

Package ‘sfsmisc’

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Title Utilities from 'Seminar fuer Statistik' ETH Zurich

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Description Useful utilities ['goodies'] from Seminar fuer Statistik ETH Zurich, some of which were ported from S-plus in the 1990s.
For graphics, have pretty (Log-scale) axes `eaxis()`, an enhanced Tukey-Anscombe plot, combining histogram and boxplot, 2d-residual plots, a `tachoPlot()`, pretty arrows, etc.
For robustness, have a robust F test and `robust range()`.
For system support, notably on Linux, provides `'Sys.*()'` functions with more access to system and CPU information.
Finally, miscellaneous utilities such as simple efficient prime numbers, integer codes, `Duplicated()`, `toLatex.numeric()` and `is.whole()`.

Depends R (>= 3.3.0)

Imports grDevices, utils, stats, tools

Suggests datasets, tcltk, cluster, lattice, MASS, Matrix, nlme,
lokern, Rmpfr, gmp

Enhances mgcv, rpart, nor1mix, polycor, sm, tikzDevice, DescTools,
e1071, Hmisc, pastecs, polynom, relevance, robustbase

EnhancesNote 2nd line: packages mentioned in Rd xrefs

Encoding UTF-8

ByteCompile yes

License GPL (>= 2)

URL <https://github.com/mmaechler/sfsmisc>

BugReports <https://github.com/mmaechler/sfsmisc/issues>

NeedsCompilation no

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Description

AsciiToInt returns [integer](#) codes in 0:255 for each (one byte) character in strings. `ichar` is an alias for it, for old S compatibility.

`strcodes` implements in R the basic engine for translating characters to corresponding integer codes.

`chars8bit()` is the *inverse* function of `AsciiToInt`, producing “one byte” characters from integer codes. Note that it (and hence `strcodes()`) depends on the locale, see [Sys.getlocale\(\)](#).

Usage

```
AsciiToInt(strings)
  ichar(strings)
chars8bit(i = 1:255)
strcodes(x, table = chars8bit(1:255))
```

Arguments

<code>strings, x</code>	character vector.
<code>i</code>	numeric (integer) vector of values in 1:255.
<code>table</code>	a vector of (unique) character strings, typically of one character each.

Details

Only codes in 1:127 make up the ASCII encoding which should be identical for all R versions, whereas the ‘*upper*’ half is often determined from the ISO-8859-1 (aka “ISO-Latin 1”) encoding, but may well differ, depending on the locale setting, see also [Sys.setlocale](#).

Note that 0 is no longer allowed since, R does not allow `\0` aka `nul` characters in a string anymore.

Value

`AsciiToInt` (and hence `ichar`) and `chars8bit` return a vector of the same length as their argument.

`strcodes(x, tab)` returns a [list](#) of the same [length](#) and [names](#) as `x` with list components of integer vectors with codes in 1:255.

Author(s)

Martin Maechler, partly in 1991 for S-plus

Examples

```

chars8bit(65:70)#-> "A" "B" .. "F"
stopifnot(identical(LETTERS, chars8bit(65:90)),
           identical(AsciiToInt(LETTERS), 65:90))

## may only work in ISO-latin1 locale (not in UTF-8):
try( strcodes(c(a= "ABC", ch="1234", place = "Zürich")) )
## in "latin-1" gives {otherwise should give NA instead of 252}:
## Not run:
$a
[1] 65 66 67

$ch
[1] 49 50 51 52

$place
[1] 90 252 114 105 99 104

## End(Not run)

myloc <- Sys.getlocale()

if(.Platform $ OS.type == "unix") withAutoprint({ # ''should work'' here
  try( Sys.setlocale(locale = "de_CH") )# "try": just in case
  strcodes(c(a= "ABC", ch="1234", place = "Zürich")) # no NA hopefully
  AsciiToInt(chars8bit()) # -> 1:255 {if setting latin1 succeeded above}

  chars8bit(97:140)
  try( Sys.setlocale(locale = "de_CH.utf-8") )# "try": just in case
  chars8bit(97:140) ## typically looks different than above
})

## Resetting to original locale .. works "mostly":
lapply(strsplit(strsplit(myloc, ";")[[1]], "="),
       function(cc) try(Sys.setlocale(cc[1], cc[2]))) -> .scratch

Sys.getlocale() == myloc # TRUE if we have succeeded to reset it

```

axTexpr

*Axis Ticks Expressions in Nice 10^{**k} Form*

Description

Produce nice $a \times 10^k$ expressions for [axis](#) labeling instead of the scientific notation " $a \times 10^k$ ".

Usage

```

axTexpr(side, at = axTicks(side, axp = axp, usr = usr, log = log),
        axp = NULL, usr = NULL, log = NULL,
        drop.1 = FALSE)

```

Arguments

side	integer in 1:4 specifying the axis side, as for axis .
at	numeric vector; with identical default as in axTicks() .
axp, usr, log	as for axTicks() .
drop.1	logical indicating if $1 \times$ should be dropped from the resulting expressions.

Details

This is just a utility with the same arguments as [axTicks](#), a wrapper [pretty10exp](#)(at, *).

Value

an expression of the same length as x, with elements of the form `a %*% 10 ^ k`.

Author(s)

Martin Maechler

See Also

[pretty10exp](#); [eaxis](#), [axis](#), [axTicks](#).

Examples

```
x <- 1e7*(-10:50)
y <- dnorm(x, m=10e7, s=20e7)
plot(x,y)## not really nice, the following is better:

## For horizontal y-axis labels, need more space:
op <- par(mar= .1+ c(5,5,4,1))
plot(x,y, axes= FALSE, frame=TRUE)
aX <- axTicks(1); axis(1, at=aX, label= axExpr(1, aX))
## horizontal labels on y-axis:
aY <- axTicks(2); axis(2, at=aY, label= axExpr(2, aY), las=2)
par(op)

### -- only 'x' and using log-scale there:
plot(x,y, xaxt= "n", log = "x")
aX <- axTicks(1); axis(1, at=aX, label= axExpr(1, aX))

## Now an "engineer's version" ( more ticks; only label "10 ^ k" ) :

axp <- par("xaxp") #-> powers of 10 *inside* 'usr'
axp[3] <- 1 # such that only 10^ are labeled
aX <- axTicks(1, axp = axp)
xu <- 10 ^ par("usr")[1:2]
e10 <- c(-1,1) + round(log10(axp[1:2])) ## exponents of 10 *outside* 'usr'
v <- c(outer(1:9, e10[1]:e10[2], function(x,E) x * 10 ^ E))
v <- v[xu[1] <= v & v <= xu[2]]
```

```
plot(x,y, xaxt= "n", log = "x", main = "engineer's version of x - axis")
axis(1, at = aX, label = axExpr(1, aX, drop.1=TRUE)) # 'default'
axis(1, at = v, label = FALSE, tcl = 2/3 * par("tcl"))
```

cairoSwd

Cairo PDF Graphics Device useful for Sweave

Description

Provides a graphics device for Sweave, based on [cairo_pdf](#). The advantage of `cairoSwd()` compared to `pdf()` is its support of Unicode characters.

Usage

```
cairoSwd(name, width, height, ...)
```

Arguments

name	file name prefix to which ‘.pdf’ will be appended.
width,height	in inches, see cairo_pdf .
...	further arguments, passed to cairo_pdf()

Note

Sweave devices need to have an argument list as above.

Usage in a Sweave chunk:

```
<<some-plot, fig=TRUE, grdevice=cairoSwd>>=
```

Author(s)

Alain Hauser

See Also

[pdf](#), [cairo_pdf](#), [Sweave](#).

capture.and.write	<i>Capture output and Write / Print First and Last Parts</i>
-------------------	--

Description

Capture output and print first and last parts, eliding middle parts. Particularly useful for teaching purposes, and, e.g., in Sweave ([RweaveLatex](#)).

By default, when `middle = NA`, `capture.output(EXPR, first, last)` basically does

```
co <- capture.output(EXPR)
writeLines(head(co, first))
cat( ... dots ... )
writeLines(tail(co, last))
```

Usage

```
capture.and.write(EXPR, first, last = 2, middle = NA,
                  i.middle, dots = " ..... ", n.dots = 2)
```

Arguments

EXPR	the (literal) expression the output of which is to be captured.
first	integer: how many lines should be printed at beginning.
last	integer: how many lines should be printed at the end.
middle	numeric (or NA logical):
i.middle	index start of middle part
dots	string to be used for elided lines
n.dots	number of dots lines added between parts.

Value

return value of [capture.output](#)(EXPR).

Author(s)

Martin Maechler, ETH Zurich

See Also

[head](#), [tail](#)

Examples

```
x <- seq(0, 10, by = .1)

## for matrix, dataframe, .. first lines include a header line:
capture.and.write( cbind(x, log1p(exp(x))), first = 5)

## first, *middle* and last :
capture.and.write( cbind(x, x^2, x^3), first = 4, middle = 3, n.dots= 1)
```

col01scale

*Matrix Scaling Utilities***Description**

col01scale and colcenter (re)scale the columns of a matrix. These are simple one-line utilities, mainly with a didactical purpose.

Usage

```
colcenter (mat)
col01scale(mat, scale.func = function(x) diff(range(x)), location.func = mean)
```

Arguments

```
mat          numeric matrix, to rescaled.
scale.func, location.func
              two functions mapping a numeric vector to a single number.
```

Value

a matrix with the same attributes as the input mat.

Author(s)

Martin Maechler

See Also

The standard R function [scale\(\)](#).

Examples

```
## See the simple function definitions:

colcenter ## simply one line

col01scale# almost as simple
```

compresid2way

*Plot Components + Residuals for Two Factors***Description**

For an analysis of variance or regression with (at least) two factors: Plot components + residuals for two factors according to Tukey's "forget-it plot". Try it!

Usage

```
compresid2way(aov, data=NULL, fac=1:2, label = TRUE, numlabel = FALSE,
              xlab=NULL, ylab=NULL, main=NULL,
              col=c(2,3,4,4), lty=c(1,1,2,4), pch=c(1,2))
```

Arguments

aov	either an <code>aov</code> object with a formula of the form $y \sim a + b$, where a and b are factors, or such a formula.
data	data frame containing a and b .
fac	the two factors used for plotting. Either column numbers or names for argument data.
label	logical indicating if levels of factors should be shown in the plot.
numlabel	logical indicating if effects of factors will be shown in the plot.
xlab, ylab, main	the usual <code>title</code> components, here with a non-trivial default constructed from <code>aov</code> and the component factors used.
col, lty, pch	colors, line types, plotting characters to be used for plotting [1] positive residuals, [2] negative residuals, [3] grid, [4] labels. If <code>pch</code> is sufficiently long, it will be used as the list of individual symbols for plotting the y values.

Details

For a two-way analysis of variance, the plot shows the additive components of the fits for the two factors by the intersections of a grid, along with the residuals. The observed values of the target variable are identical to the vertical coordinate.

The application of the function has been extended to cover more complicated models. The components of the fit for two factors are shown as just described, and the residuals are added. The result is a "component plus residual" plot for two factors in one display.

Value

Invisibly, a list with components

compy	data.frame containing the component effects of the two factors, and combined effects plus residual
coef	coefficients: Intercept and effects of the factors

Author(s)

Werner Stahel <stahel@stat.math.ethz.ch>

References

F. Mosteller and J. W. Tukey (1977) *Data Analysis and Regression: A Second Course in Statistics*. Addison-Wesley, Reading, Mass., p. 176.

John W. Tukey (1977) *Exploratory Data Analysis*. Addison-Wesley, Reading, Mass., p. 381.

See Also

[interaction.plot](#)

Examples

```
## From Venables and Ripley (2002) p.165.
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.cr <- compresid2way(yield ~ N+P+K, data=npk, fac=c("P","K"))

## Fisher's 1926 data on potatoe yield
data(potatoes)
pot.aov <- aov(yield ~ nitrogen+potash+pos, data=potatoes)
compresid2way(pot.aov, pch=as.character(potatoes$pos))

compresid2way(yield~nitrogen+potash, data=subset(potatoes, pos == 2))

## 2 x 3 design :
data(warpbreaks)
summary(fm1 <- aov(breaks ~ wool + tension, data = warpbreaks))
compresid2way(fm1)
```

cum.Vert.funkt

Kumulative Verteilung Aufzeichnen

Description

Kumulative Verteilung von x aufzeichnen, auf Wunsch auch Median und Quartile.

This is just an old German language version of [plot.ecdf\(\)](#) used for teaching at ETHZ.

Usage

```
cum.Vert.funkt(x, Quartile = TRUE, titel = TRUE, Datum = TRUE,
              rang.axis = n <= 20, xlab = "", main = "", ...)
```

Arguments

<code>x</code>	numeric vector whose empirical distribution should be plotted.
<code>Quartile</code>	logical indicating if all 3 non-trivial quartiles should be drawn.
<code>titel</code>	logical indicating if a German title should be drawn.
<code>Datum</code>	logical indicating if <code>p.datum</code> should be added.
<code>rang.axis</code>	logical indicating if all the ranks should be marked at the y-axis. Defaults to true if there are not more than 20 observations.
<code>xlab, main</code>	x-axis label and main title; default to empty.
<code>...</code>	optional further arguments, passed to <code>plotStep</code> .

Value

the return value of `plotStep()` which is called internally, *invisibly*.

Author(s)

Martin Maechler et al.

See Also

`plotStep` on which it is based; but you should really consider using `plot.ecdf()` from the **stats** package instead of this.

Examples

```
cum.Vert.funkt(runif(12))
cum.Vert.funkt(runif(20))

Z <- rnorm(50)
cum.Vert.funkt(Z)
```

Description

Compute numerical derivatives of $f()$ given observations (x, y) , using cubic smoothing splines with GCV, see `smooth.spline`. In other words, estimate $f'()$ and/or $f''()$ for the model

$$Y_i = f(x_i) + E_i, \quad i = 1, \dots, n,$$

Usage

```
D1D2(x, y, xout = x, spar.offset = 0.1384, deriv = 1:2, spl.spar = NULL)
```

Arguments

<code>x, y</code>	numeric vectors of same length, supposedly from a model $y \sim f(x)$.
<code>xout</code>	abscissa values at which to evaluate the derivatives.
<code>spar.offset</code>	numeric fudge added to the smoothing parameter, see <code>spl.par</code> below.
<code>deriv</code>	integer in 1:2 indicating which derivatives are to be computed.
<code>spl.spar</code>	direct smoothing parameter for <code>smooth.spline</code> . If it is NULL (as per default), the smoothing parameter used will be <code>spar.offset + sp\$spar</code> , where <code>sp\$spar</code> is the GCV estimated smoothing parameter, see smooth.spline .

Details

It is well known that for derivative estimation, the optimal smoothing parameter is larger (more smoothing) than for the function itself. `spar.offset` is really just a *fudge* offset added to the smoothing parameter. Note that in R's implementation of [smooth.spline](#), `spar` is really on the $\log \lambda$ scale.

When `deriv = 1:2` (as per default), both derivatives are estimated with the *same* smoothing parameter which is suboptimal for the single functions individually. Another possibility is to call `D1D2(*, deriv = k)` twice with `k = 1` and `k = 2` and use a *larger* smoothing parameter for the second derivative.

Value

a list with several components,

<code>x</code>	the abscissae values at which the derivative(s) are evaluated.
<code>D1</code>	if <code>deriv</code> contains 1, estimated values of $f'(x_i)$ where x_i are the values from <code>xout</code> .
<code>D2</code>	if <code>deriv</code> contains 2, estimated values of $f''(x_i)$.
<code>spar</code>	the smoothing parameter used in the (final) <code>smooth.spline</code> call.
<code>df</code>	the equivalent degrees of freedom in that <code>smooth.spline</code> call.

Author(s)

Martin Maechler, in 1992 (for S).

See Also

[D2ss](#) which calls `smooth.spline` twice, first on `y`, then on the $f'(x_i)$ values; [smooth.spline](#) on which it relies completely.

Examples

```
set.seed(8840)
x <- runif(100, 0, 10)
y <- sin(x) + rnorm(100)/4

op <- par(mfrow = c(2,1))
```

```

plot(x,y)
lines(ss <- smooth.spline(x,y), col = 4)
str(ss[c("df", "spar")])
plot(cos, 0, 10, ylim = c(-1.5,1.5), lwd=2,
      main = expression("Estimating f'() : "~~ frac(d,dx) * sin(x) == cos(x)))
offs <- c(-0.1, 0, 0.1, 0.2, 0.3)
i <- 1
for(off in offs) {
  d12 <- D1D2(x,y, spar.offset = off)
  lines(d12$x, d12$D1, col = i <- i+1)
}
legend(2,1.6, c("true cos()",paste("sp.off. = ", format(offs))), lwd=1,
      col = 1:(1+length(offs)), cex = 0.8, bg = NA)
par(op)

```

D2ss

Numerical Derivatives of (x,y) Data (via Smoothing Splines)

Description

Compute the numerical first or 2nd derivatives of $f()$ given observations $(x[i], y \sim f(x[i]))$.

D1tr is the *trivial* discrete first derivative using simple difference ratios, whereas D1ss and D2ss use cubic smoothing splines (see [smooth.spline](#)) to estimate first or second derivatives, respectively.

D2ss first uses `smooth.spline` for the first derivative $f'()$ and then applies the same to the predicted values $\hat{f}'(t_i)$ (where t_i are the values of `xout`) to find $\hat{f}''(t_i)$.

Usage

```
D1tr(y, x = 1)
```

```
D1ss(x, y, xout = x, spar.offset = 0.1384, spl.spar=NULL)
```

```
D2ss(x, y, xout = x, spar.offset = 0.1384, spl.spar=NULL)
```

Arguments

<code>x, y</code>	numeric vectors of same length, supposedly from a model $y \sim f(x)$. For <code>D1tr()</code> , <code>x</code> can have length one and then gets the meaning of $h = \Delta x$.
<code>xout</code>	abscissa values at which to evaluate the derivatives.
<code>spar.offset</code>	numeric fudge added to the smoothing parameter(s), see <code>spl.par</code> below. Note that the current default is there for historical reasons only, and we often would recommend to use <code>spar.offset = 0</code> instead.
<code>spl.spar</code>	direct smoothing parameter(s) for <code>smooth.spline</code> . If it is <code>NULL</code> (as per default), the smoothing parameter used will be <code>spar.offset + sp\$spar</code> , where <code>sp\$spar</code> is the GCV estimated smoothing parameter for <i>both</i> smooths, see smooth.spline .

Details

It is well known that for derivative estimation, the optimal smoothing parameter is larger (more smoothing needed) than for the function itself. `spar.offset` is really just a *fudge* offset added to the smoothing parameters. Note that in R's implementation of [smooth.spline](#), `spar` is really on the $\log \lambda$ scale.

Value

`D1tr()` and `D1ss()` return a numeric vector of the length of `y` or `xout`, respectively.

`D2ss()` returns a list with components

<code>x</code>	the abscissae values (= <code>xout</code>) at which the derivative(s) are evaluated.
<code>y</code>	estimated values of $f''(x_i)$.
<code>spl.spar</code>	numeric vector of length 2, contain the <code>spar</code> arguments to the two <code>smooth.spline</code> calls.
<code>spar.offset</code>	as specified on input (maybe <code>rep()</code> eated to length 2).

Author(s)

Martin Maechler, in 1992 (for S).

See Also

[D1D2](#) which directly uses the 2nd derivative of the smoothing spline; [smooth.spline](#).

Examples

```
## First Derivative --- spar.off = 0 ok "asymptotically" (?)
set.seed(330)
mult.fig(12)
for(i in 1:12) {
  x <- runif(500, 0,10); y <- sin(x) + rnorm(500)/4
  f1 <- D1ss(x=x,y=y, spar.off=0.0)
  plot(x,f1, ylim = range(c(-1,1,f1)))
  curve(cos(x), col=3, add= TRUE)
}

set.seed(8840)
x <- runif(100, 0,10)
y <- sin(x) + rnorm(100)/4

op <- par(mfrow = c(2,1))
plot(x,y)
lines(ss <- smooth.spline(x,y), col = 4)
str(ss[c("df", "spar")])
xx <- seq(0,10, len=201)
plot(xx, -sin(xx), type = 'l', ylim = c(-1.5,1.5))
title(expression("Estimating f'()' : " * frac(d^2,dx^2) * sin(x) == -sin(x)))
offs <- c(0.05, 0.1, 0.1348, 0.2)
i <- 1
```

```

for(off in offs) {
  d12 <- D2ss(x,y, spar.offset = off)
  lines(d12, col = i <- i+1)
}
legend(2,1.6, c("true : -sin(x)",paste("sp.off. = ", format(offs))), lwd=1,
       col = 1:(1+length(offs)), cex = 0.8, bg = NA)
par(op)

```

 Deprecated

Deprecated 'sfsmisc' Functions

Description

These functions are provided for compatibility with older versions of the **sfsmisc** package only, and may be defunct as soon as of the next release.

Usage

```

pmax.sa(scalar, arr)
pmin.sa(scalar, arr)

```

Arguments

scalar	numeric scalar.
arr	any numeric R object, typically array.

Details

`pmax.sa(s, a)` and `pmin.sa(s, a)` return (more-dimensional) arrays. These have been deprecated, because `pmax` and `pmin` do so too, **if** the array is used as *first* argument.

 diagDA

Diagonal Discriminant Analysis

Description

This function implements a simple Gaussian maximum likelihood discriminant rule, for diagonal class covariance matrices.

In machine learning lingo, this is called “Naive Bayes” (for continuous predictors). Note that naive Bayes is more general, as it models discrete predictors as multinomial, i.e., binary predictor variables as Binomial / Bernoulli.

Usage

```
dDA(x, cll, pool = TRUE)
## S3 method for class 'dDA'
predict(object, newdata, pool = object$pool, ...)
## S3 method for class 'dDA'
print(x, ...)

diagDA(ls, cll, ts, pool = TRUE)
```

Arguments

<code>x, ls</code>	learning set data matrix, with rows corresponding to cases (e.g., mRNA samples) and columns to predictor variables (e.g., genes).
<code>cll</code>	class labels for learning set, must be consecutive integers.
<code>object</code>	object of class <code>dDA</code> .
<code>ts, newdata</code>	test set (prediction) data matrix, with rows corresponding to cases and columns to predictor variables.
<code>pool</code>	logical flag. If true (by default), the covariance matrices are assumed to be constant across classes and the discriminant rule is linear in the data. Otherwise (<code>pool=FALSE</code>), the covariance matrices may vary across classes and the discriminant rule is quadratic in the data.
<code>...</code>	further arguments passed to and from methods.

Value

`dDA()` returns an object of class `dDA` for which there are `print` and `predict` methods. The latter returns the same as `diagDA()`:

`diagDA()` returns an integer vector of class predictions for the test set.

Author(s)

Sandrine Dudoit, <sandrine@stat.berkeley.edu> and
Jane Fridlyand, <janef@stat.berkeley.edu> originally wrote `stat.diag.da()` in CRAN package **sma** which was modified for speedup by Martin Maechler <maechler@R-project.org> who also introduced `dDA` etc.

References

S. Dudoit, J. Fridlyand, and T. P. Speed. (2000) Comparison of Discrimination Methods for the Classification of Tumors Using Gene Expression Data. (Statistics, UC Berkeley, June 2000, Tech Report #576)

See Also

`lda` and `qda` from the **MASS** package; `naiveBayes` from **e1071**.

Examples

```
## two artificial examples by Andreas Greutert:
d1 <- data.frame(x = c(1, 5, 5, 5, 10, 25, 25, 25, 25, 29),
                 y = c(4, 1, 2, 4, 4, 4, 6:8, 7))

n.plot(d1)
library(cluster)
(cl1P <- pam(d1,k=4)$cluster) # 4 surprising clusters
with(d1, points(x+0.5, y, col = cl1P, pch =cl1P))

i1 <- c(1,3,5,6)
tr1 <- d1[-i1,]
cl1. <- c(1,2,1,2,1,3)
cl1 <- c(2,2,1,1,1,3)
plot(tr1, cex=2, col = cl1, pch = 20+cl1)
(dd.<- diagDA(tr1, cl1., ts = d1[ i1,]))# ok
(dd <- diagDA(tr1, cl1 , ts = d1[ i1,]))# ok, too!
points(d1[ i1,], pch = 10, cex=3, col = dd)

## use new fit + predict instead :
(r1 <- dDA(tr1, cl1))
(r1.<- dDA(tr1, cl1.))
stopifnot(dd == predict(r1, new = d1[ i1,]),
          dd.== predict(r1., new = d1[ i1,]))

plot(tr1, cex=2, col = cl1, bg = cl1, pch = 20+cl1,
      xlim=c(1,30), ylim= c(0,10))
xy <- cbind(x= runif(500, min=1,max=30), y = runif(500, min=0, max=10))
points(xy, cex= 0.5, col = predict(r1, new = xy))
abline(v=c( mean(c(5,25)), mean(c(25,29))))

## example where one variable xj has Var(xj) = 0:
x4 <- matrix(c(2:4,7, 6,8,5,6, 7,2,3,1, 7,7,7,7), ncol=4)
y <- c(2,2, 1,1)
m4.1 <- dDA(x4, y, pool = FALSE)
m4.2 <- dDA(x4, y, pool = TRUE)
xx <- matrix(c(3,7,5,7), ncol=4)
predict(m4.1, xx)## gave integer(0) previously
predict(m4.2, xx)
```

diagX

The “Other” Diagonal Matrix

Description

Compute the *other* diagonal identity matrix. The result is basically a *fast* version of `diag(n)[, n:1]`.

Usage

```
diagX(n)
```

Arguments

n positive integer.

Value

a numeric $n \times n$ matrix with many zeros – apart from 1s in the *other* diagonal.

Author(s)

Martin Maechler, 1992.

See Also

[diag](#).

Examples

```
diagX(4)
for(m in 1:5)
  stopifnot(identical(diagX(m), diag(m)[, m:1, drop = FALSE]))
```

digitsBase

Digit/Bit Representation of Integers in any Base

Description

Integer number representations in other Bases.

Formally, for every element $N = x[i]$, compute the (vector of) “digits” A of the base b representation of the number N , $N = \sum_{k=0}^M A_{M-k} b^k$.

Revert such a representation to integers.

Usage

```
digitsBase(x, base = 2, ndigits = 1 + floor(1e-9 + log(max(x,1), base)))
## S3 method for class 'basedInt'
as.integer(x, ...)
## S3 method for class 'basedInt'
print(x, ...)

as.intBase(x, base = 2)
bi2int(xlist, base)
```

Arguments

<code>x</code>	For <code>digitsBase()</code> : non-negative integer (vector) whose base base digits are wanted. For <code>as.intBase()</code> : a list of numeric vectors, a character vector, or an integer matrix as returned by <code>digitsBase()</code> , representing digits in base base.
<code>base</code>	integer, at least 2 specifying the base for representation.
<code>ndigits</code>	number of bits/digits to use.
<code>...</code>	potential further arguments passed to methods, notably <code>print</code> .
<code>xlist</code>	a list of integer vectors with entries typically in $0:(base-1)$, such as resulting from <code>digitsBase()</code> .

Value

For `digitsBase()`, an object, say `m`, of class "basedInt" which is basically a (`ndigits` x `n`) [matrix](#) where `m[,i]` corresponds to `x[i]`, `n <- length(x)` and `attr(m, "base")` is the input base.

`as.intBase()` and the [as.integer](#) method for `basedInt` objects return an [integer](#) vector.

`bi2int()` is the low-level workhorse of `as.intBase()`.

Note

Some of these functions existed under names `digits` and `digits.v` in previous versions of the `sfsmisc` package.

Author(s)

Martin Maechler, Dec 4, 1991 (for S-plus; then called `digits.v`).

Examples

```
digitsBase(0:12, 8) #-- octal representation
empty.dimnames(digitsBase(0:33, 2)) # binary

## This may be handy for just one number (and default decimal):
digits <- function(n, base = 10) as.vector(digitsBase(n, base = base))
digits(128982734)      # 1 2 8 9 8 2 7 3 4
digits(128, base = 8)  # 2 0 0

## one way of pretty printing (base <= 10!)
b2ch <- function(db)
  noquote(gsub("^0+({1,})$", " \\1",
    apply(db, 2, paste, collapse = "")))
b2ch(digitsBase(0:33, 2)) #-> 0 1 10 11 100 101 ... 100001
b2ch(digitsBase(0:33, 4)) #-> 0 1 2 3 10 11 12 13 20 ... 200 201

## Hexadecimal:
i <- c(1:20, 100:106)
M <- digitsBase(i, 16)
hexdig <- c(0:9, LETTERS[1:6])
```

```

cM <- hexdig[1 + M]; dim(cM) <- dim(M)
b2ch(cM) #-> 1 2 3 4 5 6 7 8 9 A B C D E F 10 11 ... 6A

## IP (Internet Protocol) numbers coding: <n>.<n>.<n>.<n> <-> longinteger
ip_ntoa <- function(n)
  apply(digitsBase(n, base = 256), 2, paste, collapse=".")
ip_ntoa(2130706430 + (0:9))# "126.255.255.254" ... "127.0.0.7"
## and the inverse:
ip_aton <- function(a)
  bi2int(lapply(strsplit(a, "."), fixed=TRUE), as.integer), 256)

n <- 2130706430 + (0:9)
head(ip <- ip_ntoa(n))
head(ip_aton(ip))
stopifnot( n == ip_aton(ip_ntoa(n )),
  ip == ip_ntoa(ip_aton(ip)))

## Inverse of digitsBase() : as.integer method for the "basedInt" class
as.integer(M)
## or also as.intBase() working from strings:
(cb <- apply(digitsBase(0:33, 4), 2, paste, collapse = ""))
##-> "000" "001" ..... "200" "201"
all(0:33 == as.intBase(cb, base = 4))

```

Duplicated

*Counting-Generalization of duplicated()***Description**

Duplicated() generalizes the [duplicated](#) method for vectors, by returning indices of “equivalence classes” for duplicated entries and returning nomatch (NA by default) for unique entries.

Note that duplicated() is not TRUE for the first time a duplicate appears, whereas Duplicated() only marks unique entries with nomatch (NA).

Usage

```
Duplicated(v, incomparables = FALSE, fromLast = FALSE, nomatch = NA_integer_)
```

Arguments

v	a vector, often character, factor, or numeric.
incomparables	a vector of values that cannot be compared, passed to both duplicated () and match (). FALSE is a special value, meaning that all values can be compared, and may be the only value accepted for methods other than the default. It will be coerced internally to the same type as x.
fromLast	logical indicating if duplication should be considered from the reverse side, i.e., the last (or rightmost) of identical elements would correspond to duplicated=FALSE.
nomatch	passed to match (): the value to be returned in the case when no match is found. Note that it is coerced to integer.

Value

an integer vector of the same length as `v`. Can be used as a [factor](#), e.g., in [split](#), [tapply](#), etc.

Author(s)

Christoph Buser and Martin Maechler, Seminar fuer Statistik, ETH Zurich, Sep.2007

See Also

[uniqueL](#) (also in this [sfsmisc](#) package); [duplicated](#), [match](#).

Examples

```
x <- c(9:12, 1:4, 3:6, 0:7)
data.frame(x, dup = duplicated(x),
           dupL= duplicated(x, fromLast=TRUE),
           Dup = Duplicated(x),
           DupL= Duplicated(x, fromLast=TRUE))
```

eaxis

Extended / Engineering Axis for Graphics

Description

An extended [axis\(\)](#) function which labels more prettily, in particular for log-scale axes.

It makes use of [plotmath](#) or (LaTeX) [expressions](#) of the form $k \times 10^k$ for labeling a log-scaled axis and when otherwise exponential formatting would be used (see [pretty10exp](#)).

Usage

```
eaxis(side, at = if(log) axTicks(side, axp=axp, log=log, nintLog=nintLog)
      else axTicks(side, axp=axp, log=log),
      labels = NULL, log = NULL,
      use.expr = log || format.info(as.numeric(at), digits=7)[3] > 0,
      f.smalltcl = 3/5, at.small = NULL, small.mult = NULL, equidist.at.tol = 0.002,
      small.args = list(),
      draw.between.ticks = TRUE, between.max = 4,
      outer.at = TRUE, drop.1 = TRUE, sub10 = FALSE, las = 1,
      nintLog = max(12, par("lab")[2 - is.x]),
      axp = NULL, n.axp = NULL, max.at = Inf,
      lab.type = "plotmath", lab.sep = "cdot",
      ...)
```

Arguments

side	integer in 1:4, specifying side of <code>axis</code> .
at	numeric vector of (“normalized”) tick locations; by default <code>axTicks(side, .)</code> , i.e., the same as <code>axis()</code> would use.
labels	NULL (default), <code>logical</code> , character or expression, as in <code>axis()</code> ; in addition, if NA, <code>labels = TRUE</code> is passed to <code>axis()</code> , i.e. <code>pretty10exp</code> is <i>not</i> used. Use FALSE to suppress any labeling.
log	logical or NULL specifying if log-scale should be used; the default depends on the current plot’s axis.
use.expr	logical specifying if <code>pretty10exp(.)</code> should be used for constructing labels when they are NULL. The default is typically good enough, but you may occasionally <i>force</i> <code>use.expr = TRUE</code> .
f.smalltcl	factor specifying the lengths of the small ticks in proportion to the normalized, labeled ticks.
at.small	locations of <i>small</i> ticks; the default, NULL, uses <code>small.mult</code> and constructs “smart” locations.
small.mult	positive integer (or NULL), used when <code>at.small</code> is NULL to indicate which multiples of <code>at</code> (typically <code>axTicks()</code>) should be used as “small ticks”. The default NULL will use 9 in the log case and a number in 2:5 otherwise.
equidist.at.tol	a small positive number, a tolerance to be used for checking equidistant at values. Used to be hardwired at .001 which was seen to be too small; increase it when necessary.
small.args	optional <code>list</code> of further arguments to the (second) <code>axis()</code> call which draws the <i>small</i> ticks.
draw.between.ticks	(only if <code>log</code> is true): logical indicating that possible (non-small) ticks between the labeled (via <code>at</code>) ones should be drawn as well (and possibly also used for <code>at.small</code> construction), see also <code>between.max</code> .
between.max	(only if <code>log</code> and <code>draw.between.ticks</code> are true): integer indicating ticks should be drawn (approximately) between the labeled ones.
outer.at	logical specifying that <code>at.small</code> should also be constructed outside the <code>at</code> range, but still inside the corresponding <code>par("usr")</code> .
drop.1	logical specifying if $1 \times$ should be dropped from labels, passed to <code>pretty10exp()</code> .
sub10	logical, integer (of length 1 or 2) or “10”, indicating if some 10^k should be simplified to “traditional” formats, see <code>pretty10exp</code> .
nintLog	only used in R > 2.13.x, when <code>log</code> is true: approximate (lower bound on) number of intervals for log scaling.
axp	to be passed to <code>axTicks()</code> if <code>at</code> is not specified.
n.axp	to be set to <code>axp[3]</code> when <code>axp</code> and <code>at</code> are not specified, in order to tweak the <i>number</i> of (non-small) tick marks produced from <code>axTicks(.)</code> , notably when <code>log</code> is true, set <code>n.axp</code> to 1, 2, or 3: 1: will produce tick marks at 10^j for integer j ,

	2: gives marks $k10^j$ with $k \in \{1, 5\}$,
	3: gives marks $k10^j$ with $k \in \{1, 2, 5\}$
	see 'xaxp' on the par help page.
max.at	maximal number of at values to be used effectively. If you don't specify at yourself carefully, it is recommended to set this to something like 25, but this is not the default, for back compatibility reasons.
las, ...	arguments passed to (the first) axis call. Note that the default las = 1 differs from axis's default las = 0.
lab.type	string, passed to pretty10exp to choose between default plotmath or LaTeX label format.
lab.sep	separator between mantissa and exponent for LaTeX labels, see pretty10exp .

Author(s)

Martin Maechler

See Also

[axis](#), [axTicks](#), [axTexpr](#), [pretty10exp](#).

Examples

```
x <- lseq(1e-10, 0.1, length = 201)
plot(x, pt(x, df=3), type = "l", xaxt = "n", log = "x")
eaxis(1)
## without small ticks:
eaxis(3, at.small=FALSE, col="blue")

## If you like the ticks, but prefer traditional (non-"plotmath") labels:
plot(x, gamma(x), type = "l", log = "x")
eaxis(1, labels=NA)

x <- lseq(.001, 0.1, length = 1000)
plot(x, sin(1/x)*x, type = "l", xaxt = "n", log = "x")
eaxis(1)
eaxis(3, n.axp = 1)# -> xaxp[3] = 1: only 10^j (main) ticks

## non- log-scale : draw small ticks, but no "10^k" if not needed:
x <- seq(-100, 100, length = 1000)
plot(x, sin(x)/x, type = "l", xaxt = "n")
eaxis(1) # default -> {1, 2, 5} * 10^j ticks
eaxis(3, n.axp = 2)# -> xaxp[3] := 2 -- approximately two (main) ticks

x <- seq(-1, 1, length = 1000)
plot(x, sin(x)/x, type = "l", xaxt = "n")
eaxis(1, small.args = list(col="blue"))

x <- x/1000
plot(x, 1-sin(x)/x, type = "l", xaxt = "n", yaxt = "n")
eaxis(1)
```

```

eaxis(2)
## more labels than default:
op <- par(lab=c(10,5,7))
plot(x, sin(x)/x, type = "l", xaxt = "n")
eaxis(1) # maybe (depending on your canvas), there are too many,
## in that case, maybe use
plot(x, sin(x)/x, type = "l", xaxt = "n")
eaxis(1, axTicks(1)[c(TRUE,FALSE)]) # drop every 2nd label
eaxis(3, labels=FALSE)

## ore use 'max.at' which thins as well:
plot(x, sin(x)/x, type = "l", xaxt = "n")
eaxis(1, max.at=6)
par(op)

### Answering R-help "How do I show real values on a log10 histogram", 26 Mar 2013
## the data:
set.seed(1); summary(x <- rlnorm(100, m = 2, sdl = 3))
## the plot (w/o x-axis) :
r <- hist(log10(x), xaxt = "n", xlab = "x [log scale]")
## the nice axis:
axt <- axTicks(1)
eaxis(1, at = axt, labels = pretty10exp(10^axt, drop.1=TRUE))
## Additionally demo'ing 'sub10' options:
plot(r, xaxt="n")
eaxis(1, at = axt, labels = pretty10exp(10^axt, drop.1=TRUE, sub10 = 2))
## or
plot(r, xaxt="n")
eaxis(1, at = axt, labels = pretty10exp(10^axt, drop.1=TRUE, sub10 = "10"))
## or
plot(r, xaxt="n")
eaxis(1, at = axt, labels = pretty10exp(10^axt, drop.1=TRUE, sub10 = c(-2, 2)))

```

ecdf.ksCI

Plot Empirical Distribution Function With 95% Confidence Curves

Description

Plots the empirical (cumulative) distribution function (ECDF) for univariate data, together with upper and lower simultaneous 95% confidence curves, computed via Kolmogorov-Smirnov' D , see [KSd](#).

Usage

```

ecdf.ksCI(x, main = NULL, sub = NULL, xlab = deparse(substitute(x)),
          ci.col = "red", ...)

```

Arguments

`x` `x` numerical vector of observations.
`main, sub, xlab` arguments passed to [title](#).
`ci.col` color for confidence interval lines.
`...` optional arguments passed to [plot.stepfun](#).

Value

Nothing. Used for its side effect, to produce a plot.

Note

Presently, will only work if `length(x) > 9`.

Author(s)

Kjetil Halvorsen

References

Bickel and Doksum, see [KSd](#).

See Also

[ecdf](#) and [plot.stepfun](#) in standard R.

Examples

```
ecdf.ksCI( rchisq(50,3) )
```

`ellipsePoints`*Compute Radially Equispaced Points on Ellipse*

Description

Compute points on (the boundary of) an ellipse which is given by elementary geometric parameters.

Usage

```
ellipsePoints(a, b, alpha = 0, loc = c(0, 0), n = 201, keep.ab.order=FALSE)
```

Arguments

a, b	length of half axes in (x,y) direction. Note that (a, b) is equivalent to (b, a) <i>unless</i> keep.ab.order=TRUE.
alpha	angle (in degrees) giving the orientation of the ellipse, i.e., the original (x,y)-axis ellipse is rotated by angle.
loc	center (LOCation) of the ellipse.
n	number of points to generate.
keep.ab.order	logical indicating if (a, b) should be considered <i>ordered</i> . When FALSE, as per default, the orientation of the ellipse is solely determined by alpha. Note that keep.ab.order = TRUE seems a more natural default, but FALSE is there for back-compatibility.

Value

A numeric matrix of dimension $n \times 2$, each row containing the (x,y) coordinates of a point.

Author(s)

Martin Maechler, March 2002.

See Also

the **ellipse** package and **ellipsoidhull** and **ellipsoidPoints** in the **cluster** package.

Examples

```
## Simple Ellipse, centered at (0,0), x-/y- axis parallel:
ep <- ellipsePoints(5,2)
str(ep)
plot(ep, type="n",asp=1) ; polygon(ep, col = 2)
## (a,b) = (2,5) is equivalent to (5,2) :
lines(ellipsePoints(2,5), lwd=2, lty=3)
## keep.order=TRUE : Now, (2,5) are axes in x- respective y- direction:
lines(ellipsePoints(2,5, keep.ab.order=TRUE), col="blue")

## rotate by 30 degrees :
plot(ellipsePoints(5,2, alpha = 30), asp=1)
abline(h=0,v=0,col="gray")
abline(a=0,b= tan( 30 *pi/180), col=2, lty = 2)
abline(a=0,b= tan(120 *pi/180), col=3, lty = 2)

## NB: use x11(type = "Xlib") for the following if you can
if(dev.interactive(TRUE)) {
  ## Movie : rotating ellipse :
  nTurns <- 4 # #{full 360 deg turns}
  for(al in 1:(nTurns*360)) {
    ep <- ellipsePoints(3,6, alpha=al, loc = c(5,2))
    plot(ep,type="l",xlim=c(-1,11),ylim=c(-4,8),
      asp=1, axes = FALSE, xlab="", ylab="")
  }
}
```

```

    }

    ## Movie : rotating _filled_ ellipse {less nice to look at}
    for(al in 1:180) {
      ep <- ellipsePoints(3,6, alpha=al, loc = c(5,2))
      plot(ep,type="n",xlim=c(-1,11),ylim=c(-4,8),
        asp=1, axes = FALSE, xlab="", ylab="")
      polygon(ep,col=2,border=3,lwd=2.5)
    }
  }# only if interactive

```

empty.dimnames	<i>Empty Dimnames of an Array</i>
----------------	-----------------------------------

Description

Remove all dimension names from an array for compact printing.

Usage

```
empty.dimnames(a)
```

Arguments

`a` an [array](#), i.e., as special case a matrix.

Value

Returns `a` with its `dimnames` replaced by empty character strings.

Author(s)

Bill Venables / Martin Maechler, Sept 1993.

See Also

[unname](#) *removes* the `dimnames`.

Examples

```

empty.dimnames(diag(5)) # looks much nicer

(a <- matrix(-9:10, 4,5))
empty.dimnames(a) # nicer, right?

```

errbar

Scatter Plot with Error Bars

Description

Draws a scatter plot, adding vertical “error bars” to all the points.

Usage

```
errbar(x, y, yplus, yminus, cap = 0.015,
       ylim = range(y, yplus, yminus),
       xlab= deparse(substitute(x)),
       ylab= deparse(substitute(y)), ...)
```

Arguments

x	vector of x values.
y	vector of y values.
yplus	vector of y values: the tops of the error bars.
yminus	vector of y values: the bottoms of the error bars.
cap	the width of the little lines at the tops and bottoms of the error bars in units of the width of the plot. Default is 0.015.
ylim	(numeric of length 2): the y-axis extents with a sensible default.
xlab, ylab	axis labels for the plot, as in plot.default .
...	Graphical parameters (see par) may also be supplied as arguments to this function.

Author(s)

Originally Charles Geyer, U.Chicago, early 1991; then Martin Mächler.

See Also

[errbar](#) in package **Hmisc** is similar.

Examples

```
y <- rnorm(10); d <- 1 + .1*rnorm(10)
errbar(1:10, y, y + d, y - d, main="Error Bars example")
```

f.robftest*Robust F-Test: Wald test for multiple coefficients of rlm() Object*

Description

Compute a robust F-Test, i.e., a Wald test for multiple coefficients of an `rlm` object.

Usage

```
f.robftest(object, var = -1)
```

Arguments

<code>object</code>	result of <code>rlm()</code> .
<code>var</code>	variables. Either their names or their indices; the default, <code>-1</code> means all <i>but</i> the intercept.

Details

This builds heavily on `summary.rlm()`, the `summary` method for `rlm` results.

Value

An object of class `"htest"`, hence with the standard print methods for hypothesis tests. This is basically a list with components

<code>statistic</code>	the F statistic, according to ...
<code>df</code>	numerator and denominator degrees of freedom.
<code>data.name</code>	(extracted from input object.)
<code>alternative</code>	<code>"two.sided"</code> , always.
<code>p.value</code>	the P-value, using an F-test on <code>statistic</code> and <code>df[1:2]</code> .

Author(s)

Werner Stahel, July 2000; updates by Martin Maechler.

References

FIXME — Need some here !

See Also

`rlm`, `summary.aov`, etc.

Examples

```
if(require("MASS")) {
  ## same data as example(rlm)
  data(stackloss)
  summary(rsl <- rlm(stack.loss ~ ., stackloss))
  f.robftest(rsl)
} else " forget it "
```

factorize

*Prime Factorization of Integers***Description**

Compute the prime factorization(s) of integer(s) n .

Usage

```
factorize(n, verbose = FALSE)
```

Arguments

n vector of integers to factorize.
 verbose logical indicating if some progress information should be printed.

Details

works via [primes](#), currently in a cheap way, sub-optimal for large composite n .

Value

A named [list](#) of the same length as n , each element a 2-column matrix with column "p" the prime factors and column "m" their respective exponents (or multiplities), i.e., for a prime number n , the resulting matrix is `cbind(p = n, m = 1)`.

Author(s)

Martin Maechler, Jan. 1996.

See Also

[primes](#).

For factorization of moderately or really large numbers, see the [gmp](#) package, and its `factorize()`.

Examples

```
factorize(47)
factorize(seq(101, 120, by=2))
```

funEnv	<i>List-like Environment of Functions (and More)</i>
--------	--

Description

Construct a “list”, really an [environment](#) typically of functions and optionally other R objects, where the [functions](#) and [formulas](#) all share the same environment. Consequently, the functions may call each other.

On technical level, this is just a simple wrapper around [list2env\(\)](#).

Usage

```
funEnv(..., envir = NULL, parent = parent.frame(),
       hash = (...length() > 100), size = max(29L, ...length()))
```

Arguments

<code>...</code>	an arbitrary <i>named</i> “list” of R objects, typically including several functions .
<code>envir</code>	an environment or <code>NULL</code> .
<code>parent</code>	(for the case <code>envir = NULL</code>): a parent frame aka enclosing environment, see new.env and list2env .
<code>hash, size</code>	(for the case <code>envir = NULL</code>): <code>hash</code> a logical indicating if the created environment should use hashing, and <code>(size)</code> the hash size, see list2env .

Value

an [environment](#), say `E`, containing the objects from `...` (plus those in `envir`), and all function objects’ [environment\(\)](#) is `E`.

Author(s)

Martin Maechler

See Also

[list2env](#), [environment](#)

Examples

```
ee <- funEnv(f = function(x) g(2*(x+1)),
            g = function(y) hh(y+1),
            hh = function(u) u^2,
            info = "Some Information (not a function)")
ls(ee) # here the same as names(ee)
## Check that it works: i.e., that "f sees g" and "g sees hh":
stopifnot(all.equal(ee$f(pi), (2*pi+3)^2))
ee$f(0:4) # [1] 9 25 49 81 121
```

hatMat	<i>Hat Matrix of a Smoother</i>
--------	---------------------------------

Description

Compute the hat matrix or smoother matrix, of ‘any’ (linear) smoother, smoothing splines, by default.

Usage

```
hatMat(x, trace= FALSE,
      pred.sm = function(x, y, ...)
                    predict(smooth.spline(x, y, ...), x = x)$y,
      ...)
```

Arguments

<code>x</code>	numeric vector or matrix.
<code>trace</code>	logical indicating if the whole hat matrix, or only its trace, i.e. the sum of the diagonal values should be computed.
<code>pred.sm</code>	a function of at least two arguments (<code>x</code> , <code>y</code>) which returns fitted values, i.e. $\hat{y}$, of length compatible to <code>x</code> (and <code>y</code>).
<code>...</code>	optionally further arguments to the smoother function <code>pred.sm</code> .

Value

The hat matrix H (if `trace = FALSE` as per default) or a number, $tr(H)$, the *trace* of H , i.e., $\sum_i H_{ii}$.

Note that `dim(H) == c(n, n)` where `n <- length(x)` also in the case where some `x` values are duplicated (aka *ties*).

Author(s)

Martin Maechler <maechler@stat.math.ethz.ch>

References

Hastie and Tibshirani (1990). *Generalized Additive Models*. Chapman & Hall.

See Also

[smooth.spline](#), etc. Note the demo, `demo("hatmat-ex")`.

Examples

```
require(stats) # for smooth.spline() or loess()

x1 <- c(1:4, 7:12)
H1 <- hatMat(x1, spar = 0.5) # default : smooth.spline()

matplot(x1, H1, type = "l", main = "columns of smoother hat matrix")

## Example 'pred.sm' arguments for hatMat() :
pspl <- function(x,y,...) predict(smooth.spline(x,y, ...), x = x)$y
pksm <- function(x,y,...) ksmooth(sort(x),y, "normal", x.points=x, ...)$y
## Rather than ksmooth():
if(require("lokern"))
  pksm2 <- function(x,y,...) glkerns(x,y, x.out=x, ...)$est

## Explaining 'trace = TRUE'
all.equal(sum(diag((hatMat(c(1:4, 7:12), df = 4))),
             hatMat(c(1:4, 7:12), df = 4, trace = TRUE), tol = 1e-12)

## ksmooth() :
Hk <- hatMat(x1, pr = pksm, bandwidth = 2)
cat(sprintf("df = %.2f\n", sum(diag(Hk))))
image(Hk)
Matrix::printSpMatrix(as(round(Hk, 2), "sparseMatrix"))

##--> see demo("hatmat-ex") for more (and larger) examples
```

helppdf

help() Type="pdf" and View It

Description

Utility to view PDF-rendered [help](#) pages; particularly useful in case they contain mathematical formulas or otherwise sophisticated formats.

Usage

```
helppdf(topic, viewer = getOption("pdfviewer"), quiet = !interactive(), ...)
```

Arguments

topic	the topic, passed to help() .
viewer	a pdf viewer; the default is typically what you want interactively.
quiet	logical indicating that nothing should be printed to the console and the result should be returned as invisible() .
...	further optional arguments passed to help() .

Value

Returns the full path of the pdf file produced.

Author(s)

Martin Maechler

See Also

[help](#), [system](#).

Examples

```
if(interactive()) {
  ## Both calls work :
  helppdf(Normal)
  helppdf("NegBinomial")
} else if(.Platform$OS.type != "windows") { # batch mode (Windows often too slow for this)
  od <- setwd(tempdir())
  ff <- helppdf(Normal, viewer=NULL)
  stopifnot(file.exists(ff)) ; print(ff)
  setwd(od)# revert to previous dir.
}
```

histBxp

Plot a Histogram and a Boxplot

Description

Creates a histogram and a horizontal boxplot on the current graphics device.

Usage

```
histBxp(x, nclass, breaks, probability=FALSE, include.lowest=TRUE,
        xlab = deparse(substitute(x)),
        ...,
        width=0.2, boxcol=3, medcol=2, medlwd=5, whisklty=2, staplelty=1)
```

Arguments

x	numeric vector of data for histogram. Missing values (NAs) are allowed.
nclass	recommendation for the number of classes (i.e., bars) the histogram should have. The default is a number proportional to the logarithm of the length of x.
breaks	vector of the break points for the bars of the histogram. The count in the i-th bar is <code>sum(breaks[i] < x <= breaks[i+1])</code> except that if <code>include.lowest</code> is TRUE (the default), the first bar also includes points equal to <code>breaks[1]</code> . If omitted, evenly-spaced break points are determined from <code>nclass</code> and the extremes of the data.

probability	logical flag: if TRUE, the histogram will be scaled as a probability density; the sum of the bar heights times bar widths will equal 1. If FALSE, the heights of the bars will be counts.
include.lowest	If TRUE (the default), the lowest bar will include data points equal to the lowest break, otherwise it will act like the other bars (see the description of the breaks argument).
xlab	character or expression for x axis labeling.
...	additional arguments to barplot . The hist function uses the function <code>barplot</code> to do the actual plotting; consequently, arguments to the <code>barplot</code> function that control shading, etc., can also be given to <code>hist</code> . See the <code>barplot</code> documentation for arguments <code>angle</code> , <code>density</code> , <code>col</code> , and <code>inside</code> . Do not use the space or <code>histo</code> arguments.
width	width of the box relative to the height of the histogram. DEFAULT is 0.2.
boxcol	color of filled box. The default is 3.
medcol	the color of the median line. The special value, NA, indicates the current plotting color (<code>par("col")</code>). The default is 2. If <code>boxcol=0</code> and <code>medcol</code> is not explicitly specified this is set to the current plotting color (<code>par("col")</code>).
medlwd	median line width. The special value NA, is used to indicate the current line width (<code>par("lwd")</code>). The default is 5.
whisklty	whisker line type. The special value NA indicates the current line type (<code>par("lty")</code>). The default is 2 (dotted line).
staplelty	staple (whisker end cap) line type. The special value NA indicates the current line type (<code>par("lty")</code>). The default is 1 (solid line).

Graphical parameters (see [par](#)) may also be supplied as arguments to this function. In addition, the high-level graphics arguments described under `par` and the arguments to `title` may be supplied to this function.

Details

If `include.lowest` is FALSE the bottom breakpoint must be strictly less than the minimum of the data, otherwise (the default) it must be less than or equal to the minimum of the data. The top breakpoint must be greater than or equal to the maximum of the data.

This function has been called `hist.bxp()` for 17 years; in 2012, the increasingly strong CRAN policies required a new name (which could not be confused with an S3 method name).

Author(s)

S-Plus: Markus Keller, Christian Keller; port to R in 1990's: Martin Mächler.

See Also

[hist](#), [barplot](#), [boxplot](#), [rug](#) and [scat1d](#) in the [Hmisc](#) package.

Examples

```
lab <- "50 samples from a t distribution with 5 d.f."
mult.fig(2*3, main = "Hist() + Rug() and histBxp(*)")
for(i in 1:3) {
  my.sample <- rt(50, 5)
  hist(my.sample, main=lab); rug(my.sample)# for 50 obs., this is ok, too..
  histBxp(my.sample, main=lab)
}
```

integrate.xy

*Cheap Numerical Integration through Data Points***Description**

Given (x_i, f_i) where $f_i = f(x_i)$, compute a cheap approximation of $\int_a^b f(x)dx$.

Usage

```
integrate.xy(x, fx, a, b, use.spline=TRUE, xtol=2e-08)
```

Arguments

x	abscissa values.
fx	corresponding values of $f(x)$.
a, b	the boundaries of integration; these default to min(x) and max(x) respectively.
use.spline	logical; if TRUE use an interpolating spline.
xtol	tolerance factor, typically around <code>sqrt(.Machine\$double.eps)</code>(fixme)....

Details

Note that this is really not good for noisy fx values; probably a smoothing spline should be used in that case.

Also, we are not yet using Romberg in order to improve the trapezoid rule. This would be quite an improvement in equidistant cases.

Value

the approximate integral.

Author(s)

Martin Maechler, May 1994 (for S).

See Also

[integrate](#) for numerical integration of *functions*.

Examples

```
x <- 1:4
integrate.xy(x, exp(x))
print(exp(4) - exp(1), digits = 10) # the true integral

for(n in c(10, 20, 50, 100, 200)) {
  x <- seq(1, 4, len = n)
  cat(formatC(n, wid=4), formatC(integrate.xy(x, exp(x)), dig = 9), "\n")
}
```

inv.seq

*Inverse seq() – Short Expression for Index Vector***Description**

Compute a short expression for a given integer vector, typically an index, that can be expressed shortly, using `:` etc.

Usage

```
inv.seq(i)
```

Arguments

`i` vector of (usually increasing) integers.

Value

a `call` (“the inside of an [expression](#)”) to be `eval()`ed to return the original `i`.

Author(s)

Martin Maechler, October 1995; more elegant implementation from Tony Plate.

See Also

[rle](#) for another kind of integer vector coding.

Examples

```
(rr <- inv.seq(i1 <- c(3:12, 20:24, 27, 30:33)))
eval(rr)
stopifnot(eval(rr) == i1)

e2 <- expression(c(20:13, 3:12, -1:-4, 27, 30:31))
(i2 <- eval(e2))
(r2 <- inv.seq(i2))
stopifnot(all.equal(r2, e2[[1]]))
```

```
## Had {mapply()} bug in this example:
ii <- c(1:3, 6:9, 11:16)
stopifnot(identical(ii, eval(inv.seq(ii))))
```

is.whole

Test Whether a Vector or Array Consists of Whole Numbers

Description

This function tests whether a numeric or complex vector or array consists of whole numbers. The function `is.integer` is not appropriate for this since it tests whether the vector is of class `integer` (see examples).

Usage

```
is.whole(x, tolerance = sqrt(.Machine$double.eps))
```

Arguments

<code>x</code>	integer, numeric, or complex vector or array to be tested
<code>tolerance</code>	maximal distance to the next whole number

Value

The return value has the same dimension as the argument `x`: if `x` is a vector, the function returns a logical vector of the same length; if `x` is a matrix or array, the function returns a logical matrix or array of the same dimensions. Each entry in the result indicates whether the corresponding entry in `x` is whole.

Author(s)

Alain Hauser <alain@huschhus.ch>

See Also

[is.integer](#)

Examples

```
## Create a random array, matrix, vector
set.seed(307)
a <- array(runif(24), dim = c(2, 3, 4))
a[4:8] <- 4:8
m <- matrix(runif(12), 3, 4)
m[2:4] <- 2:4
v <- complex(real = seq(0.5, 1.5, by = 0.1),
             imaginary = seq(2.5, 3.5, by = 0.1))

## Find whole entries
```

```

is.whole(a)
is.whole(m)
is.whole(v)

## Numbers of class integer are always whole
is.whole(dim(a))
is.whole(length(v))

```

iterate.lin.recursion *Generate Sequence Iterating a Linear Recursion*

Description

Generate numeric sequences applying a linear recursion `nr.it` times.

Usage

```
iterate.lin.recursion(x, coeff, delta = 0, nr.it)
```

Arguments

<code>x</code>	numeric vector with <i>initial values</i> , i.e., specifying the beginning of the resulting sequence; must be of length (larger or) equal to <code>length(coeff)</code> .
<code>coeff</code>	coefficient vector of the linear recursion.
<code>delta</code>	numeric scalar added to each term; defaults to 0. If not zero, determines the linear drift component.
<code>nr.it</code>	integer, number of iterations.

Value

numeric vector, say `r`, of length `n + nr.it`, where `n = length(x)`. Initialized as `r[1:n] = x`, the recursion is `r[k+1] = sum(coeff * r[(k-m+1):k])`, where `m = length(coeff)`.

Note

Depending on the zeroes of the characteristic polynomial of `coeff`, there are three cases, of convergence, oscillation and divergence.

Author(s)

Martin Maechler

See Also

[seq](#) can be regarded as a trivial special case.

Examples

```
## The Fibonacci sequence:
iterate.lin.recursion(0:1, c(1,1), nr = 12)
## 0 1 1 2 3 5 8 13 21 34 55 89 144 233

## seq() as a special case:
stopifnot(iterate.lin.recursion(4,1, d=2, nr=20)
          == seq(4, by=2, length=1+20))

## 'Deterministic AR(2)':
round(iterate.lin.recursion(1:4, c(-0.7, 0.9), d = 2, nr=15), dig=3)
## slowly decaying :
plot(ts(iterate.lin.recursion(1:4, c(-0.9, 0.95), nr=150)))
```

KSd

Approximate Critical Values for Kolmogorov-Smirnov's D

Description

Computes the critical value for Kolmogorov-Smirnov's D_n , for sample sizes $n \geq 10$ and confidence level 95%.

Usage

KSd(n)

Arguments

n the sample size, $n \geq 10$.

Details

Based on tables values given in the reference below. For $n \leq 80$ uses interpolations from exact values, elsewhere uses asymptotic approximation.

Value

The critical value for D (two-sided) for significance level 0.05 (or confidence level 95%).

Author(s)

Kjetil Halvorsen and Martin Maechler

References

Peter J. Bickel and Kjell A. Doksum (1977), *Mathematical Statistics: Basic Ideas and Selected Topics*. Holden Day. Section 9.6 and table IX.

See Also

Is used from [ecdf.ksCI](#).

Examples

```
KSd(90)
KSd(1:9)# now works

op <- par(mfrow=c(2,1))
plot(KSd, 10, 150)# nice
abline(v = c(75,85), col = "gray")
plot(KSd, 79, 81, n = 1001)# very tiny discontinuity at 80
par(op)
```

last	<i>Get Last Elements of a Vector</i>
------	--------------------------------------

Description

Extract the last elements of a vector.

Usage

```
last(x, length.out = 1, na.rm = FALSE)
```

Arguments

x	any vector.
length.out	integer indicating how many element are desired. If positive, return the length.out last elements of x; if negative, the last length.out elements are <i>dropped</i> .
na.rm	logical indicating if the last non-missing value (if any) shall be returned. By default (it is FALSE and) the last elements (whatever its values) are returned.

Value

a vector of length `abs(length.out)` of *last* values from x.

Note

This function may eventually be deprecated for the standard R function [tail\(\)](#).

Useful for the [turnogram\(\)](#) function in package [pasteecs](#).

Author(s)

Werner Stahel (<stahel@stat.math.ethz.ch>), and independently, Philippe Grosjean (<phgrosjean@sciviews.org>),
Frédéric Ibanez (<ibanez@obs-vlfr.fr>).

See Also

[first](#), [turnogram](#)

Examples

```
a <- c(NA, 1, 2, NA, 3, 4, NA)
last(a)
last(a, na.rm=TRUE)

last(a, length = 2)
last(a, length = -3)
```

linesHyperb.lm

Plot Confidence or Prediction Hyperbolas around a Regression Line

Description

Add confidence/prediction hyperbolas for $y(x_0)$ to a plot with data or regression line.

Usage

```
linesHyperb.lm(object, c.prob=0.95, confidence=FALSE,
               k=if (confidence) Inf else 1,
               col=2, lty=2, do.abline=TRUE)
```

Arguments

object	result of <code>lm(.)</code> .
c.prob	coverage probability in (0, 1).
confidence	logical; if true, do (small) confidence band, else, realistic prediction band for the mean of k observations.
k	integer or Inf; assume k future observations; k = Inf corresponds to confidence intervals (for y).
col, lty	attributes for the lines to be drawn.
do.abline	logical; if true, the regression line is drawn as well.

Note

With `predict.lm(*, interval=)` is available, this function `linesHyperb.lm` is only slightly more general for its k argument.

Author(s)

Martin Maechler, Oct 1995

See Also

`predict.lm`(*, interval=) optionally computes prediction or confidence intervals.

Examples

```
data(swiss)
plot(Fertility ~ Education, data = swiss) # the data
(lmS <- lm(Fertility ~ Education, data = swiss))
linesHyperb.lm(lmS)
linesHyperb.lm(lmS, conf=TRUE, col="blue")
```

list_named	<i>Automatically Named list()</i>
------------	-----------------------------------

Description

A version of `list(...)`, but with “automatically” *named* list components.

Usage

```
list_(...)
```

Arguments

... components to make up the resulting `list`. Their variable names (or unevaluated expressions in the call) will become the `names(.)` of the result.

Details

The names are extracted from `sys.call()`, and the function is written to be *fast* (rather than easy to ready for the uninitiated ;-)

Value

a `list` with the components in the arguments with `names` taken from their call to `list_()`.

Author(s)

Martin Maechler

See Also

`list`, `names`

Examples

```
a <- 1:4; lett <- letters[1:9]; CH <- "Suisse"
all.equal(list(a, lett),
          list_(a, lett)) # "names for current but not for target"
str(list(a, lett, CH)) # [[1]], [[2]], .. (no names)
str(list_(a, lett, CH))#  $a  $lett ..

stopifnot(identical(
  list(a, lett, CH),
  unname(L <- list_(a, lett, CH))),
  is.list(L), names(L) == c("a", "lett", "CH"),
  identical(lett, L$lett) ## etc
)

## The function is currently defined as
function (...) `names<-`(list(...), vapply(sys.call()[-1L], as.character, ""))
```

loessDemo

Graphical Interactive Demo of loess()

Description

A graphical and interactive demonstration and visualization of how [loess](#) works. By clicking on the graphic, the user determines the current estimation window which is visualized together with the weights.

Usage

```
loessDemo(x, y, span = 1/2, degree = 1, family = c("gaussian", "symmetric"),
  nearest = FALSE, nout = 501,
  xlim = numeric(0), ylim = numeric(0), strictlim = TRUE, verbose = TRUE,
  inch.sym = 0.25, pch = 4, shade = TRUE, w.symbols = TRUE,
  sym.col = "blue", w.col = "light blue", line.col = "steelblue")
```

Arguments

x, y	numeric vectors of the same length; the demo is about loess (y ~ x).
span	the smoothing parameter α .
degree	the degree of the polynomials to be used; must be in 0, 1, 2.
family	if "gaussian" fitting is by least-squares, and if "symmetric" a re-descending M estimator is used with Tukey's biweight function. Can be abbreviated.
nearest	logical indicating how x_0 should be determined, the value at which $\hat{f}(x_0)$ is computed. If nearest is true, the closest <i>data</i> value is taken.
nout	the number of points at which to evaluate, i.e, determining $u_i, i = 1, 2, \dots, nout$, at which $\hat{f}(u_i)$ is computed.
xlim	x-range; to extend or determine (iff strictlim is true) the <i>x</i> -range for plotting.
ylim	y-range; to extend or determine (iff strictlim is true) the <i>y</i> -range for plotting.

<code>strictlim</code>	logical determining if <code>xlim</code> and <code>ylim</code> should be strict limits (as e.g., in plot.default), or just a suggestion to <i>extend</i> the data-dependent ranges.
<code>verbose</code>	logical
<code>inch.sym</code>	symbol size in inches of the maximal weight circle symbol.
<code>pch</code>	plotting character, see points .
<code>shade</code>	logical; if true, polygon (<code>..</code> , <code>density=..</code>) will be used to shade off the regions where the weights are zero.
<code>w.symbols</code>	logical indicating if the non-zero weights should be visualized by circles with radius proportional to <code>inch.sym</code> and $\sqrt{w}$ where w are the weights.
<code>sym.col, w.col, line.col</code>	colors for the symbols, weights and lines, respectively.

Author(s)

As function `loess.demo()`, written and posted to S-news, on 27 Sep 2001, by Greg Snow, Brigham Young University, it was modified by Henrik Aa. Nielsen, IMM, DTU, and subsequently spiffed up for R by Martin Maechler.

See Also

[loess](#).

Examples

```
if(dev.interactive()) {
  if(requireNamespace("lattice")) {
    data("ethanol", package = "lattice")
    attach(ethanol)
    loessDemo(E,N0x, span=.25)
    loessDemo(E,N0x, span=.25, family = "symmetric")

    loessDemo(E,N0x, degree=0)# Tricube Kernel estimate
  }

  ## Artificial Example with one outlier
  n2 <- 50; x <- 1:(1+2*n2)
  fx <- (x/10 - 5)^2
  y <- fx + 4*rnorm(x)
  y[n2+1] <- 1e4
  loessDemo(x,y, span=1/3, ylim= c(0,1000))# not robust !!
  loessDemo(x,y, span=1/3, family = "symm")
  loessDemo(x,y, span=1/3, family = "symm", w.symb = FALSE, ylim = c(0,40))
  loessDemo(x,y, span=1/3, family = "symm", ylim = c(0,40))
  ## but see warnings() --- there's a "fixup"
}
```

lseq	<i>Generate Sequences, Equidistant on Log Scale</i>
------	---

Description

Generate sequences which are equidistant on a log-scale.

Usage

```
lseq(from, to, length)
```

Arguments

from	starting value of sequence.
to	end value of the sequence.
length	desired length of the sequence.

Value

a [numeric](#) vector of length length.

See Also

[seq](#).

Examples

```
(x <- lseq(1, 990, length= 21))
plot(x, x^4, type = "b", col = 2, log = "xy")
if(with(R.version, major >= 2 && minor >= 1))
plot(x, exp(x), type = "b", col = 2, log = "xy")
```

mat2tex	<i>Produce LaTeX commands to print a matrix</i>
---------	---

Description

“Translate” an R matrix (like object) into a LaTeX table, using `\begin{tabular}`

Usage

```
mat2tex(x, file= "mat.tex", envir = "tabular",
        nam.center = "l", col.center = "c",
        append = TRUE, digits = 3, title)
```

Arguments

x	a matrix
file	names the file to which LaTeX commands should be written
envir	a string, the LaTeX environment name; default is "tabular"; useful maybe "array", or other versions of tabular environments.
nam.center	character specifying row names should be center; default "1".
col.center	character (vector) specifying how the columns should be centered; must have values from c("l", "c", "r"); defaults to "c".
append	logical; if FALSE, will destroy the file file before writing commands to it; otherwise (by default), simply adds commands at the end of file file.
digits	integer; setting of <code>options(digits=.)</code> for purpose of number representation.
title	a string, possibly using LaTeX commands, which will span the columns of the LaTeX matrix

Value

No value is returned. This function, when used correctly, only writes LaTeX commands to a file.

Author(s)

For S: Vincent Carey <vjcarey@sphunix.sph.jhu.edu>, from a post on Feb.19, 1991 to S-news.
Port to R (and a bit more) by Martin Maechler <maechler@stat.math.ethz.ch>.

See Also

`latex` in package **Hmisc** is more flexible (but may surprise by its auto-printing ..).

Examples

```
mex <- matrix(c(pi,pi/2,pi/4,exp(1),exp(2),exp(3)),nrow=2, byrow=TRUE,
              dimnames = list(c("$\\pi$", "$e$"), c("a", "b", "c")))
mat2tex(mex, file = print(tf <- tempfile("mat", , ".tex")),
        title="$\\pi$, e$, etc." )
```

```
## The last command produces the file "mat<xyz>.tex" containing
```

```
##> \begin{tabular} {| l || c| c| c|}
##> \multicolumn{ 4 }{c}{ $\\pi$, e$, etc. } \\ \hline
##> \ & a & b & c \\ \hline \hline
##> $\\pi$ & 3.14 & 1.57 & 0.785 \\ \hline
##> $e$ & 2.72 & 7.39 & 20.1 \\ \hline
##> \end{tabular}
```

```
## Now you have to properly embed the contents of this file
## in a LaTeX document -- for example, you will need a
## preamble, the \begin{document} statement, etc.
```

```
## Note that the backslash needs protection in dimnames
```

```
## or title actions.

mat2tex(mex, stdout(), col.center = c("r","r","c"))
```

missingCh

Has a Formal Argument been Set or is it Missing?

Description

missingCh can be used to test whether a value was specified as an argument to a function. Very much related to the standard R function [missing](#), here the argument is given by its name, a character string.

As missingCh() calls missing(), do consider the caveats about the latter, see [missing](#).

Usage

```
missingCh(x, envir = parent.frame())
```

Arguments

x	a character string.
envir	a (function evaluation) environment , in which the variable named x is to be “missing”.

Value

a [logical](#) indicating if the argument named x is [missing](#) in the function “above”, typically the caller of missingCh, but see the use of envir in the vapply example.

Author(s)

Martin Maechler

See Also

[missing](#)

Examples

```
tst1 <- function(a, b, dd, ...) ## does not work an with argument named 'c' !
  c(b = missingCh("b"), dd = missingCh("dd"))
tst1(2)#-> both 'b' and 'dd' are missing
tst1(,3,,3)
##      b      dd
## FALSE TRUE  -- as 'b' is not missing but 'dd' is.

Tst <- function(a,b,cc,dd,EEE, ...)
  vapply(c("a","b","cc","dd","EEE"), missingCh, NA, envir=environment())
```

```

Tst()
## TRUE ... TRUE -- as all are missing()
Tst(1,,3)
##      a      b      cc      dd      EEE
## FALSE TRUE FALSE TRUE TRUE
## .....
## as 'a' and 'cc' where not missing()

## Formal testing:
stopifnot(tst1(), !tst1(,3,3), Tst(),
          Tst(1,,3, b=2, E="bar") == c(0,0,1,0,0))
## maybe surprising that this ^^ becomes 'dd' and only 'cc' is missing

```

mpl

Simple Matrix Plots

Description

Do simple matrix plots, providing an easy interface to [matplot](#) by using a default x variable.

Usage

```

mpl(mat, ...)
p.m(mat, ...)

```

Arguments

mat	numeric matrix.
...	further arguments passed to matplot , e.g., type, xlab, etc.

Details

`p.m(m)` use the first column of `m` as *x* variable, whereas `mpl(m)` uses the integers 1, 2, ..., `nrow(m)` as coordinates and `rownames(m)` as axis labels if possible.

Note

These were really created for playing around with curves etc, and probably should be *deprecated* since in concrete examples, using `matplot()` directly is more appropriate.

Author(s)

Martin Maechler

See Also

[matplot](#), [plot.mts](#)(*, `plot.type = "single"`).

Examples

```
data(animals, package = "cluster")
mpl(animals, type = "l")
```

mult.fig

Plot Setup for MULTiple FIGures, incl. Main Title

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

```
mult.fig(nr.plots, mfrow, mfcol, marP = rep(0, 4),
        mpg = c(if(par("las") != 0) 2. else 1.5, 0.6, 0),
        mar = marP + 0.1 + c(4,4,2,1), oma = c(0,0, tit.wid, 0),
        main = NULL,
        tit.wid = if (is.null(main)) 0 else 1 + 1.5*cex.main,
        cex.main = par("cex.main"), line.main = cex.main - 1/2,
        col.main = par("col.main"), font.main = par("font.main"), ...)
```

Arguments

<code>nr.plots</code>	integer; the number of plot figures you'll want to draw.
<code>mfrow, mfcol</code>	<i>instead</i> of <code>nr.plots</code> : integer(2) vectors giving the rectangular figure layout for <code>par(mfrow = *)</code> , or <code>par(mfcol = *)</code> , respectively. The default is to use <code>mfrow = n2mfrow(nr.plots)</code> .
<code>marP</code>	numeric(4) vector of figure margins to <i>add</i> ("Plus") to default <code>mar</code> , see below.
<code>mpg</code>	argument for <code>par(mpg = .)</code> with a smaller default than usual.
<code>mar</code>	argument for <code>par(mar = .)</code> with a smaller default than usual, using the <code>marP</code> argument, see above.
<code>oma</code>	argument for <code>par(oma = .)</code> , by default for adding space for the main title if necessary.
<code>main</code>	character. The main title to be used for the whole graphic.
<code>tit.wid</code>	numeric specifying the vertical width to be used for the main title; note that this is only used for the default value of <code>oma</code> (s. above).
<code>cex.main</code>	numeric; the character size to be used for the main title.
<code>line.main</code>	numeric; the margin line at which the title is written (via <code>mtext(main, side=3, outer=TRUE, line = line.main, ...)</code>).
<code>col.main, font.main</code>	color and font for main title, passed to <code>mtext()</code> , see also <code>par(*)</code> .
<code>...</code>	further arguments to <code>mtext</code> for the main title.

Value

A [list](#) with two components that are lists themselves, a subset of [par\(\)](#),

new.par the current par settings.

old.par the par *before* the call.

Author(s)

Martin Maechler, UW Seattle, 1990 (for S).

See Also

[par](#), [layout](#).

Examples

```
opl <- mult.fig(5, main= expression("Sine Functions " * sin(n * pi * x)))
x <- seq(0, 1, len = 201)
for (n in 1:5)
  plot(x, sin(n * pi * x), ylab = "", main = paste("n = ",n))
par(opl$old.par)

rr <- mult.fig(mfrow=c(5,1), main= "Cosinus Funktionen", cex = 1.5,
              marP = - c(0, 1, 2, 0))
for (n in 1:5)
  plot(x, cos(n * pi * x), type = 'l', col="red", ylab = "")
str(rr)
par(rr$old.par)
## The *restored* par settings:
str(do.call("par", as.list(names(rr$new.par))))

## Manual setting of `tit.wid` in case subsequent code also manages par():
mult.fig(4, tit.wid = 2)$old.par -> opar
plot(lm(sr ~ pop15 + pop75 + dpi + ddpi, data = LifeCycleSavings))
par(opar) # reset
```

n.code

Convert "Round" Integers to Short Strings and Back

Description

n.code convert “round integers” to short character strings. This is useful to build up variable names in simulations, e.g.

code2n is the *inverse* function of n.code().

Usage

```
n.code(n,      ndig = 1, dec.codes = c("", "d", "c", "k"))
code2n(ncod, ndig = 1, dec.codes = c("", "d", "c", "k"))
```

Arguments

n	integer vector.
ncod	character vector, typically resulting from n.code.
ndig	integer giving number of digits before the coding character.
dec.codes	character code for 1, 10, 100, 1000 (etc).

Value

n.code(n) returns a [character](#) vector of the same length as n.
 code2n(ncod) returns a [integer](#) vector of the same length as ncod.
 Usually, code2n(n.code(n)) == n.

Author(s)

Martin Maechler

Examples

```
n10 <- c(10,20,90, 100,500, 2000,10000)
(c10 <- n.code(n10))#-> "1d" "2d" "9d" "1c" ..
stopifnot(code2n(c10) == n10)
```

n.plot

Name Plot: Names or Numbers instead of Points in Plot

Description

A utility function which basically calls [plot](#)(*, type="n") and [text](#). To have names or numbers instead of points in a plot is useful for identification, e.g., in a residual plot, see also [TA.plot](#).

Usage

```
n.plot(x, y = NULL, nam = NULL, abbr = n >= 20 || max(nchar(nam))>=8,
       xlab = NULL, ylab = NULL, log = "",
       cex = par("cex"), col = par("col"), ...)
```

Arguments

x, y	coordinates at which to plot. If y is missing, x is used for both, if it's a data.frame , list , 2-column matrix etc – via xy.coords ; formula do not work.
nam	the labels to plot at each (x,y). Per default, these taken from the data x and y; case numbers 1:n are taken if no names are available.
abbr	logical indicating if the nam labels should be abbreviated – with a sensible default.
xlab, ylab	labels for the x- and y- axis, the latter being empty by default.

log	character specifying if log scaled axes should be used, see plot.default .
cex	plotting character expansion, see par .
col	color to use for text() .
...	further arguments to be passed to the plot call.

Value

invisibly, a character vector with the labels used.

Author(s)

Martin Maechler, since 1992

See Also

[plot.default](#), [text](#).

Examples

```
n.plot(1:20, cumsum(rnorm(20)))
data(cars)
with(cars, n.plot(speed, dist, cex = 0.8, col = "forest green"))
```

nearcor

Find the Nearest Proper Correlation Matrix

Description

This function “smoothes” an improper correlation matrix as it can result from [cor](#) with use=”pairwise.complete.obs” or [hetcor](#).

It is *deprecated* now, in favor of [nearPD\(\)](#) from package **Matrix**.

Usage

```
nearcor(R, eig.tol = 1e-6, conv.tol = 1e-07, posd.tol = 1e-8,
        maxits = 100, verbose = FALSE)
```

Arguments

R	a square symmetric approximate correlation matrix
eig.tol	defines relative positiveness of eigenvalues compared to largest, default=1e-6.
conv.tol	convergence tolerance for algorithm, default=1.0e-7
posd.tol	tolerance for enforcing positive definiteness, default=1.0e-8
maxits	maximum number of iterations
verbose	logical specifying if convergence monitoring should be verbose.

Details

This implements the algorithm of Higham (2002), then forces symmetry, then forces positive definiteness using code from [posdefify](#). This implementation does not make use of direct LAPACK access for tuning purposes as in the MATLAB code of Lucas (2001). The algorithm of Knol DL and ten Berge (1989) (not implemented here) is more general in (1) that it allows constraints to fix some rows (and columns) of the matrix and (2) to force the smallest eigenvalue to have a certain value.

Value

A [list](#), with components

<code>cor</code>	resulting correlation matrix
<code>fnorm</code>	Froebenius norm of difference of input and output
<code>iterations</code>	number of iterations used
<code>converged</code>	logical

Author(s)

Jens Oehlschlägel

References

See those in [posdefify](#).

See Also

the slightly more flexible [nearPD](#) which also returns a *classed* matrix (class `dpoMatrix`). For new code, `nearPD()` is really preferred to `nearcor()`, which hence is considered deprecated.

[hetcor](#), [eigen](#); [posdefify](#) for a simpler algorithm.

Examples

```
cat("pr is the example matrix used in Knol DL, ten Berge (1989)\n")
pr <- matrix(c(1,      0.477, 0.644, 0.478, 0.651, 0.826,
0.477, 1,      0.516, 0.233, 0.682, 0.75,
0.644, 0.516, 1,      0.599, 0.581, 0.742,
0.478, 0.233, 0.599, 1,      0.741, 0.8,
0.651, 0.682, 0.581, 0.741, 1,      0.798,
0.826, 0.75, 0.742, 0.8, 0.798, 1),
  nrow = 6, ncol = 6)

ncr <- nearcor(pr)
nr <- ncr$cor

plot(pr[lower.tri(pr)],
     nr[lower.tri(nr)]); abline(0,1, lty=2)
round(cbind(eigen(pr)$values, eigen(nr)$values), 8)
```

```

cat("The following will fail:\n")
try(factanal(cov=pr, factors=2))
cat("and this should work\n")
try(factanal(cov=nr, factors=2))

if(require("polycor")) {

  n <- 400
  x <- rnorm(n)
  y <- rnorm(n)

  x1 <- (x + rnorm(n))/2
  x2 <- (x + rnorm(n))/2
  x3 <- (x + rnorm(n))/2
  x4 <- (x + rnorm(n))/2

  y1 <- (y + rnorm(n))/2
  y2 <- (y + rnorm(n))/2
  y3 <- (y + rnorm(n))/2
  y4 <- (y + rnorm(n))/2

  dat <- data.frame(x1, x2, x3, x4, y1, y2, y3, y4)

  x1 <- ordered(as.integer(x1 > 0))
  x2 <- ordered(as.integer(x2 > 0))
  x3 <- ordered(as.integer(x3 > 1))
  x4 <- ordered(as.integer(x4 > -1))

  y1 <- ordered(as.integer(y1 > 0))
  y2 <- ordered(as.integer(y2 > 0))
  y3 <- ordered(as.integer(y3 > 1))
  y4 <- ordered(as.integer(y4 > -1))

  odat <- data.frame(x1, x2, x3, x4, y1, y2, y3, y4)

  xcor <- cor(dat)
  pcor <- cor(data.matrix(odat)) # cor() no longer works for factors
  hcor <- hetcor(odat, ML=TRUE, std.err=FALSE)$correlations
  ncor <- nearcor(hcor)$cor

  try(factanal(covmat=xcor, factors=2, n.obs=n))
  try(factanal(covmat=pcor, factors=2, n.obs=n))
  try(factanal(covmat=hcor, factors=2, n.obs=n))
  try(factanal(covmat=ncor, factors=2, n.obs=n))
}

```

nr.sign.chg

*Number of Sign Changes in Sequence***Description**

Compute the number of sign changes in the sequence y .

Usage

```
nr.sign.chg(y)
```

Arguments

y numeric vector.

Value

an integer giving the number of sign changes in sequence y. Note that going from positive to 0 to positive is *not* a sign change.

Author(s)

Martin Maechler, 17 Feb 1993.

Examples

```
(y <- c(1:2,1:-1,0:-2))
nr.sign.chg(y)## = 1
```

p.arrows

Prettified Arrows Plots

Description

Draws arrows, like the [arrows](#) function, but with “nice” *filled* arrow heads.

Usage

```
p.arrows(x1, y1, x2, y2, size = 1, width, fill = 2, ...)
```

Arguments

x1, y1	coordinates of points from which to draw.
x2, y2	coordinates of points to which to draw.
size	symbol size as a fraction of a character height; default 1.
width	width of the arrow head; defaults to
fill	color for filling the arrow head.
...	further arguments passed to segments() .

Author(s)

Andreas Ruckstuhl, 19 May 1994; (cosmetic by MM).

See Also

[arrows.](#)

Examples

```
example(arrows, echo = FALSE) #-> x, y, s
plot(x,y, main="p.arrows(.)")
p.arrows(x[s], y[s], x[s+1], y[s+1], col= 1:3, fill = "dark blue")
```

p.datum

Plot 'Datum' (deutsch!) unten rechts

Description

Plot the date (and time, if required) in German, at the lower right hand margin of your plot.`date`

Usage

```
p.datum(outer = FALSE, cex = 0.75, ...)
```

Arguments

<code>outer</code>	logical; passed to mtext.
<code>cex</code>	non-negative; passed to mtext.
<code>...</code>	any arguments to u.Datumvonheute.

See Also

[u.date](#), [date.](#)

Examples

```
plot(1)
p.datum()
```

p.dnorm

*Plot Parametric Density Functions***Description**

These are utilities for pretty plotting of often used parametric densities.

Usage

```
p.dnorm(mu = 0, s = 1, h0.col = "light gray",
        ms.lines = TRUE, ms.col = "gray", ...)
p.dchisq(nu, h0.col = "light gray", ...)
p.dgamma(shape, h0.col = "light gray", ...)
```

Arguments

mu, s	numbers, the mean and standard deviation of the normal distribution.
nu	positive number, the degrees of freedom df argument for the χ^2 -density function dchisq .
shape	number, the shape parameter for the Gamma distribution.
h0.col	color specification for the line $y = 0$.
ms.lines	logical, used for the normal only: should lines be drawn at the mean and ± 1 standard deviation.
ms.col	color for the ms lines if ms.lines is TRUE.
...	further parameter passed to curve() , e.g., add = TRUE for adding to current plot.

Author(s)

Werner Stahel et al.

See Also

the underlying density functions, [dnorm](#), [dchisq](#), [dgamma](#).

Examples

```
p.dnorm()
p.dnorm(mu=1.5, add = TRUE, ms.lines = FALSE) # add to the plot above

p.dchisq(2, main= "Chi^2 Densities -- nu = 2,3,4")
p.dchisq(3, add = TRUE, col = "red")
p.dchisq(4, add = TRUE, col = "blue")

op <- par(mfrow = c(2,2), mgp = c(1.6, 0.6,0), mar = c(3,3,1,1))
for(sh in 1:4)
  p.dgamma(sh)
par(op)
```

p.hboxp

Add a Horizontal Boxplot to the Current Plot

Description

Add a horizontal boxplot to the current plot. This is mainly an auxiliary function for `histBxp`, since `boxplot`(*, horizontal = TRUE, add = TRUE) is usually much preferable to this.

Usage

```
p.hboxp(x, y.lo, y.hi, boxcol = 3,
        medcol = 2, medlwd = 5, whisklty = 2, staplelty = 1)
```

Arguments

x	univariate data set.
y.lo, y.hi	minimal and maximal <i>user</i> coordinates or y.lo = c(ylo,hyi).
boxcol, medcol	color of the box and the median line.
medlwd	line width of median line.
whisklty, staplelty	line types of the whisker and the staple, the latter being used for the outmost non-outliers.

Details

....

Author(s)

Martin Maechler building on code from Markus and Christian Keller.

See Also

`boxplot`(**, horizontal = TRUE, add= TRUE).

Examples

```
## ==> See code in 'histBxp' (.) and example(histBxp) !
##
```

p.profileTraces	<i>Plot a profile.nls Object With Profile Traces</i>
-----------------	--

Description

Displays a series of plots of the profile t function and the likelihood profile traces for the parameters in a nonlinear regression model that has been fitted with [nls](#) and profiled with [profile.nls](#).

Usage

```
p.profileTraces(x, cex = 1,
               subtitle = paste("t-Profiles and traces of ",
                               deparse(attr(x, "summary")$formula)))
```

Arguments

x	an object of class "profile.nls", typically resulting from <code>profile(nls(.))</code> , see profile.nls .
cex	character expansion, see <code>par(cex =)</code> .
subtitle	a subtitle to set for the plot. The default now includes the <code>nls()</code> formula used.

Note

the **stats**-internal `stats:::plot.profile.nls` plot method just does “the diagonals”.

Author(s)

Andreas Ruckstuhl, R port by Isabelle Flückiger and Marcel Wolbers

See Also

[profile](#), and [nls](#) (which has unexported `profile` and `stats:::plot.profile.nls` methods).

Examples

```
require(stats)
data(Puromycin)
Treat <- Puromycin[Puromycin$state == "treated", ]
fm <- nls(rate ~ T1*conc/(T2+conc), data=Treat,
         start = list(T1=207, T2=0.06))
(pr <- profile(fm)) # quite a few things..
op <- par(mfcol=1:2)
plot(pr) # -> 2 'standard' plots
par(op)
## ours:
p.profileTraces(pr)
```

p.res.2fact

*Plot Numeric (e.g. Residuals) vs 2 Factors Using Boxplots***Description**

Plots a numeric “residual like” variable against two factor covariates, using boxplots.

Usage

```
p.res.2fact(x, y, z, restricted, notch = FALSE,
            xlab = NULL, ylab = NULL, main = NULL)
```

Arguments

x, y	two factors or numeric vectors giving the levels of factors.
z	numeric vector of same length as x and y, typically residuals.
restricted	positive value which truncates the size. The corresponding symbols are marked by stars.
notch	logical indicating if the boxplots should be notched, see boxplot(*, notch) .
xlab, ylab	axis labels, see plot.default , per default the actual argument expressions.
main	main title passed to plot, defaulting to the deparsed z argument.

Details

if values *are* restricted, this make use of the auxiliar function [u.boxplot.x](#).

Author(s)

Lorenz Gygax <logyg@wild.unizh.ch> and Martin Maechler, Jan.95; starting from [p.res.2x\(\)](#).

See Also

[p.res.2x](#), [boxplot](#), [plot.lm](#), [TA.plot](#).

Examples

```
I <- 8; J <- 3; K <- 20
xx <- factor(rep(rep(1:I, rep(K,I)),J))
yy <- factor(rep(1:J, rep(I*K,J)))
zz <- rt(I*J*K, df=5) #-- Student t with 5 d.f.
p.res.2fact(xx,yy,zz, restr= 4, main= "i.i.d. t <- 5 random |.| <= 4")
mtext("p.res.2fact(xx,yy,zz, restr= 4, ..)",
      line=1, adj=1, outer=TRUE, cex=1)

## Real data
data(warpbreaks)
(fm1 <- lm(breaks ~ wool*tension, data = warpbreaks))
```

```
## call via formula method of p.res.2x():
p.res.2x(~ ., fm1) # is shorter than, but equivalent to
## p.res.2x(~ wool + tension, fm1) ## or the direct
## with(warpbreaks, p.res.2fact(wool, tension, residuals(fm1)))
##
## whereas this is "transposed":
p.res.2x(~ tension+wool, fm1)
```

p.res.2x

Stahel's Residual Plot against 2 X's

Description

Plot Residuals, e.g., of a multiple linear regression, against two (predictor) variables, using positively and negatively oriented line segments for positive and negative residuals.

This is a (S3) *generic* function with a default and a [formula](#) method.

Usage

```
p.res.2x(x, ...)

## Default S3 method:
p.res.2x(x, y, z, restricted, size = 1, slwd = 1, scol = 2:3,
        xlab = NULL, ylab = NULL, main = NULL,
        xlim = range(x), ylim = range(y), ...)

## S3 method for class 'formula'
p.res.2x(x = ~., data, main = deparse(substitute(data)),
        xlab = NULL, ylab = NULL, ...)
```

Arguments

x, y	numeric vectors of the same length specifying 2 covariates. For the formula method, x is a formula .
z	numeric vector of same length as x and y, typically residuals.
restricted	positive value which truncates the size. The corresponding symbols are marked by stars.
size	the symbols are scaled so that size is the size of the largest symbol in cm.
slwd, scol	line width and color(s) for the residual segments . If scol has length 2 as per default, the two colors are used for positive and negative z values, respectively.
xlab, ylab, main	axis labels, and title see title , each with a sensible default. To suppress, use, e.g., main = "".
xlim, ylim	the basic x- and y- axis extents, see plot.default . Note that these will be slightly extended such that segments are not cut off.
...	further arguments passed to plot, or p.res.2x.default(), respectively.
data	(for the formula method:) a data frame or a fitted " lm " object.

Details

Each residual $zz[i]$ is visualized as line segment centered at (xx_i, yy_i) , $i = 1, \dots, n$, where the *lengths* of the segments are proportional to the absolute values $\|zz_i\|$.

Positive residuals' line segments have slope $+1$, and negative ones slope -1 , and `scol` is used to use different colors for negative and positive segments.

The formula interface calls `p.res.2fact()` when *both* `x` and `y` are *factors*.

Author(s)

Andreas Ruckstuhl in June 1991 and Martin Maechler, in 1992, '94, 2003-4.

References

Stahel, W.-A. (2008) *Statistische Datenanalyse: Eine Einführung für Naturwissenschaftler*, 5. Auflage, Vieweg, Wiesbaden; Paragraph 13.8.r and 13.8.v.

See Also

`p.res.2fact`, `plot.lm`, `TA.plot`.

Examples

```
xx <- rep(1:10,7)
yy <- rep(1:7, rep(10,7))
zz <- rnorm(70)
p.res.2x(xx,yy,zz, restricted = 2, main = "i.i.d. N(0,1) random residuals")

example(lm.influence, echo = FALSE)

op <- mult.fig(2, marP=c(-1,-1,-1,0), main="p.res.2x(*,*, residuals(lm.SR))")$old.par
with(LifeCycleSavings,
  { p.res.2x(pop15, ddpi, residuals(lm.SR), scol=c("red", "blue"))
    p.res.2x(pop75, dpi, residuals(lm.SR), scol=2:1)
  })

## with formula interface:
p.res.2x(~ pop15 + ddpi, lm.SR, scol=c("red", "blue"))
p.res.2x(~ pop75 + dpi, lm.SR, scol=2:1)

par(op) # revert par() settings above
```

p.scales

Conversion between plotting scales: usr, cm, symbol

Description

Give scale conversion factors of three coordinate systems in use for traditional R graphics: use, cm, symbol.

Usage

```
p.scales(unit = relsysize * 2.54 * min(pin), relsysize = 0.05)
```

Arguments

unit	length of unit (or x and y units) of symbol coordinates in cm.
relsysize	same, as a proportion of the plotting area.

Value

A numeric 2x2 matrix, with rows named x and y, and columns, named "sy2usr" and "usr2cm" which give the scale conversion factors from 'symbol' (as given) to 'usr' coordinates and from these to 'cm', respectively.

Author(s)

Werner Stahel, 1990; simplification: M.Maechler, 1993, 2004

See Also

`par("usr")`, of also `("pin")` on which this is based.

Examples

```
p.scales()
```

p.tachoPlot

Draw Symbol on a Plot

Description

Puts a symbol (pointer) on a plot at each of the specified locations.

Usage

```
p.tachoPlot(x, y, z, angle=c(pi/4,3*pi/4), size,
  method = c("robust", "sensitive", "rank"),
  legend = TRUE, show.method = legend,
  xlab = deparse(substitute(x)), ylab = deparse(substitute(y)),
  xlim, ylim, ...)
```

Arguments

x, y, z	coordinates of points. Numeric vectors of the same length. Missing values (NAs) are allowed.
angle	numeric vector whose elements give the angles between the horizontal baseline and the minimum and maximum direction of the pointer measured clockwise in radians.
size	length of the pointers in cm.
method	string specifying the method to calculate the angle of the pointer. One of "sensitive", "robust" or "rank". Only the first two characters are necessary. The minimum and maximum direction of the pointer corresponds to min(z) and max(z) if method is "sensitive" or "rank" and to the upper and lower extreme of z if method is "robust" (see boxplot or rrange for details). The angle is proportional to z or rank(z) in case of method="rank".
legend	logical flag: if TRUE (default), a legend giving the values of the minimum and maximum direction of the pointer is drawn.
show.method	logical flag, defaulting to legend; if true, the method name is printed.
xlab, ylab	labels for x and y axis; defaults to the 'expression' used in the function call.
xlim, ylim	numeric of length 2, the limits for the x and y axis, respectively; see plot.default .
...	further arguments to plot . Graphical parameters (see par) may also be supplied as arguments to this function.

Details

A scatter plot of the variables x and y is plotted. The value of the third variable z is given by the direction of a pointer (similar to a tachometer). Observations whose z-coordinate is missing are marked by a dot.

Side Effects

A plot is created on the current graphics device.

Author(s)

Christian Keller, June 1995

See Also

[symbols](#)

Examples

```
data(state)
data(USArrests)
p.tachoPlot(state.center $x, state.center $y, USArrests[, "UrbanPop"])

data(mtcars)
par(mfrow=c(2,2))
```

```
## see the difference between the three methods (not much differ. here!)
p.tachoPlot(mtcars$hp, mtcars$disp, mtcars$mpg, method="sens")
p.tachoPlot(mtcars$hp, mtcars$disp, mtcars$mpg, method="rank")
p.tachoPlot(mtcars$hp, mtcars$disp, mtcars$mpg, method="rob")
```

p.ts

plot.ts with multi-plots and Auto-Title – on 1 page

Description

For longer time-series, it is sometimes important to spread the time-series plots over several sub-plots. `p.ts(.)` does this both automatically, and under manual control.

Actually, this is a generalization of `plot.ts` (with different defaults).

Usage

```
p.ts(x, nrplots = max(1, min(8, n %/% 400)), overlap = nk %/% 16,
     date.x = NULL, do.x.axis = !is.null(date.x), do.x.rug = FALSE,
     ax.format, main.tit = NULL, ylim = NULL, ylab = "", xlab = "Time",
     quiet = FALSE, mgp = c(1.25, .5, 0), ...)
```

Arguments

<code>x</code>	timeseries (possibly multivariate) or numeric vector.
<code>nrplots</code>	number of sub-plots. Default: in $\{1..8\}$, approximately $n/400$ if possible.
<code>overlap</code>	by how much should subsequent plots overlap. Defaults to about 1/16 of sub-length on each side.
<code>date.x</code>	a time “vector” of the same length as <code>x</code> and coercable to class “POSIXct” (see DateTimeClasses).
<code>do.x.axis</code>	logical specifying if an x axis should be drawn (i.e., tick marks and labels).
<code>do.x.rug</code>	logical specifying if rug of <code>date.x</code> values should drawn along the x axis.
<code>ax.format</code>	when <code>do.x.axis</code> is true, specify the format to be used in the call to <code>axis.POSIXct</code> .
<code>main.tit</code>	Main title (over all plots). Defaults to name of <code>x</code> .
<code>ylim</code>	numeric(2) or NULL; if the former, specifying the y-range for the plots. Defaults to a common pretty range.
<code>ylab, xlab</code>	labels for y- and x-axis respectively, see description in <code>plot.default</code> .
<code>quiet</code>	logical; if TRUE, there’s no reporting on each subplot.
<code>mgp</code>	numeric(3) to be passed to <code>mult.fig()</code> , see <code>par(mgp = .)</code> .
<code>...</code>	further graphic parameters for each <code>plot.ts(. .)</code> .

Side Effects

A page of `nrplots` subplots is drawn on the current graphics device.

Author(s)

Martin Maechler, <maechler@stat.math.ethz.ch>; July 1994 (for S).

See Also

p.ts() calls [mult.fig\(\)](#) for setup. Further, [plot.ts](#) and [plot](#).

Examples

```
stopifnot(require(stats))
## stopifnot(require(datasets))

data(sunspots)
p.ts(sunspots, nr=1) # == usual  plot.ts(..)
p.ts(sunspots)
p.ts(sunspots, nr=3, col=2)

data(EuStockMarkets)
p.ts(EuStockMarkets[, "SMI"])
## multivariate :
p.ts(log10(EuStockMarkets), col = 2:5)

## with Date - x-axis (dense random dates):
set.seed(12)
x <- as.Date("2000-02-29") + cumsum(1+ rpois(1000, lambda= 2.5))
z <- cumsum(.1 + 2*rt(1000, df=3))
p.ts(z, 4, date.x = x)
p.ts(z, 6, date.x = x, ax.format = "%b %Y", do.x.rug = TRUE)
```

paste.vec

Utility for 'Showing' S vectors

Description

A simple utility for displaying simple S vectors; can be used as debugging utility.

Usage

```
paste.vec(name, digits = options()$digits)
```

Arguments

name	string with an variable name which must exist in the current environment (R session).
digits	how many decimal digits to be used; passed to format .

Value

a string of the form "NAME = x1 x2 ..."

Author(s)

Martin Maechler, about 1992.

Examples

```
x <- 1:4
paste.vec(x) ##-> "x = 1 2 3 4"
```

pkgDesc

Version of packageDescription() as Simple Vector

Description

a simple “version”, or wrapper for `packageDescription()`, returning a named character vector, including “file”, and still has a useful `print()` method.

Usage

```
pkgDesc(pkg, lib.loc = NULL, fields = NULL, ...)
pkgBuilt(pkg, lib.loc = NULL, ...)
```

Arguments

<code>pkg</code>	a character string, name of an installed R package.
<code>lib.loc</code>	library location to find the package in; the default NULL uses the full <code>.libPaths()</code> .
<code>fields</code>	a character vector (or NULL) specifying fields to be returned.
<code>...</code>	further optional arguments passed to <code>packageDescription()</code> .

Value

a named [character](#) vector, with [names](#), the *fields*, identical to the names of the [list](#) returned by `packageDescription`, plus its “file” attribute. Additionally the resulting vector is of class “Dlist” which activates a useful `print()` method.

Note

The file is always returned; not the least that the author wants to see it quite often as his `.libPaths()` is non-trivial and typically longer than 4 entries.

Author(s)

Martin Maechler, Jan. 2021

See Also

[packageDescription](#), [.libPaths](#).

Examples

```
str(pd <- pkgDesc("sfsmisc"))
pd[c("Date", "Packaged", "Built", "file")]

pkgBuilt("sfsmisc")

## Show "Built" (and "file") for all packages whose namespaces are loaded:
lNs <- loadedNamespaces()
mLNs <- sapply(lNs, pkgBuilt)
t(mLNs) # typically prints nicely

pkgs <- c("grid", "lattice", "MASS", "Matrix", "nlme", "lme4", "sfsmisc")
pkgs <- c("foobar", "barbar", pkgs, "kitty") # + names that typically don't exist
pkgsOk <- basename(find.package(pkgs, quiet=TRUE))
mpkg <- sapply(pkgsOk, pkgBuilt)
stopifnot(is.matrix(mpkg), nrow(mpkg) == 2)
mpkg["Built",]
```

pkgLibs

R Package Compiled Code Library Dependencies (on Unix-alikes)

Description

List some system level information about the compiled code library, typically its dependencies, for R packages with compiled code; for Unix-alikes or more generally when cmd is installed locally.

Usage

```
pkgLibs(pkg,
  cmd = if(Sys.info()[["sysname"]] == "Darwin") "otool -L" else "ldd")
```

Arguments

pkg	character vector of package names of <i>installed</i> R packages.
cmd	a character string with the name of an OS / system level program (to be called via system (cmd, ...)) which gives information about the shared library (of compiled code), also known as “DLL” (dynamically loadable library) or “so” ((dynamic) shared object) library. The default, “ldd” is a standard binary utility on Unix-alike platforms such as Linux. On macOS, “oTool -L” is used by default.

Details

Note that there seems some language confusion as “DLL” on Windows is *also* used for “Dynamic-link Library” and Wikipedia warns about confusing the two concepts (“dynamically loaded ..” vs “dynamic-link ..”).

Value

a named `list` with one entry per package in `pkg`, the `names` being the directory / folder names of the corresponding `pkgs` from `pkg`.

The exact structure of such entries is currently subject to change and you should not rely on its exact format for now.

Author(s)

Martin Maechler

References

‘Dynamic Loading’ on Wikipedia, https://en.wikipedia.org/wiki/Dynamic_loading

On Windows, “DLL” is also used for Dynamic-link library, https://en.wikipedia.org/wiki/Dynamic-link_library.

`man ldd` from a terminal on a valid OS.

See Also

`dyn.load()`, `library.dynam()`, and `getLoadedDLLs()`.

Also, `.C`, `.Call` which use such DLLs.

Examples

```
# for the example only using standard R packages :
myPkgs <- c("stats", "MASS", "rpart", "Matrix")
pl <- pkgLibs(myPkgs)
pl
stopifnot(exprs = {
  is.list(pl)
  length(pl) == length(myPkgs)
  is.character(pkgD <- names(pl))
})
## Have seen this failing when a strange development version of "Matrix" was picked up:
try( stopifnot( dir.exists(pkgD)) )
```

plotDS

Plot Data and Smoother / Fitted Values

Description

For one-dimensional nonparametric regression, plot the data and fitted values, typically a smooth function, and optionally use segments to visualize the residuals.

Usage

```
plotDS(x, yd, ys, xlab = "", ylab = "", ylim = rrange(c(yd, ys)),
       xpd = TRUE, do.seg = TRUE, seg.p = 0.95,
       segP = list(lty = 2, lwd = 1, col = 2),
       linP = list(lty = 1, lwd = 2.5, col = 3),
       ...)
```

Arguments

x, yd, ys	numeric vectors all of the same length, representing (x_i, y_i) and fitted (smooth) values $\hat{y}_i$. x will be sorted increasingly if necessary, and yd and ys accordingly. Alternatively, ys can be an x-y list (as resulting from xy.coords) containing fitted values on a finer grid than the observations x. In that case, the observational values x[] must be part of the larger set; seqXtend () may be applied to construct such a set of abscissa values.
xlab, ylab	x- and y- axis labels, as in plot.default .
ylim	limits of y-axis to be used; defaults to a <i>robust</i> range of the values.
xpd	see par(xpd=.) ; by default do allow to draw outside the plot region.
do.seg	logical indicating if residual segments should be drawn, at x[i], from yd[i] to ys[i] (approximately, see seg.p).
seg.p	segment percentage of segments to be drawn, from yd to seg.p*ys + (1-seg.p)*yd.
segP	list with named components lty, lwd, col specifying line type, width and color for the residual segments, used only when do.seg is true.
linP	list with named components lty, lwd, col specifying line type, width and color for “smooth curve lines”.
...	further arguments passed to plot .

Note

Non-existing components in the lists segP or linP will result in the [par](#) defaults to be used.
 plotDS() used to be called pl.ds up to November 2007.

Author(s)

Martin Maechler, since 1990

See Also

[seqXtend](#)() to construct more smooth ys “objects”.

Examples

```
data(cars)
x <- cars$speed
yd <- cars$dist
ys <- lowess(x, yd, f = .3)$y
```

```

plotDS(x, yd, ys)

## More interesting : Version of example(Theoph)
data(Theoph)
Th4 <- subset(Theoph, Subject == 4)
## just for "checking" purposes -- permute the observations:
Th4 <- Th4[sample(nrow(Th4)), ]
fm1 <- nls(conc ~ SSfol(Dose, Time, lKe, lKa, lCl), data = Th4)

## Simple
plotDS(Th4$Time, Th4$conc, fitted(fm1),
      sub = "Theophylline data - Subject 4 only",
      segP = list(lty=1,col=2), las = 1)

## Nicer: Draw the smoother not only at x = x[i] (observations):
xsm <- unique(sort(c(Th4$Time, seq(0, 25, length = 201))))
ysm <- c(predict(fm1, newdata = list(Time = xsm)))
plotDS(Th4$Time, Th4$conc, ys = list(x=xsm, y=ysm),
      sub = "Theophylline data - Subject 4 only",
      segP = list(lwd=2), las = 1)

```

plotStep

Plot a Step Function

Description

Plots a step function $f(x) = \sum_i y_i 1_{[t_{i-1}, t_i]}(x)$, i.e., a piecewise constant function of one variable. With one argument, plots **the** empirical cumulative distribution function.

Usage

```

plotStep(ti, y,
        cad.lag = TRUE,
        verticals = !cad.lag,
        left.points= cad.lag, right.points= FALSE, end.points= FALSE,
        add = FALSE,
        pch = par('pch'),
        xlab=deparse(substitute(ti)), ylab=deparse(substitute(y)),
        main=NULL, ...)

```

Arguments

ti	numeric vector = X[1:N] or t[0:n].
y	numeric vector y[1:n]; if omitted take y = k/N for empirical CDF.
cad.lag	logical: Draw 'cad.lag', i.e., " <i>continue à droite, limite à gauche</i> ". Default = TRUE.
verticals	logical: Draw vertical lines? Default= ! cad.lag
left.points	logical: Draw left points? Default= cad.lag

<code>right.points</code>	logical: Draw right points? Default= FALSE
<code>end.points</code>	logical: Draw 2 end points? Default= FALSE
<code>add</code>	logical: Add to existing plot? Default= FALSE
<code>pch</code>	plotting character for points, see <code>par()</code> .
<code>xlab, ylab</code>	labels of x- and y-axis
<code>main</code>	main title; defaults to the call' if you do not want a title, use <code>main = ""</code> .
<code>...</code>	Any valid argument to <code>plot(..)</code> .

Value

invisibly: List with components `t` and `y`.

Side Effects

Calls `plot(..)`, `points(..)`, `segments(..)` appropriately and plots on current graphics device.

Author(s)

Martin Maechler, Seminar for Statistics, ETH Zurich, <maechler@stat.math.ethz.ch>, 1991 ff.

See Also

The `plot` methods `plot.ecdf` and `plot.stepfun` in R which are conceptually nicer.

`segments(..., method = "constant")`.

Examples

```
##-- Draw an Empirical CDF (and see the default title ..)
plotStep(rnorm(15))

plotStep(runif(25), cad.lag=FALSE)
plotStep(runif(25), cad.lag=FALSE, add=TRUE, lty = 2)

ui <- sort(runif(20))
plotStep(ui, ni <- cumsum(rpois(19, lambda=1.5) - 1.5), cad.lag = FALSE)
plotStep(ui, ni, verticals = TRUE, right.points = TRUE)

plotStep(rnorm(201), pch = '.') #- smaller points
```

polyn.eval

*Evaluate Polynomials***Description**

Evaluate one or several univariate polynomials at several locations, i.e. compute $\text{coef}[1] + \text{coef}[2]*x + \dots + \text{coef}[p+1]*x^p$ (in the simplest case where x is scalar and coef a vector).

Usage

```
polyn.eval(coef, x)
```

Arguments

coef	“numeric” vector or matrix. If a vector, x can be an array and the result matches x. If coef is a matrix it specifies several polynomials of the same degree as rows, x must be a vector, $\text{coef}[,k]$ is for x^{k-1} and the result is a matrix of dimension $\text{length}(x) * \text{nrow}(\text{coef})$. Note that coef can also be complex or bigrational (<code>as.bigq(.)</code> from gmp, or arbitrary precision (“mpfr”) from Rmpfr, or similar number-like objects for which basic arithmetic is defined.
x	“numeric” vector or array. Either x or coef must be a vector.

Details

The stable “Horner rule” is used for evaluation in any case.

When $\text{length}(\text{coef}) == 1$, `polyn.eval(coef, x)` now returns a vector of $\text{length}(x)$ whereas previously, it just gave the number coef independent of x .

Value

numeric vector or array, depending on input dimensionalities, see above.

Author(s)

Martin Maechler, ages ago.

See Also

For much more sophisticated handling of polynomials, use the **polynom** package, see, e.g., [predict.polynomial](#). For multivariate polynomials (and also for nice interface to the **orthopolynom** package), consider the **mpoly** package.

Examples

```

polyn.eval(c(1,-2,1), x = 0:3)# (x - 1)^2
polyn.eval(c(0, 24, -50, 35, -10, 1), x = matrix(0:5, 2,3))# 5 zeros!
(cf <- rbind(diag(3), c(1,-2,1)))
polyn.eval(cf, 0:5)

x <- seq(-3,7, by=1/4)
cf <- 4:1
(px <- polyn.eval(cf, x)) # is exact
if((gmpT <- "package:gmp" %in% search()) || require("gmp")) withAutoprint({
  pxq <- polyn.eval(coef = as.bigq(cf, 1), x=x)
  pxq
  stopifnot(pxq == px)
  if(!gmpT) detach("package:gmp")
})

if((RmpfrT <- "package:Rmpfr" %in% search()) || require("Rmpfr")) withAutoprint({
  pxM <- polyn.eval(coef = mpfr(cf, 80), x=x) # 80 bits accuracy
  pxM
  stopifnot(pxM == px)
  if(!RmpfrT) detach("package:Rmpfr")
})

stopifnot(identical(polyn.eval(12, x), rep(12, length(x))),
           identical(polyn.eval(7, diag(3)), matrix(7, 3,3)))

```

posdefify

Find a Close Positive Definite Matrix

Description

From a matrix *m*, construct a "close" positive definite one.

Usage

```

posdefify(m, method = c("someEVadd", "allEVadd"),
          symmetric = TRUE, eigen.m = eigen(m, symmetric= symmetric),
          eps.ev = 1e-07)

```

Arguments

<i>m</i>	a numeric (square) matrix.
<i>method</i>	a string specifying the method to apply; can be abbreviated.
<i>symmetric</i>	logical, simply passed to eigen (unless <i>eigen.m</i> is specified); currently, we do not see any reason for <i>not</i> using TRUE.
<i>eigen.m</i>	the eigen value decomposition of <i>m</i> , can be specified in case it is already available.
<i>eps.ev</i>	number specifying the tolerance to use, see Details below.

Details

We form the eigen decomposition

$$m = V\Lambda V'$$

where Λ is the diagonal matrix of eigenvalues, $\Lambda_{j,j} = \lambda_j$, with *decreasing* eigenvalues $\lambda_1 \geq \lambda_2 \geq \dots \geq \lambda_n$.

When the smallest eigenvalue λ_n are less than $\text{Eps} < -\text{eps} \cdot \text{ev} * \text{abs}(\text{lambda}[1])$, i.e., negative or “almost zero”, some or all eigenvalues are replaced by *positive* ($\geq \text{Eps}$) values, $\tilde{\Lambda}_{j,j} = \tilde{\lambda}_j$. Then, $\tilde{m} = V\tilde{\Lambda}V'$ is computed and rescaled in order to keep the original diagonal (where that is $\geq \text{Eps}$).

Value

a matrix of the same dimensions and the “same” diagonal (i.e. [diag](#)) as `m` but with the property to be positive definite.

Note

As we found out, there are more sophisticated algorithms to solve this and related problems. See the references and the [nearPD\(\)](#) function in the **Matrix** package. We consider `nearPD()` to also be the successor of this package’s [nearcor\(\)](#).

Author(s)

Martin Maechler, July 2004

References

Section 4.4.2 of Gill, P.-E., Murray, W. and Wright, M.-H. (1981) *Practical Optimization*, Academic Press.

Cheng, Sheung Hun and Higham, Nick (1998) A Modified Cholesky Algorithm Based on a Symmetric Indefinite Factorization; *SIAM J. Matrix Anal. Appl.*, **19**, 1097–1110.

Knol DL, ten Berge JMF (1989) Least-squares approximation of an improper correlation matrix by a proper one. *Psychometrika* **54**, 53–61.

Higham (2002) Computing the nearest correlation matrix - a problem from finance; *IMA Journal of Numerical Analysis* **22**, 329–343.

Lucas (2001) Computing nearest covariance and correlation matrices. A thesis submitted to the University of Manchester for the degree of Master of Science in the Faculty of Science and Engineering.

See Also

[eigen](#) on which the current methods rely. [nearPD\(\)](#) in the **Matrix** package. (Further, the deprecated [nearcor\(\)](#) from this package.)

Examples

```
set.seed(12)
m <- matrix(round(rnorm(25),2), 5, 5); m <- 1+ m + t(m); diag(m) <- diag(m) + 4
m
posdefify(m)
1000 * zapsmall(m - posdefify(m))
```

potatoes	<i>Fisher's Potato Crop Data</i>
----------	----------------------------------

Description

Fisher's potato crop data set is of historical interest as an early example of a multi-factor block design.

Usage

```
data(potatoes)
```

Format

A data frame with 64 observations on the following 5 variables.

pos a factor with levels 1:4.

treat a factor with 16 levels A to H and J to Q, i.e., LETTERS[1:17][-9].

nitrogen a factor specifying the amount of nitrogen sulfate (NH_4), with the four levels 0, 1, 2, 4.

potash a factor specifying the amount of potassium (K, 'kalium') sulfate, with the four levels 0, 1, 2, 4.

yield a numeric vector giving the yield of potatoes in ...

Source

Bennett, J. H. (1972) *Collected Papers of R. A. Fischer* vol.~II, 1925-31; The University of Adelaide.

References

T.Eden and R. A. Fisher (1929) Studies in Crop Variation. VI. Experiments on the Response of the Potato to Potash and Nitrogen. *J. Agricultural Science* **19**, 201–213. Accessible from Bennett (1972), see above.

Examples

```
data(potatoes)
## See the experimental design:
with(potatoes, {
  cat("4 blocks of experiments;",
      "each does every (nitrogen,potash) combination (aka 'treat'ment) once.",
      '', sep="\n")
  print(ftable(table(nitrogen, potash, treat)))
  print(ftable(tt <- table(pos,potash,nitrogen)))
  tt[cbind(pos,potash,nitrogen)] <- as.character(treat)
  cat("The 4 blocks pos = 1, 2, 3, 4:\n")
  ftable(tt)
})
## First plot:
with(potatoes, interaction.plot(potash,nitrogen, response=yield))

## ANOVAs:
summary(aov(yield ~ nitrogen * potash + Error(pos), data = potatoes))
# "==" can use simply
summary(aov(yield ~ nitrogen + potash + pos, data = potatoes))
# and
summary(aov(yield ~ nitrogen + potash, data = potatoes))
```

pretty10exp

Nice 10 $\times$ k Label Expressions

Description

Produce nice $a \times 10^k$ expressions to be used instead of the scientific notation " $a \times 10^k$ ".

Usage

```
pretty10exp(x, drop.1 = FALSE, sub10 = FALSE, digits = 7, digits.fuzz,
  off = pmax(10^-digits, 2^-(110x*log2(10)+1075)),
  lab.type = c("plotmath","latex"),
  lab.sep = c("cdot", "times"))
```

Arguments

x	numeric vector (e.g. axis tick locations)
drop.1	logical indicating if $1 \times$ should be dropped from the resulting expressions.
sub10	logical, "10", a non-negative integer number or an integer vector of length two, say (k_1, k_2) , indicating if some 10^j expressions for $j \in J$ should be formatted traditionally, notably e.g., $10^0 \equiv 1$. When a (non-negative) number, say k , $J = \{j; j \leq k\}$ are all simplified, when a length-2 vector, $J = \{j; k_1 \leq j \leq k_2\}$ are. Special cases: sub10 = TRUE means to use 1 instead of 10^0 and sub10 = "10" uses both 1 for 10^0 and 10 for 10^1 ; these are short forms of sub10 = c(0,0) and sub10 = c(0,1) respectively.

<code>digits</code>	number of digits for mantissa (<i>a</i>) construction; the number of <i>significant</i> digits, see <code>signif</code> .
<code>digits.fuzz</code>	the old deprecated name for <code>digits</code> .
<code>off</code>	a numeric offset in <code>eT <- floor(110x + off)</code> where <code>110x <- log10(abs(x))</code> and <code>eT</code> are the exponents k for label factors 10^k . Previously hardcoded to 10^{digits} , the new default provides better results for subnormal <code>abs(x)</code> values.
<code>lab.type</code>	a string indicating how the result should look like. By default, (<code>plotmath</code> -compatible) <code>expressions</code> are returned. Alternatively, <code>lab.type = "plotmath"</code> returns LaTeX formatted strings for labels. (The latter is useful, e.g., when using the <code>tikzDevice</code> package to generate LaTeX-processed figures.)
<code>lab.sep</code>	character separator between mantissa and exponent for LaTeX labels; it will be prepended with a backslash, i.e., <code>"\cdot"</code> will use <code>"\cdot"</code>

Value

For the default `lab.type = "plotmath"`, an expression of the same length as `x`, typically with elements of the form `a%% 10 ^ k`. Exceptions are `0` which is kept simple, if `drop.1` is true and $a = 1$, 10^k is used, and if `sub10` is not false, `a%% 10 ^ 0` as `a`, and `a%% 10 ^ k` as as the corresponding formatted number `a * 10^k` independently of `drop.1`.

Otherwise, a `character` vector of the same length as `x`. For `lab.type = "latex"`, currently the only alternative to the default, these strings are LaTeX (math mode) compatible strings.

Note

If `sub10` is set, it will typically be a small number such as 0, 1, or 2. Setting `sub10 = TRUE` will be interpreted as `sub10 = 1` where resulting exponents k will either be negative or $k \geq 2$.

Author(s)

Martin Maechler; Ben Bolker contributed `lab.type = "latex"` and `lab.sep`.

See Also

`axExpr` and `eaxis()` which build on `pretty10exp()`, notably the `eaxis()` example plots.

The new `toLatex.numeric` method which gives very similar results with option `scientific = TRUE`.

Further, `axis`, `axTicks`.

Examples

```
pretty10exp(-1:3 * 1000)
pretty10exp(-1:3 * 1000, drop.1 = TRUE)
pretty10exp(c(1,2,5,10,20,50,100,200) * 1e3)
pretty10exp(c(1,2,5,10,20,50,100,200) * 1e3, drop.1 = TRUE)

set.seed(17); lx <- rlnorm(10, m=8, s=6)
pretty10exp(lx, digits = 3)
```

```

pretty10exp(lx, digits = 3, sub10 = 2)

pretty10exp(lx, digits = 3, lab.type="latex")
pretty10exp(lx, digits = 3, lab.type="latex", lab.sep="times", sub10=2)

## use regular formatted numbers from 0.03 to 300 :
pretty10exp(3*10^(-3:4), sub10 = c(-2,2))
pretty10exp(3*10^(-3:4), sub10 = c(-2,2), lab.type = "l")

ax <- 10^(-6:0) - 2e-16
pretty10exp(ax, drop.1=TRUE) # nice for plotting
pretty10exp(ax, drop.1=TRUE, sub10=TRUE)
pretty10exp(ax, drop.1=TRUE, sub10=c(-2,2))

## in sfsmisc version <= 1.0-16, no 'digits',
## i.e., implicitly had digits := #{double precision digits} ==
(dig. <- .Machine$double.digits * log10(2)) # 15.95
pretty10exp(ax, drop.1=TRUE, digits= dig.) # 'ugly'

## Subnormal numbers
x <- sort(c(outer(10^-(323:305), 1:9))); x <- c(x[1]/2, x)
tail(x, 12) # nice
head(x, 6) # "ugly" (they are multiple's of 2^-1074):
head(x, 6) / 2^-1074 # nice

head(p0 <- pretty10exp(x, off = 10^-7), 30) # previous behavior {before 'off' existed}
str(head(pTen <- lapply(p0, `[`, 3L)))
str(exTen <- sapply(pTen, `[`, 3L)) # -324 -324 ..
head(f0 <- sapply(p0, `[`, 2L), 17)

head(p1 <- pretty10exp(x))# new default
str(head(pTen1 <- lapply(p1, `[`, 3L)))
str(exTen1 <- sapply(pTen1, `[`, 3L)) # -324 -324 ..
head(f1 <- sapply(p1, `[`, 2L), 17) #

head(cbind(x, f0, f1, exTen, exTen1), 80)
(nEQ <- which(sapply(1:length(p0), function(i) p0[[i]] != p1[[i]])))
cbind(x, f0, f1, exTen, exTen1)[nEQ,]
stopifnot(is.finite(f1), 0.5 <= f1, f1 <= 9)

```

primes

Find all Primes Less Than n

Description

Find all prime numbers aka ‘primes’ less than n .

Uses an obvious sieve method (and some care), working with [logical](#) and [integers](#) to be quite fast.

Usage

```
primes(n, pSeq = NULL)
```

Arguments

n a (typically positive integer) number.

pSeq optionally a vector of primes (2,3,5,...) as if from a `primes()` call; **must** be correct. The goal is a speedup, but currently we have not found one single case, where using a non-NULL `pSeq` is faster.

Details

As the function only uses `max(n)`, `n` can also be a *vector* of numbers.

The famous prime number theorem states that $\pi(n)$, the *number* of primes below n is asymptotically $n / \log(n)$ in the sense that $\lim_{n \rightarrow \infty} \pi(n) \cdot \log(n) / n \sim 1$.

Equivalently, the inverse of $pi()$, the n -th prime number p_n is around $n \log n$; recent results (Pierre Dusart, 1999), prove that

$$\log n + \log \log n - 1 < \frac{p_n}{n} < \log n + \log \log n \quad \text{for } n \geq 6.$$

Value

numeric vector of all prime numbers $\leq n$.

Author(s)

Bill Venables (<= 2001); Martin Maechler gained another 40% speed, carefully working with logicals and integers.

See Also

`factorize`. For large n , use the `gmp` package and its `isprime` and `nextprime` functions.

Examples

```
(p1 <- primes(100))
system.time(p1k <- primes(1000)) # still lightning fast
stopifnot(length(p1k) == 168)

system.time(p.e7 <- primes(1e7)) # still only 0.3 sec (2015 (i7))
stopifnot(length(p.e7) == 664579)
## The famous pi(n) := number of primes <= n:
pi.n <- approxfun(p.e7, seq_along(p.e7), method = "constant")
pi.n(c(10, 100, 1000)) # 4 25 168
plot(pi.n, 2, 1e7, n = 1024, log="xy", axes = FALSE,
     xlab = "n", ylab = quote(pi(n)),
     main = quote("The prime number function " ~ pi(n)))
eaxis(1); eaxis(2)
```

```
## Exploring p(n) := the n-th prime number ~~~ n * pnn(n), where
## pnn(n) := log n + log log n
pnn <- function(n) { L <- log(n); L + log(L) }
n <- 6:(N <- length(PR <- primes(1e5)))
m.pn <- cbind(l.pn = ceiling(n*(pnn(n)-1)), pn = PR[n], u.pn = floor(n*pn(n)))
matplot(n, m.pn, type="l", ylab = quote(p[n]), main = quote(p[n] ~~~
      "with lower/upper bounds" ~ n*(log(n) + log(log(n)) -(1~"or"~0))))
## (difference to the lower approximation) / n --> ~ 0.0426 (?) :
plot(n, PR[n]/n - (pnn(n)-1), type = 'l', cex = 1/8, log="x", xaxt="n")
eaxis(1); abline(h=0, col=adjustcolor(1, 0.5))
```

printTable2

Add and Print Marginals for 2-way Contingency Tables

Description

printTable2() prints a 2-way contingency table “with all bells and whistles” (currently using German labeling).

margin2table() computes marginals, adds them to the table and returns a margin2table object the print method for which adds text decorations (using “-” and “|”).

Usage

```
printTable2(table2, digits = 3)
margin2table(x, totName = "sum", name.if.empty=FALSE)
## S3 method for class 'margin2table'
print(x, digits = 3, quote = FALSE, right = TRUE, ...)
```

Arguments

table2	a matrix with non-negative integer entries, i.e. the contingency table.
x	a matrix; for print(), the result of margin2table.
digits	Anzahl Dezimalstellen, auf die die Häufigkeiten gerundet werden sollen.
quote, right	logicals passed to <code>print.default()</code> , but with different default values.
totName	string to use as row- and column- name if x has corresponding <code>dimnames</code> .
name.if.empty	logical indicating if the margin “totals” should be named in any case.
...	further potential arguments, unused currently.

Value

margin2table returns a matrix with *added marginals*, i.e., an extra row and column, and is of class “margin2table” (and “table” still) which has a nice print method.

printTable2 is just producing output.

Author(s)

Martin Maechler, Feb.1993; then Dec 2003

See Also

[table](#), [ftable](#).

Examples

```
margin2table(diag(4),,TRUE)
m <- diag(3); colnames(m) <- letters[1:3]
margin2table(m)
margin2table(m / sum(m))

data(HairEyeColor)
margin2table(HairEyeColor[, , "Male"])
printTable2(HairEyeColor[, , "Male"])
printTable2(HairEyeColor[, , "Female"])
```

prt.DEBUG

Utility Printing in DEBUG mode

Description

This is **defunct** now: The global DEBUG has been a cheap precursor to R's [options\(verbose=\)](#) (or a verbose function argument).

This function prints out its arguments as [cat\(\)](#) does, additionally printing the name of function in which it's been called — only when a global variable DEBUG exists and is [TRUE](#).

Usage

```
prt.DEBUG(..., LEVEL = 1)
```

Arguments

...	arguments to be passed to cat() for printing.
LEVEL	integer (or logical) indicating a debugging level for printing.

Author(s)

Martin Maechler, originally for S-PLUS.

ps.end	<i>Close PostScript or Acrobat Graphics Device opened by 'ps.do' / 'pdf.do'</i>
--------	---

Description

Closes the PostScript or PDF file ([postscript](#),[pdf](#)), opened by a previous [ps.do](#) (or [pdf.latex](#), or ...) call, using [dev.off](#), and additionally opens a previewer for that file, *unless* the previewer is already up. This almost provides an 'interactive' device (like [x11](#)) for [postscript](#) or [pdf](#).

Usage

```
ps.end(call.gv= NULL, command = getOption("eps_view"),
       debug = getOption("verbose"))
pdf.end(call.viewer= NULL, command = getOption("pdfviewer"),
       debug = getOption("verbose"))
```

Arguments

call.gv, call.viewer	logical, indicating if the postscript or acrobat reader (e.g., ghostview or acroread or the command given by command) should be called. By default, find out if the viewer is already running on this file and only call it if needed.
command	character, giving a system command for PostScript previewing. By default, <code>getOption("eps_view")</code> is set to <code>gv -watch -geometry -0+0 -magstep -2 -media BBox -noantialias</code> which assumes gv (aka <i>ghostview</i>) to be in your OS path.
debug	logical; if TRUE print information during execution.

Details

Depends on Unix tools, such as ps.

Author(s)

Martin Maechler

See Also

[postscript](#), [postscript pdf.do](#), [ps.do](#), ...

Examples

```
if(interactive())
) {
  myPS <- tempfile("ex", fileext = ".ps")
  ps.do(myPS)
  data(sunspots)
```

```

plot(sunspots)
ps.end()

tempfile("ex-sun", fileext = ".pdf") -> myPDF
pdf.latex(myPDF)
plot(sunspots)
pdf.end(call. = FALSE) # basically the same as dev.off()
}
ps.latex(tempfile("ex2", fileext = ".eps"))
plot(sunspots)
ps.end(call.gv = FALSE) # basically the same as dev.off()

```

ps.latex

*PostScript/PDF Preview Device with Optional 'LaTeX' Touch***Description**

All functions start a pseudo PostScript or Acrobat preview device, using [postscript](#) or [pdf](#), and further registering the file name for subsequent calls to [pdf.end\(\)](#) or [ps.end\(\)](#).

Usage

```

pdf.do(file, paper = "default", width = -1, height = -1, onefile = FALSE,
       title = NULL, version = "1.4", quiet = FALSE, ...)

pdf.latex(file, height = 5 + main.space * 1.25, width = 9.5,
          main.space=FALSE, lab.space = main.space,
          paper = "special", title = NULL,
          lab=c(10, 10, 7), mgp.lab=c(1.6, 0.7, 0), mar=c(4, 4, 0.9, 1.1), ...)

ps.do(file, width=-1, height=-1, onefile=FALSE, horizontal=FALSE,
      title = NULL, ...)

ps.latex(file, height = 5 + main.space * 1.25, width = 9.5,
          main.space=FALSE, lab.space = main.space,
          paper = "special", title = NULL,
          lab=c(10, 10, 7), mgp.lab=c(1.6, 0.7, 0), mar=c(4, 4, 0.9, 1.1), ...)

```

Arguments

file	character giving the PostScript/PDF file name to be written.
height	device height in <i>inches</i> , height * 2.54 are <i>cm</i> . The default is 5 plus 1.25 iff main.space.
width	device width in <i>inches</i> ; for this and height, see postscript .
onefile, horizontal	logicals passed to postscript(...) or pdf(...) , most probably to be left alone.

title	PostScript/PDF (not plot!) title passed to <code>postscript()</code> or <code>pdf()</code> ; by default use a title with R version and file in it.
version	a string describing the PDF version that will be required to view the output, see <code>pdf</code> ; our (high) default ensures alpha-transparency.
quiet	logical specifying that some (informative/warning) messages should not be issued.
main.space	logical; if true, leave space for a main title (unusual for LaTeX figures!).
lab.space	logical; if true, leave space for x- and y- labels (by <i>not</i> subtracting from mar).
paper	character (or missing), typically "a4" or "a4r" in non-America, see <code>postscript</code> . Only if this is "special" (or missing) are your choices of width and height completely honored (and this may lead to files that cannot print on A4) with resizing.
lab	integer of length 3, <code>lab[1:2]</code> are desired number of tick marks on x- and y-axis, see <code>par(lab=)</code> .
mgp.lab	three decreasing numbers determining space for axis labeling, see <code>par(mgp=)</code> , the default is here smaller than usual.
mar	four numbers, indicating marginal space, see <code>par(mar=)</code> , the default is here smaller than usual.
...	arguments passed to <code>ps.do()</code> or <code>pdf.do()</code> from <code>ps.latex</code> / <code>pdf.latex</code> and to <code>ps.options</code> from <code>ps.do/pdf.do</code> .

Details

`ps.latex` and `pdf.latex` have an additional LaTeX flavor, and just differ by some extra `par` settings from the `*.do` siblings: E.g., after `ps.do(..)` is called, the graphical parameters `c("mar", "mgp", "lab")` are reset (to values that typically are better than the defaults for LaTeX figures).

Whereas the defaults for paper, width, and height *differ* between `pdf` and `postscript`, they are set such as to provide very similar functionality, for the functions `ps.do()` and `pdf.do()`; e.g., by default, both use a full plot on portrait-oriented page of the default paper, as per `getOption("papersize")`. `pdf.do()` sets the default paper to "special" when both width and height are specified.

Value

A list with components

<code>old.par</code>	containing the old par values
<code>new.par</code>	containing the newly set par values

Author(s)

Martin Maechler

See Also

`ps.end`, `pdf`, `postscript`, `dev.print`.

Examples

```

if(interactive()) {

  ps.latex("ps.latex-ex.ps", main= TRUE)
  data(sunspots)
  plot(sunspots,main=paste("Sunspots Data, n=",length(sunspots)),col="red")
  ps.end()

  pdf.latex("pdf.latex-ex.pdf", main= TRUE)
  data(sunspots)
  plot(sunspots,main=paste("Sunspots Data, n=",length(sunspots)),col="red")
  pdf.end()

  ps.do("ps_do_ex.ps")
  example(plot.function)
  ps.end()

  pdf.do("pdf_do_ex.pdf", width=12, height=5)
  plot(sunspots, main="Monthly Sunspot numbers (in Zurich, then Tokyo)")
  pdf.end()
}

```

quadrant

Give the Quadrant Number of Planar Points

Description

Determine the quadrant of planar points, i.e. in which of the four parts cut by the x- and y- axis the points lie. Zero values (i.e. points on the axes) are treated as if *positive*.

Usage

```
quadrant(x, y=NULL)
```

Arguments

x, y numeric vectors of the same length, or x is an $x - y$ structure and y=NULL, see [xy.coords](#).

Value

numeric vector of same length as x (if that's a vector) with values in 1:4 indicating the quadrant number of the corresponding point.

Examples

```
xy <- as.matrix(expand.grid(x= -7:7, y= -7:7)); rownames(xy) <- NULL
(qu <- quadrant(xy))
plot(xy, col = qu+1, main = "quadrant() number", axes = FALSE)
abline(h=0, v=0, col="gray") # the x- and y- axis
text(xy, lab = qu, col = qu+1, adj = c(1.4,0))
```

QUnif

Quasi Random Numbers via Halton Sequences

Description

These functions provide quasi random numbers or *space filling* or *low discrepancy* sequences in the p -dimensional unit cube.

Usage

```
sHalton(n.max, n.min = 1, base = 2, leap = 1)
QUnif (n, min = 0, max = 1, n.min = 1, p, leap = 1, silent = FALSE)
```

Arguments

n.max	maximal (sequence) number.
n.min	minimal sequence number.
n	number of p -dimensional points generated in QUnif. By default, n.min = 1, leap = 1 and the maximal sequence number is n.max = n.min + (n-1)*leap.
base	integer ≥ 2 : The base with respect to which the Halton sequence is built.
min, max	lower and upper limits of the univariate intervals. Must be of length 1 or p.
p	dimensionality of space (the unit cube) in which points are generated.
leap	integer indicating (if > 1) if the series should be leaped, i.e., only every leapth entry should be taken.
silent	logical asking to suppress the message about enlarging the prime table for large p.

Value

sHalton(n,m) returns a numeric vector of length n-m+1 of values in $[0, 1]$.
 QUnif(n, min, max, n.min, p=p) generates n-n.min+1 p -dimensional points in $[min, max]^p$ returning a numeric matrix with p columns.

Note

For leap Kocis and Whiten recommend values of $L = 31, 61, 149, 409$, and particularly the $L = 409$ for dimensions up to 400.

Author(s)

Martin Maechler

References

James Gentle (1998) *Random Number Generation and Monte Carlo Simulation*; sec.\6.3. Springer.
 Kocis, L. and Whiten, W.J. (1997) Computational Investigations of Low-Discrepancy Sequences.
ACM Transactions of Mathematical Software **23**, 2, 266–294.

Examples

```
32*sHalton(20, base=2)

stopifnot(sHalton(20, base=3, leap=2) ==
          sHalton(20, base=3)[1+2*(0:9)])
## ----- a 2D Visualization -----

Uplot <- function(xy, axes=FALSE, xlab="", ylab="", ...) {
  plot(xy, xaxs="i", yaxs="i", xlim=0:1, ylim=0:1, xpd = FALSE,
        axes=axes, xlab=xlab, ylab=ylab, ...)
  box(lty=2, col="gray40")
}

do4 <- function(n, ...) {
  op <- mult.fig(4, main=paste("n =", n, ": Quasi vs. (Pseudo) Random"),
                marP=c(-2,-2,-1,0))$old.par
  on.exit(par(op))
  for(i in 1:2) {
    Uplot(QUnif(n, p=2), main="QUnif", ...)
    Uplot(cbind(runif(n), runif(n)), main="runif", ...)
  }
}
do4(100)
do4(500)
do4(1000, cex = 0.8, col="slateblue")
do4(10000, pch= ".", col="slateblue")
do4(40000, pch= ".", col="slateblue")
```

read.org.table

*Read.table for an Emacs Org Table***Description**

Read an emacs “Org” table (in file or text) by `read.table()`.

Usage

```
read.org.table(file, header = TRUE, skip = 0,
               encoding = "native", fileEncoding = "", text, quiet=FALSE, ...)
```

Arguments

file	a file name, a file or other connection.
header	logical indicating if the org table has header line (in the usual " "-separated org table format).
skip	integer number of initial lines to skip.
encoding	to be used in the main readLines (file, encoding=encoding) call.
fileEncoding	if file is a file name, i.e., a character string, and fileEncoding is not the empty string, file(file, "rt", encoding = fileEncoding) will be used.
text	instead of file, a character or string (of a few lines, typically).
quiet	logical to suppress the message which is signalled when no nrows=* has been specified and the automatic number of rows is smaller than 95% of the rows / non-header lines of the file.
...	further arguments passed to read.table . You should <i>not</i> use encoding (but possibly fileEncoding!) here, as we do not call read.table on file (but on a textConnection).

Value

a [data.frame](#)

Note

TODO: It should be easy to extend read.org.table() to also work for some of the proposed Markdown formats for tables. Please write to [maintainer](#)("sfsmisc") or open a github issue if you are interested.

References

Org-Mode *Manual* on tables, <https://orgmode.org/manual/Tables.html>
 Org *tutorial* for tables, <https://orgmode.org/worg/org-tutorials/tables.html>

See Also

CRAN package [ascii](#) can *write* org tables. [read.table](#)

Examples

```
t1 <-
"
| a | var2 | C |
|---+-----+-----|
| 2 | may  | 3.4 |
| 7 | feb  | 4.7 |
"

d <- read.org.table(text = t1)
d
stopifnot(dim(d) == c(2, 3),
          identical(names(d), c("a", "var2", "C")),
          d[, "a"] == c(2,7))
```

relErr	<i>Relative Error When Appropriate, Absolute Otherwise</i>
--------	--

Description

relErrV(): Compute the signed relative error componentwise (“vectorized”) between the target and current vectors, using the *absolute* error, i.e., the difference in case the relative error is not well defined, i.e., when target is zero or infinite.

relErr(): simply the *mean* absolute value of the relative errors between target and current vectors; typically the “same” as `all.equal.numeric(target, vector, tolerance=0, countEQ=TRUE)`. Currently useful only when both vectors are finite.

Usage

```
relErrV(target, current, eps0 = .Machine$double.xmin)
relErr (target, current)
```

Arguments

target	numeric, possibly scalar.
current	numeric vector of <code>length()</code> a multiple of <code>length(target)</code> ; if an <code>array</code> (incl <code>matrix</code>), dimensions are preserved; for vectors, <code>names(target)</code> are preserved.
eps0	non-negative number; values <code>abs(target) < eps0</code> should be treated as zero (and hence <i>absolute</i> instead of relative error be computed). This may be crucial when target is an “mpfr”-number vector.

Value

relErrV(): a numeric vector of the same length (or array of the same dimension) as `current`.

relErr(): a single number.

Author(s)

Martin Maechler, originally as part of **Matrix** package’s ‘test-tools.R’.

See Also

`all.equal.numeric()` is similar in spirit but returns TRUE or string containing the *mean* relative or absolute error.

Examples

```
## relErrV() test example: showing how it works fine with {NA, Inf, 0} :
eps <- 1e-4*c(-9, -8, -6, -4, 0.5, 1, 5)
target <- c(-1:1, 0, 0, NA, NaN, Inf, -Inf, Inf, 0, Inf, 1, -3:3)
current <- c(-1:1, 1e-7, NaN, NA, 0, Inf, Inf, 0, Inf, 1, Inf, -3:3+ eps)
cbind(target, current, absE = current-target,
```

```

relE = relErrV(target,current)) -> M ; M
stopifnot(exprs = {
  is.logical(isFr <- is.finite(rF <- M[, "relE"]))
  target==current | isFr == is.finite(aF <- M[, "absE"])
  identical(aF[!isFr] , rF[!isFr])
  identical(numeric(), relErrV(numeric(), integer())) # length 0 {used to fail}
})
tools::assertError(relErrV(1, numeric()), verbose=TRUE) # no longer allowed
## relErr() is pretty simple --- (possibly too simple, currently)
relErr
relErr(target, current) # NA (of course)
all.equal.numeric(target, current) ## "'is.NA' value mismatch ..."

## comparison after dropping NA's :
hasN <- is.na(target) | is.na(current)
all.equal(target[!hasN], current[!hasN], tolerance=0) # "Mean abs. diff.: Inf"
relErr(target[!hasN], current[!hasN]) # NaN (to improve?)

## comparison after only keeping cases where both are finite:
finN <- is.finite(target) & is.finite(current)
all.equal(target[finN], current[finN], tol=0) # "Mean abs.d.: 0.000279.."
all.equal(target[finN], current[finN], tol=0, countEQ=TRUE) # " " : 0.000239..
relErr(target[finN], current[finN]) # 0.0002392929

```

repChar

Make Simple String from Repeating a Character, e.g. Blank String

Description

Simple constructors of a constant character string from one character, notably a “blank” string of given string length.

M.M. is now ‘*mentally deprecating*’ `bl.string` in favor of using `repChar()` in all cases.

With R 3.3.0 (May 2016), the *new* function `strrep()` was introduced; it is faster typically, and more flexible, e.g. accepting a *vector* for the 2nd argument.

This (for now informally) deprecates all uses of `repChar()` and `bl.string()`.

Usage

```

repChar(char, no)
bl.string(no)

```

Arguments

char	single character (or arbitrary string).
no	non-negative integer.

Value

One string, i.e., `character(1)`, for `bl.string` a blank string, fulfilling `n == nchar(bl.string(n))`.

Author(s)

Martin Maechler, early 1990's (for `bl.string`).

See Also

[paste](#), [character](#), [nchar](#).

Examples

```
r <- sapply(0:8, function(n) ccat(repChar(" ",n), n))
cbind(r)

repChar("-", 4)
repChar("_", 6)
## it may make sense to a string of more than one character:
repChar("--- ", 6)

## show the very simple function definitions:
repChar
bl.string
```

rot2

Rotate Planar Points by Angle

Description

Rotate planar (xy) points by angle phi (in radians).

Usage

```
rot2(xy, phi)
```

Arguments

xy	numeric 2-column matrix, or coercable to one.
phi	numeric scalar, the angle in radians (i.e., $\phi=\pi$ corresponds to 180 degrees) by which to rotate the points.

Value

A two column matrix as xy, containing the rotated points.

Author(s)

Martin Maechler, Oct.1994

Examples

```
## Rotate three points by 60 degrees :
(xy0 <- rbind(c(1,0.5), c(1,1), c(0,1)))
(Txy <- rot2(xy0, phi = 60 * pi/180))
plot(xy0, col = 2, type = "b", asp = 1,
      xlim=c(-1,1), ylim=c(0,1.5), main = "rot2(*, pi/3) : 2d rotation by 60°")
points(Txy, col = 3, type = "b")
O <- rep(0,2); P2 <- rbind(xy0[2,], Txy[2,])
arrows(0,0,P2[,1],P2[,2], col = "dark gray")

xy0 <- .8*rbind(c(1,0), c(.5,.6), c(.7,1), c(1,1), c(.9,.8), c(1,0)) - 0.2
plot(xy0, col= 2, type="b", main= "rot2( <polygon>, pi/4 * 1:7)", asp=1,
      xlim=c(-1,1),ylim=c(-1,1), lwd= 2, axes = FALSE, xlab="", ylab="")
abline(h=0, v=0, col="thistle"); text(1.05, -.05, "x"); text(-.05,1.05, "y")
for(phi in pi/4 * 0:7)
  do.call("arrows",c(list(0,0),rot2(xy0[2,], phi), length=0.1, col="gray40"))
for(phi in pi/4 * 1:7)
  polygon(rot2(xy0, phi = phi), col = 1+phi/(pi/4), border=2, type = "b")
```

rotn

Generalized Rot13 Character Translation (Rotation)

Description

Compute generalized ‘rot13’ character translations or “rotations”

In the distant past, considered as poor man’s encryption, such rotations are way too poor nowadays and provided mainly for didactical reasons.

Usage

```
rotn(ch, n = 13)
```

Arguments

ch	a character vector; often a string (of length 1).
n	an integer in $\{1 \dots 26\}$; the default is particularly useful.

Details

Note that the default $n = 13$ makes rotn into a function that is its own inverse.

Written after having searched for it and found `seqinr::rot13()` which was generalized and rendered more transparently to my eyes.

Value

a character as ch, but with each character (which belongs to [letters](#) or [LETTERS](#) “rotated” by n (positions in the alphabet).

Author(s)

Martin Maechler

See Also[rot2](#), a completely different rotation (namely in the plane aka R^2).**Examples**

```
rotn(c("ABC", "a", "b", "c"), 1)
rotn(c("ABC", "a", "b", "c"), 2)
rotn(c("ABC", "a", "b", "c"), 26) # rotation by 26 does not change much

(ch <- paste("Hello", c("World!", "you too")))
rotn(ch)
rotn( rotn(ch) ) # rotn(*, 13) is its own inverse
```

roundfixS

*Round to Integer Keeping the Sum Fixed***Description**

Given a real numbers y_i with the particular property that $\sum_i y_i$ is integer, find *integer* numbers x_i which are close to y_i ($|x_i - y_i| < 1\forall i$), and have identical “marginal” sum, `sum(x) == sum(y)`.

As I found later, the problem is known as “Apportionment Problem” and it is quite an old problem with several solution methods proposed historically, but only in 1982, Balinski and Young proved that there is no method that fulfills three natural desiderata.

Note that the (first) three methods currently available here were all (re?)-invented by M.Maechler, without any knowledge of the litterature. At the time of writing, I have not even checked to which (if any) of the historical methods they match.

Usage

```
roundfixS(x, method = c("offset-round", "round+fix", "1greedy"))
```

Arguments

x	a numeric vector which must sum to an integer
method	character string specifying the algorithm to be used.

Details

Without hindsight, it may be surprising that all three methods give identical results (in all situations and simulations considered), notably that the idea of ‘mass shifting’ employed in the iterative “1greedy” algorithm seems equivalent to the much simpler idea used in “offset-round”.

I am pretty sure that these algorithms solve the L_p optimization problem, $\min_x \|y - x\|_p$, typically for all $p \in [1, \infty]$ *simultaneously*, but have not bothered to find a formal proof.

Value

a numeric vector, say r , of the same length as x , but with integer values and fulfilling $\text{sum}(r) == \text{sum}(x)$.

Author(s)

Martin Maechler, November 2007

References

Michel Balinski and H. Peyton Young (1982) **Fair Representation: Meeting the Ideal of One Man, One Vote**;

https://en.wikipedia.org/wiki/Apportionment_paradox

<https://www.ams.org/samplings/feature-column/fcarc-apportionii3>

See Also

[round](#) etc

Examples

```
## trivial example
kk <- c(0,1,7)
stopifnot(identical(kk, roundfixS(kk))) # failed at some point

x <- c(-1.4, -1, 0.244, 0.493, 1.222, 1.222, 2, 2, 2.2, 2.444, 3.625, 3.95)
sum(x) # an integer
r <- roundfixS(x)
stopifnot(all.equal(sum(r), sum(x)))
m <- cbind(x=x, `r2i(x)` = r, resid = x - r, `|res|` = abs(x-r))
rbind(m, c(colSums(m[,1:2]), 0, sum(abs(m[,`|res|`]))))

chk <- function(y) {
  cat("sum(y) =", format(S <- sum(y)), "\n")
  r2 <- roundfixS(y, method="offset")
  r2. <- roundfixS(y, method="round")
  r2_ <- roundfixS(y, method="1g")
  stopifnot(all.equal(sum(r2 ), S),
            all.equal(sum(r2.), S),
            all.equal(sum(r2_), S))
  all(r2 == r2. & r2. == r2_) # TRUE if all give the same result
}

makeIntSum <- function(y) {
  n <- length(y)
  y[n] <- ceiling(y[n]) - (sum(y[-n]) %% 1)
  y
}
set.seed(11)
y <- makeIntSum(rnorm(100))
chk(y)
```

```
## nastier example:
set.seed(7)
y <- makeIntSum(rpois(100, 10) + c(runif(75, min= 0, max=.2),
                                runif(25, min=.5, max=.9)))

chk(y)

## Not run:
for(i in 1:1000)
  stopifnot(chk(makeIntSum(rpois(100, 10) +
                        c(runif(75, min= 0, max=.2),
                          runif(25, min=.5, max=.9)))))

## End(Not run)
```

range

Robust Range using Boxplot 'Quartiles'

Description

Compute a robust range, i.e. the usual `range()` as long as there are no outliers, using the “whisker boundaries” of `boxplot`, i.e., `boxplot.stats`.

Usage

```
rrange(x, range=1, coef = 1.5, na.rm = TRUE)
```

Arguments

<code>x</code>	numeric vector the robust range of which shall be computed.
<code>range</code>	number for S compatibility; $1.5 * \text{range}$ is equivalent to <code>coef</code> .
<code>coef</code>	numeric multiplication factor defining the outlier boundary, see ‘Details’ below.
<code>na.rm</code>	logical indicating how <code>NA</code> values should be handled; they are simply dropped when <code>na.rm = TRUE</code> as by default.

Details

The robust range is really just what `boxplot.stats(x, coef=coef)` returns as the whisker boundaries. This is the most extreme values `x[j]` still inside median plus/minus `coef * IQR`.

Value

numeric vector `c(m, M)` with $m \leq M$ which is (not strictly) inside `range(x) = c(min(x), max(x))`.

Author(s)

Martin Maechler, 1990.

See Also

[range](#), [fivenum](#), [boxplot](#) and [boxplot.stats](#).

A more sophisticated robust range for (strongly) asymmetric data can be derived from the skewness adjusted boxplot statistics [adjboxStats](#) which is a generalization of [boxplot.stats](#).

Examples

```
stopifnot(rrange(c(1:10,1000)) == c(1,10))
```

seqXtend	<i>Sequence Covering the Range of X, including X</i>
----------	--

Description

Produce a sequence of unique values (sorted increasingly), *containing* the initial set of values `x`. This can be useful for setting prediction e.g. ranges in nonparametric regression.

Usage

```
seqXtend(x, length., method = c("simple", "aim", "interpolate"),
        from = NULL, to = NULL)
```

Arguments

<code>x</code>	numeric vector.
<code>length.</code>	integer specifying <i>approximately</i> the desired length() of the result.
<code>method</code>	string specifying the method to be used. The default, "simple" uses seq(*, length.out = length.) where "aim" aims a bit better towards the desired final length, and "interpolate" interpolates evenly <i>inside</i> each interval $[x_i, x_{i+1}]$ in a way to make all the new intervalls of approximately the same length.
<code>from, to</code>	numbers to be passed to (the default method for) seq() , defaulting to the minimal and maximal <code>x</code> value, respectively.

Value

numeric vector of increasing values, of approximate length `length.` (unless `length. < length(unique(x))` in which case, the result is simply `sort(unique(x))`), containing the original values of `x`.

From, `r <- seqXtend(x, *)`, the original values are at indices `ix <- match(x, r)`, i.e., `identical(x, r[ix])`.

Note

`method = "interpolate"` typically gives the best results. Calling [roundfixS](#), it also need more computational resources than the other methods.

Author(s)

Martin Maechler

See Also[seq](#); [plotDS](#) can make particularly good use of `seqXtend()`**Examples**

```

a <- c(1,2,10,12)
seqXtend(a, 12)# --> simply 1:12
seqXtend(a, 12, "interp")# ditto
seqXtend(a, 12, "aim")# really worse
stopifnot(all.equal(seqXtend(a, 12, "interp"), 1:12))

## for a "general" x, however, "aim" aims better than default
x <- c(1.2, 2.4, 4.6, 9.9)
length(print(seqXtend(x, 12)))      # 14
length(print(seqXtend(x, 12, "aim"))) # 12
length(print(seqXtend(x, 12, "int"))) # 12

## "interpolate" is really nice:
xt <- seqXtend(x, 100, "interp")
plot(xt, main="seqXtend(*, 100, \"interpol\")")
points(match(x,xt), x, col = 2, pch = 20)
# ... you don't even see that it's not equidistant
# whereas the cheap method shows ...
xt2 <- seqXtend(x, 100)
plot(xt2, col="blue")
points(match(x,xt2), x, col = 2, pch = 20)

## with "Date" objects
Drng <- as.Date(c("2007-11-10", "2012-07-12"))
(px <- pretty(Drng, n = 16)) # say, for the main labels
## say, a finer grid, for ticks -- should be almost equidistant
n3 <- 3*length(px)
summary(as.numeric(diff(seqXtend(px, n3))))      # wildly varying
summary(as.numeric(diff(seqXtend(px, n3, "aim")))) # (ditto)
summary(as.numeric(diff(seqXtend(px, n3, "int")))) # around 30

```

Description

Collect (and print) information about the current R session and environment, using `sessionInfo()` and more mostly low-level and platform dependent information.

`isRshared()` is a utility called from `sessionInfoX()`.

Usage

```
sessionInfoX(pkgs = NULL, list.libP = FALSE, extraR.env = TRUE)

## S3 method for class 'sessionInfoX'
print(x, locale = TRUE, RLIBS = TRUE, Renv = TRUE, ...)

isRshared(platform = .Platform)
```

Arguments

pkgs	NULL (default), TRUE or a character vector of R package names, whose packageDescription() s are wanted. No packages by default, TRUE takes all currently loaded pkgs.
list.libP	a logical indicating if for all .libPaths entries, the files should be listed via list.files .
extraR.env	logical indicating if <i>all</i> environment variables should be recorded which start with "R_" or "_R_".
x	typically the result of sessionInfoX() .
locale	logical, passed to print.sessionInfo() indicating if the locale information should be printed.
RLIBS	logical indicating if the information about R_LIBS should be printed.
Renv	logical indicating if the information about R environment variables should be printed.
...	passed to print methods.
platform	a list "like" .Platform .

Value

For [isRshared\(\)](#), a [logical](#) indicating if R has been installed as "shared", i.e., linked to 'libR*' shared library.

For [sessionInfoX\(\)](#), an object of S3 class "sessionInfoX", a [list](#) with components (there may be more, experimental and not yet listed here):

sInfo	simply the value of sessionInfo() .
sysInf	the value of Sys.info() .
capabilities	the value of capabilities() .
Machine	the value of .Machine .
compiledBy	for R 4.3.0 and newer, the value of R_compiled_by() .
extSoft	for R 3.2.0 and newer, the value of extSoftVersion() .
grSoft	for R 3.2.0 and newer, the value of grSoftVersion() .
tclVersion	for R 3.2.0 and newer and when tcltk is loaded, the Tcl version (tclVersion()).
LAPACK	for R 3.0.3 and newer, the value of La_version() .
pcre	for R 3.1.3 and newer, the value of pcre_config() .

pkgDescr	If pkgs was non-empty, a named list of packageDescription() s for each entry in pkgs.
libPath	the value of .libPaths() .
RLIBS	a character vector of entries from Sys.getenv("R_LIBS") , typically very similar to the libPaths component.
n.RLIBS	simply a normalizePath() ed version of RLIBS.
R.env	a named character vector with the “important” R environment variables R_ENVIRON, R_PROFILE, R_CHECK_ENVIRON.
xR.env	if extraR.env was true, a named character vector of “all R related” environment variables, as specified in extraR.env’s description above.
shared	(not available on Windows, where it is conceptually always true:) logical indicating if the version of R is “shared”.

Author(s)

Martin Maechler, December 2015 ff.

See Also

[sessionInfo](#), [.libPaths](#), [R.version](#), [Sys.getenv](#).

Examples

```
six0 <- sessionInfoX()
six0$shared # useful (for some, e.g., MM) on Unix alike

sixN <- sessionInfoX("nlme", list.libP = TRUE)
sixN # -> print() method for "sessionInfoX"
names(sixN)
str(sixN, max = 1) # outline of lower-level structure
str(sixN$pkgDescr) # list with one component "nlme"
```

shortRversion

Short R Version String

Description

From **base** R’s [R.version.string](#), produce a somewhat shorter version, with or without date, notably also for *patched* or *development* versions of R.

Main use is for plotting or construction of file of variable names.

Usage

```
shortRversion(Rv = R.version, Rst = Rv$status,
              Rvstring = if (!is.null(s <- Rv$version.string)) s else R.version.string,
              date = Rst != "", spaces = TRUE)
```

Arguments

Rv	a list with some of the same components as R.version .
Rst	a string specifying the <i>status</i> of R's version. For released versions of R, this is <code>""</code> ; almost always use the default <code>Rv\$status</code> .
Rvstring	a string with a default that should work even for R versions previous to 1.0.0.
date	logical specifying if the date of the R version should be included in the result; by default, this will be true only for non-released versions of R.
spaces	logical indicating if the result may contain spaces (aka 'blanks'); setting it to false, replaces the blanks by <code>"-"</code> and <code>"_"</code> .

Value

a [character](#) string, typically a shortened version of `Rvstring`.

Author(s)

Martin Maechler

See Also

[R.version](#), [R.version.string](#)

Examples

```
shortRversion() ## (including the date, typically for an R Core developer)
## but this is shorter:
(Rver <- shortRversion(date=FALSE))
shortRversion(spaces=FALSE) # e.g. for a file or even directory name
shortRversion(spaces=FALSE, date=FALSE) # even shorter, ditto

## If you want even shorter { abbreviate() will remove spaces, too }:
abbreviate(shortRversion(), 11)
abbreviate(shortRversion(date=FALSE), 13)
```

signi

Rounding to Significant Digits

Description

Rounds to significant digits similarly to [signif](#).

Usage

```
signi(x, digits = 6)
```

Arguments

`x` numeric vector to be rounded.
`digits` number of significant digits required.

Value

numeric vector “close” to `x`, i.e. by at least `digits` significant digits.

Note

This is really just `round(x, digits - trunc(log10(abs(x))))` and hence mainly of didactical use. Rather use `signif()` otherwise.

Author(s)

Martin Maechler, in prehistoric times (i.e. before 1990).

See Also

[signif](#), [round](#).

Examples

```
(x1 <- seq(-2, 4, by = 0.5))
identical(x1, signif(x1))# since 0.5 is exact in binary arithmetic
(x2 <- pi - 3 + c(-5,-1,0, .1, .2, 1, 10,100))
signif(x2, 3)
```

sourceAttach

Source and Attach an R source file

Description

Source (via [sys.source\(\)](#)) and attach ([attach](#)) an R source file.

Usage

```
sourceAttach(file, pos=2,
             name = paste(abbreviate(gsub(fsep,"", dirname(file)),
                                         12, method="both.sides"),
                           basename(file), sep=fsep),
             keep.source = getOption("keep.source.pkgs"),
             warn.conflicts = TRUE)
```

Arguments

file	file name
pos	passed to attach()
name	character, with a smart default, passed to attach() .
keep.source	logical, see sys.source() .
warn.conflicts	logical, see attach .

Value

the return value of [attach\(\)](#).

Author(s)

Martin Maechler, 29 Jul 2011

See Also

[attach](#), [sys.source](#), [source](#)

Examples

```
sourceAttach(system.file("test-tools-1.R", package="Matrix", mustWork=TRUE))
search() # shows the new "data base" at position 2
## look what it contains:
ls.str(pos = 2)
```

str_data

Overview on All Datasets in an R Package

Description

Provide an overview over all datasets available by [data\(\)](#) in a (list of) given R packages.

Usage

```
str_data(pkgs, filterFUN, ...)
```

Arguments

pkgs	character vector of names of R packages.
filterFUN	optionally a logical function for filtering the R objects.
...	potential further arguments to be passed to str ; str(utils:::str.default) gives useful list.

Value

invisibly (see [invisible](#)) a [list](#) with named components matching the `pkgs` argument. Each of these components is a named list with one entry per `data(.)` argument name. Each entry is a [character](#) vector of the names of all objects, typically only one.

The side effect is, as with [str\(\)](#), to print everything (via [cat](#)) to the console.

Author(s)

Martin Maechler

See Also

[str](#), [data](#).

Examples

```
str_data("cluster")

str_data("datasets", max=0, give.attr = FALSE)

## Filtering (and return value)
df1 <- str_data("datasets", filterFUN=is.data.frame)
str(df.d <- df1$datasets)
## dim() of all those data frames:
t(sapply(unlist(df.d), function(.) dim(get(.))))

### Data sets in all attached packages but "datasets" (and stubs):
s <- search()
(Apkgs <- sub("^package:", '', s[grepl("^package:", s)]))
str_data(Apkgs[!Apkgs %in% c("datasets", "stats", "base")])
```

Sys.cpuinfo

Provide Information about the Linux Hardware (CPU, Memory, etc)

Description

Return information about the Linux hardware, notably the CPU (the central processor unit) and memory of the computer R is running on. This is currently **only available for Linux**.

These functions exist on other unix-alike platforms, but produce an error when called.

Usage

```
Sys.procinfo(procfile)
Sys.cpuinfo()
Sys.meminfo()
Sys.memGB(kind = "MemTotal")
Sys.MIPS()
```

Arguments

procfile	name of file the lines of which give the CPU info “as on Linux”
kind	a character string specifying which <i>kind</i> of memory is desired.

Value

The Sys.*info() functions return a "simple.list", here basically a named character vector, (where the names have been filtered through [make.names](#)(*, unique=TRUE) which is of importance for multi-processor or multi-core CPUs, such that vector can easily be indexed.

Sys.memGB() returns available memory in giga bytes [GB];

Sys.MIPS() returns a number giving an approximation of the **M**illion **I**nstructions **P**er **S**econd that the CPU processes (using “bogomips”). This is a performance measure of the basic *non-numeric* processing capabilities. For single-core Linux systems, often about twice the basic clock rate in “MHz” (as available by Sys.cpuinfo()[“cpu.MHz”]); now, with multicore systems, the result is often around (but smaller than) $2 * \#\{\text{cores}\} * \text{clock.rate}$.

Note

These currently do rely on the Linux ‘/proc/’ file system, and may not easily be portable to non-Linux environments.

On multi-processor machines, Sys.cpuinfo() contains each field for each processor (i.e., names(Sys.cpuinfo()) has [duplicated](#) entries).

Conceivably, the bogoMIPS source code is open and available and could be built into R.

Author(s)

Martin Maechler

See Also

[Sys.ps](#), etc.

Examples

```
(n.cores <- parallel::detectCores())
if(substr(R.version[["os"]], 1,5) == "linux") { ##-- only on Linux
  Sys.cpuinfo() # which is often ugly; this looks much better:
  length(Sys.cpu2 <- local({I <- Sys.cpuinfo(); I[ !grepl("^flags", names(I)) ] })))
  ## may still be too much, notably if n.cores > 2:
  (Sys3 <- Sys.cpu2[!grepl("[.][0-9]+$", names(Sys.cpu2))])

  Sys.MIPS() ## just the 'bogomips' from above:
  Sys.MIPS() / as.numeric(Sys.cpuinfo()[“cpu.MHz”]) ## ~~ 2 * #{cores} ((no longer))

  ## Available Memory -- can be crucial:
  Sys.memGB() #- default "MemTotal"
  if(Sys.memGB("MemFree") > 16)
    message("Be happy! You have more than 16 Gigabytes of free memory")
}
```

Sys.ps

Return Process Status (Unix 'ps') Information

Description

These functions return process id and status information, typically about the running R process.

Usage

```
Sys.ps(process= Sys.getpid(),
       fields = c("pid", "pcpu", "time", "vsz", "comm"),
       usefile = length(fields) > 10,
       ps.cmd  = Sys.ps.cmd(),
       verbose = getOption("verbose"),
       warn.multi = verbose || any(fields != "ALL"))
```

```
Sys.sizes(process = Sys.getpid(), ps.cmd = Sys.ps.cmd())
```

Arguments

process	the process id, an integer.
fields	character strings of "ALL", specifying which process status fields are desired.
usefile	logical; if true, system writes to a temporary file and that is scanned subsequently.
ps.cmd	character string, giving the "ps" command name to be used.
verbose	logical ...
warn.multi	logical ...

Details

Use `man ps` on your respective Unix system, to see what fields are supported exactly. Unix dialects *do* differ here, and, SunOS-Solaris even has more than one ps command...

Value

Note, that `Sys.sizes()` currently returns two integers which are "common" to Solaris and Linux.

Author(s)

Martin Maechler

See Also

[Sys.info](#), [Sys.getpid](#), [proc.time](#).

Examples

```
(.pid <- Sys.getpid()) ## process ID of current process
Sys.sizes(.pid)

## The default process statistics about the running R process
try( Sys.ps() )
```

TA.plot

Tukey-Anscombe Plot (Residual vs. Fitted) of a Linear Model

Description

From a linear (or glm) model fitted, produce the so-called Tukey-Anscombe plot. Useful (optional) additions include: 0-line, lowess smooth, 2sigma lines, and automatic labeling of observations.

Usage

```
TA.plot(lm.res,
        fit= fitted(lm.res), res= residuals(lm.res, type="pearson"),
        labels= NULL, main= mk.main(), xlab = "Fitted values",
        draw.smooth= n >= 10, show.call = TRUE, show.2sigma= TRUE,
        lo.iter = NULL, lo.cex= NULL,
        par0line = list(lty = 2, col = "gray"),
        parSmooth = list(lwd = 1.5, lty = 4, col = 2),
        parSigma = list(lwd = 1.2, lty = 3, col = 4),
        verbose = FALSE,
        ...)
```

Arguments

lm.res	Result of <code>lm(.)</code> , <code>avov(.)</code> , <code>glm(.)</code> or a similar object.
fit	fitted values; you probably want the default here.
res	residuals to use. Default: Weighted ("Pearson") residuals if weights have been used for the model fit.
labels	strings to use as plotting symbols for each point. Default(NULL): extract observations' names or use its sequence number. Use, e.g., "*" to get simple * symbols.
main	main title to plot. Default: sophisticated, resulting in something like "Tukey-Anscombe Plot of : y ~ x" constructed from <code>lm.res \$ call</code> .
xlab	x-axis label for plot.
draw.smooth	logical; if TRUE, draw a lowess smoother (with automatic smoothing fraction).
show.call	logical; if TRUE, write the "call"ing syntax with which the fit was done.
show.2sigma	logical; if TRUE, draw horizontal lines at $\pm 2\sigma$ where σ is <code>mad(resid)</code> .
lo.iter	positive integer, giving the number of lowess robustness iterations. The default depends on the model and is 0 for non Gaussian <code>glm</code> 's.

lo.cex	character expansion ("cex") for lowess and other marginal texts.
par0line	a list of arguments (with reasonable defaults) to be passed to <code>abline(.)</code> when drawing the x-axis, i.e., the $y = 0$ line.
parSmooth, parSigma	each a list of arguments (with reasonable default) for drawing the smooth curve (if <code>draw.smooth</code> is true), or the horizontal sigma boundaries (if <code>show.2sigma</code> is true) respectively.
verbose	logical indicating if some construction details should be reported (<code>print()</code> ed).
...	further graphical parameters are passed to <code>n.plot(.)</code> .

Side Effects

The above mentioned plot is produced on the current graphic device.

Author(s)

Martin Maechler, Seminar fuer Statistik, ETH Zurich, Switzerland; <maechler@stat.math.ethz.ch>

See Also

`plot.lm` which also does a QQ normal plot and more.

Examples

```
data(stackloss)
TA.plot(lm(stack.loss ~ stack.x))

example(airquality)
summary(lm0 <- lm(Ozone ~ ., data= airquality))
TA.plot(lm0)
TA.plot(lm0, label = "O") # instead of case numbers

if(FALSE) {
  TA.plot(lm(cost ~ age+type+car.age, claims, weights=number, na.action=na.omit))
}

##--- for aov(.) : -----
data(Gun, package = "nlme")
TA.plot( aov(rounds ~ Method + Physique/Team, data = Gun))

##--- Not so clear what it means for GLM, but: -----
if(require(rpart)) { # for the two datasets only
  data(solder, package = "rpart")
  TA.plot(glm(skips ~ ., data = solder, family = poisson), cex= .6)

  data(kyphosis, package = "rpart")
  TA.plot(glm(Kyphosis ~ poly(Age,2) + Start, data=kyphosis, family = binomial),
    cex=.75) # smaller title and plotting characters
}
```

tapplySimpl

*More simplification in tapply() result***Description**

For the case of more than two categories or indices (in INDEX), traditional `tapply(*, simplify = TRUE)` still returns a list when an array may seem more useful and natural. This is provided by `tapplySimpl()` if the function `FUN()` is defined such as to return a vector of the same length in all cases.

Usage

```
tapplySimpl(X, INDEX, FUN, ...)
```

Arguments

X	an atomic object, typically a vector. All these arguments are as in <code>tapply()</code> and are passed to <code>tapply(. .)</code> .
INDEX	list of (typically more than one) factors, each of same length as X.
FUN	the function to be applied. For the result to be simplifiable, <code>FUN()</code> must return a vector of always the same length.
...	optional arguments to FUN.

Value

If the above conditions are satisfied, the list returned from `r <- tapply(X, INDEX, FUN, ...)` is simplified into an `array` of rank $1 + \#\{indices\}$, i.e., $1 + \text{length}(INDEX)$; otherwise, `tapplySimpl()` returns the list `r`, i.e., the same as `tapply()`.

Author(s)

Martin Maechler, 14 Jun 1993 (for S-plus).

See Also

`tapply(*, simplify=TRUE)`.

Examples

```
## Using tapply() would give a list (with dim() of a matrix);
## here we get 3-array:

data(esoph)
with(esoph, {
  mima <- tapplySimpl(ncases/ncontrols, list(agegp, alcgp), range)
  stopifnot(dim(mima) == c(2, nlevels(agegp), nlevels(alcgp)))
})
aperm(mima)
```

tkdensity

*GUI Density Estimation using Tcl/Tk***Description**

This is graphical user interface (GUI) to [density](#), allowing for dynamic bandwidth choice and a simple kind of zooming, relying on `library(tcltk)`.

Usage

```
tkdensity(y, n = 1024, log.bw = TRUE, showvalue = TRUE,
          xlim = NULL, do.rug = size < 1000, kernels = NULL,
          from.f = if (log.bw) -2 else 1/1000,
          to.f   = if (log.bw) +2.2 else 2,
          col = 2)
```

Arguments

<code>y</code>	numeric; the data the density of which we want.
<code>n</code>	integer; the number of abscissa values for density evaluation (and plotting).
<code>log.bw</code>	logical; if true (default), the gui scrollbar is on a <i>log</i> bandwidth scale, otherwise, simple interval.
<code>showvalue</code>	logical; if true, the value of the current (log) bandwidth is shown on top of the scrollbar.
<code>xlim</code>	initial <code>xlim</code> for plotting, see plot.default .
<code>do.rug</code>	logical indicating if rug(y) should be added to each plot. This is too slow for really large sample sizes.
<code>kernels</code>	character vector of kernel names as allowable for the <code>kernels</code> argument of the standard density function.
<code>from.f, to.f</code>	numeric giving the left and right limit of the bandwidth scrollbar.
<code>col</code>	color to be used for the density curve.

Details

`library(tcltk)` must be working, i.e., Tcl/Tk must have been installed on your platform, and must have been visible during R's configuration and/or installation.

You can not only choose the bandwidth (the most important parameter), but also the kernel, and you can zoom in and out (in x-range only).

Value

none.

(How could this be done? `tcltk` widgets run as separate processes!)

Author(s)

Martin Maechler, building on demo(tkdensity).

Examples

```
if (dev.interactive(TRUE)) ## does really not make sense otherwise
  if(try(require("tcltk"))){ ## sometimes (rarely) there, but broken

  data(faithful)
  tkdensity(faithful $ eruptions)

  set.seed(7)
  if(require("nor1mix"))
    tkdensity(rnorMix(1000, MW.nm9), kernels = c("gaussian", "epanechnikov"))
}
```

toLatex.numeric

LaTeX or Sweave friendly Formatting of Numbers

Description

Formats real numbers, possibly in scientific notation, with a given number of digits after the decimal point. Output can be used in LaTeX math mode, e.g., for printing numbers in a table, where each number has to be printed with the same number of digits after the decimal point, even if the last digits are zeros.

Usage

```
## S3 method for class 'numeric'
toLatex(object, digits = format.info(object)[2],
        scientific = format.info(object)[3] > 0, times = "\\cdot", ...)
```

Arguments

object	a numeric vector.
digits	number of digits <i>after the decimal point</i> (for the mantissa if scientific). The default behaves the same as R's <code>format()</code> .
scientific	logical indicating if scientific notation $a * 10^k$ should be used. The default behaves the same as R's <code>format()</code> .
times	character string indicating the LaTeX symbol to be used for the ‘times’ sign.
...	unused; for compatibility with <code>toLatex</code> .

Value

a `character` vector of the same length as `object`, containing the formatted numbers.

Note

We use digits for `round`, i.e., round after the decimal point on purpose, rather than `signif()` icant digit rounding as used by `print()` or `format()`.

Author(s)

Alain Hauser

See Also

`pretty10exp` which gives `expressions` similar to our `scientific=TRUE`. `toLatex` with other methods.

Examples

```
xx <- pi * 10^(-9:9)

format(xx)
formatC(xx)

toLatex(xx) #-> scientific = TRUE is chosen
toLatex(xx, scientific=FALSE)

sapply(xx, toLatex)
sapply(xx, toLatex, digits = 2)
```

u.assign0

'Portable' assign / get functions (R / S-plus) for 'Frame 0'

Description

R does not have S' concept of `frame = 0`, aka 'session frame'. These two function were an attempt to provide a portable way for working with frame 0, particularly when porting code *from* S.

They have been **deprecated** since August 2013.

Usage

```
u.assign0(x, value, immediate = FALSE)
u.get0(x)
```

Arguments

<code>x</code>	character string giving the <i>name</i> of the object.
<code>value</code>	any R object which is to be assigned.
<code>immediate</code>	logical, for S compatibility. No use in R.

Note

Really don't use these anymore...

Author(s)

Martin Maechler

See Also

[get](#), [assign](#).

u.boxplot.x

Utility Returning x-Coordinates of Boxplot

Description

Return the x-coordinates in an 'n-way' side-by-side boxplot. This is an auxiliary function and exists mainly for backcompatibility with S-plus.

Usage

```
u.boxplot.x(n, j = 1:n, fullrange = 100)
```

Arguments

n	number of boxplots.
j	indices of boxplots.
fullrange	x-coords as 'uniform' in $[0, fullrange]$; (f.=100, corresponds to Splus 3.x (x = 1,2)).

Value

a numeric vector of length n, with values inside $(0, M)$ where $M = fullrange$.

Author(s)

Martin Maechler

See Also

[boxplot](#).

Examples

```
u.boxplot.x(7) # == 8.93 22.62 36.3 ... 91.07
```

u.date	<i>Return Date[-Time] String in 'European' Format</i>
--------	---

Description

Return one string of the form "day/month/year", plus "hour:minutes", optionally.

Usage

```
u.date(short=FALSE)
```

Arguments

short logical; if TRUE, no time is given.

Value

String with current date (and time).

Author(s)

Martin Maechler, ca. 1992

See Also

[u.Datumvonheute.](#)

Examples

```
u.date()
u.date(short = TRUE)
```

u.datumdecode	<i>Convert "Numeric" Dates</i>
---------------	--------------------------------

Description

Daten der Form 8710230920 aufspalten in Jahr, Monat, Tag, Std, Min

Usage

```
u.datumdecode(d, YMDHMnames = c("Jahr", "Monat", "Tag", "Std", "Min"))
```

Arguments

d numeric dates in the form YYMMDDHHMM.
YMDHMnames (column) names to be used for the result.

Value

a numeric matrix (or vector) with 5 columns containing the year, month, etc.

Note

MM: This is a wrong concept, and also suffers from the “millenium bug” (by using only 2 digits for the year).

Author(s)

?? (someone at SfS ETH)

See Also

R’s *proper* date-time coding: [DateTimeClasses](#); [u.date](#) etc.

Examples

```
u.datumdecode(8710230920)
##  Jahr Monat  Tag  Std  Min
##   87    10   23    9   20

u.datumdecode(c(8710230900, 9710230920, 0210230920))
##      Jahr Monat Tag Std Min
## [1,]   87    10  23   9  00
## [2,]   97    10  23   9  20
## [3,]    2    10  23   9  20
```

u.Datumvonheute	<i>Datum und Uhrzeit (auf deutsch)</i>
-----------------	--

Description

Return current date and time as a string, possibly including day of the week in *German*.

Usage

```
u.Datumvonheute(W.tag=2, Zeit=FALSE)

C.Monatsname
C.Wochentag
C.Wochentagkurz
C.weekday
```

Arguments

- W.tag logical or integer specifying you want weekday (‘Wochentag’). 0 or FALSE gives no, 1 or TRUE gives a short and 2 the long version of the day of the week.
- Zeit logical or integer specifying if time ("Zeit") is desired. 0 or FALSE gives no, 1 or TRUE gives a hours only and 2 hours and minutes.

Value

A string with the current date/time, in the form specified by the arguments.

The C.* are [character](#) vector “constants”, the German ones actually used by u.Datumvonheute.

Author(s)

Caterina Savi, Martin Maechler

See Also

[u.date](#) for a similar English version, and [p.datum](#) which plots. For English month names, etc [month.name](#).

Examples

```
u.Datumvonheute()
u.Datumvonheute(W.tag=1, Zeit=TRUE)
u.Datumvonheute(W.tag= FALSE, Zeit=2)
```

u.log

(Anti)Symmetric Log High-Transform

Description

Compute $\log()$ only for high values and keep low ones – antisymmetrically such that $u.\log(x)$ is (once) continuously differentiable, it computes $f(x) = x$ for $|x| \leq c$ and $\text{sign}(x)c \cdot (1 + \log(|x|/c))$ for $|x| \geq c$.

Usage

```
u.log(x, c = 1)
```

Arguments

x	numeric vector to be transformed.
c	scalar, > 0

Details

Alternatively, the ‘IHS’ (inverse hyperbolic sine) transform has been proposed, first in more generality as $S_U()$ curves by Johnson(1949); in its simplest form, $f(x) = \text{arsinh}(x) = \log(x + \sqrt{x^2 + 1})$, which is also antisymmetric, continuous and once differentiable as our $u.\log(.)$.

Value

numeric vector of same length as x.

Author(s)

Martin Maechler, 24 Jan 1995

References

N. L. Johnson (1949). Systems of Frequency Curves Generated by Methods of Translation, *Biometrika* **36**, pp 149–176. doi:10.2307/2332539

See Also

Werner Stahel's sophisticated version of John Tukey's "started log" (which was $\log(x + c)$), with concave extension to negative values and adaptive default choice of c ; **logst** in his CRAN package **relevance**, or **LogSt** in package **DescTools**.

Examples

```
curve(u.log, -3, 10); abline(h=0, v=0, col = "gray20", lty = 3)
curve(1 + log(x), .01, add = TRUE, col= "brown") # simple log
curve(u.log(x, 2), add = TRUE, col=2)
curve(u.log(x, c= 0.4), add = TRUE, col=4)

## Compare with IHS = inverse hyperbolic sine == asinh
ihs <- function(x) log(x+sqrt(x^2+1)) # == asinh(x) {aka "arsinh(x)" or "sinh^{-1}(x)"}
xI <- c(-Inf, Inf, NA, NaN)
stopifnot(all.equal(xI, asinh(xI))) # but not for ihs():
cbind(xI, asinh=asinh(xI), ihs=ihs(xI)) # differs for -Inf
x <- runif(500, 0, 4); x[100+0:3] <- xI
all.equal(ihs(x), asinh(x)) #==> is.NA value mismatch: asinh() is correct {i.e. better!}

curve(u.log, -2, 20, n=1000); abline(h=0, v=0, col = "gray20", lty = 3)
curve(ihs(x)+1-log(2), add=TRUE, col=adjustcolor(2, 1/2), lwd=2)
curve(ihs(x), add=TRUE, col=adjustcolor(4, 1/2), lwd=2)
## for x >= 0, u.log(x) is nicely between IHS(x) and shifted IHS

## a log10-scale version of asinh() {aka "IHS"}: ihs10(x) := asinh(x/2) / ln(10)
ihs10 <- function(x) asinh(x/2)/log(10)
xyaxis <- function() abline(h=0, v=0, col = "gray20", lty = 3)
leg3 <- function(x = "right")
  legend(x, legend = c(quote(ihs10(x) == asinh(x/2)/log(10)),
    quote(log[10](1+x)), quote(log[10](x))),
    col=c(1,2,5), bty="n", lwd=2)
curve(asinh(x/2)/log(10), -5, 20, n=1000, lwd=2); xyaxis()
curve(log10(1+x), col=2, lwd=2, add=TRUE)
curve(log10( x ), col=5, lwd=2, add=TRUE); leg3()

## zoom out and x-log-scale
curve(asinh(x/2)/log(10), .1, 100, log="x", n=1000); xyaxis()
curve(log10(1+x), col=2, add=TRUE)
curve(log10( x ), col=5, add=TRUE) ; leg3("center")

curve(log10(1+x) - ihs10(x), .1, 1000, col=2, n=1000, log="x", ylim = c(-1,1)*0.10,
  main = "absolute difference", xaxt="n"); xyaxis(); eaxis(1, sub10=1)
```

```

curve(log10( x ) - ihs10(x), col=4, n=1000, add = TRUE)

curve(abs(1 - ihs10(x) / log10(1+x)), .1, 5000, col=2, log = "xy", ylim = c(6e-9, 2),
      main="|relative error| of approx. ihs10(x) := asinh(x/2)/log(10)", n=1000, axes=FALSE)
eaxis(1, sub10=1); eaxis(2, sub10=TRUE)
curve(abs(1 - ihs10(x) / log10(x)), col=4, n=1000, add = TRUE)
legend("left", legend = c(quote(log[10](1+x)), quote(log[10](x))),
      col=c(2,4), bty="n", lwd=1)

## Compare with Stahel's version of "started log"
## (here, for *vectors* only, and 'base', as DescTools::LogSt());

## by MM: "modularized" by providing a threshold-computer function separately:
logst_thrWS <- function(x, mult = 1) {
  lq <- quantile(x[x > 0], probs = c(0.25, 0.75), na.rm = TRUE, names = FALSE)
  if (lq[1] == lq[2])
    lq[1] <- lq[2]/2
  lq[1]^(1 + mult)/lq[2]^mult
}
logst0L <- function(x, calib = x, threshold = thrFUN(calib, mult=mult),
  thrFUN = logst_thrWS, mult = 1, base = 10)
{
  ## logical index sub-assignment instead of ifelse(): { already in DescTools::LogSt }
  res <- x # incl NA's
  notNA <- !is.na(sml <- (x < (th <- threshold)))
  i1 <- sml & notNA; res[i1] <- log(th, base) + ((x[i1] - th)/(th * log(base)))
  i2 <- !sml & notNA; res[i2] <- log(x[i2], base)
  attr(res, "threshold") <- th
  attr(res, "base") <- base
  res
}
logst0 <- function(x, calib = x, threshold = thrFUN(calib, mult=mult),
  thrFUN = logst_thrWS, mult = 1, base = 10)
{
  ## Using pmax.int() instead of logical indexing -- NA's work automatically - even faster
  xm <- pmax.int(threshold, x)
  res <- log(xm, base) + (x - xm)/(threshold * log(base))
  attr(res, "threshold") <- threshold
  attr(res, "base") <- base
  res
}

## u.log() is really using natural log() -- whereas logst() defaults to base=10
curve(u.log, -4, 10, n=1000); abline(h=0, v=0, col = "gray20", lty = 3); points(-1:1, -1:1, pch=3)
curve(log10(x) + 1, add=TRUE, col=adjustcolor("midnightblue", 1/2), lwd=4, lty=6)
curve(log10(x), add=TRUE, col=adjustcolor("skyblue3", 1/2), lwd=4, lty=7)
curve(logst0(x, threshold= 2 ), add=TRUE, col=adjustcolor("orange",1/2), lwd=2)
curve(logst0(x, threshold= 1 ), add=TRUE, col=adjustcolor(2, 1/2), lwd=2)
curve(logst0(x, threshold= 1/4), add=TRUE, col=adjustcolor(3, 1/2), lwd=2, lty=2)
curve(logst0(x, threshold= 1/8), add=TRUE, col=adjustcolor(4, 1/2), lwd=2, lty=2)

```

u.sys

'Portable' System function (R / S-plus)

Description

u.sys() is a convenient wrapper (of system()) to call to the underlying operating system. The main purpose has been to provide a function with identical UI both in S-PLUS and R. MM thinks you shouldn't use this anymore, usually.

Sys.ps.cmd() returns the 'ps' (**p**rocess **s**tatus) OS command name (as [character](#) string), and is typically usable on unix alike only.

Usage

```
u.sys(..., intern = TRUE)
```

```
Sys.ps.cmd()
```

Arguments

... any number of strings – which will be [paste\(\)](#)d together and passed to system.
intern logical – note that the default is *reversed* from the one in [system\(\)](#).

Author(s)

Martin Maechler

See Also

[system](#), really!; on non-Windows, Sys.ps() which makes use of Sys.ps.cmd().

Examples

```
u.sys # shows how simply the function is defined :
## Not run:
  function (... , intern = TRUE)
    system(paste(... , sep = " "), intern = intern)

## End(Not run)

# All *running* processes of user [sometimes only R]:
try ( u.sys(Sys.ps.cmd(), "ur") )
```

unif*Nice Uniform Points in Interval*

Description

Give regularly spaced points on interval $[-c, c]$ with mean 0 (exactly) and variance about 1 (very close for **even** `n` and larger `round.dig`). Note that c depends on `n`.

Usage

```
unif(n, round.dig = 1 + trunc(log10(n)))
```

Arguments

<code>n</code>	positive integer specifying the number of points desired.
<code>round.dig</code>	integer indicating to how many digits the result is rounded.

Value

numeric vector of length `n`, symmetric around 0, hence with exact mean 0, and variance approximately 1.

Note

It relies on the fact that $Var(1, 2, \dots, n) = n(n + 1)/12$.

Author(s)

Martin Maechler, ca 1990

See Also

[runif](#) for producing uniform *random* numbers.

Examples

```
(u <- unif(8))
var(u)

(u. <- unif(8, 12))# more digits in result, hence precision for Var :
var(u.)
```

uniqueL	<i>A Reversible Version of unique()</i>
---------	---

Description

A version of [unique](#) keeping enough information to reverse (or *invert*) to the original data.

Usage

```
uniqueL(x, isuniq = !duplicated(x), need.sort = is.unsorted(x))
```

Arguments

<code>x</code>	numeric vector, of length <code>n</code> , say.
<code>isuniq</code>	logical vector of the same length as <code>x</code> . For the reversion to work this should select at least all unique values of <code>x</code> .
<code>need.sort</code>	logical indicating if <code>x</code> is not yet sorted. Note that this argument exists only for speedup possibility when it is known, and that it <i>must be set correctly</i> .

Value

list of two components,

<code>ix</code>	integer vector of indices
<code>xU</code>	vector of values from <code>x</code>

such that both `x[isuniq] == xU` and `xU[ix] == x`.

Author(s)

Martin Maechler

See Also

[Duplicated](#) from the `sfsmisc` package in addition to the standard [unique](#) and [duplicated](#).

Examples

```
x0 <- c(1:3,2:7,8:4)
str(r0 <- uniqueL(x0))
with(r0, xU[ix]) ## == x0 !
```

Description

Concatenate vector elements or anything using `paste(*, collapse = .)`. These are simple short abbreviations I have been using in my own codes in many places.

Usage

```
vcat(vec, sep = " ")
ccat(...)
```

Arguments

<code>vec, ...</code>	any vector and other arguments to be pasted to together.
<code>sep</code>	the separator to use, see the <i>Details</i> section.

Details

The functions are really just defined as

```
vcat := function(vec, sep = " ") paste(vec, collapse = sep)
```

```
ccat := function(...) paste(..., collapse = "", sep = "")
```

Value

a character string (of length 1) with the concatenated arguments.

Author(s)

Martin Maechler, early 1990's.

See Also

`paste`, `as.character`, `format`. `cat()` is really for printing.

Examples

```
ch <- "is"
ccat("This ", ch, " it: ", 100, "%")
vv <- c(1,pi, 20.4)
vcat(vv)
vcat(vv, sep = ", ")
```

wrapFormula

*Enhance Formula by Wrapping each Term, e.g., by "s(.)"***Description**

The main motivation for this function has been the easy construction of a “full GAM formula” from something as simple as $Y \sim \dots$

The potential use is slightly more general.

Usage

```
wrapFormula(f, data, wrapString = "s(*)")
```

Arguments

f the initial [formula](#); typically something like $Y \sim \dots$
data [data.frame](#) to which the formula applies; see, [formula](#) or also [gam](#) or [lm](#).
wrapString [character](#) string, containing `"*"`, specifying the wrapping expression to use.

Value

a [formula](#) very similar to `f`; just replacing each *additive* term by its wrapped version.

Note

There are limits for this to work correctly; notably the right hand side of the formula `f` should not be nested or otherwise complicated, rather typically just $\dots$ as in the examples.

Author(s)

Martin Maechler, May 2007.

See Also

[formula](#); [gam](#) from package [mgcv](#) (or also from package [gam](#)).

Examples

```
myF <- wrapFormula(Fertility ~ . , data = swiss)
myF # Fertility ~ s(Agriculture) + s(....) + ...

if(require("mgcv")) {
  m1 <- gam(myF, data = swiss)
  print( summary(m1) )
  plot(m1, pages = 1) ; title(format(m1$call), line= 2.5)
}

## other wrappers:
```

```
wrapFormula(Fertility ~ . , data = swiss, wrap = "lo(*)")
wrapFormula(Fertility ~ . , data = swiss, wrap = "poly(*, 4)")
```

xy.grid*Produce Regular Grid Matrix*

Description

Produce the grid used by [persp](#), [contour](#), etc, as an N x 2 matrix. This is really outdated by [expand.grid\(\)](#) nowadays.

Usage

```
xy.grid(x, y)
```

Arguments

x, y any vectors of same mode.

Value

a 2-column matrix of “points” for each combination of x and y, i.e. with $\text{length}(x) * \text{length}(y)$ rows.

Author(s)

Martin Maechler, 26 Oct 1994.

See Also

[expand.grid](#) which didn’t exist when `xy.grid` was first devised.

Examples

```
plot(xy.grid(1:7, 10*(0:4)))

x <- 1:3 ; y <- 10*(0:4)
xyg <- xy.grid(x,y)

## Compare with expand.grid() :
m2 <- as.matrix(expand.grid(y,x)[, 2:1])
dimnames(m2) <- NULL
stopifnot(identical(xyg, m2))
```

xy.unique.x

*Uniqify (X,Y) Values using Weights***Description**

Given *smoother* data (x_i, y_i) and maybe weights w_i , with multiple x_i , use the unique x values, replacing the y 's by their (weighted) mean and updating the weights accordingly.

Usage

```
xy.unique.x(x, y, w, fun.mean = mean, ...)
```

Arguments

x, y	numeric vectors of same length. Alternatively, x can be a 'xy' like structure, see xy.coords .
w	numeric vector of non-negative weights – or missing which corresponds to all weights equal.
fun.mean	the mean function to use.
...	optional arguments all passed to unique .

Value

Numeric matrix with three columns, named x, y and w with unique x values and corresponding y and weights w.

Author(s)

Martin Maechler, 8 Mar 1993.

See Also

e.g., [smooth.spline](#) uses something like this internally.

Examples

```
## simple example:
x <- c(1,1,2,4,3,1)
y <- 1:6
rbind(x, y)
xy.unique.x(x, y)
#   x y w
# 1 1 3 3
# 2 2 3 1
# 3 4 4 1
# 4 3 5 1
xy.unique.x(x, y, fromLast = TRUE)
```

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