Usage}
selfStart(model, initial, parameters, template)

\section*{Arguments}
model a function object defining a nonlinear model.
initial a function object, taking three arguments: mCall, data, and LHS, representing, respectively, a matched call to the function model, a data frame in which to interpret the variables in mCall, and the expression from the left-hand side of the model formula in the call to nls. This function should return initial values for the parameters in model.

\section*{parameters, template}
arguments used by some methods..

\section*{Value}
a function object of the selfStart class.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

selfStart.default, selfStart.formula

```

\section*{Examples}
\#\# see documentation for the methods
selfStart.default Construct Self-starting Nonlinear Models

\section*{Description}

A method for the generic function 'selfStart' for formula objects.

\section*{Usage}
selfStart(model, initial, parameters, template)

\section*{Arguments}
\begin{tabular}{ll} 
model & a function object defining a nonlinear model. \\
initial & \begin{tabular}{l} 
a function object, taking three arguments: mCall, data, and LHS, repre- \\
senting, respectively, a matched call to the function model, a data frame \\
in which to interpret the variables in mCall, and the expression from the
\end{tabular} \\
left-hand side of the model formula in the call to nls. This function \\
should return initial values for the parameters in model.
\end{tabular}

\section*{Value}
a function object of class selfStart, corresponding to a self-starting nonlinear model function. An initial attribute (defined by the initial argument) is added to the function to calculate starting estimates for the parameters in the model automatically.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
selfStart.formula

\section*{Examples}
```


# 'first.order.log.model' is a function object defining a first order

# compartment model

# 'first.order.log.initial' is a function object which calculates initial

# values for the parameters in 'first.order.log.model'

# self-starting first order compartment model

SSfol <- selfStart(first.order.log.model, first.order.log.initial)

```

\section*{Description}

A method for the generic function 'selfStart' for formula objects.

\section*{Usage}
```

selfStart(model, initial, parameters, template)

```

\section*{Arguments}
\begin{tabular}{ll} 
model & a nonlinear formula object of the form ~expression. \\
initial & \begin{tabular}{l} 
a function object, taking three arguments: mCall, data, and LHS, repre- \\
senting, respectively, a matched call to the function model, a data frame \\
in which to interpret the variables in mCall, and the expression from the \\
left-hand side of the model formula in the call to nls. This function \\
should return initial values for the parameters in model.
\end{tabular} \\
parameters & \begin{tabular}{l} 
a character vector specifying the terms on the right hand side of model \\
for which initial estimates should be calculated. Passed as the namevec \\
argument to the deriv function.
\end{tabular} \\
template & \begin{tabular}{l} 
an optional prototype for the calling sequence of the returned object, \\
passed as the function.arg argument to the deriv function. By default, \\
a template is generated with the covariates in model coming first and the \\
parameters in model coming last in the calling sequence.
\end{tabular}
\end{tabular}

\section*{Value}
a function object of class selfStart, obtained by applying deriv to the right hand side of the model formula. An initial attribute (defined by the initial argument) is added to the function to calculate starting estimates for the parameters in the model automatically.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also} selfStart.default, deriv

\section*{Examples}
```


## self-starting logistic model

SSlogis <- selfStart(~ Asym/(1 + exp((xmid - x)/scal)),
function(mCall, data, LHS)
{
xy <- sortedXyData(mCall[["x"]], LHS, data)
if(nrow(xy) < 4) {
stop("Too few distinct x values to fit a logistic")
}

```
```

    z <- xy[["y"]]
    if (min(z) <= 0) { z <- z + 0.05 * max(z) } # avoid zeroes
    z<- z/(1.05 * max(z)) # scale to within unit height
    xy[["z"]] <- log(z/(1 - z)) # logit transformation
    aux <- coef(lm(x ~ z, xy))
    parameters(xy) <- list(xmid = aux[1], scal = aux[2])
    pars <- as.vector(coef(nls(y ~ 1/(1 + exp((xmid - x)/scal)),
        data = xy, algorithm = "plinear")))
    value <- c(pars[3], pars[1], pars[2])
    names(value) <- mCall[c("Asym", "xmid", "scal")]
    value
    }, c("Asym", "xmid", "scal"))

```
```

setNames
Set the Names in an Object

```

\section*{Description}

This is a convenience function that sets the names on an object and returns the object. It is most useful at the end of a function definition where one is creating the object to be returned and would prefer not to store it under a name just so the names can be assigned.

\section*{Usage}
setNames(object, nm)

\section*{Arguments}
\begin{tabular}{ll} 
object & an object for which a names attribute will be meaningful \\
nm & a character vector of names to assign to the object
\end{tabular}

\section*{Value}

An object of the same sort as object with the new names assigned.

\section*{Author(s)}

Douglas M. Bates and Saikat DebRoy

\section*{See Also}
clearNames

\section*{Examples}
```

setNames( 1:3, c("foo", "bar", "baz") )

# this is just a short form of

tmp <- 1:3
names(tmp) <- c("foo", "bar", "baz")
tmp

```
```

sortedXyData Create a sortedXyData object

```

\section*{Description}

This is a constructor function for the class of sortedXyData objects. These objects are mostly used in the initial function for a self-starting nonlinear regression model, which will be of the selfStart class.

\section*{Usage}
sortedXyData(x, y, data)

\section*{Arguments}
\(\mathrm{x} \quad\) a numeric vector or an expression that will evaluate in data to a numeric vector
y
a numeric vector or an expression that will evaluate in data to a numeric vector
data an optional data frame in which to evaluate expressions for x and y , if they are given as expressions

\section*{Value}

A sortedXyData object. This is a data frame with exactly two numeric columns, named \(x\) and y . The rows are sorted so the x column is in increasing order. Duplicate x values are eliminated by averaging the corresponding y values.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

selfStart, NLSstClosestX, NLSstLfAsymptote, NLSstRtAsymptote

```

\section*{Examples}
```

data( DNase )
DNase.2 <- DNase[ DNase\$Run == "2", ]
sortedXyData( expression(log(conc)), expression(density), DNase.2 )

```

\section*{SSasymp Asymptotic Regression Model}

\section*{Description}

This selfStart model evaluates the asymptotic regression function and its gradient. It has an initial attribute that will evaluate initial estimates of the parameters Asym, RO, and lrc for a given set of data.

\section*{Usage}

SSasymp(input, Asym, RO, lrc)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
Asym a numeric parameter representing the horizontal asymptote on the right side (very large values of input).
R0 a numeric parameter representing the response when input is zero.
\(\operatorname{lrc} \quad\) a numeric parameter representing the natural logarithm of the rate constant.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Asym+(ROAsym) \(* \exp (-\exp (\operatorname{lrc}) * i n p u t)\). If all of the arguments Asym, RO, and lrc are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

nls, selfStart

```

\section*{Examples}
```

data( Loblolly )
Lob.329 <- Loblolly[ Loblolly$Seed == "329", ]
SSasymp( Lob.329$age, 100, -8.5, -3.2 ) \# response only
Asym <- 100
resp0 <- -8.5
lrc <- -3.2
SSasymp( Lob.329\$age, Asym, resp0, lrc ) \# response and gradient
getInitial(height ~ SSasymp( age, Asym, resp0, lrc), data = Lob.329)

## Initial values are in fact the converged values

fm1 <- nls(height ~ SSasymp( age, Asym, resp0, lrc), data = Lob.329)
summary(fm1)

```

\section*{Description}

This selfStart model evaluates an alternative parameterization of the asymptotic regression function and the gradient with respect to those parameters. It has an initial attribute that creates initial estimates of the parameters Asym, 1 rc , and c 0 .

\section*{Usage}

SSasympOff(input, Asym, lrc, c0)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
Asym a numeric parameter representing the horizontal asymptote on the right side (very large values of input).
\(\operatorname{lrc} \quad\) a numeric parameter representing the natural logarithm of the rate constant.
c0 a numeric parameter representing the input for which the response is zero.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Asym*(1 \(\exp (-\exp (\operatorname{lrc}) *(\) input \(-c 0))\) ). If all of the arguments Asym, lrc, and c0 are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data( CO2 )
CO2.Qn1 <- CO2[CO2$Plant == "Qn1", ]
SSasympOff( CO2.Qn1$conc, 32, -4, 43 ) \# response only
Asym <- 32; lrc <- -4; c0 <- 43
SSasympOff( CO2.Qn1\$conc, Asym, lrc, c0 ) \# response and gradient
getInitial(uptake ~ SSasymp( conc, Asym, lrc, c0), data = C02.Qn1)

## Initial values are in fact the converged values

fm1 <- nls(uptake ~ SSasymp( conc, Asym, lrc, c0), data = CO2.Qn1)
summary(fm1)

```

SSasympOrig Asymptotic Regression Model through the Origin

\section*{Description}

This selfStart model evaluates the asymptotic regression function through the origin and its gradient. It has an initial attribute that will evaluate initial estimates of the parameters Asym and lrc for a given set of data.

\section*{Usage}

SSasympOrig(input, Asym, lrc)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
Asym a numeric parameter representing the horizontal asymptote.
\(\operatorname{lrc} \quad\) a numeric parameter representing the natural logarithm of the rate constant.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Asym*(1 \(\exp (-\exp (\operatorname{lrc}) * i n p u t))\). If all of the arguments Asym and lrc are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data( Loblolly )
Lob. }329\mathrm{ <- Loblolly[ Loblolly$Seed == "329", ]
SSasympOrig( Lob.329$age, 100, -3.2 ) \# response only
Asym <- 100; lrc <- -3.2
SSasympOrig( Lob.329\$age, Asym, lrc ) \# response and gradient
getInitial(height ~ SSasympOrig(age, Asym, lrc), data = Lob.329)

## Initial values are in fact the converged values

fm1 <- nls(height ~ SSasympOrig( age, Asym, lrc), data = Lob.329)
summary(fm1)

```

\section*{SSbiexp Biexponential model}

\section*{Description}

This selfStart model evaluates the biexponential model function and its gradient. It has an initial attribute that creates initial estimates of the parameters A1, \(\operatorname{lrc} 1\), A2, and lrc2.

\section*{Usage}

SSbiexp(input, A1, lrc1, A2, lrc2)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
A1 a numeric parameter representing the multiplier of the first exponential.
lrc1 a numeric parameter representing the natural logarithm of the rate constant of the first exponential.
A2 a numeric parameter representing the multiplier of the second exponential.
lrc2 a numeric parameter representing the natural logarithm of the rate constant of the second exponential.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression \(\mathrm{A} 1 * \exp (-\) \(\exp (\operatorname{lrc} 1) *\) input \()+A 2 * \exp (-\exp (\operatorname{lrc} 2) *\) input \()\). If all of the arguments A1, \(\operatorname{lrc} 1, A 2\), and lrc2 are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data( Indometh )
Indo.1 <- Indometh[Indometh$Subject == 1, ]
SSbiexp( Indo.1$time, 3, 1, 0.6, -1.3 ) \# response only
A1 <- 3; lrc1 <- 1; A2 <- 0.6; lrc2 <- -1.3
SSbiexp( Indo.1\$time, A1, lrc1, A2, lrc2 ) \# response and gradient
getInitial(conc ~ SSbiexp(time, A1, lrc1, A2, lrc2), data = Indo.1)

## Initial values are in fact the converged values

fm1 <- nls(conc ~ SSbiexp(time, A1, lrc1, A2, lrc2), data = Indo.1)
summary(fm1)

```

\section*{SSfol First-order Compartment Model}

\section*{Description}

This selfStart model evaluates the first-order compartment function and its gradient. It has an initial attribute that creates initial estimates of the parameters \(1 \mathrm{Ke}, 1 \mathrm{Ka}\), and 1 Cl .

\section*{Usage}

SSfol(Dose, input, \(1 \mathrm{Ke}, 1 \mathrm{Ka}, 1 \mathrm{Cl})\)

\section*{Arguments}

Dose a numeric value representing the initial dose.
input a numeric vector at which to evaluate the model.
1Ke a numeric parameter representing the natural logarithm of the elimination rate constant.

1Ka a numeric parameter representing the natural logarithm of the absorption rate constant.

1 Cl a numeric parameter representing the natural logarithm of the clearance.

\section*{Value}
a numeric vector of the same length as input, which is the value of the expression Dose \(* \exp (1 \mathrm{Ke}+1 \mathrm{Ka}-1 \mathrm{Cl}) *(\exp (-\exp (1 \mathrm{Ke}) * i n p u t)-\exp (-\exp (1 \mathrm{Ka}) * i n p u t)) /\) ( \(\exp (1 \mathrm{Ka})-\exp (1 \mathrm{Ke}))\).

If all of the arguments \(1 \mathrm{Ke}, 1 \mathrm{Ka}\), and 1 Cl are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data( Theoph )
Theoph.1 <- Theoph[ Theoph$Subject == 1, ]
SSfol( Theoph.1$Dose, Theoph.1$Time, -2.5, 0.5, -3 ) # response only
1Ke <- -2.5; 1Ка <- 0.5; 1Cl <- -3
SSfol( Theoph.1$Dose, Theoph.1\$Time, lKe, lKa, lCl ) \# response and gradient
getInitial(conc ~ SSfol(Dose, Time, 1Ke, lKa, 1Cl), data = Theoph.1)

## Initial values are in fact the converged values

fm1 <- nls(conc ~ SSfol(Dose, Time, lKe, lKa, lCl), data = Theoph.1)
summary(fm1)

```

\section*{SSfpl}

\section*{Description}

This selfStart model evaluates the four-parameter logistic function and its gradient. It has an initial attribute that will evaluate initial estimates of the parameters A, B, xmid, and scal for a given set of data.

\section*{Usage}

SSfpl(input, A, B, xmid, scal)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
A a numeric parameter representing the horizontal asymptote on the left side (very small values of input).

B a numeric parameter representing the horizontal asymptote on the right side (very large values of input).
xmid a numeric parameter representing the input value at the inflection point of the curve. The value of SSfpl will be midway between A and B at xmid.
scal a numeric scale parameter on the input axis.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression \(A+(B-\) A) \(/(1+\exp ((x m i d-i n p u t) / s c a l))\). If all of the arguments A, B, xmid, and scal are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

nls, selfStart

```

\section*{Examples}
```

data( ChickWeight )
Chick.1 <- ChickWeight[ChickWeight$Chick == 1, ]
SSfpl( Chick.1$Time, 13, 368, 14, 6 ) \# response only
A <- 13; B <- 368; xmid <- 14; scal <- 6
SSfpl( Chick.1\$Time, A, B, xmid, scal ) \# response and gradient
getInitial(weight ~ SSfpl(Time, A, B, xmid, scal), data = Chick.1)

## Initial values are in fact the converged values

fm1 <- nls(weight ~ SSfpl(Time, A, B, xmid, scal), data = Chick.1)
summary(fm1)

```

\section*{Description}

This selfStart model evaluates the Gompertz growth model and its gradient. It has an initial attribute that creates initial estimates of the parameters Asym, b2, and b3.

\section*{Usage}

SSgompertz(x, Asym, b2, b3)

\section*{Arguments}
x

Asym a numeric parameter representing the asymptote.
b2 a numeric parameter related to the value of the function at \(\mathrm{x}=0\)
b3 a numeric parameter related to the scale the x axis.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Asym*exp (\(\mathrm{b} 2 * \mathrm{~b} 3^{\wedge} \mathrm{x}\) ). If all of the arguments Asym, b2, and b3 are names of objects the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data(DNase )
DNase.1 <- subset(DNase, Run == 1)
SSlogis(log(DNase.1$conc), 4.5, 2.3, 0.7) # response only
Asym <- 4.5; b2 <- 2.3; b3 <- 0.7
SSgompertz(log(DNase.1$conc), Asym, b2, b3 ) \# response and gradient
getInitial(density ~ SSgompertz(log(conc), Asym, b2, b3),
data = DNase.1)

## Initial values are in fact the converged values

fm1 <- nls(density ~ SSgompertz(log(conc), Asym, b2, b3),
data = DNase.1)
summary(fm1)

```

SSlogis Logistic Model

\section*{Description}

This selfStart model evaluates the logistic function and its gradient. It has an initial attribute that creates initial estimates of the parameters Asym, xmid, and scal.

\section*{Usage}

SSlogis(input, Asym, xmid, scal)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
Asym a numeric parameter representing the asymptote.
xmid a numeric parameter representing the x value at the inflection point of the curve. The value of SSlogis will be Asym/2 at xmid.
scal a numeric scale parameter on the input axis.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Asym/(1+exp((xmid-input)/scal)). If all of the arguments Asym, xmid, and scal are names of objects the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
```

nls, selfStart

```

\section*{Examples}
```

data( ChickWeight )
Chick.1 <- ChickWeight[ChickWeight$Chick == 1, ]
SSlogis( Chick.1$Time, 368, 14, 6 ) \# response only
Asym <- 368; xmid <- 14; scal <- 6
SSlogis( Chick.1\$Time, Asym, xmid, scal ) \# response and gradient
getInitial(weight ~ SSlogis(Time, Asym, xmid, scal), data = Chick.1)

## Initial values are in fact the converged values

fm1 <- nls(weight ~ SSlogis(Time, Asym, xmid, scal), data = Chick.1)
summary(fm1)

```

\section*{SSmicmen Michaelis-Menten Model}

\section*{Description}

This selfStart model evaluates the Michaelis-Menten model and its gradient. It has an initial attribute that will evaluate initial estimates of the parameters Vm and K

\section*{Usage}

SSmicmen(input, Vm, K)

\section*{Arguments}
input a numeric vector of values at which to evaluate the model.
Vm a numeric parameter representing the maximum value of the response.
K a numeric parameter representing the input value at which half the maximum response is attained. In the field of enzyme kinetics this is called the Michaelis parameter.

\section*{Value}
a numeric vector of the same length as input. It is the value of the expression Vm*input/(K+input). If both the arguments Vm and K are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Jose Pinheiro and Douglas Bates

\section*{See Also}
nls, selfStart

\section*{Examples}
```

data( Puromycin )
PurTrt <- Puromycin[ Puromycin$state == "treated", ]
SSmicmen( PurTrt$conc, 200, 0.05 ) \# response only
Vm <- 200; K <- 0.05
SSmicmen( PurTrt\$conc, Vm, K ) \# response and gradient
getInitial(rate ~ SSmicmen(conc, Vm, K), data = PurTrt)

## Initial values are in fact the converged values

fm1 <- nls(rate ~ SSmicmen(conc, Vm, K), data = PurTrt)
summary(fm1 )

## Alternative call using the subset argument

fm2 <- nls(rate ~ SSmicmen(conc, Vm, K), data = Puromycin,
subset = state == "treated")
summary(fm2)

```

\section*{Description}

This selfStart model evaluates the Weibull model for growth curve data and its gradient. It has an initial attribute that will evaluate initial estimates of the parameters Asym, Drop, lrc, and pwr for a given set of data.

\section*{Usage}

SSweibull(x, Asym, Drop, lrc, pwr)

\section*{Arguments}
x
Asym a numeric parameter representing the horizontal asymptote on the right side (very small values of x ).

Drop a numeric parameter representing the change from Asym to the y intercept.
\(\operatorname{lrc} \quad\) a numeric parameter representing the natural logarithm of the rate constant.
pwr a numeric parameter representing the power to which x is raised.

\section*{Details}

This model is a generalization of the SSasymp model in that it reduces to SSasymp when pwr is unity.

\section*{Value}
a numeric vector of the same length as x . It is the value of the expression Asym-Drop*exp (\(\left.\exp (\operatorname{lrc}) * x^{\wedge} p w r\right)\). If all of the arguments Asym, Drop, lrc, and pwr are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

\section*{Author(s)}

Douglas Bates

\section*{References}

Ratkowsky, David A. (1983), Nonlinear Regression Modeling, Dekker. (section 4.4.5)

\section*{See Also}
```

nls, selfStart, SSasymp

```

\section*{Examples}
```

data(ChickWeight)
Chick.6 <- subset(ChickWeight, (Chick == 6) \& (Time > 0))
SSweibull(Chick.6$Time, 160, 115, -5.5, 2.5 ) # response only
Asym <- 160; Drop <- 115; lrc <- -5.5; pwr <- 2.5
SSweibull(Chick.6$Time, Asym, Drop, lrc, pwr) \# response and gradient
getInitial(weight ~ SSweibull(Time, Asym, Drop, lrc, pwr), data = Chick.6)

## Initial values are in fact the converged values

fm1 <- nls(weight ~ SSweibull(Time, Asym, Drop, lrc, pwr), data = Chick.6)
summary(fm1)

```

Theoph Pharmacokinetics of theophylline

\section*{Description}

The Theoph data frame has 132 rows and 5 columns of data from an experiment on the pharmacokinetics of theophylline.

\section*{Usage}
data(Theoph)

\section*{Format}

This data frame contains the following columns:
Subject an ordered factor with levels \(1, \ldots, 12\) identifying the subject on whom the observation was made. The ordering is by increasing maximum concentration of theophylline observed.

Wt weight of the subject ( kg ).
Dose dose of theophylline administered orally to the subject ( \(\mathrm{mg} / \mathrm{kg}\) ).
Time time since drug administration when the sample was drawn (hr).
conc theophylline concentration in the sample ( \(\mathrm{mg} / \mathrm{L}\) ).

\section*{Details}

Boeckmann, Sheiner and Beal (1994) report data from a study by Dr. Robert Upton of the kinetics of the anti-asthmatic drug theophylline. Twelve subjects were given oral doses of theophylline then serum concentrations were measured at 11 time points over the next 25 hours.

These data are analyzed in Davidian and Giltinan (1995) and Pinheiro and Bates (2000) using a two-compartment open pharmacokinetic model, for which a self-starting model function, SSfol , is available.

\section*{Source}

Boeckmann, A. J., Sheiner, L. B. and Beal, S. L. (1994), NONMEM Users Guide: Part V, NONMEM Project Group, University of California, San Francisco.
Davidian, M. and Giltinan, D. M. (1995) Nonlinear Models for Repeated Measurement Data, Chapman \& Hall (section 5.5, p. 145 and section 6.6, p. 176)
Pinheiro, J. C. and Bates, D. M. (2000) Mixed-effects Models in \(S\) and S-PLUS, Springer (Appendix A.29)

See Also
SSfol

\section*{Examples}
```

data(Theoph)
coplot(conc ~ Time | Subject, data = Theoph, show = FALSE)
Theoph.4 <- subset(Theoph, Subject == 4)
fm1 <- nls(conc ~ SSfol(Dose, Time, lKe, lKa, lCl),
data = Theoph.4)
summary(fm1)
plot(conc ~ Time, data = Theoph.4,
xlab = "Time since drug administration (hr)",
ylab = "Theophylline concentration (mg/L)",
main = "Observed concentrations and fitted model",
sub = "Theophylline data - Subject 4 only",
las = 1, col = 4)
xvals <- seq(0, par("usr")[2], len = 55)
lines(xvals, predict(fm1, newdata = list(Time = xvals)),
col = 4)

```

\section*{Chapter 9}

\section*{The splines package}
asVector
Coerce an Object to a Vector

\section*{Description}

This is a generic function. Methods for this function coerce objects of given classes to vectors.

\section*{Usage}
asVector (object)

\section*{Arguments}
object An object.

\section*{Details}

Methods for vector coercion in new classes must be created for the asVector generic instead of as.vector. The as.vector function is internal and not easily extended. Currently the only class with an asVector method is the xyVector class.

\section*{Value}
a vector

\section*{Author(s)}

Douglas Bates and Bill Venables
```

See Also
xyVector

```

\section*{Examples}
```

data( women )
ispl <- interpSpline( weight ~ height, women )
pred <- predict(ispl)
class(pred)
str(pred)
asVector(pred)

```
backSpline Monotone Inverse Spline

\section*{Description}

Create a monotone inverse of a monotone natural spline.

\section*{Usage}
backSpline(object)

\section*{Arguments}
object an object that inherits from class nbSpline or npolySpline. That is, the object must represent a natural interpolation spline but it can be either in the B-spline representation or the piecewise polynomial one. The spline is checked to see if it represents a monotone function.

\section*{Value}

An object of class polySpline that contains the piecewise polynomial representation of a function that has the appropriate values and derivatives at the knot positions to be an inverse of the spline represented by object. Technically this object is not a spline because the second derivative is not constrained to be continuous at the knot positions. However, it is often a much better approximation to the inverse than fitting an interpolation spline to the \(\mathrm{y} / \mathrm{x}\) pairs.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
```

interpSpline

```

\section*{Examples}
```

data( women )
ispl <- interpSpline( women$height, women$weight )
bspl <- backSpline( ispl )
plot( bspl ) \# plots over the range of the knots
points( women$weight, women$height )

```

\section*{Description}

Generate the B-spline basis matrix for a polynomial spline.

\section*{Usage}
bs(x, df = NULL, knots = NULL, degree = 3, intercept = FALSE, Boundary.knots \(=\) range \((x)\) )

\section*{Arguments}
\begin{tabular}{ll}
x & the predictor variable. \\
df & \begin{tabular}{l} 
degrees of freedom; one can specify df rather than knots; bs () then \\
chooses df-degree-1 knots at suitable quantiles of x.
\end{tabular} \\
knots & \begin{tabular}{l} 
the internal breakpoints that define the spline. The default is NULL, which \\
results in a basis for ordinary polynomial regression. Typical values are \\
the mean or median for one knot, quantiles for more knots. See also \\
Boundary.knots.
\end{tabular} \\
degree & \begin{tabular}{l} 
degree of the piecewise polynomial-default is 3 for cubic splines.
\end{tabular} \\
intercept & if TRUE, an intercept is included in the basis; default is FALSE. \\
Boundary.knots
\end{tabular}\(\quad\)\begin{tabular}{l} 
boundary points at which to anchor the B-spline basis (default the range of \\
the data). If both knots and Boundary.knots are supplied, the basis pa- \\
rameters do not depend on x. Data can extend beyond Boundary.knots.
\end{tabular}

\section*{Value}

A matrix of dimension length (x) * df, where either df was supplied or if knots were supplied, df = length(knots) \(+3+\) intercept. Attributes are returned that correspond to the arguments to bs, and explicitly give the knots, Boundary.knots etc for use by predict.bs().
bs () is based on the function spline. des() written by Douglas Bates. It generates a basis matrix for representing the family of piecewise polynomials with the specified interior knots and degree, evaluated at the values of x . A primary use is in modeling formulas to directly specify a piecewise polynomial term in a model.

Beware of making predictions with new \(x\) values when \(d f\) is used as an argument. Either use safe.predict.gam(), or else specify knots and Boundary.knots.

\section*{See Also}
```

ns, poly, smooth.spline, predict.bs, SafePrediction

```

\section*{Examples}
```

data(women)
bs(women\$height, df = 5)
summary(fm1 <- lm(weight ~ bs(height, df = 5), data = women))

## example of safe prediction

plot(women, xlab = "Height (in)", ylab = "Weight (lb)")
ht <- seq(57, 73, len = 200)
lines(ht, predict(fm1, data.frame(height=ht)))

## Consistency:

x <- c(1:3,5:6)
stopifnot(identical(bs(x), bs(x, df = 3)),
!is.null(kk <- attr(bs(x), "knots")),\# not true till 1.5.1
length(kk) == 0)

```
interpSpline Create an Interpolation Spline

\section*{Description}

Create an interpolation spline, either from x and y vectors, or from a formula/data.frame combination.

\section*{Usage}
```

interpSpline(obj1, obj2, bSpline = FALSE, period = NULL,
na.action = na.fail)

```

\section*{Arguments}
obj1 Either a numeric vector of x values or a formula.
obj2 If obj1 is numeric this should be a numeric vector of the same length. If obj1 is a formula this can be an optional data frame in which to evaluate the names in the formula.
bSpline If TRUE the b-spline representation is returned, otherwise the piecewise polynomial representation is returned. Defaults to FALSE.
period An optional positive numeric value giving a period for a periodic interpolation spline.
na.action a optional function which indicates what should happen when the data contain NAs. The default action (na.omit) is to omit any incomplete observations. The alternative action na.fail causes interpSpline to print an error message and terminate if there are any incomplete observations.

\section*{Value}

An object that inherits from class spline. The object can be in the B-spline representation, in which case it will be of class nbSpline for natural B-spline, or in the piecewise polynomial representation, in which case it will be of class npolySpline.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
splineKnots, splineOrder, periodicSpline

\section*{Examples}
```

data( women )
ispl <- interpSpline( women$height, women$weight )
ispl2 <- interpSpline( weight ~ height, women )

# ispl and ispl2 should be the same

plot( predict( ispl, seq( 55, 75, len = 51 ) ), type = "l" )
points( women$height, women$weight )
plot( ispl ) \# plots over the range of the knots
points( women$height, women$weight )
splineKnots( ispl )

```

\section*{Description}

Generate the B-spline basis matrix for a natural cubic spline.

\section*{Usage}
ns ( \(\mathrm{x}, \mathrm{df}=\mathrm{NULL}, \mathrm{knots}=\mathrm{NULL}\), intercept \(=\) FALSE, Boundary.knots \(=\) range \((x)\) )

\section*{Arguments}
\begin{tabular}{ll}
x & the predictor variable. \\
df & \begin{tabular}{l} 
degrees of freedom. One can supply df rather than knots; ns () then \\
chooses df \(-1-\) intercept knots at suitably chosen quantiles of x.
\end{tabular} \\
knots & \begin{tabular}{l} 
breakpoints that define the spline. The default is no knots; together with \\
the natural boundary conditions this results in a basis for linear regression \\
on x. Typical values are the mean or median for one knot, quantiles for \\
more knots. See also Boundary.knots.
\end{tabular} \\
intercept & if TRUE, an intercept is included in the basis; default is FALSE. \\
Boundary.knots
\end{tabular}
boundary points at which to impose the natural boundary conditions and anchor the B-spline basis (default the range of the data). If both knots and Boundary.knots are supplied, the basis parameters do not depend on x. Data can extend beyond Boundary.knots

\section*{Value}

A matrix of dimension length ( \(x\) ) * df where either df was supplied or if knots were supplied, df = length(knots) \(+1+\) intercept. Attributes are returned that correspond to the arguments to ns, and explicitly give the knots, Boundary.knots etc for use by predict.ns().
ns() is based on the function spline. des(). It generates a basis matrix for representing the family of piecewise-cubic splines with the specified sequence of interior knots, and the natural boundary conditions. These enforce the constraint that the function is linear beyond the boundary knots, which can either be supplied, else default to the extremes of the data. A primary use is in modeling formula to directly specify a natural spline term in a model.
Beware of making predictions with new \(x\) values when \(d f\) is used as an argument. Either use safe.predict.gam(), or else specify knots and Boundary.knots.

\section*{See Also}
bs, poly, predict.ns, SafePrediction

\section*{Examples}
```

data(women)
ns(women\$height, df = 5)
summary(fm1 <- lm(weight ~ ns(height, df = 5), data = women))

## example of safe prediction

plot(women, xlab = "Height (in)", ylab = "Weight (lb)")
ht <- seq(57, 73, len = 200)
lines(ht, predict(fm1, data.frame(height=ht)))

```
periodicSpline Create a Periodic Interpolation Spline

\section*{Description}

Create a periodic interpolation spline, either from x and y vectors, or from a formula/data.frame combination.

\section*{Usage}
periodicSpline(obj1, obj2, knots, period \(=2 *\) pi, ord \(=4\) )

\section*{Arguments}
obj1 either a numeric vector of x values or a formula.
obj2 if obj1 is numeric this should be a numeric vector of the same length. If obj1 is a formula this can be an optional data frame in which to evaluate the names in the formula.
knots optional numeric vector of knot positions.
period positive numeric value giving the period for the periodic spline. Defaults to 2 * pi.
ord integer giving the order of the spline, at least 2. Defaults to 4. See splineOrder for a definition of the order of a spline.

\section*{Value}

An object that inherits from class spline. The object can be in the B-spline representation, in which case it will be a pbSpline object, or in the piecewise polynomial representation (a ppolySpline object).

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
```

splineKnots, interpSpline

```

\section*{Examples}
```

xx <- seq( -pi, pi, len = 16 ) [-1]
yy <- sin( xx )
frm <- data.frame( xx, yy )
( pispl <- periodicSpline( xx, yy, period = 2 * pi ) )
pispl2 <- periodicSpline( yy ~ xx, frm, period = 2 * pi )
stopifnot(all.equal(pispl, pispl2))\# pispl and pispl2 are the same
plot( pispl ) \# displays over one period
points( yy ~ xx, col = "brown")
plot( predict( pispl, seq(-3*pi, 3*pi, len = 101) ), type = "l" )

```
```

polySpline Piecewise Polynomial Spline Representation

```

\section*{Description}

Create the piecewise polynomial representation of a spline object.

\section*{Usage}
polySpline(object, ...)
as.polySpline(object, ...)

\section*{Arguments}
object An object that inherits from class spline.
... Optional additional arguments. At present no additional arguments are used.

\section*{Value}

An object that inherits from class polySpline. This is the piecewise polynomial representation of a univariate spline function. It is defined by a set of distinct numeric values called knots. The spline function is a polynomial function between each successive pair of knots. At each interior knot the polynomial segments on each side are constrained to have the same value of the function and some of its derivatives.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
```

interpSpline, periodicSpline, splineKnots, splineOrder

```

\section*{Examples}
```

data( women )
ispl <- polySpline( interpSpline( weight ~ height, women, bSpline = TRUE ) )
print( ispl ) \# print the piecewise polynomial representation
plot( ispl ) \# plots over the range of the knots
points( women$height, women$weight )

```
predict.bs Evaluate a Spline Basis

\section*{Description}

Evaluate a predefined spline basis at given values.

\section*{Usage}
```

predict(object, newx, ...)

```

\section*{Arguments}
object the result of a call to bs or ns having attributes describing knots, degree, etc.
newx the \(x\) values at which evaluations are required.
... Optional additional arguments. Presently no additional arguments are used.

\section*{Value}

An object just like object, except evaluated at the new values of x .
These are methods for the generic function predict for objects inheriting from classes "bs" or "ns". See predict for the general behavior of this function.

\section*{See Also}
bs, ns, poly.

\section*{Examples}
```

data(women)
basis <- ns(women\$height, df = 5)
newX <- seq(58, 72, len = 51)

# evaluate the basis at the new data

predict(basis, newX)

```

\section*{Description}

The predict methods for the classes that inherit from the virtual classes bSpline and polySpline are used to evaluate the spline or its derivatives. The plot method for a spline object first evaluates predict with the x argument missing, then plots the resulting xyVector with type = "l".

\section*{Usage}
predict (object, \(x, n s e g=50\), deriv=0, ...)
plot.spline(x, ...)

\section*{Arguments}
object An object that inherits from the bSpline or the polySpline class. For plot.spline this argument is called \(x\).
\(x \quad\) A numeric vector of \(x\) values at which to evaluate the spline. If this argument is missing a suitable set of x values is generated as a sequence of nseq segments spanning the range of the knots. For plot.spline the x argument is as described under object above.
nseg A positive integer giving the number of segments in a set of equally-spaced x values spanning the range of the knots in object. This value is only used if x is missing.
deriv An integer between 0 and splineOrder (object) - 1 specifying the derivative to evaluate.
... predict: further arguments passed to or from other methods. plot: additional graphical parameters (see link\{par\}).

\section*{Value}
an xyVector with components
\(\mathrm{x} \quad\) the supplied or inferred numeric vector of x values
\(\mathrm{y} \quad\) the value of the spline (or its deriv'th derivative) at the x vector

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
xyVector, interpSpline, periodicSpline

\section*{Examples}
```

data( women )
ispl <- interpSpline( weight ~ height, women )
opar <- par(mfrow = c(2, 2), las = 1)
plot(predict(ispl, nseg = 201), \# plots over the range of the knots
main = "Original data with interpolating spline", type = "l",
xlab = "height", ylab = "weight")
points(women$height, women$weight, col = 4)
plot(predict(ispl, nseg = 201, deriv = 1),
main = "First derivative of interpolating spline", type = "l",
xlab = "height", ylab = "weight")
plot(predict(ispl, nseg = 201, deriv = 2),
main = "Second derivative of interpolating spline", type = "l",
xlab = "height", ylab = "weight")
plot(predict(ispl, nseg = 401, deriv = 3),
main = "Third derivative of interpolating spline", type = "l",
xlab = "height", ylab = "weight")
par(opar)

```
```

splineDesign Design Matrix for B-splines

```

\section*{Description}

Evaluate the design matrix for the B-splines defined by knots at the values in x .

\section*{Usage}
```

splineDesign(knots, x, ord = 4, derivs)
spline.des(knots, x, ord = 4, derivs)

```

\section*{Arguments}
knots a numeric vector of knot positions with non-decreasing values.
\(\mathrm{x} \quad\) a numeric vector of values at which to evaluate the B-spline functions or derivatives. The values in x must be between knots [ord] and knots [ length(knots) +1 - ord ].
ord a positive integer giving the order of the spline function. This is the number of coefficients in each piecewise polynomial segment, thus a cubic spline has order 4. Defaults to 4.
derivs an integer vector of the same length as \(x\) and with values between 0 and ord - 1. The derivative of the given order is evaluated at the x positions. Defaults to a vector of zeroes of the same length as \(x\).

\section*{Value}

A matrix with length ( x ) rows and length ( knots ) - ord columns. The i'th row of the matrix contains the coefficients of the B-splines (or the indicated derivative of the Bsplines) defined by the knot vector and evaluated at the i'th value of x. Each B-spline is defined by a set of ord successive knots so the total number of B-splines is length(knots)ord.

\section*{Note}

The older spline.des function takes the same arguments but returns a list with several components including knots, ord, derivs, and design. The design component is the same as the value of the splineDesign function.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{Examples}
```

splineDesign(knots = 1:10, x = 4:7)

```
```

splineKnots Knot Vector from a Spline

```

\section*{Description}

Return the knot vector corresponding to a spline object.

\section*{Usage}
splineKnots(object)

\section*{Arguments}
object an object that inherits from class "spline".

\section*{Value}

A non-decreasing numeric vector of knot positions.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{Examples}
```

data( women )
ispl <- interpSpline( weight ~ height, women )
splineKnots( ispl )

```

\section*{splineOrder Determine the Order of a Spline}

\section*{Description}

Return the order of a spline object.

\section*{Usage}
splineOrder(object)

\section*{Arguments}
object An object that inherits from class "spline".

\section*{Details}

The order of a spline is the number of coefficients in each piece of the piecewise polynomial representation. Thus a cubic spline has order 4.

\section*{Value}

A positive integer.

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{See Also}
splineKnots, interpSpline, periodicSpline

\section*{Examples}
```

data( women )
splineOrder( interpSpline( weight ~ height, women ) )

```
```

xyVector

```

Construct an xyVector Object

\section*{Description}

Create an object to represent a set of \(x-y\) pairs. The resulting object can be treated as a matrix or as a data frame or as a vector. When treated as a vector it reduces to the y component only.
The result of functions such as predict.spline is returned as an xyVector object so the x -values used to generate the y -positions are retained, say for purposes of generating plots.

\section*{Usage}
```

xyVector(x, y)

```

\section*{Arguments}
\begin{tabular}{ll}
x & a numeric vector \\
y & a numeric vector of the same length as x
\end{tabular}

\section*{Value}

An object of class xyVector with components
\begin{tabular}{ll}
\(x\) & a numeric vector \\
\(y\) & a numeric vector of the same length as \(x\)
\end{tabular}

\section*{Author(s)}

Douglas Bates and Bill Venables

\section*{Examples}
```

data( women )
ispl <- interpSpline( weight ~ height, women )
weights <- predict( ispl, seq( 55, 75, len = 51 ))
class( weights )
plot( weights, type = "l", xlab = "height", ylab = "weight" )
points( women$height, women$weight )
weights

```

\section*{Chapter 10}

\section*{The stepfun package}

\section*{ecdf \\ Empirical Cumulative Distribution Function}

\section*{Description}

Compute or plot an empirical cumulative distribution function.

\section*{Usage}
ecdf (x)
plot(..., verticals = FALSE, col.01line = "gray70")
print(x, digits= getOption("digits") - 2, ...)
summary(object, ...)

\section*{Arguments}
x
numeric vector of "observations" in ecdf; for the methods \(x\) is as object below.
... arguments to be passed to plot.stepfun, the first of which should be an R object of class "ecdf".
verticals
see plot.stepfun.
col.01line numeric or character specifying the color of the horizontal lines at \(y=0\) and 1 , see colors.
object (or x:) object of class "ecdf", typically.
digits number of significant digits to use, see print.

\section*{Details}

The e.c.d.f. (empirical cumulative distribution function) \(F_{n}\) is a step function with jump \(1 / n\) at each observation (possibly with multiple jumps at one place if there are ties).

For observations \(\mathrm{x}=\left(x_{1}, x_{2}, \ldots x_{n}\right), F_{n}\) is the fraction of observations less or equal to \(t\), i.e.,
\[
F_{n}(t)=\#\left\{x_{i} \leq t\right\} / n=\frac{1}{n} \sum_{i=1}^{n} \mathbf{1}_{\left[x_{i} \leq t\right]}
\]

The function plot.ecdf which implements the plot method for ecdf objects, is implemented via a call to plot.stepfun; see its documentation.

\section*{Value}

For ecdf, a function of class "ecdf", inheriting from the "stepfun" class.

\section*{Author(s)}

Martin Maechler, 〈maechler@stat.math.ethz.ch〉.

\section*{See Also}
stepfun, the more general class of step functions, approxfun and splinefun.

\section*{Examples}
```

\#\#-- Simple didactical ecdf example:
Fn <- ecdf(rnorm(12))
Fn; summary(Fn)
12*Fn(knots(Fn)) == 1:12 \#\# == 1:12 if and only if there are no ties !
y <- round(rnorm(12),1); y[3] <- y[1]
Fn12 <- ecdf(y)
Fn12
print(knots(Fn12), dig=2)
12*Fn12(knots(Fn12)) \#\# ~ = 1:12 if there where no ties
summary(Fn12)
summary.stepfun(Fn12)
print(ls.Fn12 <- ls(env= environment(Fn12)))
\#\#[1] "f" "method" "n" "ties" "x" "y" "yleft" "yright"
12 * Fn12((-20:20)/10)
\#\#\#------------------ Plotting
op <- par(mfrow=c(3,1), mgp=c(1.5, 0.8,0), mar= . 1+c(3,3,2,1))
F10 <- ecdf(rnorm(10))
summary(F10)
plot(F10)
plot(F10, verticals= TRUE, do.p = FALSE)
plot(Fn12)\# , lwd=2) dis-regarded
xx <- unique(sort(c(seq(-3,2, length=201), knots(Fn12))))
lines(xx, Fn12(xx), col='blue')
abline(v=knots(Fn12),lty=2,col='gray70')
plot(xx, Fn12(xx), type='b', cex=.1)\#- plot.default
plot(Fn12, col.h='red', add= TRUE) \#- plot method
abline(v=knots(Fn12),lty=2,col='gray70')
plot(Fn12, verticals=TRUE, col.p='blue', col.h='red',col.v='bisque')
par(op)
\#\#-- this works too (automatic call to ecdf(.)):

```
```

plot.ecdf(rnorm(24))

```
```

plot.stepfun

```

Plot Step Functions

\section*{Description}

Method of the generic plot for stepfun objects and utility for plotting piecewise constant functions.

\section*{Usage}
```

plot(x, xval, xlim, xlab = "x", ylab = "f(x)", main = NULL,
add = FALSE, verticals = TRUE, do.points = TRUE,
pch = par("pch"), col.points=par("col"), cex.points=par("cex"),
col.hor = par("col"), col.vert= par("col"),
lty = par("lty"), lwd = par("lwd"), ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & an R object inheriting from "stepfun". \\
xval & \begin{tabular}{l} 
numeric vector of abscissa values at which to evaluate x. . Defaults to \\
knots (x) restricted to xlim.
\end{tabular} \\
xlim & numeric(2); range of x values to use. \\
xlab, ylab & labels of x and y axis. \\
main & main title. \\
add & logical; if TRUE only add to an existing plot. \\
verticals & logical; if TRUE, draw vertical lines at steps. \\
do.points & logical; if true, also draw points at the (xlim restricted) knot locations. \\
pch & character; point character if do.points. \\
col.points & character or integer code; color of points if do.points. \\
cex.points & numeric; character expansion factor if do.points. \\
col.hor & color of horizontal lines. \\
col.vert & color of vertical lines. \\
lty, lwd & line type and thickness for all lines. \\
\(\ldots\) & further arguments of plot (.), or if (add) segments (.).
\end{tabular}

\section*{Value}

A list with two components
\(\mathrm{t} \quad\) abscissa ( x ) values, including the two outermost ones.
\(y \quad y\) values 'in between' the \(t[]\).
Author(s)
Martin Maechler 〈maechler@stat.math.ethz.ch〉, 1990, 1993; ported to R, 1997.

\section*{See Also}
ecdf for empirical distribution functions as special step functions, approxfun and splinefun.

\section*{Examples}
```

y0 <- c(1,2,4,3)
sfun0 <- stepfun(1:3, y0, f = 0)
sfun.2 <- stepfun(1:3, y0, f = .2)
sfun1 <- stepfun(1:3, y0, f = 1)
tt <- seq(0,3, by=0.1)
op <- par(mfrow=c(2,2))
plot(sfun0); plot(sfun0, xval=tt, add=TRUE, col.h="bisque")
plot(sfun.2);plot(sfun.2,xval=tt, add=TRUE, col.h="orange")
plot(sfun1); plot(sfun1, xval=tt, add=TRUE, col.h="coral")
\#\#-- This is revealing :
plot(sfun0, verticals= FALSE,
main = "stepfun(x, y0, f=f) for f = 0, .2, 1")
for(i in 1:3)
plot(list(sfun0,sfun.2,sfun1)[[i]], add=TRUE, col.h=i, col.v=i)
legend(2.5, 1.9, paste("f =", c(0,0.2,1)), col=1:3, lty=1, y.inter=1); par(op)
\#\#-- this works too (automatic call to ecdf(.)):
plot.stepfun(rt(50, df=3), col.vert = "gray20")

```
stepfun Step Functions

\section*{Description}

Given the vectors \(\left(x_{1}, \ldots, x_{n}\right)\) and ( \(y_{0}, y_{1}, \ldots, y_{n}\) ) (one value more!), stepfun( \(\mathrm{x}, \mathrm{y}, \ldots\) ) returns an interpolating "step" function, say fn. I.e., \(f n(t)=c_{i}\) (constant) for \(t \in\left(x_{i}, x_{i+1}\right)\) and \(f n\left(x_{i}\right)=y_{i}\) for \(i=1, \ldots, n\).
The value of the constant \(c_{i}\) above depends on the "continuity" parameter f . For the default, \(\mathrm{f}=0, \mathrm{fn}\) is a "cadlag" function, i.e. continuous at right, limit ("the point") at left. In general, \(c_{i}\) is interpolated in between the neighbouring \(y\) values, \(c_{i}=(1-f) y_{i}+f \cdot y_{i+1}\). Therefore, for non- 0 values of \(f, f n\) may no longer be a proper step function, since it can be discontinuous from both sides.

\section*{Usage}
```

stepfun(x, y, f = 0, ties = "ordered")
is.stepfun(x)
knots(Fn, ...)
print(x, digits= getOption("digits") - 2, ...)
summary(object, ...)

```

\section*{Arguments}
x
y
f
ties

Fn, object
digits
numeric vector giving the knots or jump locations of the step function for stepfun(). For the other functions, x is as object below.
numeric vector one longer than x , giving the heights of the function values between the x values.
a number between 0 and 1, indicating how interpolation outside the given \(x\) values should happen. See approxfun.
Handling of tied x values. Either a function or the string "ordered". See approxfun.
an R object inheriting from "stepfun".
number of significant digits to use, see print.
potentially further arguments (require by the generic).

\section*{Value}

A function of class "stepfun", say fn. There are methods available for summarizing ("summary(.)"), representing ("print(.)") and plotting ("plot(.)", see plot.stepfun) "stepfun" objects.

The environment of fn contains all the information needed;
```

"x","y" the original arguments
"n" number of knots (x values)
"f" continuity parameter
"yleft", "yright"
the function values outside the knots;
"method" (always == "constant", from approxfun(.)).

```
normal-bracket 97 bracket-normal The knots are also available by knots (fn).

\section*{Author(s)}

Martin Maechler, 〈maechler@stat.math.ethz.ch〉 with some basic code from Thomas Lumley.

\section*{See Also}
ecdf for empirical distribution functions as special step functions and plot.stepfun for plotting step functions.
approxfun and splinefun.

\section*{Examples}
```

y0 <- c(1,2,4,3)
sfun0 <- stepfun(1:3, y0, f = 0)
sfun.2 <- stepfun(1:3, y0, f = .2)
sfun1 <- stepfun(1:3, y0, f = 1)
sfun0
summary(sfun0)
summary(sfun.2)

## look at the internal structure:

print.default(sfun0)

```
ls.str(envir = environment(sfun0))
x0 <- seq(0.5,3.5, by = 0.25)
\(\operatorname{rbind}(x=x 0, f . f 0=\operatorname{sfun0}(x 0), f . f 02=\operatorname{sfun.2(x0),f.f1=} \operatorname{sfun1(x0))}\)

\section*{Chapter 11}

\section*{The tcltk package}

\section*{TclInterface Low-level Tcl/Tk Interface}

\section*{Description}

These functions and variables provide the basic glue between R and the Tcl interpreter and Tk GUI toolkit. Tk windows may be represented via R objects. Tcl variables can be accessed via objects of class tclVar and the C level interface to Tcl objects is accessed via objects of class tclObj.

\section*{Usage}
```

.Tcl(...)
.Tcl.args(...)
.Tcl.callback(...)
.Tk.ID(win)
.Tk.newwin(ID)
.Tk.subwin(parent)
.TkWin
.TkRoot
tkdestroy(win)
is.tkwin(x)
tclVar(init="")
as.character(x) \# x of class "tclVar" or "tclObj"
tclvalue(x) \# x of class "tclVar" or "tclObj"
tclvalue(x) <- value
tclObj(x) \# x of class "tclVar"
tclObj(x) <- value
as.tclObj(x)
is.tclObj(x)

```
```

as.integer(x) \# x of class "tclObj"
as.double(x) \# x of class "tclObj"
addTclPath(path = ".")
tclRequire(package, warn = TRUE)

```

\section*{Arguments}
\begin{tabular}{ll} 
win & a window structure \\
x & an object \\
ID & a window ID \\
parent & a window which becomes the parent of the resulting window \\
path & path to a directory containing Tcl packages \\
package & a Tcl package name \\
warn & logical. Warn if not found? \\
\(\ldots\) & Additional arguments. See below. \\
init & initialization value
\end{tabular}

\section*{Details}

Many of these functions are not intended for general use but are used internally by the commands that create and manipulate Tk widgets and Tcl objects. At the lowest level .Tcl sends a command as a text string to the Tcl interpreter and returns the result as an object of class tclObj (see below).
.Tcl.args converts an \(R\) argument list of tag=value pairs to the Tcl -option value style, thus enabling a simple translation between the two languages. To send a value with no preceding option flag to Tcl, just use an untagged argument. In the rare case one needs an option with no subsequent value tag=NULL can be used. Most values are just converted to character mode and inserted in the command string, but window objects are passed using their ID string, and callbacks are passed via the result of .Tcl.callback. Tags are converted to option flags simply by prepending a -
Callbacks can be either atomic callbacks handled by .Tcl.callback or expressions. An expression is treated as a list of atomic callbacks, with the following exceptions: if an element is a name, it is first evaluated in the callers frame, and likewise if it is an explicit function definition; the break expression is translated directly to the Tcl counterpart. .Tcl.callback converts R functions and unevaluated calls to Tcl command strings. The argument must be either a function closure or an object of mode "call" followed by an environment. The return value in the first case is of the form R_call \(0 \times 408 \mathrm{~b} 94 \mathrm{~d} 4\) in which the hexadecimal number is the memory address of the function. In the second case it will be of the form R_call_lang 0x8a95904 0x819bfd0. For expressions, a sequence of similar items is generated, separated by semicolons. .Tcl.args takes special precautions to ensure that functions or calls will continue to exist at the specified address by assigning the callback into the relevant window environment (see below).

Tk windows are represented as objects of class tkwin which are lists containing a ID field and an env field which is an \(R\) environments, enclosed in the global environment. The value of the ID field is identical to the Tk window name. The env environment contains a parent variable and a num.subwin variable. If the window obtains subwindows and callbacks, they are added as variables to the environment. . TkRoot is the top window with ID "."; this window is not displayed in order to avoid ill effects of closing it via window manager controls. The parent variable is undefined for . TkRoot.
.Tk.ID extracts the ID of a window, .Tk.newwin creates a new window environment with a given ID and .Tk.subwin creates a new window which is a subwindow of a given parent window.
tkdestroy destroys a window and also removes the reference to a window from its parent. is.tkwin can be used to test whether a given object is a window environment.
tclVar creates a new Tcl variable and initializes it to init. An R object of class tclVar is created to represent it. Using as.character on the object returns the Tcl variable name. Accessing the Tcl variable from R is done using the tclvalue function, which can also occur on the left sie of assignments. If tclvalue is passed an argument which is not a tclVar object, then it will assume that it is a character string explicitly naming global Tcl variable. Tcl variables created by tclVar are uniquely named and automatically unset by the garbage collector when the representing object is no longer in use.
It is possible to access Tcl's 'dual-ported' objects directly, thus avoiding parsing and deparsing of their string representation. This works by using objects of class tclObj. The string representation of such objects can be extracted (but not set) using tclvalue and conversion to vectors of mode "character", "double", or "integer". Conversely, such vectors can be converted using as.tclObj. The object behind a tclVar object is extracted using tclObj(x) which also allows an assignment form, in which the right hand side of the assignment is automatically converted using as.tclObj. There is a print method for tclObj objects; it prints <Tcl> followed by the string representation of the object.
Tcl packages can be loaded with tclRequire; it may be necessary to add the directory where they are found to the Tcl search path with addTclPath.

\section*{Note}

Strings containing unbalanced braces are currently not handled well in many circumstances.

\section*{See Also}

TkWidgets, TkCommands, TkWidgetcmds. capabilities("tcltk")

\section*{Examples}
```


## These cannot be run by example() but should be OK when pasted

## into an interactive R session with the tcltk package loaded

.Tcl("format \"%s\n\" \"Hello, World!\"")
f <- function()"HI!"
.Tcl.callback(f)
.Tcl.args(text="Push!", command=f) \# NB: Different address
xyzzy <- tclVar(7913)
tclvalue(xyzzy)
tclvalue(xyzzy) <- "foo"
as.character(xyzzy)
tkcmd("set", as.character(xyzzy))
top <- tktoplevel() \# a Tk widget, see Tk-widgets
ls(envir=top$env, all=TRUE)
ls(envir=.TkRoot$env, all=TRUE)\# .Tcl.args put a callback ref in here

```

\section*{TkCommands Tk non-widget commands}

\section*{Description}

These functions interface to Tk non-widget commands, such as the window manager interface commands and the geometry managers.

\section*{Usage}
```

tkcmd(...)
tktitle(x)
tktitle(x) <- value
tkbell(...)
tkbind(...)
tkbindtags(...)
tkfocus(...)
tklower(...)
tkraise(...)
tkclipboard.append(...)
tkclipboard.clear(...)
tkevent.add(...)
tkevent.delete(...)
tkevent.generate(...)
tkevent.info(...)
tkfont.actual(...)
tkfont.configure(...)
tkfont.create(...)
tkfont.delete(...)
tkfont.families(...)
tkfont.measure(...)
tkfont.metrics(...)
tkfont.names(...)
tkgrab(...)
tkgrab.current(...)
tkgrab.release(...)
tkgrab.set(...)
tkgrab.status(...)

## NB: some widgets also have a selection.clear command, hence the "X".

tkXselection.clear(...)
tkXselection.get(...)
tkXselection.handle(...)
tkXselection.own(...)

```
```

tkwait.variable(...)
tkwait.visibility(...)
tkwait.window(...)

## winfo actually has a large number of subcommands, but it's rarely

## used, so use tkwinfo("atom", ...) etc. instead.

tkwinfo(...)

# Window manager interface

tkwm.aspect(...)
tkwm.client(...)
tkwm.colormapwindows(...)
tkwm.command(...)
tkwm.deiconify(...)
tkwm.focusmodel(...)
tkwm.frame(...)
tkwm.geometry(...)
tkwm.grid(...)
tkwm.group(...)
tkwm.iconbitmap(...)
tkwm.iconify(...)
tkwm.iconmask(...)
tkwm.iconname(...)
tkwm.iconposition(...)
tkwm.iconwindow(...)
tkwm.maxsize(...)
tkwm.minsize(...)
tkwm.overrideredirect(...)
tkwm.positionfrom(...)
tkwm.protocol(...)
tkwm.resizable(...)
tkwm.sizefrom(...)
tkwm.state(...)
tkwm.title(...)
tkwm.transient(...)
tkwm.withdraw(...)

### Geometry managers

tkgrid(...)
tkgrid.bbox(...)
tkgrid.columnconfigure(...)
tkgrid.configure(...)
tkgrid.forget(...)
tkgrid.info(...)
tkgrid.location(...)
tkgrid.propagate(...)
tkgrid.rowconfigure(...)
tkgrid.remove(...)

```
```

tkgrid.size(...)
tkgrid.slaves(...)
tkpack(...)
tkpack.configure(...)
tkpack.forget(...)
tkpack.info(...)
tkpack.propagate(...)
tkpack.slaves(...)
tkplace(...)
tkplace.configure(...)
tkplace.forget(...)
tkplace.info(...)
tkplace.slaves(...)

## Standard dialogs

tkgetOpenFile(...)
tkgetSaveFile(...)
tkmessageBox(...)

## File handling functions

tkfile.tail(...)
tkfile.dir(...)
tkopen(...)
tkclose(...)
tkputs(...)

```

\section*{Arguments}
```

x A window object
... Handled via.Tcl.args

```

\section*{Details}
tkcmd provides a generic interface to calling any Tk or Tcl command by simply running .Tcl.args on the arguments and passing the result to .Tcl. Most of the other commands simply call tkcmd with a particular first argument and sometimes also a second argument giving the subcommand.
tktitle and its assignment form provides an alternate interface to Tk's wm title
There are far too many of these commands to describe them and their arguments in full. Please refer to the Tcl/Tk documentation for details. Except for a few exceptions, the pattern is that Tcl subcommands like pack configure are converted to function names like tkpack. configure.

\section*{See Also}

TclInterface, TkWidgets, TkWidgetcmds

\section*{Examples}
```


## into an interactive R session with the tcltk package loaded

tt <- tktoplevel()
tkpack(l1<-tklabel(tt,text="Heave"),12<-tklabel(tt,text="Ho"))
tkpack.configure(l1,side="left")

## Try stretching the window and then

tkdestroy(tt)

```
tkpager

Page file using Tk text widget

\section*{Description}

This plugs into file.show, showing files in separate windows.

\section*{Usage}
tkpager(file, header, title, delete.file)

\section*{Arguments}
\begin{tabular}{ll} 
file & character vector containing the names of the files to be displayed \\
header & headers to use at the beginning of each file \\
title & title to use for the window \\
delete.file & logical. Should file(s) be deleted after display
\end{tabular}

\section*{Note}

The "
b_" string used for underlining is currently quietly removed. The font and background colour are currently hardcoded to Courier and gray90.

\section*{See Also}
file.show

\section*{Examples}

\section*{TkWidgetcmds Tk widget commands}

\section*{Description}

These functions interface to Tk widget commands.

Usage
```

tkactivate(widget, ...)
tkadd(widget, ...)
tkaddtag(widget, ...)
tkbbox(widget, ...)
tkcanvasx(widget, ...)
tkcanvasy(widget, ...)
tkcget(widget, ...)
tkcompare(widget, ...)
tkconfigure(widget, ...)
tkcoords(widget, ...)
tkcreate(widget, ...)
tkcurselection(widget,...)
tkdchars(widget, ...)
tkdebug(widget, ...)
tkdelete(widget, ...)
tkdelta(widget, ...)
tkdeselect(widget, ...)
tkdlineinfo(widget, ...)
tkdump(widget, ...)
tkentrycget(widget, ...)
tkentryconfigure(widget, ...)
tkfind(widget, ...)
tkflash(widget, ...)
tkfraction(widget, ...)
tkget(widget, ...)
tkgettags(widget, ...)
tkicursor(widget, ...)
tkidentify(widget, ...)
tkimage.cget(widget, ...)
tkimage.configure(widget, ...)
tkimage.create(widget, ...)
tkimage.names(widget, ...)
tkindex(widget, ...)
tkinsert(widget, ...)
tkinvoke(widget, ...)
tkitembind(widget, ...)
tkitemcget(widget, ...)
tkitemconfigure(widget, ...)
tkitemfocus(widget, ...)
tkitemlower(widget, ...)
tkitemraise(widget, ...)
tkitemscale(widget, ...)

```
```

tkmark.gravity(widget, ...)
tkmark.names(widget, ...)
tkmark.next(widget, ...)
tkmark.previous(widget, ...)
tkmark.set(widget, ...)
tkmark.unset(widget, ...)
tkmove(widget, ...)
tknearest(widget, ...)
tkpost(widget, ...)
tkpostcascade(widget, ...)
tkpostscript(widget, ...)
tkscan.mark(widget, ...)
tkscan.dragto(widget, ...)
tksearch(widget, ...)
tksee(widget, ...)
tkselect(widget, ...)
tkselection.adjust(widget, ...)
tkselection.anchor(widget, ...)
tkselection.clear(widget, ...)
tkselection.from(widget, ...)
tkselection.includes(widget, ...)
tkselection.present(widget, ...)
tkselection.range(widget, ...)
tkselection.set(widget, ...)
tkselection.to(widget,...)
tkset(widget, ...)
tksize(widget, ...)
tktoggle(widget, ...)
tktag.add(widget, ...)
tktag.bind(widget, ...)
tktag.cget(widget, ...)
tktag.configure(widget, ...)
tktag.delete(widget, ...)
tktag.lower(widget, ...)
tktag.names(widget, ...)
tktag.nextrange(widget, ...)
tktag.prevrange(widget, ...)
tktag.raise(widget, ...)
tktag.ranges(widget, ...)
tktag.remove(widget, ...)
tktype(widget, ...)
tkunpost(widget, ...)
tkwindow.cget(widget, ...)
tkwindow.configure(widget, ...)
tkwindow.create(widget, ...)
tkwindow.names(widget, ...)
tkxview(widget, ...)
tkxview.moveto(widget, ...)
tkxview.scroll(widget, ...)
tkyposition(widget, ...)
tkyview(widget, ...)
tkyview.moveto(widget, ...)

```
```

tkyview.scroll(widget, ...)

```

\section*{Arguments}
widget The widget this applies to
... Handled via .Tcl.args

\section*{Details}

There are far too many of these commands to describe them and their arguments in full. Please refer to the \(\mathrm{Tcl} / \mathrm{Tk}\) documentation for details. Except for a few exceptions, the pattern is that Tcl widget commands possibly with subcommands like .a.b selection clear are converted to function names like tkselection.clear and the widget is given as the first argument.

\section*{See Also}

TclInterface, TkWidgets, TkCommands

\section*{Examples}
```


## These cannot be run by examples() but should be OK when pasted

## into an interactive R session with the tcltk package loaded

tt <- tktoplevel()
tkpack(txt.w <- tktext(tt))
tkinsert(txt.w, "0.0", "plot(1:10)")

# callback function

eval.txt <- function()
eval(parse(text=tkget(txt.w, "0.0", "end")))
tkpack(but.w <- tkbutton(tt,text="Submit", command=eval.txt))

## Try pressing the button, edit the text and when finished:

```
tkdestroy (tt)
TkWidgets Tk widgets

\section*{Description}

Create Tk widgets and associated R objects.

\section*{Usage}
tkwidget(parent, type, ...)
tkbutton(parent, ...)
tkcanvas(parent, ...)
tkcheckbutton(parent, ...)
```

tkentry(parent, ...)
tkframe(parent, ...)
tklabel(parent, ...)
tklistbox(parent, ...)
tkmenu(parent, ...)
tkmenubutton(parent, ...)
tkmessage(parent, ...)
tkradiobutton(parent, ...)
tkscale(parent, ...)
tkscrollbar(parent, ...)
tktext(parent, ...)
tktoplevel(parent=.TkRoot, ...)

```

\section*{Arguments}
parent Parent of widget window
type string describing the type of widget desired
... handled via .Tcl.args

\section*{Details}

These functions create Tk widgets. tkwidget creates a widget of a given type, the others simply call tkwidget with the respective type argument.

It is not possible to describe the widgets and their arguments in full. Please refer to the \(\mathrm{Tcl} / \mathrm{Tk}\) documentation.

\section*{See Also}

TclInterface, TkCommands, TkWidgetcmds

\section*{Examples}
```


## These cannot be run by examples() but should be OK when pasted

## into an interactive R session with the tcltk package loaded

tt <- tktoplevel()
label.widget <- tklabel(tt, text="Hello, World!")
button.widget <- tkbutton(tt, text="Push",
command=function()cat("OW!\n"))
tkpack(label.widget, button.widget) \# geometry manager
\# see Tk-commands

## Push the button and then...

tkdestroy(tt)

```

\section*{Chapter 12}

\section*{The tools package}

\author{
checkFF Check Foreign Function Calls
}

\section*{Description}

Performs checks on calls to compiled code from R code. Currently only whether the interface functions such as .C and .Fortran are called with argument PACKAGE specified, which is highly recommended to avoid name clashes in foreign function calls.

\section*{Usage}
```

checkFF(package, dir, file, lib.loc = NULL,
verbose = getOption("verbose"))

```

\section*{Arguments}
package a character string naming an installed package. If given, the installed \(R\) code of the package is checked.
dir a character string specifying the path to a package's root source directory. This should contain the subdirectory R (for R code). Only used if package is not given.
file the name of a file containing R code to be checked. Used if neither package nor dir are given.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. The specified library trees are used to to search for package.
verbose a logical. If TRUE, additional diagnostics are printed (and the result is returned invisibly).

\section*{Value}

An object of class "checkFF", which currently is a list of the (parsed) foreign function calls with no PACKAGE argument.
There is a print method for nicely displaying the information contained in such objects.

\section*{Warning}

This function is still experimental. Both name and interface might change in future versions.

\section*{See Also}
.C, .Fortran; Foreign.

\section*{Examples}
```

checkFF(package = "ts", verbose = TRUE)

```
checkTnF

Check R Packages or Code for T/F

\section*{Description}

Checks the specified \(R\) package or code file for occurrences of \(T\) or \(F\), and gathers the expression containing these. This is useful as in R T and F are just variables which are set to the logicals TRUE and FALSE by default, but are not reserved words and hence can be overwritten by the user. Hence, one should always use TRUE and FALSE for the logicals.

\section*{Usage}
checkTnF(package, dir, file, lib.loc = NULL)

\section*{Arguments}
package a character string naming an installed package. If given, the installed R code and the examples in the documentation files of the package are checked. R code installed as an image file cannot be checked.
dir a character string specifying the path to a package's root source directory. This must contain the subdirectory ' \(R\) ' (for \(R\) code), and should also contain 'man' (for documentation). Only used if package is not given. If used, the R code files and the examples in the documentation files are checked.
file the name of a file containing \(R\) code to be checked. Used if neither package nor dir are given.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. The specified library trees are used to to search for package.

\section*{Value}

An object of class "checkTnF" which is a list containing, for each file where occurences of T or F were found, a list with the expressions containing these occurrences. The names of the list are the corresponding file names.
There is a print method for nicely displaying the information contained in such objects.

\section*{Warning}

This function is still experimental. Both name and interface might change in future versions.
```

checkVignettes Check Package Vignettes

```

\section*{Description}

Check all Sweave files of a package by running Sweave and/or Stangle on them. All R source code files found after the tangling step are sourceed to check whether all code can be executed without errors.
```

Usage
checkVignettes(package, dir, lib.loc = NULL, tangle = TRUE,
weave = TRUE, workdir = c("tmp", "src", "cur"),
keepfiles = FALSE)

```

\section*{Arguments}
package a character string naming an installed package. If given, Sweave files are searched in subdirectory doc.
dir a character string specifying the path to a package's root source directory. This subdirectory inst/doc is searched for Sweave files.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. The specified library trees are used to to search for package.
tangle Perform a tangle and source the extraced code?
weave Perform a weave?
workdir Directory used as working directory while checking the vignettes. If "tmp" then a temporary directory is created, this is the default. If "src" then the directory containing the vignettes itself is used, if "cur" then the current working directory of R is used.
keepfiles Delete file in temporary directory? This option is ignored when workdir!="tmp".

\section*{Value}

An object of class "checkVignettes" which is a list with the error messages found during the tangle and weave steps. There is a print method for nicely displaying the information contained in such objects.
```

codoc Check Code/Documentation Consistency

```

\section*{Description}

Find inconsistencies between actual and documented usage of R function objects in a package, by comparing names and optionally also corresponding positions and default values of the arguments of the functions.
```

Usage
codoc(package, dir, lib.loc = NULL,
use.values = FALSE, use.positions = TRUE,
ignore.generic.functions = FALSE,
verbose = getOption("verbose"))

```

\section*{Arguments}
\[
\begin{array}{ll}
\text { package } & \text { a character string naming an installed package. } \\
\text { dir } & \begin{array}{l}
\text { a character string specifying the path to a package's root source directory. } \\
\text { This must contain the subdirectories 'man' with R documentation sources } \\
\text { (in Rd format) and ' } R \text { ' with R code. Only used if package is not given. }
\end{array} \\
\text { lib.loc } & \begin{array}{l}
\text { a character vector of directory names of R libraries, or NULL. The default } \\
\text { value of NULL corresponds to all libraries currently known. The specified } \\
\text { library trees are used to to search for package. }
\end{array} \\
\text { use.positions } \begin{array}{ll}
\text { a logical indicating whether to use the positions of function arguments } \\
\text { when comparing. }
\end{array} \\
\text { use.values } \quad \begin{array}{l}
\text { a logical indicating whether to use function default values when comparing } \\
\text { code and docs. }
\end{array} \\
\text { ignore.generic.functions } \\
\text { if TRUE, functions the body of which contains "UseMethod" are ignored. } \\
\text { verbose } & \begin{array}{l}
\text { a logical. If TRUE, additional diagnostics are printed. }
\end{array}
\end{array}
\]

\section*{Details}

The purpose of this function is to check whether the documented usage of function objects agrees with their formal arguments as defined in the R code. This is not always straightforward, in particular as the usage information for methods to generic functions often employs the name of the generic rather than the method.
The following algorithm is used. If an installed package is used, it is loaded (unless it is the base package), after possibly detaching an already loaded version of the package. Otherwise, if the sources are used, the R code files of the package are collected and sourced in a new environment. Then, the usage sections of the Rd files are extracted and manipulated in order to give function stubs corresponding to the indicated usage, which are then sourced in another new environment. For interpreted functions in both the code and docs environment, the formals are compared according to the values of the arguments use.positions and use.values.

Currently, synopsis sections are used, but multiple usage examples (such as in abline) are not combined when building the stubs. Occurences of synopsis sections are reported if verbose is true.

\section*{Value}

An object of class "codoc" which is a list the names of which are the names of the functions where an inconsistency was found. The elements of the list are lists of length 2 with elements code and docs, giving the corresponding arguments obtained from the function's code and documented usage.
There is a print method for nicely displaying the information contained in such objects.

\author{
See Also
}
undoc

\section*{QA}
\(Q A\) Checks for \(R\) Code and/or Documentation

\section*{Description}

Functions for performing various quality checks.

\section*{Usage}
```

checkAssignFuns(package, dir, lib.loc = NULL)
checkDocArgs(package, dir, lib.loc = NULL)
checkDocStyle(package, dir, lib.loc = NULL)
checkMethods(package, dir, lib.loc = NULL)

```

\section*{Arguments}
package a character string naming an installed package.
dir a character string specifying the path to a package's root source directory. This should contain the subdirectories R (for R code) and 'man' with R documentation sources (in Rd format). Only used if package is not given.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. The specified library trees are used to to search for package.

\section*{Details}
checkAssignFuns checks whether assignment functions in the package R code have their final argument named value.
checkDocArgs checks, for all Rd files in a package, whether all arguments shown in the usage sections of the Rd file are documented in its arguments section. It also reports duplicated entries in the arguments section.
checkDocStyle investigates how (S3) methods are shown in the usages of the Rd files in a package. It reports if methods are shown along with their generic, which typically causes problems for the documentation of the primary argument of the generic. It also finds the methods shown by their full name (rather than that of the generic using the Rd \(\backslash\) method markup).
checkMethods checks whether all methods defined in the package \(R\) code have all arguments of the corresponding generic, with positional arguments of the generics in the same positions for the method. The generics are sought first in the given package and then in the base package. The rules when . . . is involved are subtle: see the source code.

If using an installed package, the checks needing access to all R objects of the package will load the package (unless it is the base package), after possibly detaching an already loaded version of the package.

\section*{Value}

The functions return objects of class the same as the respective function names containing the information about problems detected. There is a print method for nicely displaying the information contained in such objects.

\section*{Warning}

These functions are still experimental. Both names, interfaces and values might change in future versions.

\section*{Rtangle \(\quad R\) Driver for Stangle}

\section*{Description}

A driver for Stangle that extracts R code chunks.

\section*{Usage}
```

Rtangle()
RtangleSetup(file, syntax, output=NULL, annotate=TRUE, split=FALSE,
prefix=TRUE, quiet=FALSE)

```

\section*{Arguments}
\begin{tabular}{ll} 
file & Name of Sweave source file. \\
syntax & An object of class SweaveSyntax. \\
output & \begin{tabular}{l} 
Name of output file, default is to remove extension '.nw', '.Rnw' or '.Snw' \\
and to add extension '. \(R^{\prime}\). Any directory names in file are also removed \\
such that the output is created in the current working directory.
\end{tabular} \\
annotate & \begin{tabular}{l} 
By default, code chunks are seperated by comment lines specifying the \\
names and numbers of the code chunks. If FALSE, only the code chunks \\
without any decorating comments are extracted.
\end{tabular} \\
split & \begin{tabular}{l} 
Split output in single files per code chunk?
\end{tabular} \\
prefix & \begin{tabular}{l} 
If split=TRUE, prefix the chunk labels by the basename of the input file \\
to get output file names?
\end{tabular} \\
quiet & If TRUE all progress messages are suppressed.
\end{tabular}

\section*{Author(s)}

Friedrich Leisch

\section*{References}

Friedrich Leisch: Sweave User Manual, 2002
http://www.ci.tuwien.ac.at/~1eisch/Sweave

\section*{See Also}

Sweave, RweaveLatex
```

RweaveLatex R/LaTeX Driver for Sweave

```

\section*{Description}

A driver for Sweave that translates R code chunks in LaTeX files.

\section*{Usage}
```

RweaveLatex()
RweaveLatexSetup(file, syntax, output=NULL, quiet=FALSE, debug=FALSE,
echo=TRUE, eval = TRUE, split=FALSE, stylepath=TRUE,
pdf=TRUE, eps=TRUE)

```

\section*{Arguments}
\begin{tabular}{|c|c|}
\hline file & Name of Sweave source file. \\
\hline syntax & An object of class SweaveSyntax. \\
\hline output & Name of output file, default is to remove extension '.nw', '.Rnw' or '.Snw' and to add extension '.tex'. Any directory names in file are also removed such that the output is created in the current working directory. \\
\hline quiet & If TRUE all progress messages are suppressed. \\
\hline debug & If TRUE, input and output of all code chunks is copied to the console. \\
\hline stylepath & If TRUE, a hard path to the file 'Sweave.sty' installed with this package is set, if FALSE, only \usepackage\{Sweave\} is written. The hard path makes the TeX file less portable, but avoids the problem of installing the current version of 'Sweave.sty' to some place in your TeX input path. The argument is ignored if a \usepackage\{Sweave\} is already present in the Sweave source file. \\
\hline echo & set default for option echo, see details below. \\
\hline eval & set default for option eval, see details below. \\
\hline split & set default for option split, see details below. \\
\hline pdf & set default for option pdf, see details below. \\
\hline eps & set default for option eps, see details below. \\
\hline
\end{tabular}

\section*{Supported Options}

RweaveLatex supports the following options for code chunks (the values in parentheses show the default values):
echo: logical (TRUE). Include \(S\) code in the output file?
eval: logical (TRUE). If FALSE, the code chunk is not evaluated, and hence no text or graphical output produced.
results: character string (verbatim). If verbatim, the output of \(S\) commands is included in the verbatim-like Soutput environment. If tex, the output is taken to be already proper latex markup and included as is. If hide then all output is completely suppressed (but the code executed during the weave).
print: logical (FALSE) If TRUE, each expression in the code chunk is wrapped into a print () statement before evaluation, such that the values of all expressions become visible.
term: logical (TRUE). If TRUE, visibility of values emulates an interactive R session: values of assignments are not printed, values of single objects are printed. If FALSE, output comes only from explicit print or cat statements.
split: logical (FALSE). If TRUE, text output is written to separate files for each code chunk.
strip.white: logical (TRUE). If TRUE, blank lines at the beginning and end of output are removed.
prefix: logical (TRUE). If TRUE generated filenames of figures and output have a common prefix.
prefix.string: a character string, default is the name of the '.Snw' source file.
include: logical (TRUE), indicating whether input statements for text output and includegraphics statements for figures should be auto-generated. Use include=FALSE if the output should appear in a different place than the code chunk (by placing the input line manually).
fig: logical (FALSE), indicating whether the code chunk produces graphical output. Note that only one figure per code chunk can be processed this way.
eps: logical (TRUE), indicating whether EPS figures shall be generated. Ignored if fig=FALSE.
pdf: logical (TRUE), indicating whether PDF figures shall be generated. Ignored if fig=FALSE.
width: numeric (6), width of figures in inch.
height: numeric (6), height of figures in inch.

\section*{Author(s)}

Friedrich Leisch

\section*{References}

Friedrich Leisch: Sweave User Manual, 2002
http://www.ci.tuwien.ac.at/~1eisch/Sweave

\section*{See Also}
```

Sweave, Rtangle

```

\section*{Sweave Automatic Generation of Reports}

\section*{Description}

Sweave provides a flexible framework for mixing text and \(S\) code for automatic report generation. The basic idea is to replace the S code with its output, such that the final document only contains the text and the output of the statistical anlysis.

\section*{Usage}
```

Sweave(file, driver=RWeaveLatex(), syntax=getOption("SweaveSyntax"), ...)
Stangle(file, driver=RTangle(), syntax=getOption("SweaveSyntax"), ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
file & Name of Sweave source file. \\
driver & The actual workhorse, see details below. \\
syntax & An object of class SweaveSyntax or a character string with its \\
name. The default installation provides SweaveSyntaxNoweb and \\
& SweaveSyntaxLatex. \\
\(\ldots\) & Further arguments passed to the driver's setup function.
\end{tabular}

\section*{Details}

Automatic generation of reports by mixing word processing markup (like latex) and S code. The \(S\) code gets replaced by its output (text or graphs) in the final markup file. This allows to re-generate a report if the input data change and documents the code to reproduce the analysis in the same file that also produces the report.

Sweave combines the documentation and code chunks together (or their output) into a single document. Stangle extracts only the code from the Sweave file creating a valid S source file (that can be run using source). Code inside \(\backslash \operatorname{Sexpr}\}\) statements is ignored by Stangle.

Stangle is just a frontend to Sweave using a simple driver by default, which discards the documentation and concatenates all code chunks the current S engine understands.

\section*{Hook Functions}

Before each code chunk is evaluated, a number of hook functions can be executed. If getOption("SweaveHooks") is set, it is taken to be a collection of hook functions. For each logical option of a code chunk (echo, print, ...) a hook can be specified, which is executed if and only if the respective option is TRUE. Hooks must be named elements of the list returned by getOption("SweaveHooks") and be functions taking no arguments. E.g., if option "SweaveHooks" is defined as list (fig = foo), and foo is a function, then it would be executed before the code in each figure chunk. This is especially useful to set defaults for the graphical parameters in a series of figure chunks.

Note that the user is free to define new Sweave options and associate arbitrary hooks with them. E.g., one could define a hook function for option clean that removes all objects in the global environment. Then all code chunks with clean=TRUE would start operating on an empty workspace.

\section*{Syntax Definition}

Sweave allows a very flexible syntax framework for marking documentation and text chunks. The default is a noweb-style syntax, as alternative a latex-style syntax can be used. See the user manual for details.

\section*{Author(s)}

Friedrich Leisch

\section*{References}

Friedrich Leisch: Sweave User Manual, 2002
http://www.ci.tuwien.ac.at/~1eisch/Sweave

Friedrich Leisch: Dynamic generation of statistical reports using literate data analysis. In W. Härdle and B. Rönz, editors, Compstat 2002 - Proceedings in Computational Statistics, pages 575-580. Physika Verlag, Heidelberg, Germany, 2002. ISBN 3-7908-1517-9.

\author{
See Also \\ RweaveLatex, Rtangle
}

\section*{Examples}
```

testfile <- file.path(.path.package("tools"),
"Sweave", "Sweave-test-1.Rnw")

## create a LaTeX file

Sweave(testfile)

## create an S source file from the code chunks

Stangle(testfile)

## which can be simply sourced

source("Sweave-test-1.R")

```

SweaveSyntConv
Convert Sweave Syntax

\section*{Description}

This function converts the syntax of files in Sweave format to another Sweave syntax definition.

\section*{Usage}

SweaveSyntConv(file, syntax, output=NULL)

\section*{Arguments}
\begin{tabular}{ll} 
file & Name of Sweave source file. \\
syntax & An object of class SweaveSyntax or a character string with its name giving \\
the target syntax to which the file is converted. \\
output & \begin{tabular}{l} 
Name of output file, default is to remove the extension from the input \\
file and to add the default extension of the target syntax. Any directory \\
names in file are also removed such that the output is created in the \\
current working directory.
\end{tabular}
\end{tabular}

\section*{Author(s)}

Friedrich Leisch

\section*{References}

Friedrich Leisch: Sweave User Manual, 2002
http://www.ci.tuwien.ac.at/~1eisch/Sweave

\section*{See Also}

RweaveLatex, Rtangle

\section*{Examples}
```

testfile <- file.path(.path.package("tools"),
"Sweave", "Sweave-test-1.Rnw")

## convert the file to latex syntax

SweaveSyntConv(testfile, SweaveSyntaxLatex)

## and run it through Sweave

Sweave("Sweave-test-1.Stex")

```
    tools-internal Internal tools functions

\section*{Description}

Internal tools functions.

\section*{Usage}
sQuote(s)
.convertFilePathToAbsolute(path)
.listFilesWithExts(dir, exts, path = TRUE)
.loadPackageQuietly(package, lib.loc)
.makeS3MethodsStopList(package)
.sourceAssignments(file, envir)

\section*{Details}

These are not to be called by the user.

\section*{undoc}

Find Undocumented Objects

\section*{Description}

Finds the objects in a package which are undocumented, in the sense that they are visible to the user (or data objects or S4 classes provided by the package), but no documentation entry exists.

\section*{Usage}
undoc(package, dir, lib.loc = NULL)

\section*{Arguments}
package a character string naming an installed package.
dir a character string specifying the path to a package's root source directory. This must contain the subdirectory 'man' with R documentation sources (in \(R d\) format), and at least one of the ' \(R\) ' or 'data' subdirectories with \(R\) code or data objects, respectively.
lib.loc a character vector of directory names of R libraries, or NULL. The default value of NULL corresponds to all libraries currently known. The specified library trees are used to to search for package.

\section*{Details}

This function is useful for package maintainers mostly. In principle, all user level R objects should be documented; note however that the precise rules for documenting methods of generic functions are still under discussion.

\section*{Value}

An object of class "undoc" which is a list of character vectors containing the names of the undocumented objects split according to documentation type. This representation is still experimental, and might change in future versions.
There is a print method for nicely displaying the information contained in such objects.

\section*{Examples}
```

undoc("eda") \# Undocumented objects in 'eda'

```

\section*{Chapter 13}

\section*{The ts package}

Auto- and Cross- Covariance and -Correlation Function Estimation

\section*{Description}

The function acf computes (and by default plots) estimates of the autocovariance or autocorrelation function. Function pacf is the function used for the partial autocorrelations. Function ccf computes the cross-correlation or cross-covariance of two univariate series.

\section*{Usage}
```

acf(x, lag.max = NULL,
type = c("correlation", "covariance", "partial"),
plot = TRUE, na.action = na.fail, demean = TRUE, ...)
pacf(x, lag.max = NULL, plot = TRUE, na.action = na.fail, ...)
ccf(x, y, lag.max = NULL, type = c("correlation", "covariance"),
plot = TRUE, na.action = na.fail, ...)

```

\section*{Arguments}
\begin{tabular}{ll}
\(\mathrm{x}, \mathrm{y}\) & \begin{tabular}{l} 
a univariate or multivariate (not ccf) time series object or a numeric \\
vector or matrix.
\end{tabular} \\
lag.max & \begin{tabular}{l} 
maximum number of lags at which to calculate the acf. Default is \\
\(10 \log _{10}(N)\) where \(N\) is the number of observations.
\end{tabular} \\
type & \begin{tabular}{l} 
character string giving the type of acf to be computed. Allowed values \\
are "correlation" (the default), "covariance" or "partial".
\end{tabular} \\
plot & \begin{tabular}{l} 
logical. If TRUE the acf is plotted.
\end{tabular} \\
na.action & \begin{tabular}{l} 
function to be called to handle missing values. na.pass can be used. \\
demean
\end{tabular} \\
logical. Should the covariances be about the sample means?
\end{tabular}

\section*{Details}

For type = "correlation" and "covariance", the estimates are based on the sample covariance.
By default, no missing values are allowed. If the na. action function passes through missing values (as na.pass does), the covariances are computed from the complete cases. This means that the estimate computed may well not be a valid autocorrelation sequence, and may contain missing values. Missing values are not allowed when computing the PACF of a multivariate time series.

The partial correlation coefficient is estimated by fitting autoregressive models of successively higher orders up to lag.max.
The generic function plot has a method for objects of class "acf".
The lag is returned and plotted in units of time, and not numbers of observations.

\section*{Value}

An object of class "acf ", which is a list with the following elements:
\begin{tabular}{ll} 
lag & A three dimensional array containing the lags at which the acf is esti- \\
mated. \\
acf & An array with the same dimensions as lag containing the estimated acf. \\
type & The type of correlation (same as the type argument). \\
n.used & The number of observations in the time series. \\
series & The name of the series \(x\). \\
snames & The series names for a multivariate time series.
\end{tabular}

The result is returned invisibly if plot is TRUE.

\section*{Author(s)}

Original: Paul Gilbert, Martyn Plummer. Extensive modifications and univariate case of pacf by B.D. Ripley.

\section*{See Also}
```

plot.acf

```

\section*{Examples}
```


## Examples from Venables \& Ripley

data(lh)
acf(lh)
acf(lh, type = "covariance")
pacf(lh)
data(UKLungDeaths)
acf(ldeaths)
acf(ldeaths, ci.type = "ma")
acf(ts.union(mdeaths, fdeaths))
ccf(mdeaths, fdeaths) \# just the cross-correlations.
data(presidents) \# contains missing values
acf(presidents, na.action = na.pass)
pacf(presidents, na.action = na.pass)

```

\section*{acf2AR}

\section*{Description}

Compute an AR process exactly fitting an autocorrelation function.

\section*{Usage}
acf 2 AR (acf)

\section*{Arguments}
acf An autocorrelation or autocovariance sequence.

\section*{Value}

A matrix, with one row for the computed \(\mathrm{AR}(\mathrm{p})\) coefficients for \(1<=\mathrm{p}<=\) length(acf).

\section*{Author(s)}
B. D. Ripley

\section*{See Also}

ARMAacf, ar. yw which does this from an empirical ACF.

\section*{Examples}
```

(Acf <- ARMAacf (c(0.6, 0.3, -0.2)))
acf2AR(Acf)

```

AirPassengers Monthly Airline Passenger Numbers 1949-1960

\section*{Description}

The classic Box \& Jenkins airline data. Monthly totals of international airline passengers, 1949 to 1960 .

\section*{Usage}
data(AirPassengers)

\section*{Format}

A monthly time series, in thousands.

\section*{Source}

Box, G. E. P., Jenkins, G. M. and Reinsel, G. C. (1976) Time Series Analysis, Forecasting and Control. Third Edition. Holden-Day. Series G.

\section*{Examples}
```


## These are quite slow and so not run by example(AirPassengers)

data(AirPassengers)

## The classic 'airline model', by full ML

(fit <- arima(log10(AirPassengers), c(0, 1, 1),
seasonal = list(order=c(0, 1 ,1), period=12)))
update(fit, method = "CSS")
update(fit, x=window(log10(AirPassengers), start = 1954))
pred <- predict(fit, n.ahead = 24)
tl <- pred$pred - 1.96 * pred$se
tu <- pred$pred + 1.96 * pred$se
ts.plot(AirPassengers, 10^tl, 10^tu, log = "y", lty = c(1,2,2))

## full ML fit is the same if the series is reversed, CSS fit is not

ap0 <- rev(log10(AirPassengers))
attributes(ap0) <- attributes(AirPassengers)
arima(ap0, c(0, 1, 1), seasonal = list(order=c(0, 1 ,1), period=12))
arima(ap0, c(0, 1, 1), seasonal = list(order=c(0, 1 ,1), period=12),
method = "CSS")

## Structural Time Series

ap <- log10(AirPassengers) - 2
(fit <- StructTS(ap, type= "BSM"))
par(mfrow=c(1,2))
plot(cbind(ap, fitted(fit)), plot.type = "single")
plot(cbind(ap, tsSmooth(fit)), plot.type = "single")

```

\section*{Description}

Fit an autoregressive time series model to the data, by default selecting the complexity by AIC.

\section*{Usage}
```

ar(x, aic = TRUE, order.max = NULL,
method=c("yule-walker", "burg", "ols", "mle", "yw"), na.action,
series, ...)
ar.burg(x, aic = TRUE, order.max = NULL,
na.action = na.fail, demean = TRUE, series, var.method = 1, ...)
ar.yw(x, aic = TRUE, order.max = NULL,
na.action = na.fail, demean = TRUE, series, ...)
ar.mle(x, aic = TRUE, order.max = NULL, na.action = na.fail, demean = TRUE,
series, ...)
predict(object, newdata, n.ahead = 1, se.fit = TRUE, ...)

```

\section*{Arguments}
x
aic
order.max Maximum order (or order) of model to fit. Defaults to \(10 \log _{10}(N)\) where \(N\) is the number of observations except for method="mle" where it is the minimum of this quantity and 12 .
method Character string giving the method used to fit the model. Must be one of the strings in the default argument (the first few characters are sufficient). Defaults to "yule-walker".
na.action function to be called to handle missing values.
demean should a mean be estimated during fitting?
series names for the series. Defaults to deparse (substitute(x)).
\(\begin{array}{ll}\text { series } & \text { names for the series. Defaults to deparse (substitute (x)). } \\ \text { var.method } & \text { the method to estimate the innovations variance (see Details). }\end{array}\)
... additional arguments for specific methods.
object
newdata data to which to apply the prediction.
n. ahead number of steps ahead at which to predict.
se.fit
A univariate or multivariate time series.
Logical flag. If TRUE then the Akaike Information Criterion is used to choose the order of the autoregressive model. If FALSE, the model of order order.max is fitted.
a fit from ar.
logical: return estimated standard errors of the prediction error?

\section*{Details}

For definiteness, note that the AR coefficients have the sign in
\[
x_{t}-\mu=a_{1}\left(x_{t-1}-\mu\right)+\cdots+a_{p}\left(x_{t-p}-\mu\right)+e_{t}
\]
ar is just a wrapper for the functions ar. yw, ar.burg, ar.ols and ar.mle.
Order selection is done by AIC if aic is true. This is problematic, as of the methods here only ar.mle performs true maximum likelihood estimation. The AIC is computed as if the variance estimate were the MLE, omitting the determinant term from the likelihood. Note that this is not the same as the Gaussian likelihood evaluated at the estimated parameter values. In ar. yw the variance matrix of the innovations is computed from the fitted coefficients and the autocovariance of x .
ar. burg allows two methods to estimate the innovations variance and hence AIC. Method 1 is to use the update given by the Levinson-Durbin recursion (Brockwell and Davis, 1991, (8.2.6) on page 242), and follows S-PLUS. Method 2 is the mean of the sum of squares of the forward and backward prediction errors (as in Brockwell and Davis, 1996, page 145). Percival and Walden (1998) discuss both. In the multivariate case the estimated coefficients will depend (slightly) on the variance estimation method.

Remember that ar includes by default a constant in the model, by removing the overall mean of \(x\) before fitting the AR model, or (ar.mle) estimating a constant to subtract.

\section*{Value}

For ar and its methods a list of class "ar" with the following elements:
order The order of the fitted model. This is chosen by minimizing the AIC if aic=TRUE, otherwise it is order.max.
ar Estimated autoregression coefficients for the fitted model.
var.pred The prediction variance: an estimate of the portion of the variance of the time series that is not explained by the autoregressive model.
\(x\).mean The estimated mean of the series used in fitting and for use in prediction.
x .intercept (ar.ols only.) The intercept in the model for \(\mathrm{x}-\mathrm{x}\). mean.
aic The value of the aic argument.
n.used The number of observations in the time series.
order.max The value of the order.max argument.
partialacf The estimate of the partial autocorrelation function up to lag order.max.
resid residuals from the fitted model, conditioning on the first order observations. The first order residuals are set to NA. If x is a time series, so is resid.
method The value of the method argument.
series The name(s) of the time series.
asy.var.coef (univariate case.) The asymptotic-theory variance matrix of the coefficient estimates.

For predict.ar, a time series of predictions, or if se.fit = TRUE, a list with components pred, the predictions, and se, the estimated standard errors. Both components are time series.

\section*{Note}

Only the univariate case of ar.mle is implemented.
Fitting by method="mle" to long series can be very slow.

\section*{Author(s)}

Martyn Plummer. Univariate case of ar.yw, ar.mle and C code for univariate case of ar. burg by B. D. Ripley.

\section*{References}

Brockwell, P. J. and Davis, R. A. (1991) Time Series and Forecasting Methods. Second edition. Springer, New York. Section 11.4.
Brockwell, P. J. and Davis, R. A. (1996) Introduction to Time Series and Forecasting. Springer, New York. Sections 5.1 and 7.6.
Percival, D. P. and Walden, A. T. (1998) Spectral Analysis for Physical Applications. Cambridge University Press.

Whittle, P. (1963) On the fitting of multivariate autoregressions and the approximate canonical factorization of a spectral density matrix. Biometrika 40, 129-134.

\section*{See Also}
ar.ols, arima0 for ARMA models.

\section*{Examples}
```

data(lh)
ar(lh)
ar(lh, method="burg")
ar(lh, method="ols")
ar(lh, FALSE, 4) \# fit ar(4)
data(sunspot)
(sunspot.ar <- ar(sunspot.year))
predict(sunspot.ar, n.ahead=25)

## try the other methods too

data(BJsales)
ar(ts.union(BJsales, BJsales.lead))

## Burg is quite different here, as is OLS (see ar.ols)

ar(ts.union(BJsales, BJsales.lead), method="burg")

```

Fit Autoregressive Models to Time Series by OLS

\section*{Description}

Fit an autoregressive time series model to the data by ordinary least squares, by default selecting the complexity by AIC.

\section*{Usage}
```

ar.ols(x, aic = TRUE, order.max = NULL, na.action = na.fail,
demean = TRUE, intercept = demean, series, ...)

```

\section*{Arguments}

\section*{\(x \quad\) A univariate or multivariate time series.}
aic Logical flag. If TRUE then the Akaike Information Criterion is used to choose the order of the autoregressive model. If FALSE, the model of order order.max is fitted.
order.max Maximum order (or order) of model to fit. Defaults to \(10 \log _{10}(N)\) where \(N\) is the number of observations.
na.action function to be called to handle missing values.
demean should the AR model be for x minus its mean?
intercept should a separate intercept term be fitted?
series names for the series. Defaults to deparse(substitute(x)).
... further arguments to be passed to or from methods.

\section*{Details}
ar.ols fits the general AR model to a possibly non-stationary and/or multivariate system of series x . The resulting unconstrained least squares estimates are consistent, even if some of the series are non-stationary and/or co-integrated. For definiteness, note that the AR coefficients have the sign in
\[
x_{t}-\mu=a_{0}+a_{1}\left(x_{t-1}-\mu\right)+\cdots+a_{p}\left(x_{t-p}-\mu\right)+e_{t}
\]
where \(a_{0}\) is zero unless intercept is true, and \(\mu\) is the sample mean if demean is true, zero otherwise.
Order selection is done by AIC if aic is true. This is problematic, as ar.ols does not perform true maximum likelihood estimation. The AIC is computed as if the variance estimate (computed from the variance matrix of the residuals) were the MLE, omitting the determinant term from the likelihood. Note that this is not the same as the Gaussian likelihood evaluated at the estimated parameter values.
Some care is needed if intercept is true and demean is false. Only use this is the series are roughly centred on zero. Otherwise the computations may be inaccurate or fail entirely.

\section*{Value}

A list of class "ar" with the following elements:
order The order of the fitted model. This is chosen by minimizing the AIC if aic=TRUE, otherwise it is order.max.
ar Estimated autoregression coefficients for the fitted model.
var.pred The prediction variance: an estimate of the portion of the variance of the time series that is not explained by the autoregressive model.
\(x . m e a n \quad\) The estimated mean (or zero if demean is false) of the series used in fitting and for use in prediction.
x .intercept The intercept in the model for \(\mathrm{x}-\mathrm{x}\).mean, or zero if intercept is false. aic The value of the aic argument.
n.used The number of observations in the time series.
order.max The value of the order.max argument.
partialacf NULL. For compatibility with ar.
resid residuals from the fitted model, conditioning on the first order observations. The first order residuals are set to NA. If x is a time series, so is resid.
method The character string "Unconstrained LS".
series The name(s) of the time series.
asy.se.coef The asymptotic-theory standard errors of the coefficient estimates.

\section*{Author(s)}

Adrian Trapletti, Brian Ripley.

\section*{References}

Luetkepohl, H. (1991): Introduction to Multiple Time Series Analysis. Springer Verlag, NY, pp. 368-370.

See Also
ar

\section*{Examples}
```

data(lh)
ar(lh, method="burg")
ar.ols(lh)
ar.ols(lh, FALSE, 4) \# fit ar(4)
data(BJsales)
ar.ols(ts.union(BJsales, BJsales.lead))
data(EuStockMarkets)
x <- diff(log(EuStockMarkets))
ar.ols(x, order.max=6, demean=FALSE, intercept=TRUE)

```

\section*{arima ARIMA Modelling of Time Series}

\section*{Description}

Fit an ARIMA model to a univariate time series.

\section*{Usage}
```

arima(x, order $=c(0,0,0)$,
seasonal = list(order $=c(0,0,0)$, period $=N A)$,
xreg = NULL, include.mean = TRUE, transform.pars = TRUE,
fixed = NULL, init = NULL, method = c("CSS-ML", "ML", "CSS"),
n.cond, optim.control = list(), kappa $=1 \mathrm{e} 6$ )

```

\section*{Arguments}
\begin{tabular}{ll}
x & a univariate time series \\
order & \begin{tabular}{l} 
A specification of the non-seasonal part of the ARIMA model: the three \\
components \((p, d, q)\) are the AR order, the degree of differencing, and the \\
MA order.
\end{tabular} \\
seasonal & \begin{tabular}{l} 
A specification of the seasonal part of the ARIMA model, plus the period \\
(which defaults to frequency (x)). This should be a list with components \\
order and period, but a specification of just a numeric vector of length \\
3 will be turned into a suitable list with the specification as the order.
\end{tabular} \\
xreg & \begin{tabular}{l} 
Optionally, a vector or matrix of external regressors, which must have the \\
same number of rows as x.
\end{tabular} \\
include.mean & \begin{tabular}{l} 
Should the ARIMA model include a mean term? The default is TRUE for \\
undifferenced series, FALSE for differenced ones (where a mean would not \\
affect the fit nor predictions).
\end{tabular}
\end{tabular}
transform.pars
Logical. If true, the AR parameters are transformed to ensure that they remain in the region of stationarity. Not used for method = "CSS".
fixed optional numeric vector of the same length as the total number of parameters. If supplied, only non-NA entries in fixed will be varied. transform.pars \(=\) TRUE will be overridden if any AR parameters are fixed.
init optional numeric vector of initial parameter values. Missing values will be filled in, by zeroes except for regression coefficients.
method Fitting method: maximum likelihood or minimize conditional sum-ofsquares. The default (unless there are missing values) is to use conditional-sum-of-squares to find starting values, then maximum likelihood.
n.cond Only used if fitting by conditional-sum-of-squares: the number of initial observations to ignore. It will be ignored if less than the maximum lag of an AR term.
optim.control List of control parameters for optim.
kappa the prior variance (as a multiple of the innovations variance) for the past observations in a differenced model. Do not reduce this.

\section*{Details}

Different definitions of ARMA models have different signs for the AR and/or MA coefficients. The definition here has
\[
X_{t}=a_{1} X_{t-1}+\cdots+a_{p} X_{t-p}+e_{t}+b_{1} e_{t-1}+\ldots+b_{q} e_{t-q}
\]
and so the MA coefficients differ in sign from those of S-PLUS. Further, if include.mean is true, this formula applies to \(X-m\) rather than \(X\). For ARIMA models with differencing, the differenced series follows a zero-mean ARMA model.

The variance matrix of the estimates is found from the Hessian of the log-likelihood, and so may only be a rough guide.
Optimization is done by optim. It will work best if the columns in xreg are roughly scaled to zero mean and unit variance, but does attempt to estimate suitable scalings.

\section*{Value}

A list of class "Arima" with components:
\begin{tabular}{|c|c|}
\hline coef & a vector of AR, MA and regression coefficients, \\
\hline sigma2 & the MLE of the innovations variance. \\
\hline var.coef & the estimated variance matrix of the coefficients coef. \\
\hline loglik & the maximized log-likelihood (of the differenced data), or the approximation to it used. \\
\hline arma & A compact form of the specification, as a vector giving the number of AR, MA, seasonal AR and seasonal MA coefficients, plus the period and the number of non-seasonal and seasonal differences. \\
\hline aic & the AIC value corresponding to the log-likelihood. Only valid for method = "ML" fits. \\
\hline residuals & the standardized residuals. \\
\hline call & the matched call. \\
\hline series & the name of the series \(x\). \\
\hline convergence & the value returned by optim. \\
\hline n. cond & the number of initial observations not used in the fitting. \\
\hline model & A list representing the Kalman Filter used in the fitting. See KalmanLike. \\
\hline
\end{tabular}

\section*{Fitting methods}

The exact likelihood is computed via a state-space representation of the ARIMA process, and the innovations and their variance found by a Kalman filter. The initialization of the differenced ARMA process uses stationarity and is based on Gardner et al. (1980). For a differenced process the non-stationary components are given a diffuse prior (controlled by kappa). Observations which are still controlled by the diffuse prior (determined by having a Kalman gain of at least 1e4) are excluded from the likelihood calculations. (This gives comparable results to arima0 in the absence of missing values, when the observations excluded are precisely those dropped by the differencing.)
Missing values are allowed, and are handled exactly in method "ML".
If transform.pars is true, the optimization is done using an alternative parametrization which is a variation on that suggested by Jones (1980) and ensures that the model is stationary. For an \(\mathrm{AR}(\mathrm{p})\) model the parametrization is via the inverse tanh of the partial autocorrelations: the same procedure is applied (separately) to the AR and seasonal AR terms. The MA terms are not constrained to be invertible during optimization, but they will be converted to invertible form after optimization if transform.pars is true.
Conditional sum-of-squares is provided mainly for expositional purposes. This computes the sum of squares of the fitted innovations from observation n.cond on, (where n.cond is at least the maximum lag of an AR term), treating all earlier innovations to be zero. Argument n. cond can be used to allow comparability between different fits. The "part log-likelihood" is the first term, half the log of the estimated mean square. Missing values are allowed, but will cause many of the innovations to be missing.

When regressors are specified, they are orthogonalized prior to fitting unless any of the coefficients is fixed. It can be helpful to roughly scale the regressors to zero mean and unit variance.

\section*{Note}

The results are likely to be different from S-PLUS's arima.mle, which computes a conditional likelihood and does not include a mean in the model. Further, the convention used by arima.mle reverses the signs of the MA coefficients.
arima is very similar to arima0 for ARMA models or for differenced models without missing values, but handles differenced models with missing values exactly. It is somewhat slower than arima0, particularly for seasonally differenced models.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Brockwell, P. J. and Davis, R. A. (1996) Introduction to Time Series and Forecasting. Springer, New York. Sections 3.3 and 8.3.
Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press.
Gardner, G, Harvey, A. C. and Phillips, G. D. A. (1980) Algorithm AS154. An algorithm for exact maximum likelihood estimation of autoregressive-moving average models by means of Kalman filtering. Applied Statistics 29, 311-322.

Harvey, A. C. (1993) Time Series Models, 2nd Edition, Harvester Wheatsheaf, sections 3.3 and 4.4.

Jones, R. H. (1980) Maximum likelihood fitting of ARMA models to time series with missing observations. Technometrics 20 389-395.

\section*{See Also}
predict.Arima, tsdiag, arima0, ar

\section*{Examples}
```

data(lh)
arima(lh, order = c(1,0,0))
arima(lh, order =c(3,0,0))
arima(lh, order = c(1,0,1))
arima(lh, order = c(3,0,0), method = "CSS")
data(USAccDeaths)
arima(USAccDeaths, order = c(0,1,1), seasonal = list(order=c(0,1,1)))
arima(USAccDeaths, order = c(0,1,1), seasonal = list(order=c(0,1,1)),
method = "CSS") \# drops first 13 observations.

# for a model with as few years as this, we want full ML

data(LakeHuron)
arima(LakeHuron, order = c(2,0,0), xreg = time(LakeHuron)-1920)
data(presidents) \# contains NAs

## graphs in example(acf) suggest order 1 or 3

(fit1 <- arima(presidents, c(1, 0, 0)))
tsdiag(fit1)
(fit3 <- arima(presidents, c(3, 0, 0))) \# smaller AIC
tsdiag(fit3)

```
```

arima.sim Simulate from an ARIMA Model

```

\section*{Description}

Simulate from an ARIMA model.

\section*{Usage}
```

arima.sim(model, $n$, rand.gen $=$ rnorm, innov $=$ rand.gen( $n, \ldots$. .
n.start = NA, ...)

```

\section*{Arguments}
model A list with component ar and/or ma giving the AR and MA coeffcients respectively. Optionally a component order can be used.
n
length of output series.
rand.gen optional: a function to generate the innovations.
innov an optional times series of innovations. If not provided, rand.gen is used.
```

n.start length of "burn-in" period. If NA, the default, a reasonable value is com-
puted.
... additional arguments for rand.gen. Most usefully, the standard deviation
of the innovations generated by rnorm can be specified by sd.

```

\section*{Details}

The ARMA model is checked for stationarity.
ARIMA models are specified via the order component of model, in the same way as for arima. Other aspects of the order component are ignored.

\section*{Value}

A time-series object of class "ts".

\section*{See Also}
arima.sim

\section*{Examples}
```

arima.sim(n = 63, list(ar = c(0.8897, -0.4858), ma = c(-0.2279, 0.2488)),
sd = sqrt(0.1796))

# mildly long-tailed

arima.sim(n = 63, list(ar=c(0.8897, -0.4858), ma=c(-0.2279, 0.2488)),
rand.gen = function(n, ...) sqrt(0.1796) * rt(n, df = 5))

# An ARIMA simulation

ts.sim <- arima.sim(list(order = c(1,1,0), ar = 0.7), n = 200)
ts.plot(ts.sim)

```
arima0

ARIMA Modelling of Time Series - Preliminary Version

\section*{Description}

Fit an ARIMA model to a univariate time series, and forecast from the fitted model.

\section*{Usage}
```

$\operatorname{arima} 0(x$, order $=c(0,0,0)$,
seasonal $=$ list (order $=c(0,0,0)$, period $=N A)$,
xreg $=$ NULL, include.mean $=$ TRUE, delta $=0.01$,
transform.pars = TRUE, fixed = NULL, init = NULL,
method = c("ML", "CSS"), n.cond, optim.control = list())
predict (object, n.ahead = 1, newxreg, se.fit = TRUE, ...)

```
```

Arguments
x a univariate time series
order A specification of the non-seasonal part of the ARIMA model: the three
components (p,d,q) are the AR order, the degree of differencing, and the
MA order.
seasonal A specification of the seasonal part of the ARIMA model, plus the period
(which defaults to frequency (x)). This should be a list with components
order and period, but a specification of just a numeric vector of length
3 will be turned into a suitable list with the specification as the order.
xreg Optionally, a vector or matrix of external regressors, which must have the
same number of rows as x.
include.mean Should the ARIMA model include a mean term? The default is TRUE for
undifferenced series, FALSE for differenced ones (where a mean would not
affect the fit nor predictions).
delta A value to indicate at which point 'fast recursions' should be used. See
the Details section.
transform.pars
Logical. If true, the AR parameters are transformed to ensure that they remain in the region of stationarity. Not used for method = "CSS".
fixed optional numeric vector of the same length as the total number of parameters. If supplied, only non-NA entries in fixed will be varied. transform.pars $=$ TRUE will be overridden if any ARMA parameters are fixed.
init optional numeric vector of initial parameter values. Missing values will be filled in, by zeroes except for regression coefficients.
method Fitting method: maximum likelihood or minimize conditional sum-ofsquares.
n.cond Only used if fitting by conditional-sum-of-squares: the number of initial observations to ignore. It will be ignored if less than the maximum lag of an AR term.
optim.control List of control parameters for optim.
object, fit The result of an arima0 fit.
newxreg New values of xreg to be used for prediction. Must have at least n. ahead rows.
n.ahead The number of steps ahead for which prediction is required.
se.fit Logical: should standard errors of prediction be returned?
gof.lag Number of lags to be used in goodness-of-fit test.
... arguments passed to or from other methods.

```

\section*{Details}

Different definitions of ARMA models have different signs for the AR and/or MA coefficients. The definition here has
\[
X_{t}=a_{1} X_{t-1}+\cdots+a_{p} X_{t-p}+e_{t}+b_{1} e_{t-1}+\ldots+b_{q} e_{t-q}
\]
and so the MA coefficients differ in sign from those of S-PLUS. Further, if include.mean is true, this formula applies to \(X-m\) rather than \(X\). For ARIMA models with differencing, the differenced series follows a zero-mean ARMA model.

The variance matrix of the estimates is found from the Hessian of the log-likelihood, and so may only be a rough guide, especially for fits close to the boundary of invertibility.
Optimization is done by optim. It will work best if the columns in xreg are roughly scaled to zero mean and unit variance, but does attempt to estimate suitable scalings.

Finite-history prediction is used. This is only statistically efficient if the MA part of the fit is invertible, so predict.arima0 will give a warning for non-invertible MA models.

\section*{Value}

For arima0, a list of class "arima0" with components:
\(\left.\begin{array}{ll}\text { coef } & \text { a vector of AR, MA and regression coefficients, } \\
\text { sigma2 } & \text { the MLE of the innovations variance. } \\
\text { var.coef } & \begin{array}{l}\text { the estimated variance matrix of the coefficients coef. } \\
\text { toglik }\end{array} \\
\text { the maximized log-likelihood (of the differenced data), or the approxima- } \\
\text { tion to it used. }\end{array}\right]\)\begin{tabular}{l} 
A compact form of the specification, as a vector giving the number of AR, \\
MA, seasonal AR and seasonal MA coefficients, plus the period and the \\
number of non-seasonal and seasonal differences. \\
aic the AIC value corresponding to the log-likelihood. Only valid for method \\
\(=\) \\
= "ML" fits. Prior to R 1.5.0 this omitted 2 for the estimation of the \\
innovations variance.
\end{tabular}

For predict.arima0, a time series of predictions, or if se.fit \(=\) TRUE, a list with components pred, the predictions, and se, the estimated standard errors. Both components are time series.

\section*{Fitting methods}

The exact likelihood is computed via a state-space representation of the ARMA process, and the innovations and their variance found by a Kalman filter based on Gardner et al. (1980). This has the option to switch to 'fast recursions' (assume an effectively infinite past) if the innovations variance is close enough to its asymptotic bound. The argument delta sets the tolerance: at its default value the approximation is normally negligible and the speed-up considerable. Exact computations can be ensured by setting delta to a negative value.

If transform.pars is true, the optimization is done using an alternative parametrization which is a variation on that suggested by Jones (1980) and ensures that the model is stationary. For an \(\operatorname{AR}(\mathrm{p})\) model the parametrization is via the inverse tanh of the partial autocorrelations: the same procedure is applied (separately) to the AR and seasonal AR terms. The MA terms are also constrained to be invertible during optimization by the same transformation if transform.pars is true. Note that the MLE for MA terms does sometimes occur for MA polynomials with unit roots: such models can be fitted by using transform.pars = FALSE and specifying a good set of initial values (often obtainable from a fit with transform. pars = TRUE).

As from R 1.5.0 missing values are allowed, but any missing values will force delta to be ignored and full recursions used. Note that missing values will be propogated by differencing, so the procedure used in this function is not fully efficient in that case.
Conditional sum-of-squares is provided mainly for expositional purposes. This computes the sum of squares of the fitted innovations from observation n. cond on, (where n. cond is at least the maximum lag of an AR term), treating all earlier innovations to be zero. Argument n. cond can be used to allow comparability between different fits. The "part log-likelihood" is the first term, half the \(\log\) of the estimated mean square. Missing values are allowed, but will cause many of the innovations to be missing.

When regressors are specified, they are orthogonalized prior to fitting unless any of the coefficients is fixed. It can be helpful to roughly scale the regressors to zero mean and unit variance.

\section*{Note}

This is a preliminary version, and will be replaced by arima.
The standard errors of prediction exclude the uncertainty in the estimation of the ARMA model and the regression coefficients.

The results are likely to be different from S-PLUS's arima.mle, which computes a conditional likelihood and does not include a mean in the model. Further, the convention used by arima.mle reverses the signs of the MA coefficients.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Brockwell, P. J. and Davis, R. A. (1996) Introduction to Time Series and Forecasting. Springer, New York. Sections 3.3 and 8.3.
Gardner, G, Harvey, A. C. and Phillips, G. D. A. (1980) Algorithm AS154. An algorithm for exact maximum likelihood estimation of autoregressive-moving average models by means of Kalman filtering. Applied Statistics 29, 311-322.
Harvey, A. C. (1993) Time Series Models, 2nd Edition, Harvester Wheatsheaf, sections 3.3 and 4.4.

Harvey, A. C. and McKenzie, C. R. (1982) Algorithm AS182. An algorithm for finite sample prediction from ARIMA processes. Applied Statistics 31, 180-187.
Jones, R. H. (1980) Maximum likelihood fitting of ARMA models to time series with missing observations. Technometrics 20 389-395.

\section*{See Also}
arima, ar, tsdiag

\section*{Examples}
```

data(lh)
arima0(lh, order = c(1,0,0))
arima0(lh, order = c(3,0,0))
arima0(lh, order = c(1,0,1))
predict(arima0(lh, order = c(3,0,0)), n.ahead = 12)

```
```

arima0(lh, order = c(3,0,0), method = "CSS")
data(USAccDeaths)

# for a model with as few years as this, we want full ML

(fit <- arima0(USAccDeaths, order = c(0,1,1),
seasonal = list(order=c(0,1,1)), delta = -1))
predict(fit, n.ahead = 6)
data(LakeHuron)
arima0(LakeHuron, order = c(2,0,0), xreg = time(LakeHuron)-1920)
data(presidents) \# contains NAs

## graphs in example(acf) suggest order 1 or 3

(fit1 <- arima0(presidents, c(1, 0, 0), delta = -1)) \# avoid warning
tsdiag(fit1)
(fit3 <- arima0(presidents, c(3, 0, 0), delta = -1)) \# smaller AIC
tsdiag(fit3)

```

ARMAacf Compute Theoretical ACF for an ARMA Process

\section*{Description}

Compute the theoretical autocorrelation function or partial autocorrelation function for an ARMA process.

\section*{Usage}

ARMAacf(ar \(=\) numeric (0), ma \(=\) numeric (0), lag.max \(=r, \operatorname{pacf}=\) FALSE)

\section*{Arguments}
\begin{tabular}{ll} 
ar & numeric vector of AR coefficients \\
ma & numeric vector of MA coefficients \\
lag.max & \begin{tabular}{l} 
integer. Maximum lage required. Defaults to max \((\mathrm{p}, \mathrm{q}+1)\), where p, q \\
are the numbers of AR and MA terms respectively.
\end{tabular} \\
pacf & logical. Should the partial autocorrelations be returned?
\end{tabular}

\section*{Details}

The methods used follow Brockwell \& Davis (1991, section 3.3). Their equations (3.3.8) are solved for the autocovariances at lags \(0, \ldots, \max (p, q+1)\), and the remaining autocorrelations are given by a recursive filter.

\section*{Value}

A vector of (partial) autocorrelations, named by the lags.
Author(s)
B. D. Ripley

\section*{References}

Brockwell, P. J. and Davis, R. A. (1991) Time Series: Theory and Methods, Second Edition. Springer.

\section*{See Also}
arima, ARMAtoMA, filter.

\section*{Examples}
```

ARMAacf(c(1.0, -0.25), 1.0, lag.max = 10)

## Example from Brockwell \& Davis (1991, pp.92-4)

## answer 2^(-n) * (32/3 + 8 * n) /(32/3)

n <- 1:10; 2^(-n) * (32/3 + 8* n) /(32/3)
ARMAacf(c(1.0, -0.25), 1.0, lag.max = 10, pacf = TRUE)
ARMAacf(c(1.0, -0.25), lag.max = 10, pacf = TRUE)

```

\section*{ARMAtoMA Convert ARMA Process to Infinite MA Process}

\section*{Description}

Convert ARMA process to infinite MA process.

\section*{Usage}

ARMAtoMA(ar \(=\) numeric(0), ma \(=\) numeric(0), lag.max)

\section*{Arguments}
\begin{tabular}{ll} 
ar & numeric vector of AR coefficients \\
ma & numeric vector of MA coefficients \\
lag.max & Largest MA(Inf) coefficient required.
\end{tabular}

Value
A vector of coefficients.

Author(s)
B. D. Ripley

\section*{References}

Brockwell, P. J. and Davis, R. A. (1991) Time Series: Theory and Methods, Second Edition. Springer.

\section*{See Also}
arima, ARMAacf.

\section*{Examples}
```

ARMAtoMA(c(1.0, -0.25), 1.0, 10)

## Example from Brockwell \& Davis (1991, p.92)

## answer (1 + 3*n)*2^(-n)

n <- 1:10; (1 + 3*n)*2^(-n)

```
austres Quarterly Time Series of the Number of Australian Residents

\section*{Description}

Numbers (in thousands) of Australian residents measured quarterly from March 1971 to March 1994. The object is of class "ts".

\section*{Usage}
data(austres)

\section*{Source}
P. J. Brockwell and R. A. Davis (1996) Introduction to Time Series and Forecasting. Springer
beavers Body Temperature Series of Two Beavers

\section*{Description}

Reynolds (1994) describes a small part of a study of the long-term temperature dynamics of beaver Castor canadensis in north-central Wisconsin. Body temperature was measured by telemetry every 10 minutes for four females, but data from a one period of less than a day for each of two animals is used there.

\section*{Usage}
data(beavers)

\section*{Format}

The beaver 1 data frame has 114 rows and 4 columns on body temperature measurements at 10 minute intervals.
The beaver 2 data frame has 100 rows and 4 columns on body temperature measurements at 10 minute intervals.
The variables are as follows:
day Day of observation (in days since the beginning of 1990), December 12-13 (beaver1) and November 3-4 (beaver2).
time Time of observation, in the form 0330 for 3:30am temp Measured body temperature in degrees Celsius. activ Indicator of activity outside the retreat.

\section*{Note}

The observation at 22:20 is missing in beaver1.

\section*{Source}
P. S. Reynolds (1994) Time-series analyses of beaver body temperatures. Chapter 11 of Lange, N., Ryan, L., Billard, L., Brillinger, D., Conquest, L. and Greenhouse, J. eds (1994) Case Studies in Biometry. New York: John Wiley and Sons.

\section*{Examples}
```

data(beavers)
(yl <- range(beaver1$temp, beaver2$temp))
beaver.plot <- function(bdat, ...) {
nam <- deparse(substitute(bdat))
attach(bdat)
\# Hours since start of day:
hours <- time %/% 100 + 24*(day - day[1]) + (time %% 100)/60
plot (hours, temp, type = "l", ...,
main = paste(nam, "body temperature"))
abline(h = 37.5, col = "gray", lty = 2)
is.act <- activ == 1
points(hours[is.act], temp[is.act], col = 2, cex = . 8)
}
op <- par(mfrow = c(2,1), mar = c(3,3,4,2), mgp = .9* 2:0)
beaver.plot(beaver1, ylim = yl)
beaver.plot(beaver2, ylim = yl)
par(op)

```

BJsales Sales Data with Leading Indicator.

\section*{Description}

The sales time series BJsales and leading indicator BJsales.lead each contain 150 observations. The objects are of class "ts".

\section*{Usage}
```

data(BJsales)

```

\section*{Source}

The data are given in Box \& Jenkins (1976). Obtained from the Time Series Data Library at http://www-personal.buseco.monash.edu.au/~ \({ }^{\text {hyndman/TSDL/ }}\)

\section*{References}
G. E. P. Box and G. M. Jenkins (1976): Time Series Analysis, Forecasting and Control, Holden-Day, San Francisco, p. 537.
P. J. Brockwell and R. A. Davis (1991): Time Series: Theory and Methods, Second edition, Springer Verlag, NY, pp. 414.

Box.test
Box-Pierce and Ljung-Box Tests

\section*{Description}

Compute the Box-Pierce or Ljung-Box test statistic for examining the null hypothesis of independence in the time series x is computed.

\section*{Usage}

Box.test (x, lag = 1, type=c("Box-Pierce", "Ljung-Box"))

\section*{Arguments}
\(x \quad\) a numeric vector or univariate time series.
lag the statistic will be based on lag autocorrelation coefficients.
type test to be performed: partial matching is used.

\section*{Value}

A list with class "htest" containing the following components:
statistic the value of the test statistic.
parameter the degrees of freedom of the approximate chi-squared distribution of the test statistic.
p.value the p-value of the test.
method a character string indicating which type of test was performed.
data.name a character string giving the name of the data.

\section*{Note}

Missing values are not handled.

\section*{Author(s)}
A. Trapletti

\section*{References}

Box, G. E. P. and Pierce, D. A. (1970) Distribution of residual correlations in autoregressiveintegrated moving average time series models. Journal of the American Statistical Association 65, 1509-1526.
Ljung, G. M. and Box, G. E. P. (1978) On a measure of lack of fit in time series models. Biometrika 65, 553-564.
Harvey, A. C. (1993) Time Series Models, 2nd Edition, Harvester Wheatsheaf, NY, pp. 44, 45.

\section*{Examples}
```

x <- rnorm (100)
Box.test (x, lag = 1)
Box.test (x, lag = 1, type="Ljung")

```

\section*{Description}

Plots a cumulative periodogram.

\section*{Usage}
```

cpgram(ts, taper=0.1, main=
paste("Series: ", deparse(substitute(ts))), ci.col="blue")

```

\section*{Arguments}
\begin{tabular}{ll} 
ts & a univariate time series \\
taper & proportion tapered in forming the periodogram \\
main & main title \\
ci.col & colour for confidence band.
\end{tabular}

\section*{Value}

None.

\section*{Side Effects}

Plots the cumulative periodogram in a square plot.

\section*{Note}

From package 'MASS'.

\section*{Author(s)}
B.D. Ripley

\section*{Examples}
```

par(pty = "s", mfrow = c(1,2))
data(lh)
cpgram(lh)
lh.ar <- ar(lh, order.max = 9)
cpgram(lh.ar\$resid, main = "AR(3) fit to lh")
data(UKLungDeaths)
cpgram(ldeaths)

```
decompose Classical Seasonal Decomposition by Moving Averages

\section*{Description}

Decompose a time series into seasonal, trend and irregular components using moving averages. Deals with additive or multiplicative seasonal component.

\section*{Usage}
decompose(x, type = c("additive", "multiplicative"))

\section*{Arguments}
x
A time series.
type
The type of seasonal component.

\section*{Details}

The additive model used is:
\(\mathrm{Y}[\mathrm{t}]=\mathrm{T}[\mathrm{t}]+\mathrm{S}[\mathrm{t}]+\mathrm{e}[\mathrm{t}]\)
The multiplicative model used is:
\(\mathrm{Y}[\mathrm{t}]=\mathrm{T}[\mathrm{t}] * \mathrm{~S}[\mathrm{t}]+\mathrm{e}[\mathrm{t}]\)

Value
An object of class "decomposed.ts" with following components:
seasonal The seasonal component (i.e., the repeated seasonal figure)
figure The estimated seasonal figure only
trend The trend component
random The remainder part
type The value of type

\section*{Note}

The function stl provides a much more sophisticated decomposition.

\section*{Author(s)}

David Meyer 〈david.meyer@ci.tuwien.ac.at〉

\section*{See Also}
stl

\section*{Examples}
```

data(co2)
m <- decompose(co2)
m\$figure
plot(m)

```
diffinv Discrete Integrals: Inverse of Differencing

\section*{Description}

Computes the inverse function of the lagged differences function diff.

\section*{Usage}
```

diffinv(x, lag = 1, differences = 1,
xi = rep(0.0, lag*differences*NCOL(x)), ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & a numeric vector, matrix, or time series. \\
lag & a scalar lag parameter. \\
differences & an integer representing the order of the difference. \\
xi & \begin{tabular}{l} 
a numeric vector, matrix, or time series containing the initial values for \\
the integrals.
\end{tabular} \\
\(\ldots\) & arguments passed to or from other methods.
\end{tabular}

\section*{Details}
diffinv is a generic function with methods for class "ts" and default for vectors and matrices.

Missing values are not handled.

\section*{Value}

A numeric vector, matrix, or time series representing the discrete integral of x .

\section*{Author(s)}
A. Trapletti

\section*{See Also}
diff

\section*{Examples}
```

s <- 1:10
d <- diff(s)
diffinv(d, xi = 1)

```
```

embed Embedding a Time Series

```

\section*{Description}

Embeds the time series x into a low-dimensional Euclidean space.

\section*{Usage}
embed ( x , dimension \(=1\) )

\section*{Arguments}
\(\mathrm{x} \quad\) a numeric vector, matrix, or time series.
dimension a scalar representing the embedding dimension.

\section*{Details}

Each row of the resulting matrix consists of sequences \(x[t], x[t-1], \ldots, x[t-\) dimension +1 ], where \(t\) is the original index of \(x\). If \(x\) is a matrix, i.e., \(x\) contains more than one variable, then \(\mathrm{x}[\mathrm{t}]\) consists of the tth observation on each variable.

\section*{Value}

A matrix containing the embedded time series x .

\section*{Author(s)}
A. Trapletti, B.D. Ripley

\section*{Examples}
\(\mathrm{x}<-1: 10\)
embed ( \(x, 3\) )

EuStockMarkets Daily Closing Prices of Major European Stock Indices, 19911998.

\section*{Description}

Contains the daily closing prices of major European stock indices: Germany DAX (Ibis), Switzerland SMI, France CAC, and UK FTSE. The data are sampled in business time, i.e., weekends and holidays are omitted.

\section*{Usage}
data(EuStockMarkets)

\section*{Format}

A multivariate time series with 1860 observations on 4 variables. The object is of class "mts".

\section*{Source}

The data were kindly provided by Erste Bank AG, Vienna, Austria.
```

filter Linear Filtering on a Time Series

```

\section*{Description}

Applies linear filtering to a univariate time series or to each series separately of a multivariate time series.

\section*{Usage}
```

filter(x, filter, method = c("convolution", "recursive"),
sides = 2, circular = FALSE, init)

```

\section*{Arguments}
\begin{tabular}{|c|c|}
\hline x & a univariate or multivariate time series. \\
\hline filter & a vector of filter coefficients in reverse time order (as for AR or MA coefficients). \\
\hline method & Either "convolution" or "recursive" (and can be abbreviated). If "convolution" a moving average is used: if "recursive" an autoregression is used. \\
\hline sides & for convolution filters only. If sides=1 the filter coefficients are for past values only; if sides \(=2\) they are centred around lag 0 . In this case the length of the filter should be odd, but if it is even, more of the filter is forward in time than backward. \\
\hline circular & for convolution filters only. If TRUE, wrap the filter around the ends of the series, otherwise assume external values are missing (NA). \\
\hline init & for recursive filters only. Specifies the initial values of the time series just prior to the start value, in reverse time order. The default is a set of zeros. \\
\hline
\end{tabular}

\section*{Details}

Missing values are allowed in \(x\) but not in filter (where they would lead to missing values everywhere in the output).
Note that there is an implied coefficient 1 at lag 0 in the recursive filter, which gives
\[
y_{i}=x_{i}+f_{1} y_{i-1}+\cdots+f_{p} y_{i-p}
\]

No check is made to see if recursive filter is invertible: the output may diverge if it is not. The convolution filter is
\[
y_{i}=f_{1} x_{i+o}+\cdots+f_{p} x_{i+o-p-1}
\]
where o is the offset: see sides for how it is determined.

\section*{Value}

A time series object.

\section*{Note}
convolve(, type="filter") uses the FFT for computations and so may be faster for long filters on univariate series, but it does not return a time series (and so the time alignment is unclear), nor does it handle missing values. filter is faster for a filter of length 100 on a series of length 1000, for example.

\section*{Author(s)}
B.D. Ripley

\section*{See Also}
convolve, arima.sim

\section*{Examples}
```

x <- 1:100
filter(x, rep(1, 3))
filter(x, rep(1, 3), sides = 1)
filter(x, rep(1, 3), sides = 1, circular = TRUE)
data(presidents)
filter(presidents, rep(1,3))

```
```

HoltWinters Holt-Winters Filtering

```

\section*{Description}

Computes Holt-Winters Filtering of a given time series. Unknown parameters are determined by minimizing the squared prediction error.

\section*{Usage}
```

HoltWinters(x, alpha = NULL, beta = NULL, gamma = NULL,
seasonal = "additive", start.periods = 3,
l.start = NULL, b.start = NULL, s.start = NULL)

```

\section*{Arguments}
\begin{tabular}{ll}
x & An object of class ts \\
alpha & alpha parameter of Holt-Winters Filter \\
beta & \begin{tabular}{l} 
beta parameter of Holt-Winters Filter. If set to 0, the function will do \\
exponential smoothing.
\end{tabular} \\
gamma & \begin{tabular}{l} 
gamma parameter used for the seasonal component. If set to 0, an non- \\
seasonal model is fitted.
\end{tabular} \\
seasonal & \begin{tabular}{l} 
Selects an "additive" or "multiplicative" seasonal model. (Only takes \\
effect if gamma is non-zero).
\end{tabular}
\end{tabular}
start.periods Start periods used in the autodetection of start values. Must be at least 3.
1.start Start value for level (a[0]).
b.start Start value for trend (b[0]).
s.start Vector of start values for the seasonal component \(\left(s_{1}[0] \ldots s_{p}[0]\right)\)

\section*{Details}

The additive Holt-Winters prediction function (for time series with period length p) is
\[
\hat{Y}[t+h]=a[t]+h b[t]+s[t+1+(h-1) \bmod p],
\]
where \(a[t], b[t]\) and \(s[t]\) are given by
\[
\begin{gathered}
a[t]=\alpha(Y[t]-s[t-p])+(1-\alpha)(a[t-1]+b[t-1]) \\
b[t]=\beta(a[t]-a[t-1])+(1-\beta) b[t-1] \\
s[t]=\gamma(Y[t]-a[t])+(1-\gamma) s[t-p]
\end{gathered}
\]

The multiplicative Holt-Winters prediction function (for time series with period length p ) is
\[
\hat{Y}[t+h]=(a[t]+h b[t]) \times s[t+1+(h-1) \bmod p] .
\]
where \(a[t], b[t]\) and \(s[t]\) are given by
\[
\begin{gathered}
a[t]=\alpha(Y[t] / s[t-p])+(1-\alpha)(a[t-1]+b[t-1]) \\
b[t]=\beta(a[t]-a[t-1])+(1-\beta) b[t-1] \\
s[t]=\gamma(Y[t] / a[t])+(1-\gamma) s[t-p]
\end{gathered}
\]

The function tries to find the optimal values of \(\alpha\) and/or \(\beta\) and/or \(\gamma\) by minimizing the squared one-step prediction error if they are omitted.
For seasonal models, start values for \(\mathrm{a}, \mathrm{b}\) and s are detected by performing a simple decomposition in trend and seasonal component using moving averages (see function decompose) on the start.periods first periods (a simple linear regression on the trend component is used for starting level and trend.). For level/trend-models (no seasonal component), start values for a and b are x[2] and x[2] - x[1], respectively. For level-only models (ordinary exponential smoothing), the start value for \(a\) is \(x[1]\).

\section*{Value}

An object of class "HoltWinters", a list with components:
\begin{tabular}{ll} 
fitted & The filtered time series \\
x & The original series \\
alpha & \begin{tabular}{l} 
alpha used for filtering \\
beta
\end{tabular} \\
\begin{tabular}{l} 
beta used for filtering
\end{tabular} \\
coefficients & \begin{tabular}{l} 
A vector with named components a, b, s1, . . . sp containing the es- \\
timated values for the level, trend and seasonal components
\end{tabular} \\
seasonal & The specified seasonal-parameter \\
SSE & The final sum of squared errors achieved in optimizing \\
call & The call used
\end{tabular}

\section*{Author(s)}

David Meyer 〈david.meyer@ci.tuwien.ac.at〉

\section*{References}
C.C Holt (1957) Forecasting seasonals and trends by exponentially weighted moving averages, ONR Research Memorandum, Carnigie Institute 52.
P.R Winters (1960) Forecasting sales by exponentially weighted moving averages, Management Science 6, 324-342.

\section*{See Also}
predict.HoltWinters

\section*{Examples}
```

library(ts)
data(co2)
(m <- HoltWinters(co2))
plot(m)
data(AirPassengers)
(m <- HoltWinters(AirPassengers, seasonal = "mult"))
plot(m)
data(uspop)
x <- uspop + rnorm(uspop, sd = 5)
m <- HoltWinters(x, gamma = 0)
plot(m)
m2 <- HoltWinters(x, gamma = 0, beta = 0)
lines(fitted(m2), col = 3)

```

JohnsonJohnson Quarterly Earnings per Johnson \& Johnson Share

\section*{Description}

Quarterly earnings (dollars) per Johnson \& Johnson share 1960-80.

\section*{Usage}
data(JohnsonJohnson)

\section*{Format}

A quarterly time series

\section*{Source}

Shumway, R. H. and Stoffer, D. S. (2000) Time Series Analysis and its Applications. Second Edition. Springer. Example 1.1.

\section*{Examples}
```

data(JohnsonJohnson)
JJ <- log10(JohnsonJohnson)
plot(JJ)
(fit <- StructTS(JJ, type="BSM"))
tsdiag(fit)
sm <- tsSmooth(fit)
plot(cbind(JJ, sm[, 1], sm[, 3]-0.5), plot.type = "single",
col = c("black", "green", "blue"))
abline(h = -0.5, col = "grey60")
monthplot(fit)

```
KalmanLike Kalman Filtering

\section*{Description}

Use Kalman Filtering to find the (Gaussian) log-likelihood, or for forecasting or smoothing.

\section*{Usage}
```

KalmanLike(y, mod, nit = 0)
KalmanRun(y, mod, nit = 0)
KalmanSmooth(y, mod, nit = 0)
KalmanForecast(n.ahead = 10, mod)
makeARIMA(phi, theta, Delta, kappa = 1e6)

```

\section*{Arguments}
\(\mathrm{y} \quad \mathrm{a}\) univariate time series.
mod A list describing the state-space model: see Details.
nit The time at which the initialization is computed. nit \(=0\) implies that the initialization is for a one-step prediction, so Pn should not be computed at the first step.
n.ahead The number of steps ahead for which prediction is required.
phi, theta numeric vectors of length \(\geq 0\) giving AR and MA parameters.
Delta vector of differencing coefficients, so an ARMA model is fitted to y[t] -Delta[1]*y[t-1] - . ...
kappa the prior variance (as a multiple of the innovations variance) for the past observations in a differenced model.

\section*{Details}

These functions work with a general univariate state-space model with state vector a, transitions a <- \(\mathrm{T} \mathrm{a}+\mathrm{e}, e N(0, \kappa Q)\) and observation equation \(\mathrm{y}=\mathrm{Z}\) 'a +R eta, eta \(N(0, \kappa h)\). The likelihood is a profile likelihood after estimation of \(\kappa\).
The model is specified as a list with at least components
T the transition matrix

Z the observation coeficients
\(h\) the observation variance
V RQR'
a the current state estimate
P the current estimate of the state uncertainty matrix
Pn the estimate at time \(t-1\) of the state uncertainty matrix

KalmanSmooth is the workhorse function for tsSmooth. makeARIMA constructs the state-space model for an ARIMA model.

\section*{Value}

For KalmanLike, a list with components Lik (the log-likelihood less some constants) and s2, the estimate of of \(\kappa\).

For KalmanRun, a list with components values, a vector of length 2 giving the output of KalmanLike, resid (the residuals) and states, the contemporaneous state estimates, a matrix with one row for each time.

For KalmanSmooth, a list with two components. Component smooth is a n by p matrix of state estimates based on all the observations, with one row for each time. Component smooth is a \(n\) by \(p\) by \(p\) array of variance matrices.

For KalmanForecast, a list with components pred, the predictions, and var, the unscaled variances of the prediction errors (to be muliplied by s2).

For makeARIMA, a model list including components for its arguments.

\section*{Warning}

These functions are designed to be called from other functions which check the validity of the arguments passed, so very little checking is done.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press.

\author{
See Also \\ arima, StructTS. tsSmooth.
}
```

kernapply Apply Smoothing Kernel

```

\section*{Description}
kernapply computes the convolution between an input sequence and a specific kernel.

\section*{Usage}
```

kernapply(x, k, circular = FALSE, ...)
kernapply(k1, k2)

```

\section*{Arguments}
k, k1, k2
x
circular a logical indicating whether the input sequence to be smoothed is treated as circular, i.e., periodic.
... arguments passed to or from other methods.

\section*{Value}

A smoothed version of the input sequence.

\section*{Author(s)}
A. Trapletti

\section*{See Also}
kernel, convolve, filter, spectrum

\section*{Examples}
```


## see 'kernel' for examples

```
kernel Smoothing Kernel Objects

\section*{Description}

The "tskernel" class is designed to represent discrete symmetric normalized smoothing kernels. These kernels can be used to smooth vectors, matrices, or time series objects.

\section*{Usage}
```

kernel(coef, m, r, name)
df.kernel(k)
bandwidth.kernel(k)
is.tskernel(k)
print(x, digits = max(3,getOption("digits")-3), ...)
plot(x, ...)

```

\section*{Arguments}
coef the upper half of the smoothing kernel coefficients (inclusive of coefficient zero) or the name of a kernel (currently "daniell", "dirichlet", "fejer" or "modified.daniell".
m the kernel dimension. The number of kernel coefficients is \(2 * \mathrm{~m}+1\).
name the name of the kernel.
r the kernel order for a Fejer kernel.
digits the number of digits to format real numbers.
k, x
a "tskernel" object.
... arguments passed to or from other methods.

\section*{Details}
kernel is used to construct a general kernel or named specific kernels. The modified Daniell kernel halves the end coefficients (as used by S-PLUS).
df.kernel returns the "equivalent degrees of freedom" of a smoothing kernel as defined in Brockwell and Davies (1991), page 362, and bandwidth.kernel returns the equivalent bandwidth as defined in Bloomfield (1991), p. 201, with a continuity correction.

\section*{Value}
kernel returns a list with class "tskernel", and components the coefficients coef and the kernel dimension m. An additional attribute is "name".

\section*{Author(s)}
A. Trapletti; modifications by B.D. Ripley

\section*{References}

Bloomfield, P. (1976) Fourier Analysis of Time Series: An Introduction. Wiley.
Brockwell, P.J. and Davis, R.A. (1991) Time Series: Theory and Methods. Second edition. Springer, pp. 350-365.

\section*{See Also}
```

kernapply

```

\section*{Examples}
```

data(EuStockMarkets) \# Demonstrate a simple trading strategy for the
x <- EuStockMarkets[,1] \# financial time series German stock index DAX.
k1 <- kernel("daniell", 50) \# a long moving average
k2 <- kernel("daniell", 10) \# and a short one
plot(k1)
plot(k2)
x1 <- kernapply(x, k1)
x2 <- kernapply(x, k2)
plot(x)
lines(x1, col = "red") \# go long if the short crosses the long upwards
lines(x2, col = "green") \# and go short otherwise
data(sunspot) \# Reproduce example 10.4.3 from Brockwell and Davies (1991)
spectrum(sunspot.year, kernel=kernel("daniell", c(11,7,3)), log="no")

```

\section*{Description}

Computed a lagged version of a time series, shifting the time base back by a given number of observations.

\section*{Usage}
\(\operatorname{lag}(\mathrm{x}, \mathrm{k}=1, \ldots\) )

\section*{Arguments}
\begin{tabular}{ll}
x & A vector or matrix or univariate or multivariate time series \\
k & The number of lags (in units of observations). \\
\(\ldots\) & further arguments to be passed to or from methods.
\end{tabular}

\section*{Details}

Vector or matrix arguments x are coerced to time series.
lag is a generic function; this page documents its default method.

\section*{Value}

A time series object.

\section*{Note}

Note the sign of k : a series lagged by a positive k starts earlier.

Author(s)
B.D. Ripley

\section*{See Also}
```

diff, deltat

```

\section*{Examples}
```

data(UKLungDeaths)
lag(ldeaths, 12) \# starts one year earlier

```
lag.plot Time Series Lag Plots

\section*{Description}

Plot time series against lagged versions of themselves. Helps visualizing "auto-dependence" even when auto-correlations vanish.

\section*{Usage}
```

lag.plot(x, lags = 1, layout = NULL, set.lags = 1:lags,
main = NULL, asp = 1,
font.main=par("font.main"), cex.main=par("cex.main"),
diag = TRUE, diag.col="gray", type="p", oma =NULL, ask =NULL,
do.lines = n <= 150, labels = do.lines, ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & time-series (univariate or multivariate) \\
lags \\
layout & \begin{tabular}{l} 
number of lag plots desired, see arg set.lags. \\
the layout of multiple plots, basically the mfrow par() argument. The \\
default uses about a square layout (see n2mfrow such that all plots are on \\
one page.
\end{tabular} \\
set.lags & \begin{tabular}{l} 
positive integer vector allowing to specify the set of lags used; defaults to \\
\(1: l a g s . ~\)
\end{tabular} \\
main & \begin{tabular}{l} 
character with a main header title to be done on the top of each page.
\end{tabular} \\
asp & Aspect ratio to be fixed, see plot. default. \\
font.main, cex.main \\
attributes for the title, see par(). \\
diag & \begin{tabular}{l} 
logical indicating if the x=y diagonal should be drawn.
\end{tabular} \\
diag.col & \begin{tabular}{l} 
color to be used for the diagonal if (diag).
\end{tabular} \\
type & \begin{tabular}{l} 
plot type to be used, but see plot.ts about its restricted meaning. \\
oma
\end{tabular} \\
outer margins, see par.
\end{tabular}

\section*{Note}

It is more flexible and has different default behaviour than the S version. We use main \(=\) instead of head \(=\) for internal consistency.

\section*{Author(s)}

Martin Maechler

\section*{See Also}
plot.ts which is the basic work horse.

\section*{Examples}
```

data(nhtemp)
lag.plot(nhtemp, 8, diag.col = "forest green")
lag.plot(nhtemp, 5, main="Average Temperatures in New Haven")

## ask defaults to TRUE when we have more than one page:

lag.plot(nhtemp, 6, layout = c(2,1), asp = NA,
main = "New Haven Temperatures", col.main = "blue")

## Multivariate (but non-stationary! ...)

data(freeny)
lag.plot(freeny.x, lag = 3)
data(sunspots) \# no lines for long series :
lag.plot(sqrt(sunspots), set = c(1:4, 9:12), pch = ".", col = "gold")

```

LakeHuron

\section*{Description}

Annual measurements of the level, in feet, of Lake Huron 1875-1972.

\section*{Usage}
data(LakeHuron)

\section*{Format}

A time series of length 98 .

\section*{Source}

Brockwell, P. J. \& Davis, R. A. (1991). Time Series and Forecasting Methods. Second edition. Springer, New York. Series A, page 555.

Brockwell, P. J. \& Davis, R. A. (1996). Introduction to Time Series and Forecasting. Springer, New York. Sections 5.1 and 7.6.
lh
Luteinizing Hormone in Blood Samples

\section*{Description}

A regular time series giving the luteinizing hormone in blood samples at 10 mins intervals from a human female, 48 samples.

\section*{Usage}
data(lh)

\section*{Source}
P.J. Diggle (1990) Time Series: A Biostatistical Introduction. Oxford, table A.1, series 3
lynx Annual Canadian Lynx trappings 1821-1934

\section*{Description}

Annual numbers of lynx trappings for 1821-1934 in Canada. Taken from Brockwell \& Davis (1991), this appears to be the series considered by Campbell \& Walker (1977).

\section*{Usage}
data(lynx)

\section*{Source}

Brockwell, P. J. and Davis, R. A. (1991) Time Series and Forecasting Methods. Second edition. Springer. Series G (page 557).

\section*{References}

Campbell, M. J.and A. M. Walker (1977). A Survey of statistical work on the Mackenzie River series of annual Canadian lynx trappings for the years 1821-1934 and a new analysis. Journal of the Royal Statistical Society series A, 140, 411-431.

\section*{Description}

These functions plot seasonal (or other) subseries of a time series. For each season (or other category), a time series is plotted.

\section*{Usage}
monthplot(x, labels = NULL, times, phase, base, choice, ...)

\section*{Arguments}
x
Time series or related object.
labels Labels to use for each "season".
times Time of each observation.
phase Indicator for each "season".
base Function to use for reference line for subseries.
choice Which series of an stl or StructTS object?
... Graphical parameters.

\section*{Details}

These functions extract subseries from a time series and plot them all in one frame. The ts, stl, and StructTS methods use the internally recorded frequency and start and finish times to set the scale and the seasons. The default method assumes observations come in groups of 12 (though this can be changed).

If the labels are not given but the phase is given, then the labels default to the unique values of the phase. If both are given, then the phase values are assumed to be indices into the labels array, i.e. they should be in the range from 1 to length(labels).

\section*{Value}

These functions are executed for their side effect of drawing a seasonal subseries plot on the current graphical window.

Author(s)
Duncan Murdoch

\section*{See Also}
```

ts, stl, StructTS

```

\section*{Examples}
```


## The CO2 data

data(co2)
fit <- stl(log(co2), s.window = 20, t.window = 20)
plot(fit)
op <- par(mfrow = c(2,2))
monthplot(co2, ylab = "data", cex.axis = 0.8)
monthplot(fit, choice = "seasonal", cex.axis = 0.8)
monthplot(fit, choice = "trend", cex.axis = 0.8)
monthplot(fit, choice = "remainder", type = "h", cex.axis = 0.8)
par(op)

## The CO2 data, grouped quarterly

quarter <- (cycle(co2) - 1) %/% 3
monthplot(co2, phase = quarter)

## see also JohnsonJohnson

```
```

na.contiguous NA Handling Routines for Time Series

```

\section*{Description}

Find the longest consecutive stretch of non-missing values in a time series object. (In the event of a tie, the first such stretch.)

\section*{Usage}
na.contiguous(frame)

\section*{Arguments}
frame a univariate or multivariate time series.

\section*{Value}

A time series without missing values. The class of frame will be preserved.

\section*{Author(s)}
B. D. Ripley

\section*{See Also}
na.omit and na.omit.ts; na.fail

\section*{Examples}
data(presidents)
na.contiguous(presidents)

\section*{Description}

Measurements of the annual flow of the river Nile at Ashwan 1871-1970.

\section*{Usage}
data(Nile)

\section*{Format}

A time series of length 100 .

\section*{Source}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press. http://www.ssfpack.com/dkbook/

\section*{References}

Balke, N. S. (1993) Detecting level shifts in time series. Journal of Business and Economic Statistics 11, 81-92.
Cobb, G. W. (1978) The problem of the Nile: conditional solution to a change-point problem. Biometrika 65, 243-51.

\section*{Examples}
```

data(Nile)
par(mfrow = c(2,2))
plot(Nile)
acf(Nile)
pacf(Nile)
ar(Nile) \# selects order 2
cpgram(ar(Nile)\$resid)
par(mfrow = c(1,1))
arima(Nile, c(2, 0, 0))

## Now consider missing values, following Durbin \& Koopman

NileNA <- Nile
NileNA[c(21:40, 61:80)] <- NA
arima(NileNA, c(2, 0, 0))
plot(NileNA)
pred <- predict(arima(window(NileNA, 1871, 1890), c(2, 0, 0)), n.ahead = 20)
lines(pred$pred, lty = 3, col = "red")
lines(pred$pred + 2*pred$se, lty=2, col="blue")
lines(pred$pred - 2*pred$se, lty=2, col="blue")
pred <- predict(arima(window(NileNA, 1871, 1930), c(2, 0, 0)), n.ahead = 20)
lines(pred$pred, lty = 3, col = "red")
lines(pred$pred + 2*pred$se, lty=2, col="blue")
lines(pred$pred - 2*pred$se, lty=2, col="blue")

## Structural time series models

```
```

data(Nile, package = "ts")
par(mfrow = c(3, 1))
plot(Nile)

## local level model

(fit <- StructTS(Nile, type = "level"))
lines(fitted(fit), lty = 2) \# contempareneous smoothing
lines(tsSmooth(fit), lty = 2, col = 4) \# fixed-interval smoothing
plot(residuals(fit)); abline(h = 0, lty = 3)

## local trend model

(fit2 <- StructTS(Nile, type = "trend")) \#\# constant trend fitted
pred <- predict(fit, n.ahead = 30)

## with 50% confidence interval

ts.plot(Nile, pred$pred, pred$pred + 0.67*pred$se, pred$pred -0.67*pred\$se)

## Now consider missing values

plot(NileNA)
(fit3 <- StructTS(NileNA, type = "level"))
lines(fitted(fit3), lty = 2)
lines(tsSmooth(fit3), lty = 3)
plot(residuals(fit3)); abline(h = 0, lty = 3)

```
nottem Average Monthly Temperatures at Nottingham, 1920-1939

\section*{Description}

A time series object containing average air temperatures at Nottingham Castle in degrees Fahrenheit for 20 years.

\section*{Usage}
data(nottem)

\section*{Source}

Anderson, O. D. (1976) Time Series Analysis and Forecasting: The Box-Jenkins approach. Butterworths. Series R.

\section*{Examples}
```

data(nottem)
nott <- window(nottem, end=c(1936,12))
fit <- arima(nott,order=c(1,0,0), list(order=c(2,1,0), period=12))
nott.fore <- predict(fit, n.ahead=36)
ts.plot(nott, nott.fore$pred, nott.fore$pred+2*nott.fore$se,
    nott.fore$pred-2*nott.fore\$se, gpars=list(col=c(1,1,4,4)))

```

\section*{plot.acf Plotting Autocovariance and Autocorrelation Functions}

\section*{Description}

Plotting method for objects of class "acf".

\section*{Usage}
```

plot(x, ci = 0.95, type = "h", xlab = "Lag", ylab = NULL,
ylim = NULL, main = NULL, ci.col="blue",
ci.type = c("white", "ma"),
max.mfrow = 6,
ask = Npgs > 1 \&\& dev.interactive(),
mar = if(nser > 2) c(3,2,2,0.8) else par("mar"),
oma = if(nser > 2) c(1,1.2,1,1) else par("oma"),
mgp = if(nser > 2) c(1.5,0.6,0) else par("mgp"),
xpd = par("xpd"),
cex.main = if(nser > 2) 1 else par("cex.main"),
verbose = getOption("verbose"),
...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & an object of class "acf". \\
ci & \begin{tabular}{l} 
coverage probability for confidence interval. Plotting of the confidence \\
interval is suppressed if ci is zero or negative.
\end{tabular} \\
type & \begin{tabular}{l} 
the type of plot to be drawn, default to histogram like vertical lines. \\
xlab \\
ylab \\
ylim
\end{tabular} \\
\begin{tabular}{ll} 
the \(x\) label of the plot.
\end{tabular} \\
main & the label of the plot.
\end{tabular}

\section*{Note}

The confidence interval plotted in plot.acf is based on an uncorrelated series and should be treated with appropriate caution. Using ci.type = "ma" may be less potentially misleading.

\section*{See Also}
acf which calls plot.acf by default.

\section*{Examples}
```

z4 <- ts(matrix(rnorm(400), 100, 4), start=c(1961, 1), frequency=12)
z7 <- ts(matrix(rnorm(700), 100, 7), start=c(1961, 1), frequency=12)
acf(z4)
acf(z7, max.mfrow = 7)\# squeeze on 1 page
acf(z7) \# multi-page

```
plot.HoltWinters Plot function for HoltWinters objects

\section*{Description}

Produces a chart of the original time series along with the fitted values. Optionally, predicted values (and their confidence bounds) can also be plotted.

\section*{Usage}
```

plot(x, predicted.values = NA, intervals = TRUE,
separator = TRUE, col = 1, col.predicted = 2,
col.intervals = 4, col.separator = 1, lty = 1,
lty.predicted = 1, lty.intervals = 1, lty.separator = 3,
ylab = "Observed / Fitted",
main = "Holt-Winters filtering", ...)

```

\section*{Arguments}
x Object of class "HoltWinters"
predicted.values
Predicted values as returned by predict. HoltWinters
intervals If TRUE, the prediction intervals are plotted (default).
separator If TRUE, a separating line between fitted and predicted values is plotted (default).
col, lty Color/line type of original data (default: black solid).
col.predicted, lty.predicted
Color/line type of fitted and predicted values (default: red solid).
col.intervals, lty.intervals
Color/line type of prediction intervals (default: blue solid).
col.separator, lty.separator
Color/line type of observed/predicted values separator (default: black dashed).
ylab Label of the y-axis.
main Main title.
... Other graphics parametes

\section*{Author(s)}

David Meyer 〈david.meyer@ci.tuwien.ac.at〉

\section*{References}
C.C Holt (1957) Forecasting seasonals and trends by exponentially weighted moving averages, ONR Research Memorandum, Carnigie Institute 52.
P.R Winters (1960) Forecasting sales by exponentially weighted moving averages, Management Science 6, 324-342.

\section*{See Also}

HoltWinters, predict.HoltWinters
```

plot.spec Plotting Spectral Densities

```

\section*{Description}

Plotting method for objects of class "spec". For multivariate time series it plots the marginal spectra of the series or pairs plots of the coherency and phase of the cross-spectra.

\section*{Usage}
```

plot(x, add = FALSE, ci = 0.95, log = c("yes", "dB", "no"),
xlab = "frequency", ylab = NULL, type = "l", ci.col = "blue",
main = NULL, sub = NULL,
plot.type = c("marginal", "coherency", "phase"),
ci.lty = 3, ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & an object of class "spec". \\
add & \begin{tabular}{l} 
logical. If TRUE, add to already existing plot. \\
ci \\
Coverage probability for confidence interval. Plotting of the confidence \\
bar is omitted unless ci is strictly positive.
\end{tabular} \\
log & \begin{tabular}{l} 
If "dB", plot on log10 (decibel) scale (as S-PLUS), otherwise use conven- \\
tional log scale or linear scale. Logical values are also accepted. The de- \\
fault is "yes" unless options(ts.S.compat = TRUE) has been set, when \\
it is "dB".
\end{tabular} \\
xlab & the x label of the plot. \\
ylab & the y label of the plot. \\
type & the type of plot to be drawn, defaults to lines.
\end{tabular}
```

ci.col Colour for plotting confidence bar or confidence intervals for coherency
and phase.
main overall title for the plot.
sub a sub title for the plot.
plot.type For multivariate time series, the type of plot required. Only the first
character is needed.
ci.lty line type for confidence intervals for coherency and phase.
... Further graphical parameters.

```
```

See Also
spectrum

```

\section*{PP.test Phillips-Perron Unit Root Test}

\section*{Description}

Computes the Phillips-Perron test for the null hypothesis that x has a unit root against a stationary alternative.

\section*{Usage}

PP.test(x, lshort = TRUE)

\section*{Arguments}
\(\mathrm{x} \quad\) a numeric vector or univariate time series.
lshort a logical indicating whether the short or long version of the truncation lag parameter is used

\section*{Details}

The general regression equation which incorporates a constant and a linear trend is used and the corrected t-statistic for a first order autoregressive coefficient equals one is computed. To estimate sigma^2 the Newey-West estimator is used. If lshort is TRUE, then the truncation lag parameter is set to \(\operatorname{trunc}\left(4 *(\mathrm{n} / 100)^{\wedge} 0.25\right)\), otherwise \(\operatorname{trunc}\left(12 *(\mathrm{n} / 100)^{\wedge} 0.25\right)\) is used. The \(p\)-values are interpolated from Table 4.2, page 103 of Banerjee et al. (1993). Missing values are not handled.

\section*{Value}

A list with class "htest" containing the following components:
statistic the value of the test statistic.
parameter the truncation lag parameter.
p.value the \(p\)-value of the test.
method a character string indicating what type of test was performed.
data.name a character string giving the name of the data.

\section*{Author(s)}
A. Trapletti

\section*{References}
A. Banerjee, J. J. Dolado, J. W. Galbraith, and D. F. Hendry (1993) Cointegration, Error Correction, and the Econometric Analysis of Non-Stationary Data, Oxford University Press, Oxford.
P. Perron (1988) Trends and random walks in macroeconomic time series. Journal of Economic Dynamics and Control 12, 297-332.

\section*{Examples}
```

x <- rnorm(1000)
PP.test(x)
y <- cumsum(x) \# has unit root
PP.test(y)

```
predict.Arima Forecast from ARIMA fits

\section*{Description}

Forecast from models fitted by arima.

\section*{Usage}
predict(object, n.ahead = 1, newxreg = NULL, se.fit = TRUE, ...)

\section*{Arguments}
object The result of an arima fit.
n . ahead The number of steps ahead for which prediction is required.
newxreg New values of xreg to be used for prediction. Must have at least n. ahead rows.
se.fit Logical: should standard errors of prediction be returned?
... arguments passed to or from other methods.

\section*{Details}

Finite-history prediction is used, via KalmanForecast. This is only statistically efficient if the MA part of the fit is invertible, so predict. Arima will give a warning for non-invertible MA models.

The standard errors of prediction exclude the uncertainty in the estimation of the ARMA model and the regression coefficients. According to Harvey (1993, pp. 58-9) the effect is small.

\section*{Value}

A time series of predictions, or if se.fit = TRUE, a list with components pred, the predictions, and se, the estimated standard errors. Both components are time series.

\section*{Author(s)}
B. D. Ripley

\section*{References}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press.

Harvey, A. C. and McKenzie, C. R. (1982) Algorithm AS182. An algorithm for finite sample prediction from ARIMA processes. Applied Statistics 31, 180-187.

Harvey, A. C. (1993) Time Series Models, 2nd Edition, Harvester Wheatsheaf, sections 3.3 and 4.4

\section*{See Also}
arima

\section*{Examples}
```

data(lh, package="ts")
predict(arima(lh, order = c(3,0,0)), n.ahead = 12)
data(USAccDeaths, package="ts")
(fit <- arima(USAccDeaths, order = c(0,1,1),
seasonal = list(order=c(0,1,1))))
predict(fit, n.ahead = 6)

```
predict.HoltWinters prediction function for fitted Holt-Winters models

\section*{Description}

Computes predictions and prediction intervals for models fitted by the Holt-Winters method.

\section*{Usage}
```

predict(object, n.ahead=1, prediction.interval = FALSE,
quantile = qnorm(0.975), ...)

```

\section*{Arguments}
\begin{tabular}{ll} 
object & An object of class HoltWinters \\
n.ahead & Number of future periods to predict \\
prediction.interval
\end{tabular}
logical. If TRUE, the lower and upper bounds of the corresponding prediction intervals are computed
quantile Quantile used for each one-sided prediction interval (default: the 0.975 quantile to give a two-sided 0.95 interval)
... arguments passed to or from other methods

\section*{Value}

A time series of the predicted values. If prediction intervals are requested, a multiple time series is returned with columns fit, lwr and upr for the predicted values and the lower and upper bounds respectively.

\section*{Author(s)}

David Meyer 〈david.meyer@ci.tuwien.ac.at〉

\section*{References}
C.C Holt (1957) Forecasting seasonals and trends by exponentially weighted moving averages, ONR Research Memorandum, Carnigie Institute 52.
P.R Winters (1960) Forecasting sales by exponentially weighted moving averages, Management Science 6, 324-342.

\section*{See Also}

HoltWinters

\section*{Examples}
```

data(co2)
m <- HoltWinters(co2)
p <- predict(m, 50, prediction.interval = TRUE)
plot(m, p)

```
spec.ar

Estimate Spectral Density of a Time Series from AR Fit

\section*{Description}

Fits an AR model to x (or uses the existing fit) and computes (and by default plots) the spectral density of the fitted model.

\section*{Usage}
```

spec.ar(x, n.freq, order = NULL, plot = TRUE, na.action = na.fail,
method = "yule-walker", ...)

```

\section*{Arguments}
x ar.
n.freq The number of points at which to plot.
order The order of the AR model to be fitted. If omitted, the order is chosen by AIC.
plot Plot the periodogram?
na.action NA action function.
method method for ar fit.
... Graphical arguments passed to plot.spec.

\section*{Value}

An object of class "spec". The result is returned invisibly if plot is true.

\section*{Warning}

Some authors, for example Thomson (1990), warn strongly that AR spectra can be misleading.

\section*{Note}

The multivariate case is not yet implemented.

\section*{Author(s)}
B.D. Ripley

\section*{References}

Thompson, D.J. (1990) Time series analysis of Holocene climate data. Phil. Trans. Roy. Soc. A 330, 601-616.

Venables, W.N. and Ripley, B.D. (1997) Modern Applied Statistics with S-PLUS. Second edition. Springer. (Especially page 448.)

\section*{See Also}
ar, spectrum.

\section*{Examples}
```

data(lh)
spec.ar(lh)
data(UKLungDeaths)
spec.ar(ldeaths)
spec.ar(ldeaths, method="burg")

```
spec.pgram Estimate Spectral Density of a Time Series from Smoothed Periodogram

\section*{Description}
spec.pgram calculates the periodogram using a fast Fourier transform, and optionally smooths the result with a series of modified Daniell smoothers (moving averages giving half weight to the end values).

\section*{Usage}
```

spec.pgram(x, spans = NULL, kernel, taper = 0.1,
pad = 0, fast = TRUE, demean = FALSE, detrend =TRUE,
plot = FALSE, na.action = na.fail, ...)

```

\section*{Arguments}
\(\mathrm{x} \quad\) univariate or multivariate time series.
spans vector of odd integers giving the widths of modified Daniell smoothers to be used to smooth the periodogram.
kernel alternatively, a kernel smoother of class "tskernel".
taper proportion of data to taper. A split cosine bell taper is applied to this proportion of the data at the beginning and end of the series.
pad proportion of data to pad. Zeros are added to the end of the series to increase its length by the proportion pad.
fast logical; if TRUE, pad the series to a highly composite length.
demean logical. If TRUE, subtract the mean of the series.
detrend logical. If TRUE, remove a linear trend from the series. This will also remove the mean.
plot plot the periodogram?
na.action NA action function.
graphical arguments passed to plot.spec.

\section*{Details}

The raw periodogram is not a consistent estimator of the spectral density, but adjacent values are asymptotically independent. Hence a consistent estimator can be derived by smoothing the raw periodogram, assuming that the spectral density is smooth.
The series will be automatically padded with zeros until the series length is a highly composite number in order to help the Fast Fourier Transform. This is controlled by the fast and not the pad argument.
The periodogram at zero is in theory zero as the mean of the series is removed (but this may be affected by tapering): it is replaced by an interpolation of adjacent values during smoothing, and no value is returned for that frequency.

\section*{Value}

A list object of class "spec" (see spectrum) with the following additional components:
kernel The kernel argument, or the kernel constructed from spans.
df The distribution of the spectral density estimate can be approximated by a chi square distribution with df degrees of freedom.
bandwidth The equivalent bandwidth of the kernel smoother as defined by Bloomfield (1976, page 201).
taper The value of the taper argument.
pad The value of the pad argument.
detrend The value of the detrend argument.
demean The value of the demean argument.
The result is returned invisibly if plot is true.

\section*{Author(s)}

Originally Martyn Plummer; kernel smoothing by Adrian Trapletti, synthesis by B.D. Ripley

\section*{References}

Bloomfield, P. (1976) Fourier Analysis of Time Series: An Introduction. Wiley.
Brockwell, P.J. and Davis, R.A. (1991) Time Series: Theory and Methods. Second edition. Springer
Venables, W.N. and Ripley, B.D. (1997) Modern Applied Statistics with S-PLUS. Second edition. Springer. (Especially pp. 437-442.)

\section*{See Also}

\section*{Examples}
```


## Examples from Venables \& Ripley

data(UKLungDeaths)
spectrum(ldeaths)
spectrum(ldeaths, spans = c(3,5))
spectrum(ldeaths, spans = c(5,7))
spectrum(mdeaths, spans = c(3,3))
spectrum(fdeaths, spans = c(3,3))

## bivariate example

mfdeaths.spc <- spec.pgram(ts.union(mdeaths, fdeaths), spans = c(3,3))

# plots marginal spectra: now plot coherency and phase

plot(mfdeaths.spc, plot.type = "coherency")
plot(mfdeaths.spc, plot.type = "phase")

## now impose a lack of alignment

mfdeaths.spc <- spec.pgram(ts.intersect(mdeaths, lag(fdeaths, 4)),
spans = c(3,3), plot = FALSE)
plot(mfdeaths.spc, plot.type = "coherency")
plot(mfdeaths.spc, plot.type = "phase")

```
```

data(EuStockMarkets)
stocks.spc <- spectrum(EuStockMarkets, kernel("daniell", c(30,50)),
plot = FALSE)
plot(stocks.spc, plot.type = "marginal") \# the default type
plot(stocks.spc, plot.type = "coherency")
plot(stocks.spc, plot.type = "phase")
data(BJsales)
sales.spc <- spectrum(ts.union(BJsales, BJsales.lead),
kernel("modified.daniell", c(5,7)))
plot(sales.spc, plot.type = "coherency")
plot(sales.spc, plot.type = "phase")

```
spec.taper Taper a Time Series

\section*{Description}

Apply a cosine-bell taper to a time series.

\section*{Usage}
spec.taper ( \(\mathrm{x}, \mathrm{p}=0.1\) )

\section*{Arguments}
\(\mathrm{x} \quad\) A univariate or multivariate time series
p The total proportion to be tapered, either a scalar or a vector of the length of the number of series.

\section*{Details}

The cosine-bell taper is applied to the first and last \(p[i] / 2\) observations of time series \(x[\), i].

\section*{Value}

A new time series object.

\section*{Note}

From package 'MASS'.

\section*{Author(s)}

Kurt Hornik, B.D. Ripley

\section*{See Also}
spec.pgram, cpgram
```

spectrum Spectral Density Estimation

```

\section*{Description}

The spectrum function estimates the spectral density of a time series.

\section*{Usage}
```

spectrum(x, method = c("pgram", "ar"), plot = TRUE, na.action = na.fail,
...)

```

\section*{Arguments}
\begin{tabular}{ll}
x & A univariate or multivariate time series. \\
method & \begin{tabular}{l} 
String specifying the method used to estimate the spectral density. Al- \\
lowed methods are "pgram" (the default) and "ar".
\end{tabular} \\
plot & \begin{tabular}{l} 
logical. If TRUE then the spectral density is plotted.
\end{tabular} \\
na.action & NA action function. \\
\(\ldots\) & Further arguments to specific spec methods or plot.spec.
\end{tabular}

\section*{Details}
spectrum is a wrapper function which calls the methods spec.pgram and spec.ar.
The spectrum here is defined with scaling \(1 / f r e q u e n c y(x)\), following S-PLUS. This makes the spectral density a density over the range (-frequency \((x) / 2\), +frequency \((x) / 2]\), whereas a more common scaling is \(2 \pi\) and range \((-0.5,0.5\) ] (e.g., Bloomfield) or 1 and range \((-\pi, \pi]\).
If available, a confidence interval will be plotted by plot.spec: this is asymmetric, and the width of the centre mark indicates the equivalent bandwidth.

\section*{Value}

An object of class "spec", which is a list containing at least the following components:
freq vector of frequencies at which the spectral density is estimated. (Possibly approximate Fourier frequencies.) The units are the reciprocal of cycles per unit time (and not per observation spacing): see Details below.
spec Vector (for univariate series) or matrix (for multivariate series) of estimates of the spectral density at frequencies corresponding to freq.
coh NULL for univariate series. For multivariate time series, a matrix containing the squared coherency between different series. Column \(i+(j-1) *\) \((j-2) / 2\) of coh contains the squared coherency between columns \(i\) and \(j\) of x , where \(i<j\).
phase NULL for univariate series. For multivariate time series a matrix containing the cross-spectrum phase between different series. The format is the same as coh.
series The name of the time series.
snames For multivariate input, the names of the component series.
method The method used to calculate the spectrum.

The result is returned invisibly if plot is true.

\section*{Note}

The default plot for objects of class "spec" is quite complex, including an error bar and default title, subtitle and axis labels. The defaults can all be overridden by supplying the appropriate graphical parameters.

\section*{Author(s)}

Martyn Plummer, B.D. Ripley

\section*{References}

Bloomfield, P. (1976) Fourier Analysis of Time Series: An Introduction. Wiley.
Brockwell, P. J. and Davis, R. A. (1991) Time Series: Theory and Methods. Second edition. Springer.

Venables, W. N. and Ripley, B. D. (1997) Modern Applied Statistics with S-PLUS. Second edition. Springer. (Especially pages 437-442.)

\section*{See Also}
```

spec.ar, spec.pgram; plot.spec.

```

\section*{Examples}
```


## Examples from Venables \& Ripley

## spec.pgram

par(mfrow=c(2,2))
data(lh)
spectrum(lh)
spectrum(lh, spans=3)
spectrum(lh, spans=c (3,3))
spectrum(lh, spans=c (3,5))
data(UKLungDeaths)
spectrum(ldeaths)
spectrum(ldeaths, spans=c(3,3))
spectrum(ldeaths, spans=c (3,5))
spectrum(ldeaths, spans=c(5,7))
spectrum(ldeaths, spans=c(5,7), log="dB", ci=0.8)

# for multivariate examples see the help for spec.pgram

## spec.ar

spectrum(lh, method="ar")
spectrum(ldeaths, method="ar")

```

\section*{Description}

Decompose a time series into seasonal, trend and irregular components using loess, acronym STL.

\section*{Usage}
```

stl(x, s.window, s.degree = 0,
t.window = NULL, t.degree = 1,
l.window = nextodd(period), l.degree = t.degree,
s.jump = ceiling(s.window/10),
t.jump = ceiling(t.window/10),
l.jump = ceiling(l.window/10),
robust = FALSE,
inner = if(robust) 1 else 2,
outer = if(robust) 15 else 0,
na.action = na.fail)

```

\section*{Arguments}
\(\mathrm{x} \quad\) univariate time series to be decomposed. This should be an object of class "ts" with a frequency greater than one.
s.window either the character string "periodic" or the span (in lags) of the loess window for seasonal extraction, which should be odd. This has no default.
s.degree degree of locally-fitted polynomial in seasonal extraction. Should be zero or one.
t.window the span (in lags) of the loess window for trend extraction, which should be odd. If NULL, the default, nextodd(ceiling((1.5*period) / (1( \(1.5 / \mathrm{s}\).window))) ), is taken.
t.degree degree of locally-fitted polynomial in trend extraction. Should be zero or one.
1.window the span (in lags) of the loess window of the low-pass filter used for each subseries. Defaults to the smallest odd integer greater than or equal to frequency ( x ) which is recommended since it prevents competition between the trend and seasonal components. If not an odd integer its given value is increased to the next odd one.
l.degree degree of locally-fitted polynomial for the subseries low-pass filter. Must be 0 or 1 .
s.jump, t.jump, l.jump
integers at least one to increase speed of the respective smoother. Linear interpolation happens between every \(*\). jumpth value.
robust logical indicating if robust fitting be used in the loess procedure.
inner integer; the number of 'inner' (backfitting) iterations; usually very few (2) iterations suffice.
outer integer; the number of 'outer' robustness iterations.
na.action action on missing values.

\section*{Details}

The seasonal component is found by loess smoothing the seasonal sub-series (the series of all January values, ...); if s.window = "periodic" smoothing is effectively replaced by taking the mean. The seasonal values are removed, and the remainder smoothed to find the trend. The overall level is removed from the seasonal component and added to the trend component. This process is iterated a few times. The remainder component is the residuals from the seasonal plus trend fit.
Several methods for the resulting class "stl" objects, see, plot.stl.

\section*{Value}
```

stl returns an object of class "stl" with components
time.series a multiple time series with columns seasonal, trend and remainder.
weights the final robust weights (all one if fitting is not done robustly).
call the matched call.
win integer (length 3 vector) with the spans used for the "s", "t", and "l"
smoothers.
deg integer (length 3) vector with the polynomial degrees for these smoothers.
jump integer (length 3) vector with the "jumps" (skips) used for these smoothers.
ni number of inner iterations
no number of outer robustness iterations

```

\section*{Note}

This is similar to but not identical to the stl function in S-PLUS. The remainder component given by S-PLUS is the sum of the trend and remainder series from this function.

\section*{Author(s)}
B.D. Ripley; Fortran code by Cleveland et al. (1990) from 'netlib'.

\section*{References}
R. B. Cleveland, W. S. Cleveland, J.E. McRae, and I. Terpenning (1990) STL: A SeasonalTrend Decomposition Procedure Based on Loess. Journal of Official Statistics, 6, 3-73.

\section*{See Also}
plot.stl for stl methods; loess in package 'modreg' (which is not actually used in stl).

\section*{Examples}
```

data(nottem)
plot(stl(nottem, "per"))
plot(stl(nottem, s.win = 4, t.win = 50, t.jump = 1))
data(co2)
plot(stllc <- stl(log(co2), s.window=21))
summary(stllc)

## linear trend, strict period.

plot(stl(log(co2), s.window="per", t.window=1000))

```
```


## Two STL plotted side by side :

data(UKLungDeaths)
stmd <- stl(mdeaths, s.w = "per") \# un-robust
summary(stmR <- stl(mdeaths, s.w = "per", robust = TRUE))
op <- par(mar = c(0, 4, 0, 3), oma = c(5, 0, 4, 0), mfcol = c(4, 2))
plot(stmd, set.pars=NULL, labels = NULL,
main = "stl(mdeaths, s.w = \"per\", robust = FALSE / TRUE )")
plot(stmR, set.pars=NULL)

# mark the 'outliers' :

(iO <- which(stmR \$ weights < 1e-8)) \# 10 were considered outliers
sts <- stmR\$time.series
points(time(sts)[i0], .8* sts[,"remainder"][i0], pch = 4, col = "red")
par(op)\# reset

```
stlmethods
Methods for STL Objects

\section*{Description}

Methods for objects of class stl, typically the result of stl. The plot method does a multiple figure plot with some flexibility.

\section*{Usage}
```

plot(x, labels = colnames(X),
set.pars = list(mar = c(0, 6, 0, 6), oma = c(6, 0, 4, 0),
tck = -0.01, mfrow = c(nplot, 1)),
main = NULL, range.bars = TRUE, ...)
print(x, ...)
summary(object, digits = getOption("digits"), ...)

```

\section*{Arguments}
\begin{tabular}{ll}
x, object & stl object. \\
labels & character of length 4 giving the names of the component time-series. \\
set.pars & settings for par(.) when setting up the plot. \\
main & plot main title. \\
range.bars & \begin{tabular}{l} 
logical indicating if each plot should have a bar at its right side which are \\
of equal heights in user coordinates.
\end{tabular} \\
digits & \begin{tabular}{l} 
significant figures used in printing.
\end{tabular} \\
\(\ldots\) & further arguments passed to or from other methods.
\end{tabular}

\section*{See Also}
plot.ts and stl, particularly for examples.

StructTS Fit Structural Time Series

\section*{Description}

Fit a structural model for a time series by maximum likelihood.

\section*{Usage}
```

StructTS(x, type = c("level", "trend", "BSM"), init = NULL,
fixed = NULL, optim.control = NULL)

```

\section*{Arguments}

\section*{X}
type
init
fixed
a univariate time series. Missing values are allowed.
the class of structural model. If omitted, a BSM is used for a time series with frequency ( x ) > 1, and a local trend model otherwise.
initial values of the variance parameters.
optional numeric vector of the same length as the total number of parameters. If supplied, only non-NA entries in fixed will be varied. Probably most useful for setting variances to zero.
optim. control List of control parameters for optim. Method "L-BFGS-B" is used.

\section*{Details}

Structural time series models are (linear Gaussian) state-space models for (univariate) time series based on a decomposition of the series into a number of components. They are specified by a set of error variances, some of which may be zero.
The simplest model is the local level model specified by type = "level". This has an underlying level \(\mu_{t}\) which evolves by
\[
\mu_{t+1}=\mu_{t}+\xi_{t}, \quad \xi_{t} \sim N\left(0, \sigma_{\xi}^{2}\right)
\]

The observations are
\[
x_{t}=\mu_{t}+\epsilon_{t}, \quad \epsilon_{t} \sim N\left(0, \sigma_{\epsilon}^{2}\right)
\]

There are two parameters, \(\sigma_{\xi}^{2}\) and \(\sigma_{\epsilon}^{2}\). It is an \(\operatorname{ARIMA}(0,1,1)\) model, but with restrictions on the parameter set.
The local linear trend model, type = "trend", has the same measurement equation, but with a time-varying slope in the dynamics for \(\mu_{t}\), given by
\[
\begin{gathered}
\mu_{t+1}=\mu_{t}+\nu_{t}+\xi_{t}, \quad \xi_{t} \sim N\left(0, \sigma_{\xi}^{2}\right) \\
\nu_{t+1}=\nu_{t}+\zeta_{t}, \quad \zeta_{t} \sim N\left(0, \sigma_{\zeta}^{2}\right)
\end{gathered}
\]
with three variance parameters. It is not uncommon to find \(\sigma_{\zeta}^{2}=0\) (which reduces to the local level model) or \(\sigma_{\xi}^{2}=0\), which ensures a smooth trend. This is a restricted ARIMA( \(0,2,2\) ) model.
The basic structural model, type \(=\) "BSM", is a local trend model with an additional seasonal component. Thus the measurement equation is
\[
x_{t}=\mu_{t}+\gamma_{t}+\epsilon_{t}, \quad \epsilon_{t} \sim N\left(0, \sigma_{\epsilon}^{2}\right)
\]
where \(\gamma_{t}\) is a seasonal component with dynamics
\[
\gamma_{t+1}=-\gamma_{t}+\cdots+\gamma_{t-s+2}+\omega_{t}, \quad \omega_{t} \sim N\left(0, \sigma_{\omega}^{2}\right)
\]

The boundary case \(\sigma_{\omega}^{2}=0\) corresponds to a deterministic (but arbitrary) seasonal pattern. (This is sometimes known as the 'dummy variable' version of the BSM.)

\section*{Value}

A list of class "StructTS" with components:
coef the estimated variances of the components.
loglik the maximized log-likelihood. Note that as all these models are nonstationary this includes a diffuse prior for some observations and hence is not comparable with arima nor different types of structural models.
data the time series x .
residuals the standardized residuals.
fitted a multiple time series with one component for the level, slope and seasonal components, estimated contemporaneously (that is at time \(t\) and not at the end of the series.
call the matched call.
series the name of the series \(x\).
convergence the value returned by optim.
model, model0 Lists representing the Kalman Filter used in the fitting. See KalmanLike. model0 is the initial state of the filter, model its final state.
xtsp the tsp attributes of x .

\section*{Note}

Optimization of structural models is a lot harder than many of the references admit. For example, the AirPassengers data are considered in Brockwell \& Davis (1996): their solution appears to be a local maximum, but nowhere near as good a fit as that produced by StructTS. It is quite common to find fits with one or more variances zero, and this can include \(\sigma_{\epsilon}^{2}\).

\section*{Author(s)}
B. D. Ripley

\section*{References}

Brockwell, P. J. \& Davis, R. A. (1996). Introduction to Time Series and Forecasting. Springer, New York. Sections 8.2 and 8.5.
Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press.
Harvey, A. C. (1989) Forecasting, Structural Time Series Models and the Kalman Filter. Cambridge University Press.
Harvey, A. C. (1993) Time Series Models. 2nd Edition, Harvester Wheatsheaf.

\section*{See Also}

KalmanLike, tsSmooth

\section*{Examples}
```


## see also JohnsonJohnson, Nile and AirPassengers

data(treering)
trees <- window(treering, start=0)
(fit <- StructTS(trees, type = "level"))
plot(trees)
lines(fitted(fit), col = "green")
tsdiag(fit)
data(UKgas)
(fit <- StructTS(log10(UKgas), type = "BSM"))
par(mfrow = c(4, 1))
plot(log10(UKgas))
plot(cbind(fitted(fit), resids=resid(fit)), main = "UK gas consumption")

```
sunspot Yearly Sunspot Data, 1700-1988. Monthly Sunspot Data, 1749- 1997.

\section*{Description}

Monthly and yearly number of sunspots.

\section*{Usage}
data(sunspot)

\section*{Format}

The univariate time series sunspot.year and sunspot.month contain 289 and 2988 observations, respectively. The objects are of class "ts".

\section*{Source}

Monthly data: Sunspot Index Data Center, World Data Center-C1 For Sunspot Index Royal Observatory of Belgium, Av. Circulaire, 3, B-1180 BRUSSELS http://www.oma. be/KSB-ORB/SIDC/sidc_txt.html

Yearly data: H. Tong (1996) Non-Linear Time Series. Clarendon Press, Oxford, p. 471.

\section*{See Also}
sunspot.month is a longer version of sunspots in base R, that runs until 1988.

\section*{Examples}
```


## Compare the monthly series from 'base' and 'ts':

data(sunspots, package = base)
data(sunspot, package = ts)
plot (sunspot.month, main = "sunspot.month [ts]", col = 2)
lines(sunspots)\# ''very barely') see something

## Now look at the difference :

```
```

all(tsp(sunspots) [c(1,3)] ==
tsp(sunspot.month)[c(1,3)]) \#\# Start \& Periodicity are the same
n1 <- length(sunspots)
table(eq <- sunspots == sunspot.month[1:n1]) \#> 132 are different !
i <- which(!eq)
rug(time(eq)[i])
s1 <- sunspots[i] ; s2 <- sunspot.month[i]
cbind(i = i, sunspots = s1, ss.month = s2,
perc.diff = round(100*2*abs(s1-s2)/(s1+s2), 1))

```
    toeplitz Form Symmetric Toeplitz Matrix

\section*{Description}

Forms a symmetric Toeplitz matrix given its first row.

\section*{Usage}
toeplitz (x)

\section*{Arguments}
\(\mathrm{x} \quad\) the first row to form the Toeplitz matrix.

\section*{Value}

The Toeplitz matrix.

\section*{Author(s)}
A. Trapletti

\section*{Examples}
```

x <- 1:5
toeplitz (x)

```
treering Yearly Treering Data, -6000-1979.

\section*{Description}

Contains normalized tree-ring widths in dimensionless units. Each tree ring corresponds to one year. Tree: Methuselah Walk, Pilo; Location: California, Gt Basin B C pine 2805M, 3726-11810; Author: Donald A. Graybill, 1980.

\section*{Usage}
```

data(treering)

```

\section*{Format}

A univariate time series with 7981 observations. The object is of class "ts".

\section*{Source}

Time Series Data Library: http://www-personal.buseco.monash.edu.au/~hyndman/ TSDL/
ts.plot Plot Multiple Time Series

\section*{Description}

Plot several time series on a common plot. Unlike plot.ts the series can have a different time bases, but they should have the same frequency.

\section*{Usage}
ts.plot(..., gpars = list())

\section*{Arguments}
\begin{tabular}{ll}
\(\ldots\). & one or more univariate or multivariate time series. \\
gpars & list of named graphics parameters to be passed to the plotting functions. \\
& Those commonly used can be supplied directly in \(\ldots\)
\end{tabular}

\section*{Value}

None.

\section*{Note}

Although this can be used for a single time series, plot is easier to use and is preferred.

\section*{Author(s)}
B.D. Ripley

\section*{See Also}
```

plot.ts

```

\section*{Examples}
data(UKLungDeaths)
ts.plot(ldeaths, mdeaths, fdeaths,
gpars=list(xlab="year", ylab="deaths", lty=c(1:3)))

\section*{Description}

Bind time series which have a common frequency. ts.union pads with NAs to the total time coverage, ts.intersect restricts to the time covered by all the series.

\section*{Usage}
```

ts.intersect(..., dframe = FALSE)
ts.union(..., dframe = FALSE)

```

\section*{Arguments}
\begin{tabular}{ll}
\(\ldots\). & two or more univariate or multivariate time series, or objects which can \\
coerced to time series.
\end{tabular}

\section*{Details}

As a special case, ... can contain vectors or matrices of the same length as the combined time series of the time series present, as well as those of a single row.

\section*{Value}

A time series object if dframe is FALSE, otherwise a data frame.

\section*{Author(s)}
B. D. Ripley

\section*{See Also}
cbind.

\section*{Examples}
```

data(UKLungDeaths)
ts.union(mdeaths, fdeaths)
cbind(mdeaths, fdeaths) \# same as the previous line
ts.intersect(window(mdeaths, 1976), window(fdeaths, 1974, 1978))
data(BJsales)
sales1 <- ts.union(BJsales, lead = BJsales.lead)
ts.intersect(sales1, lead3 = lag(BJsales.lead, -3))

```

\section*{tsdiag Diagnostic Plots for Time-Series Fits}

\section*{Description}

A generic function to plot time-series diagnostics.

\section*{Usage}
tsdiag(object, gof.lag, ...)

\section*{Arguments}
object a fitted time-series model
gof.lag the maximum number of lags for a Portmanteau goodness-of-fit test
... further arguments to be passed to particular methods

\section*{Details}

This is a generic function. It will generally plot the residuals, often standadized, the autocorrelation function of the residuals, and the p-values of a Portmanteau test for all lags up to gof.lag.
The methods for arima and StructTS objects plots residuals scaled by the estimate of their (individual) variance, and use the Ljung-Box version of the portmanteau test.

\section*{Value}

None. Diagnostics are plotted.

\section*{Note}
arima0.diag was an earlier version using the Box-Pierce test which is now deprecated.

\section*{Author(s)}
B. D. Ripley

\section*{See Also}
```

arima, StructTS, Box.test

```

\section*{Examples}
```

data(lh)
fit <- arima(lh, c(1,0,0))
tsdiag(fit)

## see also examples(arima)

data(JohnsonJohnson)
(fit <- StructTS(log10(JohnsonJohnson), type="BSM"))
tsdiag(fit)

```

\section*{tsSmooth Use Fixed-Interval Smoothing on Time Series}

\section*{Description}

Performs fixed-interval smoothing on a univariate time series via a state-space model. Fixedinterval smoothing gives the best estimate of the state at each time point based on the whole observed series.
```

Usage
tsSmooth(object, ...)

```

\section*{Arguments}
object a time-series fit. Currently only class "StructTS" is supported ... possible arguments for future methods.

\section*{Value}

A time series, with as many dimensions as the state space and results at each time point of the original series. (For seasonal models, only the current seasonal component is returned.)

\section*{Author(s)}
B. D. Ripley

\section*{References}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press.

\section*{See Also}

KalmanSmooth, StructTS.
For examples consult AirPassengers, JohnsonJohnson and Nile.

UKDriverDeaths Road Casualties in Great Britain 1969-84

\section*{Description}

UKDriverDeaths is a time series giving the monthly totals of car drivers in Great Britain killed or seriously injured Jan 1969 to Dec 1984. Compulsory wearing of seat belts was introduced on 31 Jan 1983.

Seatbelts is more information on the same problem.

\section*{Usage}
data(UKDriverDeaths)
data(Seatbelts)

\section*{Format}

Seatbelts is a multiple time series, with columns

DriversKilled car drivers killed.
drivers same as UKDriverDeaths.
front front-seat passengers killed or seriously injured.
rear rear-seat passengers killed or seriously injured.
kms distance driven.
PetrolPrice petrol price.
VanKilled number of van ('light goods vehicle') drivers.
law \(0 / 1\) : was the law in effect that month?

\section*{Source}

Harvey, A.C. (1989) Forecasting, Structural Time Series Models and the Kalman Filter. Cambridge University Press, pp. 519-523.

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press. http://www.ssfpack.com/dkbook/

\section*{References}

Harvey, A. C. and Durbin, J. (1986) The effects of seat belt legislation on British road casualties: A case study in structural time series modelling. Journal of the Royal Statistical Society series B, 149, 187-227.

\section*{Examples}
```

data(UKDriverDeaths)

## work with pre-seatbelt period to identify a model, use logs

work <- window(log10(UKDriverDeaths), end = 1982+11/12)
par(mfrow = c(3,1))
plot(work); acf(work); pacf(work)
par(mfrow = c(1,1))
(fit <- arima(work, c(1,0,0), seasonal = list(order= c(1,0,0))))
z <- predict(fit, n.ahead = 24)
ts.plot(log10(UKDriverDeaths), z$pred, z$pred+2*z$se, z$pred-2*z\$se,
lty = c(1,3,2,2), col = c("black", "red", "blue", "blue"))

## now see the effect of the explanatory variables

data(Seatbelts)
X <- Seatbelts[, c("kms", "PetrolPrice", "law")]
X[, 1] <- log10(X[, 1]) - 4
arima(log10(Seatbelts[, "drivers"]), c(1,0,0),
seasonal = list(order= c(1,0,0)), xreg = X)

```

UKgas UK Quarterly Gas Consumption

\section*{Description}

Quarterly UK gas consumption from 1960Q1 to 1986Q4, in millions of therms.

\section*{Usage}
data(UKgas)

\section*{Format}

A quarterly time series of length 108.

\section*{Source}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press. http://www.ssfpack.com/dkbook/

\section*{Examples}
```

data(UKgas)

## maybe str(UKgas) ; plot(UKgas) ...

```
UKLungDeaths Monthly Deaths from Lung Diseases in the UK

\section*{Description}

Three time series giving the monthly deaths from bronchitis, emphysema and asthma in the UK, 1974-1979, both sexes (ldeaths), males (mdeaths) and females (fdeaths).

\section*{Usage}
data(UKLungDeaths)

\section*{Source}
P. J. Diggle (1990) Time Series: A Biostatistical Introduction. Oxford, table A. 3

\section*{Examples}
```

data(UKLungDeaths)
plot(ldeaths)
plot(mdeaths, fdeaths)

## Better labels:

yr <- floor(tt <- time(mdeaths))
plot(mdeaths, fdeaths,
xy.labels = paste(month.abb[12*(tt - yr)], yr-1900, sep=")"))

```

\section*{USAccDeaths Accidental Deaths in the US 1973-1978}

\section*{Description}

A time series giving the monthly totals of accidental deaths in the USA. The values for the first six months of 1979 are 779874068363846092179316.

\section*{Usage}
data(USAccDeaths)

\section*{Source}
P. J. Brockwell and R. A. Davis (1991) Time Series: Theory and Methods. Springer, New York.

\section*{Description}

A time series of the numbers of users connected to the Internet through a server every minute.

\section*{Usage}
data(WWWusage)

\section*{Format}

A time series of length 100 .

\section*{Source}

Durbin, J. and Koopman, S. J. (2001) Time Series Analysis by State Space Methods. Oxford University Press. http://www.ssfpack.com/dkbook/

\section*{References}

Makridakis, S., Wheelwright, S. C. and Hyndman, R. J. (1998) Forecasting: Methods and Applications. Wiley.

\section*{Examples}
```

data(WWWusage)
work <- diff(WWWusage)
par(mfrow = c(2,1)); plot(WWWusage); plot(work)
aics <- matrix(, 6, 6, dimnames=list(p=0:5, q=0:5))
for(q in 1:5) aics[1, 1+q] <- arima(WWWusage, c(0,1,q),
optim.control = list(maxit = 500))$aic
for(p in 1:5)
    for(q in 0:5) aics[1+p, 1+q] <- arima(WWWusage, c(p,1,q),
        optim.control = list(maxit = 500))$aic
round(aics - min(aics, na.rm=TRUE), 2)

```

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[^1]:    x
    object from which to extract elements
    $\mathrm{i}, \mathrm{j}, \ldots$, name elements to extract or replace
    drop For data frames, matrices, and arrays. If TRUE the result is coerced to the lowest possible dimension (see examples below).

[^2]:    write
    Write Data to a File

