

Package ‘ymse’

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Title Ymse (Various)

Version 0.6.7

Description Supplies a number of more or less useful functions and methods suitable for, eg. estimating dice roll probabilities, calculate latin squares, perform binary search, adjust colours in HSV space, produce prime numbers, find maximum acf/pacf/ccf, convert floats to simple ratio, produce averaged shifted histogram drop variables from formulae using regex, flatten a nested list, compute the similarity between two character vectors, plot a simple loess smooth, and other assorted tasks.

Depends R (>= 3.5.0)

Imports stats, utils, graphics, grDevices

License GPL (>= 2)

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LazyData true

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acf_max	<i>Maximum ACF, PACF and CCF</i>
---------	----------------------------------

Description

Find lag that maximizes correlation

Usage

```
acf_max(
  x,
  ...,
  plot = FALSE,
  show = plot,
  ci = 0.95,
  ma.ci = TRUE,
  max.type = c("pos", "neg", "abs"),
  most.signif = FALSE
)

pacf_max(
  x,
  ...,
  plot = FALSE,
  show = plot,
  ci = 0.95,
  max.type = c("pos", "neg", "abs")
)

ccf_max(
  x,
  y,
  ...,
  plot = FALSE,
  show = plot,
  ci = 0.95,
  max.type = c("pos", "neg", "abs")
)
```

Arguments

<code>x, y</code>	univariate numeric vector or time series
<code>...</code>	further arguments passed to <code>acf</code> , <code>pacf</code> , <code>ccf</code>
<code>plot</code>	logical; return a plot
<code>show</code>	indicate on the plot the maximum correlation
<code>ci</code>	confidence interval used, by default 95%
<code>ma.ci</code>	should the confidence limits assume an MA input (TRUE, the default), or white noise as is default for <code>plot.acf</code> ?
<code>max.type</code>	what maximum should be returned, the positive (default), negative, or absolute maximum?
<code>most.signif</code>	should the most significant correlation be returned. Only applicable if <code>ma.ci=TRUE</code>

Examples

```
x <- c(5, 5, 3, 6, 3, 6, 9, 6, 3, 1, 3, 2, 8, 9, 4, 3, 6, 6,
      6, 7, 5, 2, 5, 1, 5, 5, 0, 3, 7, 3, 6, 6, 2, 2, 6, 5)
y <- c(8, 9, 7, 5, 3, 5, 6, 9, 6, 3, 4, 5, 9, 7, 8, 5, 5, 7,
      4, 7, 7, 2, 5, 6, 5, 7, 5, 3, 5, 6, 7, 0, 5, 3, 8, 4)
```

```
acf_max(x, plot=TRUE, max.type="abs")
acf_max(x, max.type="neg")
acf_max(x, max.type="neg", most.signif=TRUE)
```

```
pacf_max(x, plot=TRUE)
pacf_max(x, max.type="abs")
```

```
ccf_max(x, y, plot=TRUE)
ccf_max(x, y, max.type="neg")
```

```
# Same plot
plot(acf(x, plot=FALSE), ci.type="ma")
acf_max(x, plot=TRUE)
```

```
acf_max(x, ci=0.99, plot=TRUE)
ccf_max(x, y, ci=0, max.type="pos", plot=TRUE)
```

addrows

Add rows to a data.frame

Description

An "rbind for data.frames", sort of.

Usage

```
addrows(dtf, nrw, top = FALSE)
```

Arguments

dtf	data.frame; original data.frame
nrv	data.frame; the new row(s) to be added
top	logical; should the new rows be added to the top or the bottom (default)?

Details

Can only bind two objects at a time, but will bind data.frames with non-matching column names and -classes. In such cases the original data.frame will serve as template.

Examples

```
dtf <- data.frame(A=letters[1:5],
                  B=1:5,
                  C=as.factor(5:1),
                  D=as.Date(0:4, origin="2000-01-01"),
                  stringsAsFactors=FALSE)

nrv <- data.frame(A=letters[1:5],
                  B=4:8,
                  C=5:1,
                  D=as.Date(5:1, origin="1990-01-01"),
                  stringsAsFactors=FALSE)

str(dtf)

dtf.a <- addrows(dtf, nrv, top=FALSE)
str(dtf.a)

# adding a single row with little concern for data types and column names
b <- type.convert(beaver1[80:90,])
b$activ <- as.logical(b$activ)

addrows(b, data.frame(350, 1200, 37.02, 1))
```

adjustcolorHSV

Adjust Colors in One or More Directions Conveniently.

Description

Adjust or modify a vector of colors by "turning knobs" on one or more coordinates in (h,s,v, α) space, typically by up or down scaling them.

Usage

```
adjustcolorHSV(
  col,
  alpha.f = 1,
  h.f = 1,
  s.f = 1,
  v.f = 1,
  offset = c(0, 0, 0, 0),
  transform = diag(c(h.f, s.f, v.f, alpha.f)),
```

```

    h = NULL,
    s = NULL,
    v = NULL,
    alpha = NULL
  )

```

Arguments

<code>col</code>	vector of colors, in any format that <code>col2rgb()</code> accepts
<code>alpha.f</code> , <code>h.f</code> , <code>s.f</code> , <code>v.f</code>	factors scaling the opacity, hue, saturation and value of <code>col</code>
<code>offset</code>	a length 4 numeric vector specifying the linear offset applied to the <i>hue</i> , <i>saturation</i> , <i>value</i> and <i>alpha</i> values
<code>transform</code>	a 4x4 diagonal matrix specifying the scaling applied to the <i>hue</i> , <i>saturation</i> , <i>value</i> and <i>alpha</i> values
<code>h</code> , <code>s</code> , <code>v</code> , <code>alpha</code>	fixed vlues for hue, saturation, value and alpha. Overrides any corresponding scaling factor or offset

Details

Essentially an HSV version of the RGB-based [adjustcolor](#). One important distinction is that the `h.f` value wraps around to fit the [0, 1] range, rather than simply "clamping" it between 0 and 1.

Value

A character vector the same length as `col` containng color data in standard hexadecimal RGBA format.

Examples

```

# Halve the saturation and value of the default palette colours
plot(2:8, cex=5, lwd=4, pch=21, bg=2:8,
     col=adjustcolorHSV(2:8, s.f=0.5, v.f=0.6))

# Offset the hue of the default palette colours by 0.5, inverting the colours
plot(2:8, cex=5, lwd=4, pch=21, bg=2:8,
     col=adjustcolorHSV(2:8, offset=c(0.5, 0, 0, 0)))

```

ahist

Average shifted histogram

Description

Create a smoothed histogram by averaging several histograms shifted by fractions of a bin-width

Usage

```
ahist(
  x,
  n.breaks = nclass.FD(x),
  n.shifts = 3,
  type = c("histogram", "polygon", "line", "table"),
  freq = FALSE,
  plot = TRUE,
  add = FALSE,
  ...
)
```

Arguments

<code>x</code>	a vector of values for which the histogram is desired
<code>n.breaks</code>	an integer giving the number of bins to be used
<code>n.shifts</code>	an integer giving the number of shifts to be performed
<code>type</code>	if <code>plot=TRUE</code> , the type of plot to be used
<code>freq</code>	should frequency counts be used, or density (default)
<code>plot</code>	logical; if <code>TRUE</code> (default), a graphical output will be returned
<code>add</code>	logical; if <code>TRUE</code> the plot will be added to the current plot
<code>...</code>	further graphical parameters to <code>ymse::plot.histogram</code> , <code>polygon</code> , or <code>lines</code>

Value

an object of class "histogram"

Examples

```
set.seed(1)
n <- 6

x <- sample(sample(0:20, 8), 6*n, replace=TRUE) + rnorm(6*n, -8, 0.5)
x <- c(x, rgamma(5*n, 3, 0.5), rnorm(4*n, 15, 2))
x <- round(x*5)/5

hist(x, freq=FALSE, breaks="FD", col="lightblue")
ahist(x, type="hist", border=2, col=7, freq=FALSE, lwd=2)
ahist(x, type="poly", border=2, col=7, freq=FALSE, lwd=2)
ahist(x, type="line", col=2, freq=FALSE, lwd=2)
ahist(x, type="table", col=2, freq=FALSE, lwd=2)
ahist(x, plot=FALSE)
```

align_char

Align character strings

Description

Align character strings

Usage

```
align_char(x, pattern = ".", ..., lpad = " ", rpad = " ")
```

Arguments

<code>x</code>	numeric or character vector
<code>pattern</code>	pattern, passed to <code>regexpr</code> , whose match the character strings will be aligned by
<code>...</code>	further arguments passed to <code>regexpr</code>
<code>lpad</code> , <code>rpadd</code>	character strings used for padding. Repeated to length

See Also

[align_num](#), for alignment more aimed at numeric strings

Examples

```
x <- c("Tom und Jerry", "Abbott og Costello", "Milo och Stich", "et alii")
cat(align_char(x, pat="[:alpha:]"), sep="\n")
cat(align_char(x, pat=" "), sep="\n")
cat(align_char(x, pat=" [A-Z]"), sep="\n")
cat(align_char(x, pat=" [a-z]"), sep="\n")
cat(align_char(x, pat="t", ignore.case=TRUE), sep="\n")
cat(align_char(x, pat="x"), sep="\n")
```

align_num

Align numbers

Description

Align numbers for neat vertical printing

Usage

```
align_num(x, lpad = " ", rpad = "0", dec = ".", min.dec = 0, rm.dec = TRUE)
```

Arguments

<code>x</code>	numeric or character vector
<code>lpad</code> , <code>rpadd</code>	character strings used for padding. Repeated to length
<code>dec</code>	decimal separator, or any other character to align by
<code>min.dec</code>	pad with zeros to reach a minimum number of decimal points
<code>rm.dec</code>	remove zeros at the end of whole numbers

See Also

[align_char](#), for alignment more aimed at character strings

Examples

```
x <- c(22100, 100, 1015, 13.018, 0.1, 0.01234)
cat(align_num(x), sep="\n") # Default
cat(format(x, scientific=FALSE), sep="\n")

cat(align_num(x, rpad=" "), sep="\n")
cat(align_num(x, rpad=" ", rm.dec=FALSE), sep="\n")
cat(align_num(x, rpad=" ", min.dec=1), sep="\n")
cat(align_num(x, rpad=" ", min.dec=2), sep="\n")

cat(align_num(x, rpad="\U00b7", min.dec=0), sep="\n")
cat(align_num(x, lpad="' ", rpad="'", min.dec=0), sep="\n")

cat(align_num(c("1.000.000", "10.000.000", "1.000,85"), dec=","), sep="\n")

# corner cases
x <- c("100.", "1.2", ".1111")
cat(align_num(x, rpad=" ", rm.dec=TRUE), sep="\n")
cat(align_num(round(as.numeric(x))), rpad=" ", rm.dec=TRUE), sep="\n")

# matching on more than one character
# so far not much more advanced than this
# working on align_num2 more suited for character strings
s <- c("cataract", "hematology", "pancreatic")
cat(align_num(s, dec="a", rpad=" "), sep="\n")
cat(align_num(s, dec="at", rpad=" "), sep="\n")

a <- c("Tom and Jerry", "Milo and Stich", "Abbott and Costello")
cat(align_num(a, dec="and", rpad=" "), sep="\n")
```

arfilter

AR filter

Description

Filter a time series using AR coefficients

Usage

```
arfilter(x, mod, x.mean = mod$x.mean, init = "focb")
```

Arguments

x	a time series
mod	an AR model
x.mean	the mean used. By default the mean of the original model. Set to zero for no demeaning
init	how the initial values should be chosen. First observation carried backwards (default), mean of the first values, or the first values in reverse.

See Also

[armodel](#)

Examples

```

set.seed(1)
arap <- ar(AirPassengers)
spec.ar(arap)
spec.pgram(arfilter(rnorm(10000), arap), span=21, na.action=na.omit)

arm <- armodel(c(1.3, -0.4))
spec.ar(arm)
plot(x <- rnorm(200), type="l")
lines(scale(arfilter(x, arm), center=FALSE), col="red", lwd=2)

```

arfit	<i>AR model fit</i>
-------	---------------------

Description

Fit a specified AR model to a univariate time series

Usage

```
arfit(x, mod, x.mean = mod$x.mean)
```

Arguments

x	a time series
mod	an AR model
x.mean	the mean used. By default the mean of the original model. Set to zero for no demeaning

See Also

[armodel](#) for examples

Examples

```

set.seed(1)
x <- runif(50) + sin(1:50/10)
plot(x); lines(arfilter(x, armodel(c(1.5, -0.5, 0.5)), x.mean=mean(x)))

```

arimpulse	<i>Impulse response of an AR model</i>
-----------	--

Description

Get and plot the impulse response of an AR model

Usage

```
arimpulse(mod, pulse = 1, n.ahead = 20, plot = TRUE, ...)
```

Arguments

mod	an AR model
pulse	numeric vector; the initial pulse. Magnitude is added to the model mean
n.ahead	the length of the computed response
plot	logical; should the result be plotted?
...	further arguments to plot

See Also

[armodel](#) for examples

armodel	<i>Create an AR model object</i>
---------	----------------------------------

Description

Specify the characteristics of an AR model

Usage

```
armodel(  
  coefs,  
  mean = 0,  
  intercept = 0,  
  var.pred = 1,  
  frequency = 1,  
  x.name = "Synthetic AR model"  
)
```

Arguments

coefs	a vector of model coefficients
mean	the mean of the process
intercept	the intercept in the model
var.pred	the portion of the variance not explained by this model
frequency	the sampling frequency of the process
x.name	name of the series

See Also

[arimpulse](#)

Examples

```

# short decay
ar.mod <- armodel(c(0.5))
arimpulse(ar.mod, pulse=1)

# long decay
ar.mod <- armodel(c(0.8))
arimpulse(ar.mod, pulse=1)

# negative second coefficient reduce damping, signal returns to normal
# more quickly
ar.mod <- armodel(c(0.8, -0.1))
arimpulse(ar.mod, pulse=1)

# second coefficient reduce damping too much, overdamping, oscillations
ar.mod <- armodel(c(0.8, -0.5))
arimp <- arimpulse(ar.mod, pulse=1, n.ahead=40)$pred
polyroot(c(1, -ar.mod$ar)) # complex conjugate roots
acf(arimp) # period ~= 6?
phi1 <- ar.mod$ar[1]
phi2 <- ar.mod$ar[2]
f <- (1/(2*pi)) * acos((phi1*(phi2-1))/(4*phi2))
1/f # period = 6.78
sp <- spec.ar(ar.mod, plot=FALSE)
1/sp$freq[which.max(sp$spec)] # period = 6.79

# decaying oscillations
ar.mod1 <- armodel(c(0.8, -0.6, -0.5, 0.2, -0.2))
arimpulse(ar.mod1, n.ahead=100)
Mod(1/polyroot(c(1, -ar.mod1$ar))) # barely inside the unit circle

# growing oscillations
ar.mod2 <- armodel(c(0.8, -0.7, -0.5, 0.2, -0.2))
arimpulse(ar.mod2, n.ahead=100)
Mod(1/polyroot(c(1, -ar.mod2$ar))) # barely outside the unit circle

ar.mod3 <- armodel(c(1.8, -1.1, 0.2, -0.2, 0.2))
arimpulse(ar.mod3, n.ahead=100)
spec.ar(ar.mod3)

resid(arfit(rnorm(10), armodel(c(0.5, -0.1), frequency=2)))

```

as.array.list

Coerce a list to an array

Description

Coerce a list consisting of data.frames or matrices of equal size to a 3d array

Usage

```

## S3 method for class 'list'
as.array(x, ...)

```

Arguments

`x` a list of equal sized data.frames or matrices
`...` (not used)

Value

A list of length l with elements of m rows and n columns wix result in an $m \times n \times l$ array.

Examples

```
df1 <- data.frame(x=c(1, 2, 3), y=c(2, 3, 4), z=c(3, 4, 5))
df2 <- data.frame(x=c(4, 2, 3), y=c(2, 5, 4), z=c(3, 4, 6))
df3 <- data.frame(x=c(1, 4, 2), y=c(3, 3, 8), z=c(4, 3, 5))

l <- list(df1, df2, df3)

as.array(l)

llm <- list(matrix(LETTERS[1:6], 2),
            matrix(LETTERS[7:12], 2))

as.array(llm)

as.array(speedskate)
```

aug_median

*Centre weighted mean***Description**

Offering a continuous link between median and arithmetic mean

Usage

```
aug_median(x, p = 1, rank = FALSE, na.rm = FALSE)
```

Arguments

`x` numeric vector
`p` positive numeric narrowness of the weight. 1 gives triangular weighting. Higher values gives narrower weights, approaching meadian, lower values gives broader weights, approaching arithmetic mean
`rank` logical. Should should ranks or numeric values determine relative weights?
`na.rm` logical. Should missing values be removed?

Details

A weighted arithmetic mean is calculated over the input vector, where most weight is given to the median value(s), and monotonically less towards either extreme. Falloff depends on p , with small values resulting in a gentler falloff and less difference between minimum and maximum weights.

Examples

```
x <- c(0, 8, 8, 8, 9)

aug_median(x)

# 0 and 9 are considered equidistant from 8
aug_median(x, rank=TRUE)

# Nearly a point weight placed at the median
aug_median(x, 100)
median(x)

# Nearly uniform weights
aug_median(x, 0.001)
mean(x)
```

bartlett

Maurice Stevenson Bartlett's car data

Description

This is an example data set Bartlett used for a lecture course on stochastic processes, Statistics Department, University College, London. The data represents the times, in seconds, when cars passed an observation point by a road.

Bartlett attributes the data to a Dr A. J. Miller who supplied them as a class example. According to Adery C. A. Hope the data was recorded on a rural Swedish road.

Usage

```
bartlett
```

Format

A numeric vector representing time points in seconds

M. S. Bartlett's notes

Analyse the above data with a view to examining:

- i** whether the times of passing constitute a Poisson process;
- ii** if not, whether some form of "bunching" or "clustering" seems to be present.

Possible analyses include:

- a** testing the homogeneity of the consecutive random time-intervals, by means of a partitioning of the degrees of freedom for the total (approximate) χ^2 ;
- b** testing the homogeneity of counts in consecutive fixed time-intervals, choosing an appropriate interval, and partitioning the degrees of freedom corresponding to the total dispersion by means of an analysis of variance;
- c** testing the correlation between the consecutive random time-intervals;
- d** examining the overall distribution of counts in fixed time-intervals;

e examining the overall distribution of the consecutive random time-intervals

You should undertake at least sufficient of these to answer the questions asked.

Source

The Spectral Analysis of Point Processes (p. 280), M. S. Bartlett, 1963

Also mentioned in:

Statistical Estimation of Density Functions (p. 252), M. S. Bartlett, 1963

A Simplified Monte Carlo Significance Test Procedure (p. 583), Adery C. A. Hope, 1968

Examples

```
cpgram(diff(bartlett))

bartlett2 <- bartlett - bartlett[1]

x <- rep(0, tail(bartlett2, 1)*10)
x[bartlett2*10] <- 1

par(mfrow=c(2, 1), mar=c(2, 3, 1, 1))
plot(x, type="l", ann=FALSE)
lines(cumsum(x)/sum(x), col="red", lwd=2)

sp <- spectrum(x, main="", xlim=c(0, 0.1), ylim=c(1e-3, 0.04))
spec <- predict(loess(sp$spec[1:3000] ~ sp$freq[1:3000], span=0.15), se=TRUE)
lines(sp$freq[1:3000], spec$fit, col="red", lwd=2)
lines(sp$freq[1:3000], spec$fit - qt((0.99 + 1)/2, spec$df)*spec$se,
      lty=1, col="lightblue")
lines(sp$freq[1:3000], spec$fit + qt((0.99 + 1)/2, spec$df)*spec$se,
      lty=1, col="lightblue")
```

binsearch

Binary search

Description

Find the position of a given value in a sorted array

Usage

```
binsearch(val, arr, L = 1L, H = length(arr))
```

```
binclosest(val, arr, L = 1L, H = length(arr))
```

Arguments

<code>val</code>	the value to search for
<code>arr</code>	a sorted array to make the search in
<code>L</code>	a lower bound
<code>H</code>	an upper bound

Details

While both `val` and `arr` can be either integer or double, the algorithm is limited by integer storage in how long the array can be. `L` and `H` can be used to limit the range of indices to be search within. `binsearch` will return either the index of the exact match, or the index just below if no exact match is found. This means that if `val` is less than the lowest value in `arr` (and `L=1`), a `0` will be returned, which can lead to issues as such an index does not exist in `R`. An array indexed by `0` will return a zero length object. `binclosest` will return the index of the closest match, and therefore a `1` in the situation where `binsearch` returns a `0`. If there is a tie the lower index will be returned. In either case, if there are duplicate matches, the lower index will be returned.

Value

A single integer representing an index on the input array.

Examples

```
binsearch(15, (1:9)*3.333)
binsearch(2, (1:9)*3.333)
binclosest(2, (1:9)*3.333)

binsearch(18, seq_len(2e9))
## Not run:
binsearch(18, seq_len(3e9))
## End(Not run)
binsearch(18, seq_len(3e9), H=2e9)
binsearch(2000, seq_len(3e7)*100 + 0.1)

set.seed(1)
x <- sort(sample(1:300, 30))
r <- sort(sample(1:300, 30))

plot(sapply(r, binsearch, x), type="l")
lines(sapply(r, binclosest, x), col="red")

x <- c(1, 2, 3, 5, 8, 9)
binclosest(6, x)
binclosest(7, x)
binclosest(5, x)
```

bix

Bix attributes

Description

`bix` provides access to the `bix` attribute of a variable. The first form returns the value of the levels of its argument and the second sets the attribute.

Usage

```
bix(d)

bix(d) <- value
```


Arguments

d	a "dice" object
value	value to begin index at

Examples

```
d <- dice(6)
d
bix(d)
bix(d) <- 3
d
expand(d)
```

caleidoscope	<i>Caleidoscopic effect on a matrix</i>
--------------	---

Description

Flip a matrix vertically and horizontally before recombining into a new large matrix

Usage

```
caleidoscope(m, odd = TRUE)
```

Arguments

m	a matrix
odd	logical; should the resulting matrix have odd dimensions?

Details

Three copies of m will be made. One flipped horizontally, one flipped vertically, and one flipped both horizontally and vertically. Then they are recombined with the original matrix in the upper right corner, and the flipped copies in the upper left, lower right and lower left corners, respectively.

Value

A matrix of either $2 \times$ or 2×-1 the number of rows and columns of the input matrix.

Examples

```
caleidoscope(matrix(1:4, 2), odd=FALSE)

image(caleidoscope(1:9 %o% 1:9))

image(caleidoscope(matrix(runif(180*200)^2, 180)), col=rainbow(256, start=0.58))
```

cbapply

Apply function to contents of clipboard

Description

Read in clipboard contents as lines, apply a function on them, and write results back to the clipboard

Usage

```
cbapply(FUN, ..., collapse = FALSE, write = TRUE, eval = FALSE)
```

Arguments

FUN	function to be applied
...	optional arguments to FUN
collapse	collapse the lines into a single string separated by newlines
write	write the results back to the clipboard
eval	parse and evaluate the results

Examples

```
## Not run:
# Copy to clipboard
a <- 10
b <- 20
s <- a + b
s
# end

# Run
ev <- cbapply(FUN=function(x) paste(x, "+ 2"), eval=TRUE)
ev; a; b; s

# Clipboard contents changed to
a <- 10 + 2
b <- 20 + 2
s <- a + b + 2
s + 2
# end

# Copy to clipboard
One Two
Three
# end

# Run
cbapply(FUN=toupper, write=FALSE)

# Clipboard contents unchanged
One Two
Three
```

```
# end
## End(Not run)
```

central.tendency	<i>Central tendency measures</i>
------------------	----------------------------------

Description

Central tendency measures

Usage

```
pseudomedian(x, na.rm = TRUE)

cmode(x, single = TRUE, ...)

dmode(x, single = TRUE, na.rm = FALSE)

midrange(x, na.rm = FALSE)

srmean(x, na.rm = FALSE)
```

Arguments

x	numeric vector
na.rm	remove NAs before starting calculations
single	return a single value (for cmode and dmode)
...	send further arguments to underlying function, e.g. density for cmode

See Also

[means](#)

Examples

```
xx <- c(1, 3, 4, 5, 7, 8, 9, 9, 7, 5, 4, 5, 3, 8)
median(xx)
pseudomedian(xx)

# Discrete mode
dmode(c(2, 3, 3, 4, 5))
dmode(c(2, 3, 3, 2, 5))
dmode(c(2, 3, 3, 2, 5), single=FALSE)
dmode(c(2, 1, 3, NA, 1))
dmode(c(2, 1, 3, NA, NA))

# Continuous mode
cmode(c(2, 3, 3, 4, 5))
cmode(c(2, 3, 3, 4, 5))
cmode(c(2, 3, 3, 4, 4, 5), n=512)
cmode(c(2, 2, 3, 3, 6, 6, 6, 7), single=FALSE, adjust=0.5)
```

```
# Slightly robust mean
set.seed(1)
r <- round(rexp(12)*c(-100, 100))
mean(r)
srmean(r)
weighted.mean(sort(r), c(0.5, rep(1, length(r)-2), 0.5))
```

combodice

Combine dice

Description

Generate probability density functions for combinations of dice.

Usage

```
combodice(x, FUN, ..., method = c("outer", "expand.grid", "convolve"), name)
```

Arguments

x	a list of dice objects, or objects that can be interpreted as such
FUN	function passed on to outer or apply, depending on method
...	further arguments passed to FUN
method	method for computation. One of outer, expand.grid or convolve
name	name used for the resulting PDF. Will use x object if none is given

Details

Each of the methods have their advantages and disadvantages. Outer and expand.grid work with roughly the same speed and memory, and can take the same kind of input, but FUN is interpreted differently, reflecting their use of outer and apply respectively. Convolve is much quicker than the other two, but is restricted to only summing distributions. While the first two can handle non-integer values, but only integer probabilities, the third can handle non-integer probabilities, but only integer values.

Value

A table giving the relative probability of each value

See Also

[dusd](#)

Examples

```

# Fudge dice
dF.2 <- as.table(c("-1"=2, "0"=2, "1"=2))
dF.1 <- as.table(c("-1"=1, "0"=4, "1"=1))
fudgedice2221 <- list(dF.2, dF.2, dF.2, dF.1)

combodice(fudgedice2221)

# Heterogeneous-class list and non-integer values
die1 <- as.table(c("2.6"=2, "3"=1, "5"=1))
die2 <- c(0, 1.4)
die3 <- as.dice(as.table(c("1"=2, "2"=2, "3"=2)))
diel <- list(die1, die2, die3)

combodice(diel)

# Regular d6 pair
re <- combodice(list(1:6, 1:6))

# Sichermann pair
si <- combodice(list(c(1, 2, 2, 3, 3, 4), c(1, 3, 4, 5, 6, 8)))
re; si # Identical

# One regular and one "average" d6
combodice(list(1:6, c(2, 3, 3, 4, 4, 5)))

# One 1/2 coin, one D4 and one d6, multiplied together
combodice(list(1:2, 1:4, 1:6), "*")

# Probability of getting n 1s throwing 1d4, 1d6 and 2d8
f <- function(x) sum(x == 1)
combodice(list(1:4, 1:6, 1:8, 1:8), FUN=f, method="exp")

# 3d6, discarding the lowest
discard_lowest <- function(x) sum(x[-which.min(x)])
combodice(list(1:6, 1:6, 1:6), discard_lowest, method="exp")

# 1d4, 2d6 and 1d20, discarding lowest and highest
olympic <- function(x) sum(x[-c(which.min(x), which.max(x))])
combodice(list(1:4, 1:6, 1:6, 1:20), olympic, method="exp")

# Dice pool. 3 d10 with target value 7
f <- function(x) sum(x >= 7)
combodice(lapply(rep(1, 3), seq, 10), f, method="ex")/10^3

# Equivalent using binomial PDF
dbinom(0:3, 3, 0.4)

# I have a d20 with a slight bump at the 4 and 10 facets,
# which makes 16 and 11 less likely, but the nearby 3, 18, 19 and 20
# correspondingly more likely. How does this affect the PDF?
d20l <- dice(20)
d20l[c(16, 11)] <- 0.6
d20l[c(3, 20, 18, 19)] <- 1.2
mean(d20l)

```

```

c0 <- combodice(list(dice(6), dice(10), dice(20)), method="conv", name="fair")
c1 <- combodice(list(dice(6), dice(10), d201), method="conv", name="uneven")

set_mar()
plot(c0, type="o", pch=16, col="grey")
points(c1, col=2, type="o", lwd=1, pch=16, cex=0.6)
legend("topright", c("fair", "bumpy"), bty="n", col=c("grey", "red"), lwd=2:1)

```

comparison_with_ties *Comparison with ties*

Description

Compare numeric values, returning an inbetween value for ties

Usage

```

x %tgt% y

tgt(x, y, bias = 0.5)

x %tlt% y

tlt(x, y, bias = 0.5)

```

Arguments

x, y	numeric values to be compared
bias	what bias should be given to ties? 0.5, the default, is considered neutral as it's halfway between 1 and 0 (true and false).

See Also

[Comparison](#), [tied_triple_test](#)

Examples

```

1:5 %tlt% 3
1:5 %tgt% 3

c(1, 4, 3, 1) %tlt% c(1, 3, 3, 2)
c(1, 4, 3, 1) %tgt% c(1, 3, 3, 2)

# Calculate MannWhitney U statistic
set.seed(1)
x <- sort(round(runif(20)*13, 1))
y <- sort(round(runif(15)*10, 1))
o <- outer(x, y, "%tgt%")

sum(o)
wilcox.test(x, y, exact=FALSE)$statistic

```

default_par	<i>Default par</i>
-------------	--------------------

Description

Sets par settings to their default values

Usage

```
default_par()
```

Details

Default par settings can be retrieved by `data(.def.par)`. A new default can be specified by editing `def.par` or making a `def.par <- par(no.readonly=TRUE)` type call.

See Also

Other `par_and_plot_margins_functions`: [revert_par\(\)](#), [set_mar\(\)](#)

dice	<i>Create, modify or convert from/to dice objects</i>
------	---

Description

Create, modify or convert from/to dice objects

Usage

```
dice(dval)

is.dice(x, ...)

as.dice(x, ...)

## S3 method for class 'dice'
print(x, ...)

## S3 method for class 'dice'
as.table(x, ...)
```

Arguments

dval	an integer vector
x	an arbitrary R object
...	further arguments passed to methods

See Also

[expand](#), [table](#)

Examples

```
# Regular d6 dice
dice(6)

# d4 dice with sides 0, 1, 2, 4
dice(c(0:3))

# d4 dice with two 2s and two 5s
dice(c(2, 2, 5, 5))
```

dput2

*Write an Object to console***Description**

Writes an ASCII text representation of an R object to the console for easy copy/paste sharing

Usage

```
dput2(
  x,
  width = 65,
  assign = c("front", "end", "none"),
  breakAtParen = FALSE,
  compact = TRUE,
  exdent = NULL
)
```

Arguments

x	an object
width	integer; column width
assign	character; should assignment be included?
breakAtParen	logical; should lines break at parenthesis begins
compact	remove spaces around ' = ' assignments
exdent	a non-negative integer specifying the exdentation of lines after the first. default 2 if assign="front", else 0.

Details

This is similar to the way dput is used to print ASCII representations of objects to the console. The differences are that dput2 lets you specify the width of the resulting column, and assignment of the object to the name used in the call will by default be included. Line breaks are by default only done on whitespace, but can be set to happen at parenthesis begins as well. This should not break code and can make for a more compact representation, but it can also make the code harder to read.

See Also

[dput](#), [deparse](#), [explode](#)

Examples

```
xmpl <- faithful[sort(sample(1:nrow(faithful), 50)), ]
dput(xmpl)
cat(deparse(xmpl, width.cutoff=65), sep='\n')
dput2(xmpl, compact=FALSE)
dput2(xmpl)
dput2(xmpl, assign="end")
dput2(xmpl, assign="none")
dput2(xmpl, 80)

# no line breaks on whitespaces or parens within character strings
xmpl <- mtcars[1:5, ]
rownames(xmpl) <- c("bbbb (hhhhhhh\u00A0hhhhhhhh)",
                    " rrrrrrrr ( bbbbbb )",
                    "v v v v v v v v v v",
                    "( g-god, d-god, _-_-_-)",
                    "100*(part)/(total)")
dput2(xmpl, 15)
dput2(xmpl, 15, breakAtParen=TRUE)
```

drop_pattern

*Drop predictors***Description**

Drop predictor variables according to a (regex) pattern

Usage

```
drop_pattern(form, pattern, ...)
```

Arguments

form	a formula object
pattern	predictors matching this pattern will be dropped
...	further arguments passed on to grep1

Details

form is divided into its individual terms, any term matching pattern is removed, before form is updated and returned. In case no match is made, form is returned unmodified. In case all predictors match, only the intercept is retained. In any case the response variable(s) are kept as is.

Value

A formula object

See Also

[drop_randfx](#)

Examples

```
f6 <- y ~ aa*bb + aa + ac + cc + acab

drop_pattern(f6, "a") # Drop all containing a
drop_pattern(f6, "a{2}") # Drop all containing exactly 2 consecutive as
drop_pattern(f6, "^^[a]*a^[a]*$") # All containing exactly 1 a
drop_pattern(f6, ":") # Drop interaction
drop_pattern(f6, "^[^:]*a^[^:]*$") # Drop all containg a, but not interaction
drop_pattern(f6, "^(?!a).*$", perl=TRUE) # Drop all not containing a

# Degenerate cases
drop_pattern(f6, "[abc]") # Drop all
drop_pattern(f6, "q") # Drop none
```

drop_randfx	<i>Drop random effects</i>
-------------	----------------------------

Description

Drop random effects from a mixed effects model formula

Usage

```
drop_randfx(form)
```

Arguments

form a formula object

Details

form is divided into its individual terms, any term containg a vertical bar (|) is removed, before form is updated and returned. In case form has no random effect terms, form is returned unmodified. In case all effects are random, only the intercept is retained. In any case the response variable(s) are kept as is.

Value

A formula object

See Also

[drop_pattern](#)

Examples

```
f1 <- Reaction ~ (1 + Days | Subject)
f2 <- Reaction ~ (1 | mygrp/mysubgrp) + (1 | Subject)
f3 <- Reaction ~ x1 + x2 + (1 + Days | Subject)
f4 <- Reaction ~ x1 * x2 + (1 | mygrp/mysubgrp) + (1 | Subject)
f5 <- Reaction ~ x1 + x2
```

```
sapply(list(f1, f2, f3, f4, f5), drop_randfx)
```

dtf_clean

Data cleanup

Description

Create a data.frame from a messy table

Usage

```
dtf_clean(
  x,
  header = TRUE,
  na.strings = c("NA", "N/A"),
  stringsAsFactors = FALSE,
  ...
)
```

Arguments

x a messy table the form of a character string

header does the table include headers? (default TRUE)

na.strings a vector of character strings which will be interpreted as missing values

stringsAsFactors should strings be read as factors? (default FALSE)

... further arguments passed to `read.table`

Examples

```
## Not run:
```

```
x1 <- "
```

```
+-----+-----+-----+-----+-----+
| Date | Emp1 | Case | Priority | PriorityCountinLast7days |
+-----+-----+-----+-----+-----+
| 2018-06-01 | A | A1 | 0 | 0 |
| 2018-06-03 | A | A2 | 0 | 1 |
| 2018-06-02 | B | B2 | 0 | 2 |
| 2018-06-03 | B | B3 | 0 | 3 |
+-----+-----+-----+-----+-----+
```

```
"
```

```
x2 <- '
```

```
+-----+-----+-----+-----+-----+
| Date | Emp1 | Case | Priority | PriorityCountinLast7days |
+-----+-----+-----+-----+-----+
| 2018-06-01 | A | "A 1" | 0 | 0 |
| 2018-06-03 | A | "A 2" | 0 | 1 |
| 2018-06-02 | B | "B 2" | 0 | 2 |
| 2018-06-03 | B | "B 3" | 0 | 3 |
+-----+-----+-----+-----+-----+
```

```
'  
  
x3 <- "  
  
    Date      | Emp1 | Case | Priority | PriorityCountinLast7days  
  
    2018-06-01 | A    | A|1  |      0 |      0  
    2018-06-03 | A    | A|2  |      0 |      1  
    2018-06-02 | B    | B|2  |      0 |      2  
    2018-06-03 | B    | B|3  |      0 |      3  
  
"  
  
x4 <- "  
    Maths | English | Science | History | Class  
  
    0.1   | 0.2     | 0.3     | 0.2     | Y2  
  
    0.9   | 0.5     | 0.7     | 0.4     | Y1  
  
    0.2   | 0.4     | 0.6     | 0.2     | Y2  
  
    0.9   | 0.5     | 0.2     | 0.7     | Y1  
"  
  
x5 <- "  
    Season    | Team  | W | AHWO  
-----  
    1 | 2017/2018 | TeamA | 2 | 1.75  
    2 | 2017/2018 | TeamB | 1 | 1.85  
    3 | 2017/2018 | TeamC | 1 | 1.70  
    4 | 2017/2018 | TeamD | 0 | 3.10  
    5 | 2016/2017 | TeamA | 1 | 1.49  
    6 | 2016/2017 | TeamB | 3 | 1.51  
    7 | 2016/2017 | TeamC | 2 | 1.90  
    8 | 2016/2017 | TeamD | 0 | N/A  
"  
  
lapply(c(x1, x2, x3, x4), dtf_clean)  
  
## End(Not run)
```

dusd	<i>Discrete (Uniform) Sum Distributions</i>
------	---

Description

Generate distributions of the sum of discrete (uniform) random variables. Two different approaches.

Usage

```
dusd1(xr = 1:6, n = 2, FUN = "+")  
  
dusd2(xi = rep(1, 6), n = 2, bix = 1, round, limit = 1e-13)
```

Arguments

<code>xr</code>	numeric vector; a vector of equiprobable values
<code>n</code>	integer; the number of distributions to be summed
<code>FUN</code>	function passed on to <code>outer</code>
<code>xi</code>	numeric vector; a vector of probabilities, with indices representing values
<code>bix</code>	logical; where does the index of <code>xi</code> start?
<code>round</code>	integer; number of digits to round to after each convolution
<code>limit</code>	numeric; values (frequencies or counts) less than this will be omitted.

Details

`dusd1` works by recursively taking the outer sum of `xr`, while `dusd2` recursively convolves `xi`. Although convolution is more efficient, it can introduce small errors, and with repeated convolutions those errors can compound. By rounding to a slightly lower precision after each convolution the generation of spurious singletons and general imprecisions can be mitigated.

Value

`dusd1` returns an array of size $\text{length}(\text{xr})^n$ representing every possible outcome. `dusd2` returns a probability mass function in the form of a table.

See Also

[combodice](#) for a more flexible implementation of the same ideas

Examples

```
# five coin flips
plot(table(dusd1(0:1, 5)))
plot(dusd2(c(1, 1), 5, bix=0))
plot(as.table(dbinom(0:5, 5, 0.5)))

# ten flips with a loaded coin
plot(table(dusd1(c(1, 1, 2), 10)))
plot(dusd2(c(2, 1), 10))
plot(dbinom(0:10, 10, 1/3), type="h", lwd=2)

# sample from a multi-roll d4 distribution
sample(dusd1(1:4, 5), 20, replace=TRUE)
plot(ecdf(dusd1(1:4, 5)))

tt <- dusd2(xi=rep(1, 4), n=3)
plot(tt)
tt <- tt/sum(tt)
rr <- replicate(50000, sample(names(tt), prob=tt))
barplot(apply(rr, 1, table), beside=TRUE)

# distribution of the sum of three d6 rolls
plot(table(dusd1(xr=1:6, 3)))
plot(dusd2(xi=rep(1, 6), n=3))

# d6 die with faces 2, 3, 5, 7, 11, 13 (prime numbers)
plot(table(dusd1(xr=c(2, 3, 5, 7, 11, 13), 3)))
```

```

# Probability of getting 7 or 8 with an 8-sided die in n out of 5 throws
l <- 6/8
h <- 1-l
d <- as.dice(c(l, h), bix=0)

dusd2(d, 5)
# need integer "probabilities" for dusd1
table(dusd1(d*4, 5))/(4^5)
# or an equivalent die
table(dusd1(c(0, 0, 0, 1), 5))/(4^5)

# Loaded die
p <- c(0.5, 1, 1, 1, 1, 1.5); sum(p)
plot(dusd2(xi=p, n=2))

# A loaded die with prime number faces
s <- vector(length=13)
s[c(2, 3, 5, 7, 11, 13)] <- c(0.5, 1, 1, 1, 1, 1.5)
plot(dusd2(xi=s, n=3))

# tricky to do with dusd2
plot(table(dusd1(xr=c(0.1105, 2, exp(1)), 10)))

# Demonstrating CLT
# dusd1 struggles with many iterations
# remember it returns an array of size length(xr)^n
plot(table(dusd1(xr=c(1, 2, 9), 12)))

s <- vector(length=9)
s[c(1, 2, 9)] <- 1
plot(dusd2(xi=s, 12, round=9)) # much quicker
plot(dusd2(xi=s/sum(s), 12)) # for frequencies instead of counts

# Impossible with dusd1
clt <- dusd2(xi=s, 15, round=9)
plot(clt, lwd=0.5, col="#00000088")

# small floating-point errors from convolution.
tail(dusd2(xi=s, 15))

# dusd2 isn't always quicker
## Not run:
plot(table(dusd1(xr=c(1, 220, 3779), 12)), lwd=1)
s2 <- vector(length=3779)
s2[c(1, 220, 3779)] <- 1
plot(dusd2(xi=s2, 12, round=8), lwd=1)

# making sure the length of xi is highly composite (or more precicely 'smooth')
# improves speed
# 3779 is prime, 3780 == 2*2*3*3*3*5*7
s3 <- vector(length=3780)
s3[c(1, 220, 3779)] <- 1
plot(dusd2(xi=s3, 12, round=9), lwd=1)

## End(Not run)

```

Elo_rating

Elo rating

Description

Calculate updated Elo ratings based on existing rating and results from matches

Usage

```
elo_upd(ra, rb, score, k = 16, sum = length(rb) > 1)
```

```
elo_upd_pw(ra, rb, score, k = 16)
```

Arguments

ra	rating of player A
rb	rating of player B
score	score respective to player A. Numeric or character; 1, 0.5, 0; win, tie, loss
k	a measure of how big the correction should be
sum	calculate the new rating based on sum of scores and opponents ratings

Examples

```
# as in example from
# https://en.wikipedia.org/wiki/Elo_rating_system#Mathematical_details
# per 2019-10-04
ra <- 1613
rb <- c(1609, 1477, 1388, 1586, 1720)
score <- c(0, 0.5, 1, 1, 0)
elo_upd(ra, rb, score, k=32, sum=FALSE)
elo_upd(ra, rb, score, k=32, sum=TRUE)

elo_upd_pw(c(1400, 1500), c(1300, 1400), c("w", "t"))

results <- read.table(text="
Player1  Player2  Result
Alice    Bob       Win
Charlie  Dennis    Loss
Elena    Frank     Loss
June     Rashida   Tie", header=TRUE, stringsAsFactors=FALSE)

scores <- read.table(text="
Player  Score
Alice   1150
Charlie 1150
Frank   1150
Bob      800
Dennis  800
Elena   800
June    900
Rashida 1100", header=TRUE, stringsAsFactors=FALSE)
```

```

rownames(scores) <- scores$Player

r2 <- results
r2[,1:2] <- scores[as.matrix(r2[,1:2]), 2]

r2u <- elo_upd_pw(r2)

scores.new <- data.frame(Score=c(r2u))
rownames(scores.new) <- as.matrix(results[,1:2])

scores.new <- round(scores.new[rownames(scores),, drop=FALSE])
scores.new$diff <- scores.new$Score - scores$Score
scores.new

```

entropy

Information entropy

Description

Computes the information entropy (also called Shannon entropy) of a set of discrete values, or a tabulated such set.

Usage

```

entropy(x, ...)

## S3 method for class 'table'
entropy(x, base = 2, ...)

## S3 method for class 'data.frame'
entropy(x, base = 2, ...)

## S3 method for class 'matrix'
entropy(x, base = 2, ...)

## Default S3 method:
entropy(x, base = 2, ...)

```

Arguments

x	a vector, table, data.frame or matrix. In the case of table, data.frame and matrix each row is treated as a separate set of counts or proportions, with columns representing species, types, categories etc.
...	further arguments passed to methods
base	the log base to be used.

Examples

```

entropy(c(5, 5, 4, 4, 2, 3, 5)) # default is unit bits
entropy(c(5, 5, 4, 4, 2, 3, 5), base=exp(1)) # unit nats

```



```

entropy(rep(1:4, 1:4), 4)
entropy(rep(1:4, 1), 4)

entropy(as.factor(c(1, 1, 2, 3, 4, 4)))
entropy(as.character(c(1, 1, 2, 3, 4, 4)))

mtctab <- table(mtcars$cyl, mtcars$carb)
entropy(mtctab, 6)

xx <- data.frame(bee=c(0, 0, 1, 2, 3, 2, 0, 3),
                  wasp=c(1, 3, 2, 0, 1, 1, 2, 1),
                  fly=c(1, 2, 4, 2, 1, 0, 1, 0),
                  beetle=c(1, 0, 0, 1, 2, 2, 0, 2),
                  butterfly=c(0, 0, 0, 0, 3, 1, 0, 1))

entropy(xx)

```

every_nth	<i>Select every n'th element</i>
-----------	----------------------------------

Description

Select every second, third, fourth etc. element (or slice/hyperplane) of an object

Usage

```

every_nth(...)

## Default S3 method:
every_nth(x, n = 2, start = 1, ...)

## S3 method for class 'matrix'
every_nth(x, n = 2, start = 1, margin = 1, ...)

## S3 method for class 'array'
every_nth(x, n = 2, start = 1, margin = 1, ...)

## S3 method for class 'data.frame'
every_nth(x, n = 2, start = 1, margin = 1, ...)

## S3 method for class 'list'
every_nth(x, n = 2, start = 1, ...)

```

Arguments

...	further arguments passed to methods
x	an object to be selected from
n	selection "step size"
start	integer in [1:n] specifying the start of selection
margin	what margin to select along

Examples

```

m <- matrix(1:64, 8)
every_nth(m, n=3, start=3, margin=2)

d <- data.frame(A=1:8, B=2:9, Q=letters[rep(1:3, length.out=8)])
every_nth(d, start=2)

a <- array(1:6^4, rep(6, 4))
every_nth(a)

l <- list(a=1:3, b=2:6, c=8:5, d=9:7, e=list(ea=1:2, eb=1), f=2:6)
every_nth(l, n=2, start=2)

```

expand	<i>Expand</i>
--------	---------------

Description

Expand a "table", a "table"-like object, or a list of "table"-like objects

Usage

```
expand(x, ...)
```

Arguments

x	an object to be expanded
...	further arguments passed to or from methods

Value

A vector with values and their repetitions specified by x

See Also

[dice](#), [table](#)

Examples

```

x <- c(4, 2, 2, 2, 3, 3, 2, 4, 6, 6)
(xt <- table(x))
(xd <- dice(x))

expand(xt)
expand(xd)

expand(list(xt, xd, x))

xn <- as.table(1:4)
names(xn) <- LETTERS[1:length(xn)]
expand(xn)

```

explode	<i>Explode object</i>
---------	-----------------------

Description

Presents an R object in an exploded, or expanded, form

Usage

```
explode(x, indent = 2)
```

Arguments

x	an R object, or a character string describing an R object
indent	how many spaces for indentation (and exdentation) at each level

Details

If x is an R object it is first deparsed and converted into a character string describing the object. This string is then unwrapped, or exploded, according to these rules: newline and exdentation after each open parenthesis, newline and indentation after each close parenthesis, and newline after each comma. Parentheses and commas forming part of character strings are ignored.

Value

An exploded representation of the object is printed to console, and returned invisibly. The output is in most cases a complete and reproducible representation of the object, similarly to `dput`, but less compact and more revealing of its inner structure.

See Also

[dput](#), [dput2](#)

Examples

```
xc <- 'list(v=1, A=c("abv", "bom"), B=c(1:3, 31, 28), list("foo", "bar", 1))'
explode(xc)

x1 <- list(0=NA, R=list(j=1:3, h="(a)", q=structure(list(a=1:2, b=c("A, K",
  "B, L")), class="data.frame", row.names=c(NA, -2L))), N=1, L=FALSE)
explode(x1)

mt <- 'coplot(mpg ~ disp | as.factor(cyl), data = mtcars,
  panel = panel.smooth, rows = 1)'
explode(mt)
```

factorise

*Factorise***Description**

Find the prime factors of a given integer

Usage

```
factorise(x)
```

Arguments

x integer

Value

An integer vector

See Also

[factors](#) for unique prime factors or all integer factors

Examples

```
factorise(320)
factorise(2 * 2 * 2 * 3 * 3 * 5)

prod(factorise(5641324))

## Not run:
factorise(nextn(60000000, c(2, 3)))
factorise(72*999983)

## End(Not run)
```

factors

*Factors***Description**

Find the integers a given number is divisible by

Usage

```
factors(x, prime = FALSE)
```

Arguments

x an integer
prime should only prime factors be returned?

Value

An integer vector

Note

The trivial factors 1 and x itself are not included.

See Also

[factorise](#) for prime factorisation

Examples

```
factors(210)
factors(210, prime=TRUE)
```

file_ext	<i>File extension</i>
----------	-----------------------

Description

Separate file name and extension from a file path

Usage

```
file_ext(x)

file_name(x)

file_name_ext(x)
```

Arguments

x a character vector

Details

If the supplied file name has several extensions, f.ex. like foobar.tar.bz, only the last extension will be considered.

Value

file_ext returns the file extension of each file path. file_name returns the file name, no extension, of each file path. file_name_ext returns both name and extension, but arranged in separate columns of a matrix.

See Also

[basename](#)

Examples

```
x <- c("/hg/.git.tar.gz", "ff/hg/hh.pdf", "git", ".History", ".History.log")

file_ext(x)
file_name(x)
file_name_ext(x)
```

fingerknit

Render R example

Description

Render example from R code stored on the clipboard

Usage

```
fingerknit()
```

Details

Similar to `reprex::reprex()` or `knitr::spin(text=, envir=new.env(), report=FALSE)`, but stripped down to the very basics. Input is plain valid R code taken from the clipboard. It is run in a fresh environment and both commands and results are captured. Commands are kept as is, but results are commented out. Instead of using three backticks to indicate code for markdown, each line has four whitespaces prepended. `fingerknit` output is also valid R code.

Value

The clipboard is used for both input and output, but the output is also returned invisibly as a character string. Warnings and errors are not captured, but printed to console as normal. If an error is encountered nothing is returned and the clipboard data remains unchanged.

fitrange

Fit to a range

Description

Linearly shift and scale a numeric vector so that it fits to a given range.

Usage

```
fitrange(x, lower = -1, upper = 1)
```

Arguments

<code>x</code>	a numeric vector
<code>lower</code>	the lower bound of the new vector
<code>upper</code>	the upper bound of the new vector

See Also[norma](#)**Examples**

```

range(fitrangle(runif(10, -2, 1.5), 0, 1))

fitrange(c(2, 3, 5, 7, 4), 1, 0)
# same, but without warning
1 - fitrange(c(2, 3, 5, 7, 4), 0, 1)

```

flatten	<i>Flatten list</i>
---------	---------------------

Description

Flatten a (nested) list to a list of its leaves

Usage

```
flatten(x, flatten.df = FALSE, keep.order = TRUE)
```

Arguments

x	a list object
flatten.df	should data.frames also be flattened?
keep.order	keep the order of the original list, same as seen when using str

Details

The nodes of the supplied list is traversed from root to leaf and successively unlisted until no lists are left (except possibly for data.frames).

Value

A single level list of x's leaves.

Examples

```

x1 <- list(
  0=NA,
  R=list(
    j=1:3,
    h="(a)",
    q=data.frame(
      a=1:2,
      b=c("A", "K", "B", "L"),
      stringsAsFactors=FALSE
    )
  ),
  N=1,
  L=FALSE
)

```

```
flatten(x1, flatten.df=TRUE, keep.order=FALSE)
flatten(x1, flatten.df=TRUE, keep.order=TRUE)
str(x1)
```

gcd	<i>Greatest common divisor</i>
-----	--------------------------------

Description

Find the largest integer, that when two numbers are divided by it, returns an integer in both cases

Usage

```
gcd(x, y)
```

Arguments

x, y integers whose greatest common divisor is to be found

Examples

```
gcd(sequence(10:16), rep(10:16, 10:16))
```

incdiff	<i>Increase difference</i>
---------	----------------------------

Description

Rearrange a sorted numeric sequence so that the difference between subsequent elements is increased

Usage

```
incdiff(x, step = 2)
```

Arguments

x a numeric sequence
 step how long a step the difference is considered for.

Details

With step=2 (default) only the difference between immediate neighbours are considered; the difference between every second element will remain small, or rather reduced, compared to the original sequence. With step=3 say, differences of both lag 1 and 2 is increased, but the difference of lag 1 will be less than if a step of 2 was used.

Examples

```

x <- 1:100
diff(x)

diff(incdiff(x, 2))
diff(incdiff(x, 3))

diff(incdiff(x, 2), 2)
diff(incdiff(x, 3), 2)

# incdiff will introduce a periodicity equal to the step length
acf(incdiff(x, 10))

# useful for making a sequence of colours more distinct
y <- seq(0.4, 1, l=18)
cols1 <- hsv(y, 1, y)
cols2 <- hsv(y, 1, incdiff(y, 3))

plot(y, col=cols1, pch=16, cex=5, ylim=c(0.4, 1.5))
points(y+0.5, col=cols2, pch=16, cex=5)

```

indexvalue

*Index-value representation of arrays***Description**

Represent an array as columns of dimensional indices and value

Usage

```
indexvalue(x, reverse = FALSE)
```

Arguments

x	an array or something that can be coerced into an array
reverse	logical; convert from Index-value representation to regular array representation?

Details

An n-dimensional array will be unfolded to a n+1-column data.frame where the first n columns represent the indices of the n dimensions, and the last column gives the value found at each index tuple. The reverse process can also be performed.

See Also

[latin_sq](#)

Examples

```

arr <- array(1:(2*3*4), dim=c(2, 3, 4))
arr.is <- indexvalue(arr)

# can be used to permute an array
indexvalue(arr.is[,c(2, 1, 3, 4)], rev=TRUE)
aperm(arr, c(2, 1, 3))

# can interpret values (symbols) as dimensional indices and permute them as well
arr2 <- array(rep(1:6, 4), dim=c(2, 3, 4))
arr2.is <- indexvalue(arr2)
indexvalue(arr2.is[,c(1, 2, 4, 3)], rev=TRUE)

# a latin square will produce an "orthogonal array"
set.seed(1)
lsq <- latin_sq(5)
iv <- indexvalue(lsq)
iv

# any permutation of a latin square is also a latin square
indexvalue(iv[, c(1, 3, 2)], reverse=TRUE)

```

intsect

*Intersect***Description**

Performs set intersection on a list of vectors

Usage

```
intsect(x)
```

Arguments

x list of sets (vectors of same mode or factors)

Details

The intersection between the sets in the list is found. This means no duplicate values are returned, whether or not there were any in the input.

Value

A vector of same mode as input, or a single factor object if input was factor.

Examples

```

intsect(list(0:6, c(2, 4, 6, 8), 3:8))

fc <- factor(LETTERS[sample(1:5, 20, rep=TRUE)])
fc1 <- split(fc, sample(1:3, 20, rep=TRUE))

intsect(fc1)

```

in_range	<i>Test if values is in a given range</i>
----------	---

Description

Checks either whether both extrema are in the given range, or if each individual value is.

Usage

```
in_range(x, lower, upper, inc = c(TRUE, TRUE), na = NA)
```

Arguments

x	any atomic or vector-like object
lower	lower range
upper	upper range
inc	logical vector of length one or two; should the lower and upper ranges, respectively, be considered inclusive?
na	should NAs in x return TRUE, FALSE, NA, or something else?

Examples

```
in_range(c(1:3, NA), 1, 4, na=NA)

in_range(1:4, 1, 4)
in_range(matrix(1:4, 2), 1, 4)

in_range(1:4, 1, 4, inc=1:0)
in_range(1:4, 1, 4, inc=0:1)
in_range(1:4, 1, 4, inc=0)

x <- as.Date(0:3, origin="2000-01-01")
in_range(x, "2000-01-01", "2000-01-04")

in_range(letters[1:4], "a", "d", inc=1:0)
in_range(letters[1:4], "a", "da", inc=1:0)

# no upper range
in_range(c(10^rnorm(9), NA), 0, NA)
in_range(c(10^rnorm(9), NA), 0, NA, na=TRUE)
```

is_coprime	<i>Coprimality check</i>
------------	--------------------------

Description

Test whether two integers are coprime, that is, have no factors in common

Usage

```
is_coprime(x, y)
```

Arguments

`x`, `y` integers to be tested for coprimality

Value

A logical vector

Examples

```
is_coprime(sequence(10:16), rep(10:16, 10:16))
is_coprime(2*3*5*7, 11*13)
```

<code>is_prime</code>	<i>Primality check</i>
-----------------------	------------------------

Description

Test integers for whether they are prime or not

Usage

```
is_prime(x)
```

Arguments

`x` vector of integers

See Also

[primes](#)

<code>keep_finite</code>	<i>Keep finite values</i>
--------------------------	---------------------------

Description

Remove NAs codeNaNs and codeInfs from data

Usage

```
keep_finite(x, ...)

## Default S3 method:
keep_finite(x, ...)

## S3 method for class 'matrix'
keep_finite(x, margin = 1, keep = c("any", "complete"), ...)

## S3 method for class 'data.frame'
keep_finite(x, margin = 1, keep = c("any", "complete"), ...)
```

Arguments

x	a vector or matrix
...	further arguments passed to methods
margin	if x is matrix, which margin to keep finites by
keep	if x is matrix, keep rows/columns with any finite values, or keep only complete rows/columns.

Value

If x is a matrix and margin is 1 or 2, a matrix is returned. Else a vector.

Examples

```
m1 <- matrix(c(10, 20, 30, 43,
               10, NA, 32, 50,
               NA, NA, NA, NA,
               13, 22, 70, 81,
               NA, 29, NA, 41), 5, byrow=TRUE,
             dimnames=list(letters[1:5], LETTERS[1:4]))

keep_finite(m1)
matplot(keep_finite(apply(m1, 2, sort, na.last=TRUE)), type="l")

m1[complete.cases(m1),]
keep_finite(m1, 1, "c") #same
keep_finite(m1, 2, "complete") #no complete columns

m1.df <- as.data.frame(t(m1))
keep_finite(m1.df, 2, "complete")
```

lag_vector	<i>Lag an arbitrary vector</i>
------------	--------------------------------

Description

Lag an arbitrary vector

Usage

```
lag_vector(x, k, type = c("cycle", "na.fill", "trim"), ...)
```

Arguments

x	vector to be lagged
k	integer vector specifying the number of lags
type	how to deal with non-overlapping sections
...	further arguments passed to methods

Examples

```
x <- 1:9

lag_vector(x, c(0, 1, -2, 3))
lag_vector(x, c(0, 1, -2, 3), "na")
lag_vector(x, c(0, 1, -2, 3), "trim")
```

latin_sq	<i>Latin square</i>
----------	---------------------

Description

Generate latin squares, either randomly or ordered

Usage

```
latin_sq(n, random = TRUE, reduce = TRUE)
```

Arguments

n	integer; number of unique values (aka. symbols)
random	logical; should the square be generated randomly?
reduce	logical; should the square be in reduced form?

Details

Computation time increases rapidly with n. On my computer generating a random square with n=12 takes about ten minutes, marking the upper limit of practicability, or even stretching it a little. A latin square in reduced form will have elements in the first row and the first column in a sorted order. By setting reduced=TRUE the first row and the first column will always be 1:n.

Value

A square integer matrix of size n^2

See Also

[indexvalue](#)

Examples

```
set.seed(1)
ls <- latin_sq(9, reduce=TRUE)
image(ls, col=randcolours(ncol(ls)))

# The more "classic" representation with latin capital letters
ls[] <- LETTERS[ls]
ls
```

markov_seq	<i>Discrete markov sequence</i>
------------	---------------------------------

Description

Generate a random discrete markov sequence

Usage

```
markov_seq(n = 100, tmat = rbind(1:3, 3:1, 2:0), init = 1)
```

Arguments

n	length of the sequence
tmat	a transition matrix
init	the initial state

Examples

```
m <- matrix(c(0.5, 0.3, 0.2,
              0.2, 0.6, 0.2,
              0.2, 0.3, 0.5), 3, byrow=TRUE)

set.seed(1)
ms <- markov_seq(n=1000, tmat=m)

colMeans(m)
prop.table(table(ms))
round(prop.table(table(head(ms, -1), tail(ms, -1), dnn=c("n", "n+1")), 1), 2)
```

math_constants	<i>Mathematical constants</i>
----------------	-------------------------------

Description

Various mathematical constants available as global variables

Format

An object of class `numeric` of length 1.

Details

e Euler's number
 pi Archimedes' number, the circle constant
 phi Golden ratio
 feig1 Feigenbaum's first constant, δ ; bifurcation velocity
 feig2 Feigenbaum's second constant, α ; reduction parameter
 eu.ma Euler–Mascheroni constant
 khin Khintchine's constant
 glai.kin Glaisher-Kinkelin constant

math_constants_char	<i>High precision mathematical constants</i>
---------------------	--

Description

Character strings representing various mathematical constants to ~100 decimal points

Format

An object of class character of length 1.

Details

e.char Euler's number
 pi.char Archimedes' number, the circle constant
 phi.char Golden ratio
 feig1.char Feigenbaum's first constant, δ ; bifurcation velocity
 feig2.char Feigenbaum's second constant, α ; reduction parameter
 eu.ma.char Euler–Mascheroni constant
 khin.char Khintchine's constant
 glai.kin.char Glaisher-Kinkelin constant

means	<i>Generalized means</i>
-------	--------------------------

Description

Harmonic, geometric, quadratic, cubic, power and Lehmer means.

Usage

```
harm(x, na.rm = TRUE)

geom(x, zero.rule = c("1p", "rm", "1"), na.rm = TRUE)

quad(x, na.rm = TRUE)

cubi(x, na.rm = TRUE)

powr(x, p = 1.5, na.rm = TRUE)

lehm(x, p = 2, na.rm = TRUE)
```


Arguments

x	numeric vector of values whose *mean is to be computed
na.rm	logical; should NA values be removed? (default TRUE)
zero.rule	for the geometric mean, how should zeros be dealt with? Add one before, and subtract one after the calculation (see <code>lop1p</code>), remove all zeros, or replace all zeros with 1.
p	exponential power. For the power mean $p=-1$, $p=2$ and $p=3$ gives the harmonic, quadratic and cubic means, respectively. For the Lehmer mean $p=0$, $p=1$ and $p=2$ gives the harmonic, arithmetic and contraharmonic means, respectively.

Notice

For some of these means zeros and/or negative values are undefined, or make otherwise little sense in context. Workarounds are given for the geometric mean, but if you end up using it on data ≤ 0 , the wise call would be to reconsider whether using a geometric mean really makes sense in that case.

See Also

[central.tendency](#)

Examples

```
fun1 <- substitute(c(harm, geom, mean, quad, cubi))

x1 <- list(c( 1, 2, 3, 5),
          c(-1, 1, 2, 3, 5),
          c( 0, 1, 2, 3, 5),
          c(-1, 0, 1, 2, 3, 5))

m <- sapply(x1, function(x) sapply(eval(fun1), function(f) f(x)))
rownames(m) <- as.character(fun1)[-1]
colnames(m) <- c("posi", "1neg", "zero", "1ngz")
round(m, 3)

harm(x1[[1]]); powr(x1[[1]], -1); lehm(x1[[1]], 0)

y <- c(0, 1, 5, 0, 6, 5, 9)

geom(y, zero.rule="1p")
geom(y, zero.rule="rm")
geom(y, zero.rule="1")
```

merge_multiple

Merge multiple data.frames

Description

Merge multiple data.frames

Usage

```
merge_multiple(
  x,
  by,
  all = FALSE,
  sort = TRUE,
  incomparables = NULL,
  include = NULL,
  exclude = NULL
)
```

Arguments

<code>x</code>	a list of data.frames with at least one column in common
<code>by</code>	name of the column to be merged by, by default the full intersect of column names between data.frames
<code>all</code>	include all rows, including those with no match
<code>sort</code>	sort the output on the by column(s)
<code>incomparables</code>	values which cannot be matched.
<code>include, exclude</code>	numeric, logical or character vector specifying which columns to include in or exclude from the merge

Details

If there are duplicate columns that aren't being used to merge by, one of two things will happen. If the parent data.frames of the duplicate columns are named, then that name will be appended to the relevant column names. If the data.frames aren't named, then the data.frames indices in the parent list are appended to the relevant column names.

Inclusion and exclusion are performed in sequence, so that if both `include` and `exclude` are specified, `exclude` acts on the result from `include`.

Examples

```
dtf1 <- data.frame(ast=1:4, bar=1:4, kat=c("A", "B", "C", "D"))
dtf2 <- data.frame(ast=1:6, bar=1:6, jun=9:4)
dtf3 <- data.frame(ast=2:6, bar=2:6, kat=c("A", "B", "C", "D", "E"))
dtf4 <- data.frame(ast=3:4, bar=3:4)
dtf5 <- data.frame(ast=1:-3, bar=0:4, git=0:4)

ll <- list(d1=dtf1, d2=dtf2, dtf3, A=dtf4, dtf5)

merge_multiple(ll, by="bar")
merge_multiple(ll, by="bar", all=TRUE, include=1:2)
merge_multiple(ll, by="bar", all=TRUE, exclude="kat")
merge_multiple(x=ll, by=c("bar", "ast"), all=TRUE)
```

multidensity	<i>Plot multiple kernel density estimates</i>
--------------	---

Description

Plot multiple kernel density estimates in the same window, together with a legend

Usage

```
multidensity(
  x,
  main,
  xlab = "",
  ylab = "Density",
  xlim,
  ylim,
  col = 1:9,
  lty = 1:2,
  lwd = 1,
  add = FALSE,
  frame.plot = TRUE,
  legend = TRUE,
  x.legend = "topleft",
  y.legend = NULL,
  bty = "o",
  box.col = "#FFFFFF00",
  bg.legend = "#FFFFFFAA",
  cex.legend = 0.7,
  x.intersp = 1,
  y.intersp = 1.5,
  inset = 0,
  xpd.legend = NA,
  horiz = FALSE,
  ...
)
```

Arguments

<code>x</code>	a list or data.frame of numeric values
<code>main</code>	a main title for the plot. Defaults to the call made to density
<code>xlab</code> , <code>ylab</code>	labels for the x and y axes
<code>xlim</code> , <code>ylim</code>	the x and y limits of the plot
<code>col</code> , <code>lty</code> , <code>lwd</code>	the line colours, types and widths for lines appearing in plot and legend
<code>add</code>	if TRUE, add to the current plot
<code>frame.plot</code>	an integer indicating whether a box should be drawn around the plot before the legend (1), after the legend (2), or not at all (0). Logical values are coerced to integer, so TRUE implies 1, and FALSE implies 0
<code>legend</code>	logical; if TRUE (the default) a legend is included with the plot

<code>x.legend, y.legend</code>	the x and y co-ordinates to be used to position the legend. They can be specified by keyword or in any way which is accepted by <code>xy.coords</code>
<code>bty</code>	legend box type
<code>box.col</code>	line colour for the legend box
<code>bg.legend</code>	background colour for the legend box
<code>cex.legend</code>	character expansion faftor for legend
<code>x.intersp, y.intersp</code>	horizontal and vertical character interspacing for legend
<code>inset</code>	the legends inset distance from the margins as a fraction of the plot region
<code>xpd.legend</code>	the value of <code>xpd</code> to be used while drawing the legend
<code>horiz</code>	logical; if TRUE, set the legend horizontally rather than vertically
<code>...</code>	further arguments passed to <code>density</code>

Value

An invisible list of the "density" objects the plot is based on.

See Also

[density](#), [ahist](#)

Examples

```
set.seed(1)
dl <- list("Unif-1"=runif(80, -2.1, 2.1),
          "Unif-2"=runif(70, -1.5, 1.5),
          "Normal-1"=rnorm(50, 0, 0.866),
          "Normal-2"=rnorm(90, 0, 1))

# sqrt((sd^2)*12) # sd to unif range

md <- multidensity(dl)
head(md, 2)

multidensity(dl, adj=1.2, x.leg="topright", frame=FALSE, inset=-0.02, lty=1)
multidensity(dl, x.legend="top", horiz=TRUE, cex.legend=0.5,
             inset=-0.05, bg.legend="white")
```

narm

Remove NAs

Description

Remove NAs from vector or matrix

Usage

```
narm(x, ...)
```

```
## Default S3 method:
narm(x, ...)
```

```
## S3 method for class 'matrix'
narm(x, margin = 1, keep = c("any", "complete"), ...)
```

```
## S3 method for class 'data.frame'
narm(x, margin = 1, keep = c("any", "complete"), ...)
```

Arguments

<code>x</code>	a vector or matrix
<code>...</code>	further arguments passed to methods
<code>margin</code>	if <code>x</code> is matrix, which margin to remove NAs by
<code>keep</code>	if <code>x</code> is matrix, keep rows/columns with any non-NA values, or keep only complete rows/columns.

Value

If `x` is a matrix and `margin` is 1 or 2, a matrix is returned. Else a vector.

Examples

```
m1 <- matrix(c(10, 20, 30, 43,
               10, NA, 32, 50,
               NA, NA, NA, NA,
               13, 22, 70, 81,
               NA, 29, NA, 41), 5, byrow=TRUE,
             dimnames=list(letters[1:5], LETTERS[1:4]))
```

```
narm(m1)
```

```
matplot(narm(apply(m1, 2, sort, na.last=TRUE))), type="l")
```

```
m1[complete.cases(m1),]
narm(m1, 1, "c") #same
narm(m1, 2, "complete") #no complete columns
```

```
m1.df <- as.data.frame(t(m1))
narm(m1.df, 2, "complete")
```

 norma

Normalize

Description

Linearly shift and scale a numeric vector so that it has a given range, about a given centre.

Usage

```
norma(x, c = 0, r = 2)
```

Arguments

x	a numeric vector
c	the centre (as in the midrange) for the new vector
r	the range of the new vector

See Also

[fitrange](#)

Examples

```
range(norma(runif(9, -2, 0.1), 0, 2))
```

pairwise

Apply function to columns/elements pairwise

Description

Pairwise application of a function to the columns of a matrix/data.frame or elements of a list

Usage

```
pairwise(x, FUN, ..., comm = FALSE)
```

Arguments

x	a matrix or data.frame
FUN	any function that takes two vectors as input and returns a single value
...	further arguments passed to FUN
comm	logical; is FUN commutative? If true, only the lower triangle, including the diagonal, is computed

Value

An $n \times n$ square matrix with n the number of columns of x.

See Also

[similarity](#) for a few more examples

Examples

```
dtf <- data.frame(aa=c(1, 1, 2, 2, 3, 2, 4),
                  bb=c(1, 1, 2, 3, 3, 3, 4),
                  cc=c(3, 3, 2, 1, 1, 1, 1),
                  dd=c(1, 2, 2, 2, 1, 1, 2))

# Root Mean Square Deviation
pairwise(dtf, function(x, y) sqrt(mean((x-y)^2)))

# using with cor.test() to accompany cor()
pv <- pairwise(dtf, function(x, y) cor.test(x, y)$p.val)
pvn <- 6^(1.1-pv)-5
pvn[pvn<1] <- 1

set_mar(1, 1, 1, 1)
plot(0, xlim=c(0.5, 4.5), ylim=c(0.5, 4.5), cex=0, ann=FALSE, xaxt="n", yaxt="n")
text(rep(1:4, 4), rep(4:1, each=4), t(round(cor(dtf), 2)), cex=pvn,
     col=c("black", "darkgrey")[(pv>0.1)+1])
```

pcamean	<i>PCA mean</i>
---------	-----------------

Description

Takes the average of several PCA objects

Usage

```
pcamean(...)
```

Arguments

... prcomp, princomp or factanal objects, or a single list of such objects

Details

I don't know if this kind of calculation has any sort of merit. It was written more as an impromptu challenge than as a solution to any problem

See Also

[prcomp](#), [princomp](#), [factanal](#)

Examples

```
xx <- data.frame(bee=c(0, 0, 1, 2, 3, 2, 0, 3),
                 wasp=c(1, 3, 2, 0, 1, 1, 2, 1),
                 fly=c(1, 2, 4, 2, 1, 0, 1, 0),
                 beetle=c(1, 0, 0, 1, 2, 2, 0, 2))

set.seed(1)
r <- 1000
```

```

xxs <- replicate(r, {
  xx$random <- sample(c(0:1, 0:4), 8, r=TRUE)
  xx
}, simplify=FALSE)

xxm <- Reduce("+", xxs) / r
xxl <- lapply(xxs, princomp)

biplot(pcamean(xxl))
biplot(princomp(xxm))

```

plot.histogram	<i>Plot histogram object</i>
----------------	------------------------------

Description

A very minor modification of `graphics::plot.histogram`.
Only difference is that `lwd` now specifies the width of the histogram bars' outline.

See Also

[plot.histogram](#), [plot.stl](#), [ahist](#)

plot.stl	<i>Plot stl object</i>
----------	------------------------

Description

A very minor modification of `stats::stl`.
Only difference is that the distance between the plotting window and the x and y labels is set by `par("mgp")[1]`, as it is for regular plots.

See Also

[plot.stl](#), [plot.histogram](#)

primes	<i>Prime number generator</i>
--------	-------------------------------

Description

Prime generator based on the sieve of Eratosthenes

Usage

`primes(n)`

Arguments

`n` integer; all prime numbers up to this will be returned

Details

Effective for primes up to ~100,000,000.

On my lightweight laptop: 1e7 -> 0.32s, 5e7 -> 1.7s, 1e8 -> 3.7s, 2e8 -> 7.6s, 3e8 -> 15s

Source

<https://stackoverflow.com/questions/3789968/generate-a-list-of-primes-up-to-a-certain-number/3791284#3791284>

See Also

[is_prime](#)

quartz.png

Save as PNG

Description

Save the contents of the current Quartz window as PNG file

Usage

```
quartz.png(file = "%Y%m%d_%H", width = 550, dir, force = FALSE)
```

Arguments

<code>file</code>	file name. If it contains any <code>"%"</code> it is passed on as a format string to <code>format(Sys.time(), file)</code> . A <code>.png</code> file extension is added automatically.
<code>width</code>	pixel width of the PNG file
<code>dir</code>	directory to save to. Defaults to current working directory
<code>force</code>	force overwriting of existing file with same name. By default duplicate path names are resolved by appending <code>_N</code> , using successive integers, to the end of the file name.

Value

A PNG file is written to disk and a message is written to the console, giving the new file's path and pixel dimensions. The file path is also returned invisibly.

Examples

```
## Not run:
set.seed(1)
i_h100 <- round(runif(100, 2, 30), 2)
i_cd <- rexp(100, 1/i_h100)
mydata <- data.frame(i_cd, i_h100)

mydata$i_h100_2m <- cut(mydata$i_h100, seq(2, 30, by=2))

i_cd_2m <- aggregate(i_cd ~ i_h100_2m, mydata, mean)

set_mar(x=2.5)
plot.default(i_cd_2m, xaxt="n", main="Groupwise means", xlab="", cex.main=1)
axis(1, i_cd_2m[,1], as.character(i_cd_2m[,1]), cex.axis=0.6, las=2)

quartz.png()
p <- quartz.png("test", 550)
file.info(p)

## End(Not run)
```

quick_table

*Tabulate data***Description**

Quick and simple function for creating contingency tables

Usage

```
quick_table(x, na.rm = FALSE, order = c("frequency", "value", "none"))
```

Arguments

x	a vector or factor object
na.rm	should NAs be included
order	how should the results be ordered, if any?

Value

A data.frame with columns val (the original values and class of x) and freq (the count, or frequency, of each value in x, integer). The rows are sorted by frequency in descending order.

Examples

```
set.seed(1)
m <- sample(c(rep(NA, 5), rpois(45, 3)))
quick_table(m)

x <- LETTERS[c(2, 2, 2, 2, 3, 1, 1)]
quick_table(x, order="freq")
quick_table(x, order="value")
quick_table(x, order="none")
```

rainbowHCL	<i>HCL rainbow palette</i>
------------	----------------------------

Description

HCL version of rainbow. Create a vector of n contiguous colours by specifying a range of Hues, and fixed Chroma and Luminance

Usage

```
rainbowHCL(
  n,
  c = 100,
  l = 75,
  start = 0,
  end = max(1, n - 1)/n,
  alpha = 1,
  s = NULL,
  v = NULL
)
```

Arguments

<code>n</code>	number of colours
<code>c</code> , <code>l</code>	the ‘chroma’ and ‘luminance’ to be used to complete the HSV color descriptions
<code>start</code> , <code>end</code>	the hue in $[0,1]$ at which the rainbow begins/ends
<code>alpha</code>	the alpha transparency, a number in $[0,1]$, see argument <code>alpha</code> in hsv
<code>s</code> , <code>v</code>	‘saturation’ and ‘value’ passed to adjustcolorHSV . Overrides ‘chroma’ and ‘luminance’ if specified.

Examples

```
mat2grid <- function(x) {
  eg <- expand.grid(1:NCOL(x), NROW(x):1)
  gd <- data.frame(eg, c(t(x)), stringsAsFactors=FALSE)
  colnames(gd) <- c("x", "y", "z")
  gd
}

n <- 25
hcl0 <- rainbowHCL(n)
hcl1 <- rainbowHCL(n, c=150, l=85)
hcl2 <- rainbowHCL(n, s=1, v=1)
hsv0 <- rainbow(n)

cols <- rbind(hcl0, hcl1, hcl2, hsv0)

pos <- mat2grid(cols)
plot(pos[,1:2], pch=17, cex=3.5, col=pos[,3], ylim=c(0.5, 4.5))
```

randcolours*Random colours*

Description

Generate a randomly selected colour palette

Usage

```
randcolours(  
  n,  
  l = c(0.2, 0.9),  
  c1 = c(0, 1),  
  c2 = c(0, 1),  
  alpha = 1,  
  space = c("Luv", "Lab")  
)
```

Arguments

n	number of colours
l	lightness range
c1	colour channel one range
c2	colour channel two range
alpha	alpha channel range
space	should the parameters be interpreted as Luv or Lab components?

Details

The range of l, c1, c2 and alpha, will be interpreted as the wanted range of each colour component, whether their length is 1, 2, or more. Although they all should nominally lie within [0, 1], only alpha must do so to achieve a valid output. The others can exceed this range, at an increased risk of clipping.

Examples

```
set.seed(3)  
n <- 20  
plot(1:n, col=randcolours(n), pch=16, cex=5)
```

resolve_dup	<i>Resolve duplicate</i>
-------------	--------------------------

Description

Resolve duplicate names by appending successive integers

Usage

```
resolve_dup(x, candidates, ignore.extension = FALSE)
```

Arguments

x	character string; name to be resolved
candidates	character vector; possible duplicate names
ignore.extension	logical; append to the end of x, even if it has something that can be interpreted as an extension

Examples

```
x <- c("my.var", "aaa.png", "aaa.jpg", "aaa_1.png", "doc-folder")
resolve_dup("aaa.jpg", x)
resolve_dup("aaa.png", x)
resolve_dup("aaa_1.png", x)
resolve_dup("doc-folder", x)
resolve_dup("New Document", x)
resolve_dup("my.var", x, ignore.ext=TRUE)
x <- c(x, resolve_dup("aaa.png", x))
resolve_dup("aaa.png", x)
```

revert_par	<i>Revert par</i>
------------	-------------------

Description

Reverts par settings back to old.par

Usage

```
revert_par()
```

See Also

Other par_and_plot_margins_functions: [default_par\(\)](#), [set_mar\(\)](#)

rle2

*Run Length Encoding***Description**

Compute the lengths and values of runs of equal values in a vector

Usage

```
rle2(
  x,
  na.unique = FALSE,
  output = c("data.frame", "rle", "named vector", "lengths", "values")
)
```

Arguments

x	a numeric or character vector
na.unique	should every NA be considered unique?
output	what form of output

Value

Return value depends on output.

`data.frame` A data.frame with lengths and values columns

`rle` An object of class "rle"

`named vector` A vector of lengths with values as names

`lengths` The lengths as a single vector

`values` The values as a single vector

Examples

```
x <- c(NA, NA, 1, 2, 3, 3, NA, NA, NA, 2, 2, 2, NA, 1, 1, NA, NA)
rle2(x)

m <- matrix(c(
  0.7, 0.2, 0.1,
  0.2, 0.6, 0.2,
  0.1, 0.2, 0.7
), 3, byrow=TRUE)

set.seed(1)
y <- LETTERS[markov_seq(n=100, m)]
rle2(y, out="named")

# Same result as rle
rle2(x, na.unique=TRUE, output="rle")
rle(x)
```

```
# inverse.rle works as long as output is "rle"
inverse.rle(rle2(x, output="rle"))
```

seq_range	<i>Generate a sequence spanning a given range</i>
-----------	---

Description

Generate a sequence spanning a given range

Usage

```
seq_range(x, ..., spread)
```

Arguments

x	a single numeric, a range, or a sequence
...	further arguments passed to seq
spread	use spread_seq to spread out the range by a given factor

Details

If x is a single number, the range is interpreted to be $[0, x]$. If x is length two, the numbers are interpreted as the left and right extrema of the sequence interval. If x is longer than two, the sequence is based upon its range.

Examples

```
seq_range(c(1, 4), by=0.5)
seq_range(c(1, 4), by=0.5, spread=2)
seq_range(4)

x <- sample(1:10, 3)
seq_range(x)
```

set_mar	<i>Set plot margins</i>
---------	-------------------------

Description

Moves axis titles and labels closer to the plotting window and shrinks the margins

Usage

```
set_mar(x = 1.8, y = 1.8, main = 1, right = 1, cex.main = 1, ...)
```

Arguments

x	margin width for the x axis, default 2
y	margin width for the x axis, default 2
main	margin width for the main title, default 1
right	margin width for the right edge, default 1
cex.main	The magnification to be used for main titles relative to the current setting of cex, default 1
...	further arguments passed to par

Details

Old par settings are stored in `.old.par` before a call to `par` of the form `par(mar=c(x, y, main, right), mgp=c(1.9, 0.5, 0.5))` is made.

See Also

[par](#)
Other `par_and_plot_margins_functions`: [default_par\(\)](#), [revert_par\(\)](#)

Examples

```
ymse:::.old.par
get("old.par", envir=ymse::ymseEnv)
ls(envir=ymse::ymseEnv)

par(col.axis=2)
plot(1:4)

set_mar()
plot(1:4)

default_par()
plot(1:4)

revert_par()
plot(1:4)

ymse:::.old.par
head(get("old.par", envir=ymse::ymseEnv))
```

similarity	<i>Similarity measure</i>
------------	---------------------------

Description

Calculate the similarity between two character vectors based on a similarity matrix

Usage

```
similarity(x, y, sm = smat(x, y), sfun = sum, ...)
```


Arguments

<code>x</code>	a character vecor or two-column data.frame/matrix
<code>y</code>	a character vector. Ignored if <code>x</code> is data.frame/matrix
<code>sm</code>	a similarity matrix. By default a unit matrix
<code>sfun</code>	function used to summarise the elementwise similarities
<code>...</code>	further arguments passed to <code>sfun</code>

See Also

[smat](#)

Examples

```
# In its most basic form similarity() gives the Hamming distance
similarity(c(1, 0, 1, 0), c(1, 1, 0, 0))

# Symmetry not required.
bef <- c(1, 2, 3, 1, 2, 3, 1, 2, 3)
aft <- c(0, 2, 2, 1, 2, 2, 1, 1, 2)

# Here a decrease in value of 1 is considered
# more similar than an increase in value of 1.
sm1 <- t(structure(c(
  3, 0, 0, 0,
  2, 3, 0, 0,
  0, 2, 3, 0,
  0, 0, 2, 3),
  .Dim=c(4L, 4L),
  .Dimnames=list(c("0", "1", "2","3"), c("0", "1", "2", "3"))))

# Symmetric version
sm2 <- t(structure(c(
  3, 1, 0, 0,
  1, 3, 1, 0,
  0, 1, 3, 1,
  0, 0, 1, 3),
  .Dim=c(4L, 4L),
  .Dimnames=list(c("0", "1", "2","3"), c("0", "1", "2", "3"))))

similarity(bef, aft, sm1)
similarity(bef, aft, sm2)

# Pre-aligned fragments of insulin genes
data(insulin)

# Transition-transversion matrix
data(smt)

# Using pairwise() to run similarity() over all column pairs
pairwise(insulin, similarity, smt, sfun=mean)

# Imagined result from questionnaire
qu <- data.frame(
  Alice=c("happy", "sad", "angry", "unsure", "happy", "sad", "happy", "angry"),
```

```

Bob=c("happy", "sad", "angry", "angry", "happy", "angry", "angry", "sad"),
Charlie=c("sad", "sad", "unsure", "unsure", "happy", "sad", "angry", "sad"),
stringsAsFactors=FALSE
)

# Similarity matrix describing the relative similitudes of the moods
emsm <- as.matrix(read.table(text="
      happy  sad  angry unsure
happy    5    0    1    1
sad      0    5    2    1
angry    1    2    4    2
unsure   1    1    2    3",
header=TRUE))

pairwise(qu, similarity, sm=emsm/5, sfun=mean)

```

simple_loess

Simplified Local Polynomial Regression Fitting

Description

A simplified interface to the loess and predict.loess combo.

Usage

```

simple_loess(...)

## Default S3 method:
simple_loess(
  y,
  x = seq_along(y),
  xout = sort(x),
  span = 0.75,
  periodic = FALSE,
  ...
)

## S3 method for class 'data.frame'
simple_loess(df, xout = sort(df[, 1]), ...)

```

Arguments

...	further arguments passed to loess
y	the response values to be regressed
x	the regressor, by default an integer sequence along y
xout	values used for prediction, unless it is an integer of length 1. In that case xout specifies the number of equally spaced values on the interval of x to be used. By default the same as x
span	parameter controlling the degree of smoothing
periodic	should the input be treated as periodic?
df	a data.frame with x-values in the first column and y-vlues in the second

Value

A data.frame with columns *xout* and *y.predicted*

Examples

```
# Simple equally spaced vector
h <- c(-0.63, 0.2, -0.44, 1.6, 0.33, -0.74, -0.82, 0.29, 0.74, 0.58, -0.3)

plot(h)
lines(simple_loess(h))

# More complicated unequally space x-values
x <- c(4, 3, 2, 5, 6, 7, 9, 10, 12, 13, 14, 15, 16, 17, 18, 19)
y <- c(3, 2, 4, 5, 6, 5, 5, 3, 4, 7, 10, 10, 8, 9, 7, 8)

plot(x, y)
lines(simple_loess(y, x), col="gray40")
points(simple_loess(y=y, x=x, xout=5L), col=2, cex=2)
points(simple_loess(y=y, x=x, xout=17), col=3, cex=2)
points(simple_loess(y=y, x=x, xout=seq(8, 12, 0.3)), col=3, pch=16)
lines(simple_loess(y=y, x=x, xout=50L), col=4, lty=2)

# data.frame input
dtf <- data.frame(x, y)
simple_loess(dtf)
```

simple_table

Read a simple table

Description

Read tables given in more or less elaborate human-readable formats

Usage

```
simple_table(
  x,
  header = TRUE,
  rem.dup.header = header,
  na.strings = c("NA", "N/A"),
  stringsAsFactors = FALSE,
  ...
)
```

Arguments

<code>x</code>	a teble represented as a character string
<code>header</code>	are the table columns named? By default TRUE
<code>rem.dup.header</code>	remove duplicated headers.
<code>na.strings</code>	a character vector of strings which are to be interpreted as NA values

```

stringsAsFactors
    should character vectors be converted to factors? By default FALSE
...
    further arguments passed to read.table

```

Value

A `data.frame` containing a representation of the data.

Examples

```

x1 <- "
+-----+-----+-----+-----+-----+
|   Date   | Emp1 | Case | Priority | PriorityCountinLast7days |
+-----+-----+-----+-----+-----+
| 2018-06-01 | A    | A1   | 0        | 0 |
| 2018-06-03 | A    | A2   | 0        | 1 |
| 2018-06-02 | B    | B2   | 0        | 2 |
| 2018-06-03 | B    | B3   | 0        | 3 |
+-----+-----+-----+-----+-----+
"

x2 <- "
      Date   | Emp1 | Case | Priority | PriorityCountinLast7days

2018-06-01 | A    | A|1  | 0        | 0
2018-06-03 | A    | A|2  | 0        | 1
2018-06-02 | B    | B|2  | 0        | 2
2018-06-03 | B    | B|3  | 0        | 3

"

x3 <- "
Maths | English | Science | History | Class

0.1 | 0.2 | 0.3 | 0.2 | Y2

0.9 | 0.5 | 0.7 | 0.4 | Y1

0.2 | 0.4 | 0.6 | 0.2 | Y2

0.9 | 0.5 | 0.2 | 0.7 | Y1
"

x4 <- "
      Season | Team | W | AHWO
-----+-----+-----+-----+-----+
1 | 2017/2018 | TeamA | 2 | 1.75
2 | 2017/2018 | TeamB | 1 | 1.85
3 | 2017/2018 | TeamC | 1 | 1.70
4 | 2017/2018 | TeamD | 0 | 3.10
5 | 2016/2017 | TeamA | 1 | 1.49
6 | 2016/2017 | TeamB | 3 | 1.51
7 | 2016/2017 | TeamC | 2 | 1.90
8 | 2016/2017 | TeamD | 0 | N/A
"

```

```

x5 <- "
  A   T   G   C
-----
A | 6 | 0 | 4 | 0 |
|---:---:---:---
T | 0 | 6 | 0 | 4 |
|---:---:---:---
G | 4 | 0 | 6 | 0 |
|---:---:---:---
C | 0 | 4 | 0 | 6 |
-----

"

x6 <- "
-----
|date           |Material          |Description      |
|-----|-----|
|10/04/2013     |WM.5597394        |PNEUMATIC       |
|11/07/2013     |GB.D040790        |RING            |
|-----|-----|

-----
|date           |Material          |Description      |
|-----|-----|
|08/06/2013     |WM.4M01004A05     |TOUCHEUR        |
|08/06/2013     |WM.4M010108-1     |LEVER           |
|-----|-----|

"

lapply(c(x1, x2, x3, x4, x5, x6), simple_table)

```

smat

*Similarity matrix***Description**

Create a similarity matrix

Usage

```
smat(x, y, s, byrow = FALSE)
```

Arguments

x	an object containing the values the similarity matrix should be computed for
y	same as x. If given the union of values in x and y are used, if not the unique values of x are used
s	a vector for filling the matrix. By default producing an identity matrix
byrow	should s fill the matrix by row?

Value

A square matrix with the values of s and row-/colnames of the unique values in {x, y}.

See Also

[similarity](#)

Examples

```
smat(1:3)

smat(c("f", "e", "d"), s=c(
4, 1, 1,
1, 3, 2,
1, 2, 3
))
```

speedskate

2018 MarbleLympics speed skating times

Description

Intermediate and total times for all 16 runs, arranged by lane and heat number.

Usage

```
speedskate
```

Format

A list containing two data.frames, one for each lane. Columns are heat and rows are time checks in seconds.

Source

https://www.youtube.com/watch?v=fA-O6f_jArk

Examples

```
tt <- t(do.call(cbind, speedskate))
pairs(tt)
cor(tt)
outer(
  colnames(tt),
  colnames(tt),
  Vectorize(function(i,j) cor.test(tt[,i],tt[,j])$p.value)
)
```

spread_seq	<i>Spread sequence</i>
------------	------------------------

Description

Spread out the values of a numeric vector

Usage

```
spread_seq(
  x,
  f = 1.1,
  ...,
  node = c("midrange", "first", "last", "mean", "median", "min", "max")
)
```

Arguments

x	a numeric vector
f	numeric or item matching a function. If numeric, a multiplicative factor applied to the distance between points, otherwise a function to be applied to differences
...	further arguments passed to f if matching a function
node	the location of x that will remain unchanged

Examples

```
x <- c(-1, 0, 2, NA, 4, 5, 6, 8, 9)

spread_seq(c(-1, 1))
spread_seq(x, 1.5, "midrange")
spread_seq(x, 1.5, "first")
spread_seq(x, 1.5, "last")
spread_seq(x, 1.5, "mean")
spread_seq(x, 1.5, "median")

spread_seq(c(3, 4, 1, 9, 6))
spread_seq(c(3, 4, 1, 9, 6), 2, "min")
spread_seq(c(3, 4, 1, 9, 6), 2, "max")

spread_seq(c(3, 4, 1, 9, 6), "/", 2)
spread_seq(c(3, 4, 1, 9, 6), 0.5)

y <- sort(c(3, 4, 1, 9, 6))
plot(y)
lines(spread_seq(y, "log1p"))

f <- function(x) sqrt(abs(x)*2)*sign(x)
y <- c(3, 4, 1, 9, 6)
plot(y)
lines(spread_seq(y, f=f))
```

summary.stl	<i>Summarizing seasonal decomposition</i>
-------------	---

Description

Summary method for class "stl".

Usage

```
## S3 method for class 'stl'
summary(object, digits = getOption("digits"), ...)
```

Arguments

object	an object of class "stl"
digits	the number of significant digits to use when printing
...	further arguments passed to or from other methods

Details

This function is a slight modification to `stats::summary.stl`, the main change being the addition of the variance statistic, which can be considered a parametric (normal) compliment to the existing IQR statistic.

Examples

```
set.seed(1)
x <- ts(rnorm(1e4, sd=1), frequency=12)
a <- stl(x, s.window="periodic")
stats::summary.stl(a)
summary(a)

b <- stl(co2, s.window="periodic")
summary(b)
```

tied_triple_test	<i>Tied triple test</i>
------------------	-------------------------

Description

Compare numeric values, returning an inbetween value for ties

Usage

```
x %ttt% y

ttt(x, y)

is.ttt(x)

## S3 method for class 'ttt'
print(x, symbols = TRUE, ...)

## S3 method for class 'ttt'
table(...)
```

Arguments

<code>x, y</code>	numeric values to be compared
<code>symbols</code>	should symbols be used instead of numeric values?
<code>...</code>	further arguments passed to methods

See Also

[Comparison](#), [comparison_with_ties](#)

Examples

```
1:5 %ttt% 3

ttt(1:3, 2)
print(ttt(1:3, 2), FALSE)

c(1, 6, 3, 0) %ttt% c(1, 3, 3, 2)

# Equivalent
as.integer(c(1, 6, 3, 0) %ttt% c(1, 3, 3, 2))
sign(c(1, 6, 3, 0) - c(1, 3, 3, 2))

# Demonstrating table method
dtf <- data.frame(x=1:5, y=3)
dtf$`?'` <- ttt(dtf$x, dtf$y)
dtf

x <- c(8, 4, 6, 8, 9, 6, 5, 7, 0, 3, 2, 1, 5, 6, 4, 7, 6,
      3, 1, 9, 5, 6, 7, 7, 4, 5, 8, 6, 2, 5, 9, 5, 4, 8)
y <- c(1, 3, 2, 4, 6, 0, 5, 3, 7, 5, 7, 4, 5, 6, 0, 1, 4,
      2, 4, 3, 1, 5, 3, 9, 2, 2, 4, 7, 5, 6, 8)

ou <- outer(sort(x), sort(y), "%ttt%")
ta <- table(ou)

pa <- capture.output(ta)

par(mar=c(1, 2, 3, 2))
image(ou, col=topo.colors(length(ta)), axes=FALSE)
title(pa)
```

box()

var_th	<i>Theoretical variance</i>
--------	-----------------------------

Description

Use Calculate the theoretical variance of base probability distributions

Usage

```
var_th(
  p,
  distribution = c("uniform", "exponential", "gamma", "t", "students-t", "bates",
    "binomial", "nbinom", "negative binomial", "beta", "f", "geometric", "hypergeometric",
    "lognormal", "log-normal", "weibull", "signed-rank", "rank-sum", "logistic")
)
```

Arguments

p	a named vector of parameter values, or a single unnamed numeric if only one parameter. Use a data.frame with appropriately named columns to calculate several variances of the same distribution.
distribution	the name of the distribution to calculate the variance of

Details

The parameters and their names are the same as used in their respective density function. In some cases, like gamma, (negative) binomial etc. more than one convention is followed.

See Also

[Distributions](#)

Examples

```
var_th(p=data.frame(min=1:2, max=5:6), dist="unif")
var(runif(1e5, 1, 5))

var_th(p=2:3, dist="exp")
var(rexp(1e5, 2))

var_th(p=data.frame(shape=3:2, scale=c(0.8, 1)), dist="gamma")
var(rgamma(1e5, shape=3, scale=0.8))

var_th(p=c(shape=3, rate=1.25), dist="gamma")
var(rgamma(1e5, shape=3, rate=1.25))

var_th(p=18:20, dist="t")
var(rt(1e5, 18))

var_th(p=c(a=1, b=2, n=3), dist="bates")
```

```

var(rbates(1e5, a=1, b=2, nr=3))

var_th(p=c(size=10, prob=0.8), dist="binom")
var(rbinom(1e5, 10, 0.8))

var_th(p=c(size=10, prob=0.8), dist="nbinom")
var(rnbinom(1e5, size=10, prob=0.8))

var_th(p=c(size=10, mu=2), dist="nbinom")
var(rnbinom(1e5, size=10, mu=2))

var_th(p=data.frame(shape1=c(1, 2), shape2=c(1.5, 1)), dist="beta")
var(rbeta(1e5, shape1=1, shape2=1.5))
var(rbeta(1e5, shape1=2, shape2=1))

var_th(p=c(df1=6, df2=11), dist="f")
var(rf(1e5, 6, 11))

var_th(p=c(m=3, n=3, k=2), dist="hypergeom")
var(rhyper(1e5, m=3, n=3, k=2))

var_th(p=c(meanlog=0, sdlog=1), dist="log-normal")
var(rlnorm(1e5, meanlog=0, sdlog=1))

var_th(p=c(shape=2, scale=1), dist="weibull")
var(rweibull(1e5, shape=2, scale=1))

var_th(p=20, dist="signed-rank")
var(rsignrank(1e5, n=20))

var_th(p=c(m=13, n=10), dist="rank-sum")
var(rwilcox(1e5, m=13, n=10))

```

weekday	<i>Week-day names</i>
---------	-----------------------

Description

Convert numeric, character, factor and date-time vectors to week-day names

Usage

```

weekday(x, ...)

## Default S3 method:
weekday(
  x,
  short = TRUE,
  language = c("english", "nn norwegian", "bm norwegian"),
  ...
)

## S3 method for class 'Date'
weekday(x, ...)

```

```
## S3 method for class 'POSIXt'
weekday(x, ...)
```

Arguments

<code>x</code>	a vector
<code>...</code>	further arguments passed to methods
<code>short</code>	if TRUE the names will be returned in shortened form
<code>language</code>	what language the names should be returned in

Details

This function follows the ISO 8601 standard, meaning that Monday is considered the first day of the week.

Examples

```
weekday(c("c", "b", "a"))
weekday(c("3", "2", "1"))
weekday(3:1)

weekday(Sys.Date())
weekday(Sys.Date(), short=FALSE, lang="nn nor")
```

 ymse

ymse: A collection of more or less useful functions

Description

There is no grand "theme" to ymse, other than that none of the functions, and in some cases function groups and classes, seemed to fit too well in any other package or merit their own package entirely.

ymse functions

[addrows](#) Add rown to a data.frame [ahist](#) Create an average shifted histogram

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