Package ‘Ecfun’

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Arrows

Draw arrows between pairs of points.

Description

Generalizes graphics::arrows to allow all arguments to be vectors. (As of R 3.1.0, only the first component of the length argument is used by graphics::arrows; others are ignored without a warning.)

Usage

`Arrows(x0, y0, x1 = x0, y1 = y0, length = 0.25, angle = 30, code = 2, col = par("fg"), lty = par("lty"), lwd = par("lwd"), warnZeroLength=FALSE, ...)`

Arguments

- `x0, y0, x1, y1, length, angle, code, col, lty, lwd, ...` as for `arrows`.
- `warnZeroLength` - Issue a warning for zero length arrow? `arrow` does; skip if FALSE.

Details

1. Put all arguments in a `data.frame` to force them to shared length.
2. Call `arrows` once for each row.

Author(s)

Spencer Graves

See Also

`arrows`

Examples

```r
## 1. Simple example:
## 3 arrows, the first with length 0 is suppressed
##
plot(1:3, type='n')
Arrows(1, 1, c(1, 2, 2), c(1, 2:3), col=1:3, length=c(1, .2, .6))
```

```r
## 2. with an NA
##
plot(1:3, type='n')
Arrows(1, 1, c(1, 2, 2), c(1, 2, NA), col=1:3, length=c(1, .2, .6))
```
as.Date1970

Date from a number of days since the start of 1970.

Description

as.Date.numeric requires origin to be specified. The present function assumes that this origin is January 1, 1970.

Usage

as.Date1970(x, ...)

Arguments

x

a numeric vector of dates in days since the start of 1970.

...

optional arguments to pass to as.Date.

Value

Returns a vector of Dates

Author(s)

Spencer Graves

See Also

as.Date as.POSIXct1970

Examples

days <- c(0, 1, 365)
Dates <- as.Date1970(days)

all.equal(c('1970-01-01', '1970-01-02', '1971-01-01'),
          as.character(Dates))

all.equal(days, as.numeric(Dates))
asNumericDF  

Coerce to numeric dropping commas and info after a blank

Description

Delete a leading dollar sign plus commas (thousand separators) and drop information after a blank, then coerce to numeric.

For a data.frame, apply to all columns, drop non-numeric columns, and order the rows by the orderBy. Some Excel imports include commas as thousand separators; this replaces any commas with char(0), ".

Similarly, if "%" is found as the last character in any field, drop the percent sign and divide the resulting numeric conversion by 100 to convert to proportion.

Also, some character data includes footnote references following the year.

Table F-1 from the US Census Bureau needs all three of these features: It needs orderBy, because the most recent year appears first, just the opposite of most other data sets where the most recent year appears last. It has footnote references following a character string indicating the year. And it includes commas as thousand separators.

Usage

asNumericChar(x)
asNumericDF(x, keep=function(x)any(!is.na(x)), orderBy=NA)

Arguments

x            For asNumericChar, this is a character vector to be converted to numeric after gsub(\',\', ' ', x). For asNumericDF, this is a data.frame with all character columns to be converted to numerics.
keep         something to indicate which columns to keep
orderBy      Which columns to order the rows of x[, keep] by. Default is to keep the input order.

Details

1. Replace commas by nothing
2. strsplit on ' ' and take only the first part, thereby eliminating the footnote references.
3. Replace any blanks with NAs
4. as.numeric
5. lapply(x, 1-4)
6. order the rows

Value

all numeric data.frame
Author(s)

Spencer Graves

See Also

scan gsub Quotes

Examples

```r
## 1. simple example

fakeF1 <- data.frame(yr=c('1948', '1947 (1)'),
                      q1=c('1 2 3', ''),
                      duh=rep(NA, 2),
                      dol=c('$1.23', ''),
                      pct=c('1%', '2%'))
nF1 <- asNumericDF(fakeF1)
	nF1. <- data.frame(yr=asNumericChar(fakeF1$yr),
                      q1=asNumericChar(fakeF1$q1),
                      duh=rep(NA, 2),
                      dol=asNumericChar(fakeF1$dol),
                      pct=c(.01, .02))

ncF1c <- data.frame(yr=1948:1947, q1=c(1234, NA),
                    duh=rep(NA, 2),
                    dol=asNumericChar(fakeF1$dol),
                    pct=c(.01, .02))

all.equal(nF1, nF1.)

all.equal(nF1., nF1c)

## 2. orderBy=1:2

nF. <- asNumericDF(fakeF1, orderBy=1:2)

all.equal(nF., nF1c[2:1])
```

BoxCox

**Box-Cox power transformation and its inverse**

Description

Box and Cox (1964) considered the following family of transformations indexed by \( \lambda \):

\[
w = \frac{(y^\lambda - 1)}{\lambda} \text{ if } \lambda \neq 0 \text{ or } \log(y) \text{ if } \lambda = 0.
\]
They estimate \( \lambda \) assuming \( w \) follows a normal distribution. This raises a theoretical problem in that \( y \) must be positive, which means that \( w \) must follow a truncated normal distribution conditioned on \( \lambda w > (-1) \).

Bickel and Doksum (1981) removed the restriction to positive \( y \), i.e., to \( w > (-1/\lambda) \) by modifying the transformation as follows:

\[
\begin{align*}
w & \left\{ \begin{array}{ll}
-\text{sign}(y) \times \text{abs}(y)^{\lambda-1}/\lambda & \text{if } \lambda \neq 0 \\
\text{sign}(y)^* \log(\text{abs}(y)) & \text{if } \lambda = 0
\end{array} \right.
\end{align*}
\]

In practice, we assume that \( y > 0 \), so this distinction has little practical value. However, the boxcox function computes the Bickel-Doksum version if \( \text{rescale} = \text{FALSE} \).

Box and Cox further noted that proper estimation of \( \lambda \) should include the Jacobian of the transformation in the log(likelihood). Doing this can be achieved by rescaling the transformation with the \( n \)th root of the Jacobian, which can be written as follows:

\[
j(y, \lambda) = J(y, \lambda)^{(1/n)} = \text{GeometricMean}(y)^{\lambda-1}.
\]

With this the rescaled power transformation is as follows:

\[
z = (y^\lambda)^{-1}/(\lambda^{n}) \times j(y, \lambda) \left\{ \begin{array}{ll}
\text{if } \lambda \neq 0 \\
\text{GeometricMean}(y)^{\lambda-1} \log(y) & \text{if } \lambda = 0
\end{array} \right.
\]

In addition to facilitating estimation of \( \lambda \), rescaling has the advantage that the units of \( z \) are the same as the units of \( y \).

The output has class ‘BoxCox’, which has attributes that allow the input to be recovered using invBoxCox. The default values of the arguments of invBoxCox are provided by the corresponding attributes of \( z \).

**Usage**

```
boxcox(y, lambda, rescale=TRUE, na.rm=rescale)
invBoxCox(z, lambda, sign.y, GeometricMean, rescale)
```

**Arguments**

- **y**: a numeric vector for which the power transform is desired
- **lambda**: A numeric vector of length 1 or 2. The first component is the power. If the second component is provided, \( y \) is replaced by \( y^\lambda \cdot \lambda[2] \).
- **rescale**: logical or numeric. If logical:
  - For boxcox, this is TRUE to the power transform with rescale, \( z \), above, and FALSE to return the power transform without the \( n \)th root of the Jacobian, \( w \), above. This defaults to TRUE, because this will give \( z \) the same units as \( y \).
  - For invBoxCox, this is TRUE if the input argument \( z \) is assumed to have been rescaled by the \( n \)th root of the Jacobian of the transformation. This defaults to a rescale attribute of \( z \) if present or to TRUE if absent.
- **na.rm**: logical: TRUE to remove NAs from \( y \) before computing the geometric mean. FALSE to compute NA for the geometric mean if any(is.na(y)).
- **sign.y**: logical: TRUE to compute NA for the geometric mean if any(is.na(y)).

**Usage**

```
boxcox(y, lambda, rescale=TRUE, na.rm=rescale)
invBoxCox(z, lambda, sign.y, GeometricMean, rescale)
```

**Arguments**

- **y**: a numeric vector for which the power transform is desired
- **lambda**: A numeric vector of length 1 or 2. The first component is the power. If the second component is provided, \( y \) is replaced by \( y^\lambda \cdot \lambda[2] \).
- **rescale**: logical or numeric. If logical:
  - For boxcox, this is TRUE to the power transform with rescale, \( z \), above, and FALSE to return the power transform without the \( n \)th root of the Jacobian, \( w \), above. This defaults to TRUE, because this will give \( z \) the same units as \( y \).
  - For invBoxCox, this is TRUE if the input argument \( z \) is assumed to have been rescaled by the \( n \)th root of the Jacobian of the transformation. This defaults to a rescale attribute of \( z \) if present or to TRUE if absent.
- **na.rm**: logical: TRUE to remove NAs from \( y \) before computing the geometric mean. FALSE to compute NA for the geometric mean if any(is.na(y)).
- **sign.y**: logical: TRUE to compute NA for the geometric mean if any(is.na(y)).

**NOTE**: If \( na.rm = \text{FALSE} \), the output will be all NA if \( \text{rescale} = \text{TRUE} \). This could produce non-useable answers in most cases. To avoid that, the default for \( na.rm \) is \text{TRUE} whenever \( \text{rescale} = \text{TRUE} \). Conversely, applications using \( na.rm = \text{FALSE} \) will likely also want \( \text{rescale} = \text{FALSE} \) to avoid returning a non-answer in these cases. This explains the default \( na.rm = \text{rescale} \).
BoxCox

\[
z \quad \text{a numeric vector or an object of class BoxCox for which the inverse Box-Cox transform is desired.}
\]

\[
\text{sign.y} \quad \text{an optional logical vector giving sign(y-lambda[2]) of the data values that presumably generated z.Defaults to an sign.y attribute of z or to rep(1, length(z)) if no such attribute is present.}
\]

\[
\text{GeometricMean} \quad \text{an optional numeric scalar giving the geometric mean of the data values that presumably generated z. Defaults to a GeometricMean attribute of z or to 1 if no such attribute is present.}
\]

**Details**

Box and Cox (1964) discussed

\[
w(y, \lambda) = \frac{(y^\lambda - 1)}{\lambda}.\]

They noted that \( w \) is continuous in \( \lambda \) with \( w(y, \lambda) = \log(y) \) if \( \lambda = 0 \) (by l'Hopital's rule).

They also discussed

\[
z(y, \lambda) = \frac{(y^\lambda - 1)}{(\lambda g^{(\lambda - 1)})}.
\]

where \( g \) is the geometric mean of \( y \).

They noted that proper estimation of \( \lambda \) should include the Jacobian of \( w(y, \lambda) \) with the likelihood. They further showed that a naive normal likelihood using \( z(y, \lambda) \) as the response without a Jacobian is equivalent to the normal likelihood using \( w(y, \lambda) \) adjusted appropriately using the Jacobian. See Box and Cox (1964) or the Wikipedia article on "Power transform".

Bickel and Doksum (1981) suggested adding \( \text{sign}(y) \) to the transformation, as discussed above.

**NUMERICAL ANALYSIS:**

Consider the Bickel and Doksum version described above:

\[
w = \text{code(sign(y)*abs(y)^lambda-1)/lambda}
\]

Let \( ly = \log(\text{abs}(y)) \). Then with \( la = \lambda \),

\[
w = \text{code(sign(y)*exp(la*ly)-1)/la}
\]

\[= \text{sign(y)*ly*(1+(la*ly/2)*(1+(la*ly/3)*(1+(la*ly/4)*(1+0(la*ly)))) + (sign(y)-1)/la}
\]

For \( y>0 \), the last term is zero. `boxcox` ignores cases with \( y<=0 \) and uses this formula (ignoring the final \( O(la*ly) \)) whenever \( \text{abs}(la) <= \text{eps} = 1/50 \). That form is used here also.

For `invBoxCox` a complementary analysis is as follows:

\[
\text{abs(y+lambda[2]) = abs(1+la*w)^*(1/la)
\]

\[= \exp(\log(\text{lp}(la*w)/la) \text{ for abs}(la*w)<1
\]

\[= w*(1-la*w*((1/2)-la*w*((1/3)-la*w*(1/4-...))))
\]

**Value**

`BoxCox` returns an object of class `BoxCox`, being a numeric vector of the same length as \( y \) with the following optional attributes:

- \( \lambda \) the value of \( \lambda \) used in the transformation
• sign.y sign(y) (or sign(y-lambda[2]) lambda[2] is provided and if any of these quantities are negative. Otherwise, this is omitted and all are assumed to be positive.

• rescale logical: TRUE if z(y, lambda) is returned rescaled by g^(lambda-1) with g = the geometric mean of y and FALSE if z(y, lambda) is not so rescaled.

• GeometricMean If rescale is numeric, attr(. , 'GeometricMean') <- rescale. Otherwise, attr(. , 'GeometricMean') is the Geometric mean of abs(y) = exp(mean(log(abs(y)))) or of abs(y+lambda[2]) if(length(lambda)>1).

invBoxCox returns a numeric vector, reconstructing y from BoxCox(y, ...).

Source


References

Wikipedia, "Power transform"

See Also

boxcox, quine boxcox boxcoxCensored boxcox boxcox.drc boxCox

Examples

```r
##
## 1. A simple example to check the two algorithms
##
Days <- 0:9
bc1 <- BoxCox(Days, c(0.01, 1))
# Taylor expansion used for obs 1:7; expm1 for 8:10

# check
GM <- exp(mean(log(abs(Days+1))))
bc0 <- (((Days+1)^0.01)-1)/0.01
bc1. <- (bc0 / (GM^0.01))
# log(Days+1) ranges from 0 to 4.4
# lambda = 0.01 will invoke both the obvious
# algorithm and the alternative assumed to be
# more accurate for (lambda(log(y)) < 0.02).
attr(bc1., 'lambda') <- c(0.01, 1)
attr(bc1., 'rescale') <- TRUE
attr(bc1., 'GeometricMean') <- GM
class(bc1.) <- 'BoxCox'

all.equal(bc1, bc1.)
```
## 2. The "boxcox" function in the MASS package

computes a maximum likelihood estimate with
BoxCox(Days+1, lambda=0.21)
with a 95 percent confidence interval of
approximately (0.08, 0.35)

bcDays1 <- BoxCox(MASS::quine$Days, c(0.21, 1))

# check
GeoMean <- exp(mean(log(abs(MASS::quine$Days+1))))
bcDays1. <- (((MASS::quine$Days+1)^0.21)-1) / (0.21*GeoMean^(0.21-1))

# log(Days+1) ranges from 0 to 4.4
attr(bcDays1., 'lambda') <- c(0.21, 1)
attr(bcDays1., 'rescale') <- TRUE
attr(bcDays1., 'GeometricMean') <- GeoMean
class(bcDays1.) <- 'BoxCox'

all.equal(bcDays1, bcDays1.)

iDays <- invBoxCox(bcDays1)

all.equal(iDays, MASS::quine$Days)

## 3. Easily computed example

bc2 <- BoxCox(c(1, 4), 2)

# check
bc2. <- (c(1, 4)^2-1)/4
attr(bc2., 'lambda') <- 2
attr(bc2., 'rescale') <- TRUE
attr(bc2., 'GeometricMean') <- 2
class(bc2.) <- 'BoxCox'

all.equal(bc2, bc2.)

all.equal(invBoxCox(bc2), c(1, 4))

## 4. plot(BoxCox())
Split a character string where a capital letter follows a lowercase letter.

Usage

camelParse(x, except=c(‘De’, ’Mc’, ’Mac’))

Arguments

x a character vector

except character vector giving exceptions: If any of these are found, ignore and look for the next one

Details

Find all places where a lowercase letter is followed by a capital.

Split on those points

Value

list of character vectors
checkNames

Author(s)
Spencer Graves

See Also
strsplit

Examples

tst <- c('Smith, JohnJohn Smith',
        'EducationNational DefenseOther Committee',
        'McCain, JohnJohn McCain')
tst. <- camelParse(tst)

all.equal(tst., list(c('Smith, John', 'John Smith'),
                    c('Education', 'National Defense', 'Other Committee'),
                    c('McCain, John', 'John McCain') ) )

checkNames  Check and return names

Description
Check and return names. If names are not provided or are not unique, write a message and return make.names consistent with warn and unique.

Usage
checkNames(x, warn=0, unique=TRUE,
          avoid=character(0),
          message=NULL, head(deparse(substitute(x)), 25), 2), ...)

Arguments

  x      an R object suitable for names

  warn   Numeric code for how to treat problems, consistent with the argument warn in options: Negative to ignore, 0 to save and print later, 1 to print as they occur, 2 or greater to convert to errors.

  unique logical: TRUE to check that names(x) are unique. Fix any duplicates with make.names.

  avoid  a vector of regular expressions to avoid adding to the output of make.names with a companion replacement when found.

Thus, length(avoid) must be a nonnegative even integer, with avoid[2*j-1] providing the pattern for regexpr and sub, and avoid[2*j] providing the replacement. See the second example.
checkNames

message0 Base to prepend to any message
... optional arguments for make.names

Details

1. namex <- names(x)
2. Check per warn and unique
3. Return an appropriate version of namex

Value

a character vector of the same length as x. If any problem is found, this character vector will have an attribute message describing the problem found. Message checking considers unique but ignores warn.

Author(s)

Spencer Graves

See Also

names make.names options for warn

Examples

##
## 1. standard operation with no names
##
tst1 <- checkNames(1:2)

# check
tst1. <- make.names(character(2), unique=TRUE)
attr(tst1., 'message') <- paste("1:2: names = NULL; returning",
"make.names(character(length(x))), TRUE")

all.equal(tst1, tst1.)

##
## 2. avoid=c(’\n0$’, ’\n1$’)
##
tst2 <-checkNames(1:2,
  avoid=c(‘\n0$’, ‘.2’,
  ‘\n1$’, ‘.3’))

# check
tst2. <-c(‘X’, ‘X.3’)
attr(tst2., 'message') <- paste("1:2: names = NULL; returning",
"make.names(character(length(x))), TRUE")


```
all.equal(tst2, tst2.)
```

---

### classIndex

**Convert class to an integer 1-8 and vice versa**

The `classIndex` function converts the class of `x` to an integer:

1. NULL
2. logical
3. integer
4. numeric
5. complex
6. raw
7. character
8. other

`index2class` converts an integer back to the corresponding class.

### Usage

```
classIndex(x)  
index2class(i, otherCharacter=TRUE)
```

### Arguments

- `x`: an object whose class index is desired.
- `i`: an integer to be converted to the name of the corresponding class
- `otherCharacter`: logical: TRUE to convert 8 to "character"; FALSE to convert 8 to "other".

### Details

The *Writing R Extensions* lists six different kinds of "atomic vectors": logical, integer, numeric, complex, character, and raw: See also Wickham (2013, section on "Atomic vectors" in the chapter on "Data structures"). These form a standard hierarchy, except for "raw", in that standard operations combining objects with different atomic classes will create an object of the higher class. For example, `TRUE + 2 + pi` returns a numeric object (approximately 6.141593). Similarly, `paste(1, 'a')` returns the character string "1 a".

For "interpolation", we might expect users interpolating between objects of class "raw" (i.e., bytes) might most likely prefer "Numeric" to "Character" interpolation, coerced back to type "raw".

The index numbers for the classes run from 1 to 8 to make it easy to convert them back from integers to character strings.
Value

`classIndex` returns an integer between 1 and 7 depending on `class(x)`. `index2class` returns a character string for the inverse transformation.

Author(s)

Spencer Graves

References

Wickham, Hadley (2014) *Advanced R*, especially Wickham (2013, section on "Atomic vectors" in the chapter on "Data structures").

See Also

`interpChar`

Examples

```r
##
## 1. classIndex
##
x1 <- classIndex(NULL)
x2 <- classIndex(logical(0))
x3 <- classIndex(integer(1))
x4 <- classIndex(numeric(2))
x5 <- classIndex(complex(3))
x6 <- classIndex(raw(4))
x7 <- classIndex(character(5))
x8 <- classIndex(list())

# check
all.equal(c(x1, x2, x3, x4, x5, x6, x7, x8), 1:8)

##
## 2. index2class
##
c1 <- index2class(1)
c2 <- index2class(2)
c3 <- index2class(3)
c4 <- index2class(4)
c5 <- index2class(5)
c6 <- index2class(6)
c7 <- index2class(7)
c8 <- index2class(8)
c8o <- index2class(8, FALSE)

# check
```
all.equal(c(c1, c2, c3, c4, c5, c6, c7, c8, c8o),
    c('NULL', 'logical', 'integer', 'numeric',
    'complex', 'raw', 'character', 'character',
    'other'))

compareLengths(x, y,    # Compare the lengths of two objects
    name.x=deparse(substitute(x), width.cutoff, nlines=1, ...),
    name.y=deparse(substitute(y), width.cutoff, nlines=1, ...),
    message0='', compFun=c('NROW', 'length'),
    action=c('compatible', 'incompatible', 'warning'),
    length0=c('compatible', 'incompatible', 'stop'),
    width.cutoff=20, ...)

Arguments

- x, y: objects whose lengths are to be compared
- name.x, name.y: names of x and y to use in a message. Default = deparse(substitute(.), width.cutoff, nlines=1).
- message0: character string to be included with name.x and name.y in a message.
- compFun: function to use in the comparison.
- action: A character vector of length 2 giving the names of functions to call if the lengths are not equal but are either 'compatible' or 'incompatible'; "" means no action.
- length0: If length(x) or length(y) = 0 (but not both), treat this case as specified by length0.
- width.cutoff: width.cutoff argument to pass to deparse. This gives the maximum number of characters to use in a name in error and warning messages.
- ...: optional arguments for deparse

Details

1. If nchar(name.x) = 0 = nchar(name.y), set name.x <- 'x', name.y <- 'y', and append 'in compareLengths:' to message0 for more informative messaging.
2. lenx <- do.call(compFun, list(x)); leny <- do.call(compFun, list(y))
3. if(lenx==leny) return(c('equal', '))
4. Compatible?
5. Compose the message.
6. "action", as indicated
**compareLengths**

**Value**

A character vector of length 2. The first element is either 'equal', 'compatible' or 'incompatible'. The second element is the message composed.

**Author(s)**

Spencer Graves with help from Duncan Murdoch

**See Also**

interpChar

**Examples**

```r
## 1. equal
##
all.equal(compareLengths(1:3, 4:6), c("equal", "))

## 2. compatible
##
a <- 1:2
b <- letters[1:6]
comp.ab <- compareLengths(a, b, message = 'Chk: ')
comp.ba <- compareLengths(b, a, message = 'Chk: ')
# check
chk.ab <- c('compatible',
             'Chk: length(b) = 6 is 3 times length(a) = 2')
all.equal(comp.ab, chk.ab)

all.equal(comp.ba, chk.ab)

## 3. incompatible
##
Z <- LETTERS[1:3]
comp.aZ <- compareLengths(a, Z)
# check
chk.aZ <- c('incompatible',
             ' length(Z) = 3 is not a multiple of length(a) = 2')
all.equal(comp.aZ, chk.aZ)

## 4. problems with name.x and name.y
##
```
createMessage

Compose a message as a single substring from a character vector

Description

This is a utility function to make it easier to automatically compose informative error and warning messages without using too many characters.

Usage

createMessage(x, width.cutoff=45, default='x', collapse='; ',
endchars='...')

Arguments

x input for paste

width.cutoff maximum number of characters from x to return in a single string. This differs from the width.cutoff argument in deparse in that the output include here considers endchars, not part of deparse.

default character string to return if nchar(x) = 0.

collapse collapse argument for paste

d default character string to indicate that part of the input string(s) was truncated.

Details

x <- paste(..., collapse=': ') nchx <- nchar(x) maxch <- (maxchar-nchar(endchar)) if(nchx>maxch)
x2 <- substring(x, 1, maxch) x. <- paste0(x2, endchar)

Value

a character string with at most width.cutoff characters.
createX2matchY

Author(s)
Spencer Graves

See Also
paste substr nchar

Examples

```r
## 1. typical use
##
tstVec <- c('Now', 'is', 'the', 'time')
msg <- createMessage(tstVec, 9, collapse=':',
                     endchars='//')

all.equal(msg, 'Now:is://')

## 2. in a function
##
tstFn <- function(cl) createMessage(deparse(cl), 9)
Cl <- quote(plot(1:3, y=4:6, col='red', main='Title'))
msg0 <- tstFn(Cl)
# check
msg. <- 'plot(1...'

all.equal(msg0, msg.)

## 3. default
##
y <- createMessage(character(3), default='y')

all.equal(y, 'y')
```

description

Return a default object of class index2class(max(classIndex(x), classIndex(y))) and length = length(y).

For example, suppose class(x) == 'numeric', for which classIndex = 4. If class(y) = 'integer', then an object of class 'numeric' is returned. However, if class(y) = 'character', then an object of class 'character' is returned.
Usage
createX2matchY(x, y)

Arguments
x, y  objects of possibly different classes and lengths.

Value
A vector of the same length as y whose class is index2class(max(classIndex(x), classIndex(y))).

Author(s)
Spencer Graves

See Also
interpPairs

Examples
```r
##
## 1.  NULL
##  -
nul <- createX2matchY(NULL, NULL)
# check
all.equal(nul, NULL)

##
## 2.  logical
##  #
# lgl3 <- createX2matchY(NULL,
#  c(FALSE, TRUE, FALSE))
# check
all.equal(lgl3, logical(3))

##
## 3.  integer
##  #
# int3 <- createX2matchY(integer(0),
#  c(FALSE, TRUE, FALSE))
# check
all.equal(int3, integer(3))

##
## 4.  list -> character
```
financialCrisisFiles

Files containing financial crisis data

Description

FinancialCrisisFiles in Ecdat is an object of class financialCrisisFiles created by the
financialCrisisFiles function to describe files containing data on financial crises downloadable
from http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/.

Usage

financialCrisisFiles(files=c("22_data.xls", "23_data.xls",
"Varieties_Part_III.xls", "25_data.xls"), ...)

Arguments

files character vector of file names
... arguments to pass with file and sheet name to read.xls when reading a sheet
of an MS Excel file. This is assumed to be the same for all sheets of all files. If
this is not the case, the resulting financialCrisisFiles object will have to be
edited manually before using it to read the data.

Details

Reinhart and Rogoff (http://www.reinhartandrogoff.com) provide numerous data sets ana-
yzed in their book, "This Time Is Different: Eight Centuries of Financial Folly". Of interest here
are data on financial crises of various types for 70 countries spanning the years 1800 - 2010, down-
loadable from http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/.

The function financialCrisisFiles produces a list of class financialCrisisFiles describ-
ing four different Excel files in very similar formats with one sheet per Country and a few extra
descriptor sheets. The data object FinancialCrisisFiles is the default output of that function.

It does this in several steps:
1. Read the first sheet of each file
2. Extract the names of the Countries from that first sheet.
3. Eliminate any blank spaces in the names to convert, e.g., "Costa Rica" to "CostaRica".
4. Find the sheets corresponding to each of the compressed names.
5. Construct the output list.
Value

The function `financialCrisisFiles` returns a list of class `financialCrisisFiles`. This is a list with components carrying the names of files to be read. Each component is a list of optional arguments to pass to `do.call(read.xls, ...) to read the sheet with name = name of that component.

The default value returned by `financialCrisisFiles` is the data object `FinancialCrisisFiles`. This corresponds to the files downloaded from [http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/](http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/) in January 2013 (except for the fourth, which was not available there because of an error with the web site but instead was obtained directly from Prof. Reinhart).

Author(s)

Spencer Graves

Source

[http://www.reinhartandrogoff.com](http://www.reinhartandrogoff.com)

References


See Also

`read.xls`

Examples

```r
Ecdat.demoFiles <- system.file('demoFiles', package='Ecdat')
Ecdat.xls <- dir(Ecdat.demoFiles, pattern='xls$',
                 full.names=TRUE)
if(require(gdata)){
  tst <- financialCrisisFiles(Ecdat.xls)
}
## Not run:
# check
\dontshow(stopifnot())
all.equal(tst, data(FinancialCrisisFiles))
\dontrun())

## End(Not run)
```
getElement2

Extract a named element from an object with a default

Description

Get element name of object. If object does not have an element name, return default.

If the name element of object is NULL the result depends on warn.NULL: If TRUE, issue a warning and return default. Otherwise, return NULL.

Usage

gElement2(object, name=1, default=NA, warn.NULL=TRUE, 
envir=list(), returnName)

Arguments

object object from which to extract component name.
name Name or index of the element to extract
default default value if name is not part of object.
warn.NULL logical to decide how to treat cases where object has a component name: If TRUE, return default with a warning. Otherwise, return NULL.
envir Supplemental list beyond object in which to look for names in case object[[name]] is a language object that must be evaluated.
returnName logical: TRUE to return as.character of any name found as an element of object. FALSE to eval any name found in the environment of object. Default = TRUE if name == 1 or a character string matching the name of the first element of object.

Details

1. If is.numeric(name) In <- (1 <= name <= length(object))
2. else In <- if(name %in% names(object))
3. El <- if(In) object[[name]] else default
4. warn.NULL?
5. if(returnName) return(as.character(El)) else return(eval(El, envir=object))

Value

an object of the form of object[[name]]; if object does not have an element or slot name, return default.

Author(s)

Spencer Graves with help from Marc Schwartz and Hadley Wickham
See Also

`getElement`, which also can return slots from S4 objects.

Examples

```r
## 1. name in object, return
##
e1 <- getElement2(list(ab=1), 'ab', 2) # 1
# check

all.equal(e1, 1)

## 2. name not in object, return default
##
eNA <- getElement2(list(), 'ab') # default default = NA
# check

all.equal(eNA, NA)

e0 <- getElement2(list(), 'ab', 2) # name not in object

all.equal(e0, 2)

e2 <- getElement2(list(ab=1), 'a', 2) # partial matching not used

all.equal(e2, 2)

## 3. name NULL in object, return default
##
ed <- getElement2(list(a=NULL), 'a', 2) # 2 with a warning

all.equal(ed, 2)

e. <- getElement2(list(a=NULL), 'a', 2, warn=NULL=NULL(FALSE)) # NULL

all.equal(e., NULL)

eNULL <- getElement2(list(a=NULL), 'a', NULL) # NULL

all.equal(eNULL, NULL)
```

##
grepNonStandardCharacters

grep for nonstandard characters

Description

Return the indices of elements of \( x \) containing characters that are not in \texttt{standardCharacters}.

Usage

\[
grepNonStandardCharacters(x, value=FALSE, standardCharacters=c(letters, LETTERS, ' ', '.', ',', 0:9, '\', '", ' - ', '_ ', '(', ')', '[', ']', '\n'), \ldots)\]

Arguments

\begin{itemize}
  \item \texttt{x}: character vector in which it is desired to identify elements containing characters not in \texttt{standardCharacters}.
  \item \texttt{value}: logical: \texttt{TRUE} to return the values found in \texttt{x}, \texttt{FALSE} to return their indices.
  \item \texttt{standardCharacters}: Characters to overlook in \texttt{x} to identify anything not in \texttt{standardCharacters}.
  \item \ldots: optional arguments for \texttt{regexp}
\end{itemize}

Details

1. \( x \sim \texttt{strsplit(x, "")}: \) convert the input character vector to a list of vectors of character vectors with \( \text{nchar}(x[i]) = 1 \) for \( i \in 1: \text{length}(x) \).
2. \texttt{sapply(x, \ldots)} to identify all elements for which any element of \( x[i] \) is not in \texttt{standardCharacters}.
Interp

**Value**

an integer vector identifying all elements of x containing a character not in standardCharacters.

**Author(s)**

Spencer Graves

**See Also**

stringi-package grep, rexp, subNonStandardCharacters, showNonASCII

**Examples**

```r
Names <- c('Raul', 'Ra\'l', 'Torres,Raul', 'Torres, Raul')
# confusion in character sets can create
# names like Names[2]
chk <- grepNonStandardCharacters(Names)

all.equal(chk, 2)

chkv <- grepNonStandardCharacters(Names, TRUE)

all.equal(chkv, 'Ra\'l')
```

---

**Interp**

*Interpolate between numbers or numbers of characters*

**Description**

Numeric interpolation is defined in the usual way:

```
xOut <- x*(1-proportion) + y*proportion
```

Character interpolation does linear interpolation on the number of characters of x and y. If `length(proportion) == 1`, interpolation is done on `cumsum(nchar(x))`. If `length(proportion) > 1`, interpolation is based on `nchar`. In either case, the interpolant is rounded to an integer number of characters. Interp then returns `substring(y, ...)` unless `nchar(x) > nchar(y)`, when it returns `substring(x, ...)`. Character interpolation is used in two cases: (1) At least one of x and y is character. (2) At least one of x and y is neither logical, integer, numeric, complex nor raw, and `class(unclass(.))` is either integer or character.

In all other cases, numeric interpolation is used.

**NOTE:** This seems to provide a relatively simple default for what most people would want from the six classes of atomic vectors (logical, integer, numeric, complex, raw, and character) and most
other classes. For example, class(unclass(factor)) is integer. The second rule would apply to this converting it to character. The coredata of an object of class zoo could be most anything, but this relatively simple rule would deliver what most people want in most case. An exception would be an object with integer coredata. To handle this as numeric, a Interp.zoo function would have to be written.

Usage

```r
Interp(x, ...)  
## Default S3 method:  
Interp(x, y, proportion,  
   argnames=character(3), message0=character(0), ...)  
InterpChkArgs(x, y, proportion,  
   argnames=character(3), message0=character(0), ...)  
InterpChar(argsChk, ...)  
InterpNum(argsChk, ...)  
```

Arguments

- `x, y` two vectors of the same class or to be coerced to the same class.
- `proportion` A number or numeric vector assumed to be between 0 and 1.
- `argnames` a character vector of length 3 giving args name.x, name.y, and proportion to pass to compareLengths to improve the value of any diagnostic message in case lengths are not compatible.
- `message0` A character string to be passed with argnames to compareLengths to improve the value of any diagnostic message in case lengths are not compatible.
- `argsChk` a list as returned by interpChkArgs
- `...` optional arguments for compareLengths

Details

Interp is an S3 generic function to allow users to easily modify the behavior to interpolate between special classes of objects.

Interp has two basic algorithms for "Numeric" and "Character" interpolation.

The computations begin by calling InterpChkArgs to dispose quickly of simple cases (e.g, x or y missing or length 0 or if proportion is <= 0 or >= 1 or missing). It returns a list.

If the list contains a component named "xout", Interp returns that value with no further computations.

Otherwise, the list returned by InterpChkArgs includes components "algorithm", "x", "y", "proportion", "pLength1" (defined below), "raw", and "outclass". The "algorithm" component must be either "Numeric" or "Character". That algorithm is then performed as discussed below using arguments "x", "y", and "proportion"; all three will have the same length. The class of "x" and "y" will match the algorithm. The list component "raw" is logical: TRUE if the output will be raw or such that class(unclass(.)) of the output will be raw. In that case, a "Numeric" interpolation will be transformed back into "raw". "outclass" will either be a list of attributes to apply to the output or NA. If a list, "xout" will be added as component ".Data" to the list "outclass" and then then processed as do.call('structure', outclass) to produce the desired output.
These two basic algorithms ("Numeric" and "Character") are the same if proportion is missing or not numeric: In that case Interp throws an error.

We now consider "Character" first, because it's domain of applicability is easier to describe. The "Numeric" algorithm is used in all other cases

1. "CHARACTER"

* 1.1. The "CHARACTER" algorithm is used when at least one of x and y is neither logical, integer, numeric, complex nor raw and satisfies one of the following two additional conditions:
** 1.1.1. Either x or y is character.
** 1.1.2. class(unclass(.)) for at least one of x and y is either character or integer.

NOTE: The strengths and weaknesses of 1.1.2 can be seen in considering factors and integer vectors of class zoo: For both, class(unclass(.)) is integer. For factors, we want to use as.character(.). For zoo objects with coredata of class integer, we would want to use numeric interpolation. This is not allowed with the current code but could be easily implemented by writing interp.zoo.

* 1.2. If either x or y is missing or has length 0, the one that is provided is returned unchanged.
* 1.3. Next determine the class of the output. This depends on whether neither, one or both of x and y have one of the six classes of atomic vectors (logical, integer, numeric, complex, raw, character):
** 1.3.1. If both x and y have one of the six atomic classes and one is character, return a character object.
** 1.3.2. If only one of x and y have an atomic class, return an object of the class of the other.
** 1.3.3. If neither of x nor y have a basic class, return an object with the class of y.
* 1.4. Set pLength1 <- (length(proportion) == 1):
** 1.4.1. If(pLength1) do the linear interpolation on cumsum(nchar(.)).
** 1.4.2. Else do the linear interpolation on nchar.
* 1.5. Next check x, y and proportion for comparable lengths: If all have length 0, return an object of the appropriate class. Otherwise, call compareLengths(x, proportion), compareLengths(y, proportion), and compareLengths(x, y).
* 1.6. Extend x, y, and proportion to the length of the longest using rep.
* 1.7. nchOut <- the number of characters to output using numeric interpolation and rounding the result to integer.
* 1.8. Return substring(y, 1, nchOut) except when the number of characters from x exceed those from y, in which case return substring(x, 1, nchOut). [NOTE: This meets the naive end conditions that the number of characters matches that of x when proportion is 0 and matches that of y when proportion is 1. This can be used to "erase" characters moving from one frame to the next in a video. See the examples.

2. "NUMERIC"

* 2.1. Confirm that this does NOT satisfy the condition for the "Character" algorithm.
* 2.2. If either x or y is missing or has length 0, return the one provided.
* 2.3. Next determine the class of the output. As for "Character" described in section 1.3, this depends on whether neither, one or both of x and y have a basic class other than character (logical, integer, numeric, complex, raw):
** 2.3.1. If proportion <= 0, return x unchanged. If proportion >= 1, return y unchanged.
** 2.3.2. If neither x nor y has a basic class, return an object of class equal that of y.
** 2.3.3. If exactly one of x and y does not have a basic class, return an object of class determined by class(unclass(.)) of the non-basic argument.
** 2.3.4. When interpolating between two objects of class raw, convert the interpoland back to class raw. Do this even when 2.3.2 or 2.3.3 applies and class(unclass(.)) of both x and y are of class raw.
* 2.4. Next check x, y and proportion for comparable lengths: If all have length 0, return an object of the appropriate class. Otherwise, call compareLengths(x, proportion), compareLengths(y, proportion), and compareLengths(x, y).
* 2.5. Compute the desired interpolation and convert it to the required class per step 2.3 above.

Value

Interp returns a vector whose class is described in "* 1.3" and "* 2.3" in "Details" above.
InterpChkArgs returns a list or throws an error as described in "Details" above.

Author(s)

Spencer Graves

References

The *Writing R Extensions* manual (available via help.start()) lists six different classes of atomic vectors: logical, integer, numeric, complex, raw and character. See also Wickham, Hadley (2014) *Advanced R*, especially Wickham (2013, section on "Atomic vectors" in the chapter on "Data structures").

See Also

`classIndex` `interpPairs`

Many other packages have functions with names like "interp", "interp1", and "interpolate". Some do one-dimensional interpolation. Others do two-dimensional interpolation. Some offer different kinds of interpolation beyond linear. At least one is a wrapper for `approx`.

Examples

```r
## 1. numerics
##
# 1.1. standard
xNum <- interpChar(1:3, 4:5, 0:3)/4
# answer
xN. <- c(1, 2.75, 3.5, 4)

all.equal(xNum, xN.)
```
# 1.2. with x but not y:
# return that vector with a warning

xN1 <- Interp(1:4, p=.5)
# answer
xN1. <- 1:4

all.equal(xN1, xN1.)

##
## 2. Single character vector
##

i.5 <- Interp(c('a', 'bc', 'def'), character(0), p=0.3)
# with y = NULL or character(0),
# Interp returns x

all.equal(i.5, c('a', 'bc', 'def'))

i.5b <- Interp('', c('a', 'bc', 'def'), p=0.3)
# Cumulative characters (length(proportion)=1):
# 0.3*(total 6 characters) = 1.2 characters
i.5. <- c('a', 'b', '')

all.equal(i.5b, i.5.)

##
## 3. Reverse character example
##

i.5c <- Interp(c('a', 'bc', 'def'), '', 0.3)
# check: 0.7*(total 6 characters) = 4.2 characters
i.5c. <- c('a', 'bc', 'd')

all.equal(i.5c, i.5c.)

##
## 4. More complicated example
##

xCh <- Interp('', c('Do it', 'with R.'),
    c(0, .5, .9))
# answer
xCh. <- c('', 'with', 'Do i')

all.equal(xCh, xCh.)

##
## 5. Still more complicated
##
```
xC2 <- Interp(c('a', 'fabulous', 'bug'),
              c('bigger or', 'just', 'big'),
              c(.3, .3, 1))
x.y.longer <- c('bigger or', 'fabulous', 'big')
# use y with ties
# nch smaller   1  4  3
# nch larger    9  8  3
# d.char        8, 4, 0
# prop          .3, .7, 1
# prop*d.char   2.4, 2.8, 0
# smaller+p*d   3, 7, 3
xC2. <- c('big', 'fabulous', 'big')
all.equal(xC2, xC2.)

##
## 6. with one NULL
##
null1 <- Interp(NULL, 1, .3)
all.equal(NULL, 1)

null2 <- Interp('abc', NULL, .3)
all.equal('abc')

##
## 7. length=0
##
log0 <- interpChar(logical(0), 2, .6)
all.equal(log0, 1.2)

##
## 8. Date
##
Jan1.1980 <- as.Date('1980-01-01')
Jan1.1972i <- Interp(0, Jan1.1980, .2)
# check
Jan1.1972 <- as.Date('1972-01-01')
all.equal(Jan1.1972, round(Jan1.1972i))

##
## 9. POSIXct
##
Jan1.1980c <- as.POSIXct(Jan1.1980)
```
Jan1.1972ci <- Interp(0, Jan1.1980c, .2)
# check
Jan1.1972ct <- as.POSIXct(Jan1.1972)

abs(diffTime(Jan1.1972ct, Jan1.1972ci,
  units="days"))<0.5

interpChar

Interpolate between numbers or numbers of characters

Description

For x and y logical, integer, numeric, Date or POSIX:

xOut <- x*(1-.proportion) + y*.proportion

Otherwise, coerce to character and return a substring of x or y with number of characters interpolating linearly between nchar(x) and nchar(y); see details.

*** NOTE: This function is currently in flux. The results may not match the documentation and may change in the future.

The current version does character interpolation on the cumulative number of characters with defaults with only one argument that may not be easy to understand and use. Proposed:

old: interpolate on number of characters in each string with the default for a missing argument being character(length(x)) [or character(length(y)) or numeric(length(x)) or ...]

2014-08-08: default with either x or y missing should be to set the other to the one we have, so interpChar becomes a no op – except that values with .proportion outside ("validProportion" = [0, 1] by default) should be dropped.

Usage

interpChar(x, ...)
## S3 method for class 'list'
interpChar(x, .proportion,
  argnames=character(3), message0=character(0), ...)
## Default S3 method:
interpChar(x, y, .proportion,
  argnames=character(3), message0=character(0), ...)

Arguments

x either a vector or a list. If a list, pass the first two elements as the first two arguments of interpChar.default.

y a vector

.proportion A number or numeric vector assumed to be between 0 and 1.
argnames  a character vector of length 3 giving args name. x, name. y, and .proportion to pass to `compareLengths` to improve the value of any diagnostic message in case lengths are not compatible.

messageNULL  A character string to be passed with argnames to `compareLengths` to improve the value of any diagnostic message in case lengths are not compatible.

...  optional arguments for `compareLengths`

Details

1. `x`, `y` and `.proportion` are first compared for compatible lengths using `compareLengths`. A warning is issued if the lengths are not compatible. They are then all extended to the same length using `rep`.

2. If `x` and `y` are both numeric, `interpChar` returns the standard linear interpolation (described above).

3. If `x`, `y`, and `.proportion` are all provided with at least one of `x` and `y` not being numeric or logical, the algorithm does linear interpolation on the difference in the number of characters between `x` and `y`. It returns characters from `y` except when `nchar(x) > nchar(y)`, in which case it returns characters from `x`. This meets the end conditions that the number of characters matches that of `x` when `.proportion` is 0 and matches that of `y` when `.proportion` is 1. This can be used to "erase" characters moving from one frame to the next in a video. See the examples.

4. If either `x` or `y` is missing, it is replaced by a default vector of the same type and length; for example, if `y` is missing and `x` is numeric, `y = numeric(length(x))`. (If the one supplied is not numeric or logical, it is coerced to character.)

Value

A vector: Numeric if `x` and `y` are both numeric and character otherwise. The length = max length of `x`, `y`, and `.proportion`.

Author(s)

Spencer Graves

See Also

`interpPairs`, which calls `interpChar`

`classIndex`, which is called by `interpChar` to help decide the class of the interpoland.

Examples

```r
##
## 1. numerics
##
# 1.1. standard
xNum <- interpChar(1:3, 4:5, (0:3)/4)
# answer
xN. <- c(1, 2.75, 3.5, 4)
```
all.equal(xNum, xN.)

# 1.2. list of length 1 with a numeric vector:
# return that vector with a warning
xN1 <- interpChar(list(a.0=1:4), .5)
# answer
xN1. <- 1:4
all.equal(xN1, xN1.)

##
## 2. Single character vector
##
i.5 <- interpChar(list(c('a', 'bc', 'def')), .p=0.3)
# If cumulative characters:
# 0.3*(total 6 characters) = 1.8 characters
#
# However, the current code does something different,
# returning "a", "bc", "d" <- like using 1-.p?
# This is a problem with the defaults with a single
# argument; ignore this issue for now.
# 2014-06-04
i.5. <- c('a', 'b', '')

#all.equal(i.5, i.5.)

##
## 3. Reverse character example
##
i.5c <- interpChar(c('a', 'bc', 'def'), '', 0.3)
# check: 0.7*(total 6 characters) = 4.2 characters
i.5c. <- c('a', 'bc', 'd')

all.equal(i.5c, i.5c.)

# The same thing specified in a list
i.5d <- interpChar(list(c('a', 'bc', 'def'), ''), 0.3)

all.equal(i.5d, i.5c.)

##
## 4. More complicated example
##
xCh <- interpChar(list(c('Do it', 'with R.')),
                 c(0, .5, .9))
# answer
xCh. <- c('', 'with', 'Do ')
# With only one input, it's assumed to be y.
# It is replicated to length(.proportion),
# With nchar = 5, 7, 5, cum = 5, 12, 17.

all.equal(xCh, xCh.)

##
## 5. Still more complicated
##
xC2 <- interpChar(c('a', 'fabulous', 'bug'),
                  c('bigger or', 'just', 'big'),
                  c(.3, .3, 1))

# answer
x.y.longer <- c('bigger or', 'fabulous', 'big')
# use y with ties
# nch smaller 1 4 3
# nch larger 9 8 3
# d.char 8, 4, 0
# cum characters 8, 12, 12
# prop .3, .7, 1
# prop*12 3.6, 8.4, 12
# cum.sm 1, 5, 8
# cum.sm+prop*12 5, 13, 20
# -cum(larger[-1]) 5, 4, 3
xC2. <- c('bigge', 'fabu', 'big')

all.equal(xC2, xC2.)

##
## 6. with one NULL
##
null1 <- interpChar(NULL, 1, 1)

all.equal(null1, 1)

null2 <- interpChar('abc', NULL, .3)

all.equal(null2, 'ab')

##
## 7. length=0
##
log0 <- interpChar(logical(0), 2, .6)

all.equal(log0, 1.2)

##
## 8. Date
##
interpPairs

interpolate between pairs of vectors in a list

Description

This does two things:

1. Computes a proportion interpolation between pairs by passing each pair with proportion to interpChar. interpChar does standard linear interpolation with numerics and interpolates based on the number of characters with non-numerics.

2. Discards rows of interpolants for which proportion is outside validProportion. If object is a list, corresponding rows of other vectors of the same length are also discarded.

NOTE: There are currently discrepancies between the documentation and the code over defaults when one but not both elements of a pair are provided. The code returns an answer. If that's not acceptable, provide the other half of the pair. After some experience is gathered, the question of defaults will be revisited and the code or the documentation will change.

Usage

interpPairs(object, ...)
## S3 method for class 'call'
interpPairs(object,
nFrames=1, iFrame=nFrames,
endFrames=round(0.2*nFrames),
envir = parent.frame(),
pairs=c('1'='\\.0$', '2'='\\.1$', replace0='',
       replace1='.'2', replace2='.'3'),
validProportion=0:1, message0=character(0), ...)
## S3 method for class 'function'
interpPairs(object,
nFrames=1, iFrame=nFrames,
endFrames=round(0.2*nFrames),
envir = parent.frame(),
pairs=c('1'='\\.0$', '2'='\\.1$', replace0='',
       replace1='.'2', replace2='.'3'),
validProportion=0:1, message0=character(0), ...)
## S3 method for class 'list'
interpPairs(object,
.proportion, envir=list(),
pairs=c('1'='\\.0$', '2'='\\.1$', replace0='',
       replace1='.'2', replace2='.'3'),
validProportion=0:1, message0=character(0), ...)
interpPairs

Arguments

object  A call, function, list or data.frame with names possibly matching pairs[1:2]. When names matching both of pairs[1:2], they are converted to potentially common names using sub(pairs[1], pairs[3], ...). When matches are found among the potentially common names, they are passed with .proportion to interpChar to compute an interpolation. The matches are removed and replaced with the interpolant, shortened by excluding any rows for which .proportion is outside validProportion.

Elements with "common names" that do not have a match are replaced by elements with the common names that have been shortened by omitting rows with .proportion outside validProportion. Thus, if x.0 is found without x.1, x.0 is removed and replaced by x.

nFrames  number of distinct plots to create.

iframe  integer giving the index of the single frame to create. Default = nFrames.

An error is thrown if both iframe and .proportion are not NULL.

dendFrames  Number of frames to hold constant at the end.

.proportion  a numeric vector assumed to lie between 0 and 1 specifying how far to go from suffixes[1] to suffixes[2]. For example, if x.0 and x.1 are found and are numeric, x = x.0 + .proportion * (x.1 - x.0). Rows of x and any other element of object of the same length are dropped for any .proportion outside validProportion.

An error is thrown if both iframe and .proportion are not NULL.

eenvi  environment / list to use with codeobject, which can optionally provide other variables to compute what gets plotted; see the example below using this argument.

pairs  a character vector of two regular expressions to identify elements of object between which to interpolate and three replacements.

(1) The first of the three replacements is used in sub to convert each pairs[1:2] name found to the desired name of the interpolate. Common names found are then passed with .proportion to interpChar, which does the actual interpolation.

(2,3) interpPairs also calls checkNames(object, avoid = pairs[c(1, 3, 2, 5)]). This confirms that object has names, and all such names are unique. If object does not have names or has some duplicate names, the make.names is called to fix that problem, and any new names that match pairs[1:2] are modified using sub to avoid creating a new match. If the modification still matches pairs[1:2], it generates an error.

validProportion  Range of values of .proportion to retain, as noted with the discussion of the object argument.

message0  a character string passed to interpChar to improve the value of diagnostic messages

...  optional arguments for sub
Details

*** FUNCTION ***

First interpPairs function looks for arguments firstFrame, lastFrame, and Keep. If any of these are found, they are stored locally and removed from the function. If iframe is provided, it is used with with these arguments plus nframes and endFrames to compute .proportion.

If .proportion is outside validProportion, interpPairs does nothing, returning enquote(NULL).

If any (.proportion) is inside validProportion, interpPairs function next uses grep to look for arguments with names matching pairs[1:2]. If any are found, they are passed with .proportion to interpChar. The result is stored in the modified object with the common name obtained from sub(pairs[i], pairs[3], ...), i = 1, 2.

The result is then evaluated and then returned.

*** LIST ***

1. ALL.OUT: if (none(0<=.proportion<=1)) return 'no.op' = list(fun='return', value=NULL)

2. FIND PAIRS: Find names matching pairs[1:2] using grep. For example, names like x.0 match the default pairs[1], and names like x.1 match the default pairs[1].

3. MATCH PAIRS: Use sub(pairs[i], pairs[3], ...) for i = 1:2, to translate each name matching pairs[1:2] into something else for matching. For example, he default pairs thus translates, e.g., x.0 and x.1 both into x. In the output, x.0 and x.1 are dropped, replaced by x = interpChar(x.0, x.1, .proportion, ...). Rows with .proportion outside validProportion are dropped in x. Drop similar rows of any numeric or character vector or data.frame with the same number of rows as x or .proportion.

4. Add component .proportion to envir to make it available to eval any language component of object in the next step.

5. Loop over all elements of object to create outList, evaluating any expressions and computing the desired interpolation using interpChar. Computing xleft in this way allows xright to be specified later as quote(xleft + xinch(0.6)), for example. This can be used with a call to rasterImageAdj.

6. Let N = the maximum number of rows of elements of outList created by interpolation in the previous step. If .proportion is longer, set N = length(.proportion). Find all vectors and data.frames in outList with N rows and delete any rows for which .proportion is outside validProportion.

7. Delete the raw pairs found in steps 1-3, retaining the element with the target name computed in steps 4 and 5 above. For other elements of object modified in the previous step, retain the shortened form. Otherwise, retain the original, unevaluated element.

Value

a list with elements containing the interpolation results.

Author(s)

Spencer Graves
See Also

interpchar for details on interpolation. compareLengths for how lengths are checked and messages composed and written.
enquote

Examples

```r
###
### 1. interpPairs.function
###
###
## 1.1. simple
##
plot0 <- quote(plot(0))
plot0. <- interpPairs(plot0)
# check
all.equal(plot0, plot0.)

##
## 1.2. no op
##
noop <- interpPairs(plot0, iframe=-1)
# check
all.equal(noop, enquote(NULL))

##
## 1.3. a more typical example
## example function for interpPairs

tstPlot <- function(){
  plot(1:2, 1:2, type='n')
  lines(firstFrame=1:3,
        lastFrame=4,
        x.l=seq(1, 2, .5),
        y.l=x,
        z.0=0, z.1=1,
        txt.l=c('CRAN is', 'good', '...'),
        col='red')
}
tstbo <- body(tstPlot)
iPlot <- interpPairs(tstbo[[2]])
# check
iP <- quote(plot(1:2, 1:2, type='n'))
all.equal(iPlot, iP)
```
iLines <- interpPairs(tstbo[[3]], nFrames=5, iFrame=2)
# check:
#.proportion = (iFrame-firstFrame)/(lastFrame-firstFrame)
# = c(1/3, 0, -1/3)

# if x.0 = 0 and y.0 = 0 by default:
il <- quote(line(x=c(1/3, 0), y=c(1/9, 0), z=c(1/3, 0),
               tst=c('CR', '{')

## This example seems to give the wrong answer
## 2014-06-03: Ignore for the moment
##
#all.equal(iLines, il)

##
## 1.4. Don't throw a cryptic error with NULL
##
ip0 <- interpPairs(quote(text(labels.1=NULL)))

###
### 2. interpPairs.list
###
###
###
### 2.1. (x.0, y.0, x.1, y.1) -> (x,y)
###
tstList <- list(x.0=1:5, y.0=5:9, y.1=9:5, x.1=9,
                 ignore=letters, col=1:5)
xy <- interpPairs(tstList, 0.1)
# check
xy. <- list(ignore=letters, col=1:5,
             x=1:5 + 0.1*(9-1:5),
             y=5:9 + 0.1*(9:5-5:9) )
# New columns, 'x' and 'y', come after
# columns 'col' and 'ignore' already in tstList

all.equal(xy, xy.)

##
## 2.2. Select the middle 2:
##
x=(1-(0,1))3:4+0:1*0=(3,0)
##
xy0 <- interpPairs(tstList[-4], c(-Inf, -1, 0, 1, 2) )
# check
xy0. <- list(ignore=letters, col=3:4, x=c(3,0), y=7:6)

all.equal(xy0, xy0.)

##
## 2.3. Null interpolation because of absence of y.1 and x.0
##
## xy02 <- interpPairs(tstList[c(2, 4)], 0.1)
## check
##### NOT the current default answer; revisit later.
xy02. <- list(y=5:9, x=9)

# NOTE: length(x) = 1 = length(x.1) in testList
#all.equal(xy02, xy02.)

##
## 2.4. Select an empty list (make sure this works)
##
## x0 <- interpPairs(list(), 0:1)
## check
## x0. <- list()
names(x0.) <- character(0)

all.equal(x0, x0.)

##
## 2.5. subset one vector only
##
## xyz <- interpPairs(list(x=1:4), c(-1, 0, 1, 2))
## check
xyz. <- list(x=2:3)

all.equal(xyz, xyz.)

##
## 2.6. with elements of class call
##
## xc <- interpPairs(list(x=1:3, y=quote(x+sin(pi*x/6))), 0:1)
## check
xc. <- list(x=1:3, y=quote(x+sin(pi*x/6)))

all.equal(xc, xc.)

##
## 2.7. text
##
# 2 arguments
j.5 <- interpPairs(list(x.0='', x.1=c('a', 'bc', 'def')), 0.5)
# check
j.5. <- list(x=c('a', 'bc', ''))
all.equal(j.5, j.5.)

##
## 2.8. text, 1 argument as a list
##
j.50 <- interpPairs(list(x.1=c('a', 'bc', 'def')), 0.5)
# check
all.equal(j.50, j.5.)

##
## 2.9. A more complicated example with elements to eval
##
logo.jpg <- paste(R.home(), "doc", "html", "logo.jpg",
sep = .Platform$file.sep)
if(require(jpeg)){
  Rlogo <- readJPEG(logo.jpg)
  # argument list for a call to rasterImage or rasterImageAdj
  RlogoLoc <- list(image=Rlogo,
                   xleft.0 = c(NZ=176.5, CH=172, US=171, CN=177, RU=9.5, UK=8),
                   xleft.1 = c(NZ=176.5, CH=9, US=-73.5, CN=125, RU=37, UK=2),
                   ybottom.0=c(NZ=-37, CH=-34, US=-34, CN=-33, RU=48, UK=47),
                   ybottom.1=c(NZ=-37, CH=47, US=46, CN=32, RU=55.6, UK=55),
                   xright=quote(xleft+xinch(0.6)),
                   ytop = quote(ybottom+yinch(0.6)),
                   angle.0 =0,
                   angle.1 =c(NZ=0, CH=3*360, US=5*360, CN=2*360, RU=360, UK=360)
  )
  RlogoInterp <- interpPairs(RlogoLoc,
                           .proportion=rep(c(0, -1), c(2, 4))
  )
  # check
  all.equal(names(RlogoInterp),
            c('image', 'xright', 'ytop', 'xleft', 'ybottom', 'angle'))

  # NOTE: 'xleft', and 'ybottom' were created in interpPairs,
  # and therefore come after 'xright' and 'ytop', which were
  # already there.

##
## 2.10. using envir
##
RlogoDiag <- list(x0=quote(Rlogo.$xleft),
                   y0=quote(Rlogo.$ybottom),
                   x1=quote(Rlogo.$xright),
                   y1=quote(Rlogo.$ytop),
                   xright=quote(xleft+xinch(0.6)),
                   ytop = quote(ybottom+yinch(0.6)),
                   angle.0 =0,
                   angle.1 =c(NZ=0, CH=3*360, US=5*360, CN=2*360, RU=360, UK=360)
match.data.frame

Identify the row of \( y \) best matching each row of \( x \)

```r

yl=quote(Rlogo.$ytop )

RlogoD <- interpPairs(RlogoDiag, .p=1,
                    envir=list(Rlogo.=RlogoInterp )

all.equal(RlogoD, RlogoDiag)
}
##
## 2.11. assign; no interp but should work
##
tstAsgn <- as.list(quote(op <- (1:3)*2))
tstAsgn <- interpPairs(tstAsgn, 1)

# check
intA. <- tstAsgn
names(intA.) <- c(‘X’, ‘X.3’, ‘X.2’)

all.equal(intAsgn, intA.)

# op <- par(…)
tstP <- quote(op <- par(mar=c(5, 4, 2, 2)+0.1))
tstP <- as.list(tstP)
intPar <- interpPairs(tstPar, 1)

# check
intP. <- list(quote(‘<-’), quote(op),
              quote(par(mar=c(5, 4, 2, 2)+0.1)) )
names(intP.) <- c(‘X”, ‘X.3’, ‘X.2’)

all.equal(intPar, intP.)

intP. <- interpPairs(tstP)
all.equal(intP., tstP)

##
## NULL
##

all.equal(interpPairs(NULL), quote(NULL))
```
**Description**

For each row of \(x\[, by.x\], find the best matching row of \(y\[, by.y\], with the best match defined by grep. and split.

grep. and split must either be `missing` or have the same length as by.x and by.y. If grep.\([i\] and split\([i\) are NA, do a complete match of \(x\[, by.x[i]\] and \(y\[, by.y[i]\]. Otherwise, for each row \(j\), look for a match for `strsplit(x[j, by.x[i]], split[i][1][1]` among `strsplit(y[, by.y[i]], split[i]).` See details.

**Usage**

```r
match.data.frame(x, y, by, by.x=by, by.y=by, grep., split, sep=':')
```

**Arguments**

- \(x\), \(y\) : data.frames
- by, by.x, by.y : names of columns of \(x\) and \(y\) to match.
- grep. : a character vector of the type of match for each element of by.x and by.y. If NA, require a perfect match. Alternatives are `grep` and `agrep` to find a match for the first segment in strsplit(x, split=split\([i\]) among any of the segments of strsplit(y, split=split\([i\]). Use `fixed=TRUE` with the calls to these functions.
- NOTE: These alternatives are not examined if a unique match is found between \(x\[, by.x[is.na(grep.) & is.na(split)]\) and the corresponding columns of \(y\).
- split : A character vector of split characters to pass to `strsplit`; `strsplit` is not called if is.na(split).
- sep : a sep argument to use with `paste` to produce a matching key for the columns of \(x\) and \(y\) for which perfect matches are required. If(missing(sep) && !is.na(grep.)) sep <- `'` except where grep. = NA.

**Details**

1. Check by.x, by.y, grep. and split. If((missing(by.x) | missing(by.y)) && missing(by)) by <- names(x)
2. fullMatch <- (is.na(grep.) & is.na(split)). Create keyfx and keyfy by by pasting columns of \(x\[, by.x[fullMatch]\] and \(y\[, by.y[fullMatch]\]. Also create x. and y. = `strsplit` of \(x\[, by.x[fullMatch]\] and \(y\[, by.y[fullMatch]\].
3. Iterate over rows of \(x\) looking for the best match. This includes an inner loop over columns of \(x\[, by.x[fullMatch]\], stopping on the first unique match. Return (-1) if no unique match is found.

**Value**

an integer vector of length nrow(x) containing the index of the best matching row of \(y\) or NA if no adequate match was found.

**Author(s)**

Spencer Graves
See Also

`strsplit`, `is.na`, `grep`, `agrep`, `match`, `row.match`, `join`, `match_df`, `classify`

Examples

```r
newdata <- data.frame(state=c("AL", "MI","NY"),
  surname=c("Rogers", "Rogers", "Smith"),
  givenName=c("Mike R.", "Mike K.", "Al"),
stringsAsFactors=FALSE)
reference <- data.frame(state=c("NY", "NY", "MI", "AL", "NY", "MI"),
  surname=c("Smith", "Rogers", "Rogers (MI)",
  "Rogers (AL)", "Smith", 'Jones'),
  givenName=c("John", "Mike", "Mike", "Mike",
  "T. Albert", 'Al Thomas'),
stringsAsFactors=FALSE)
newInRef <- match.data.frame(newdata, reference,
grep.=c(NA, 'agrep', 'agrep'))

all.equal(newInRef, c(4, 3, 5))
```

Description

Use `parsename` to split a name into `surname` and `givenName`, the look for matches in table.

Usage

```r
matchName(x, data, Names=1:2,
  nicknames=matrix(character(0), 0, 2),
  namesNotFound="attr.replacement", ...)
matchName1(x1, data, name=data[, 1],
  nicknames=matrix(character(0), 0, 2), ...)
```

Arguments

- **x**
  - One of the following:
    - A character matrix or `data.frame` with the same number of rows as `data`. The best partial match is sought in `Names`. The algorithm stops when a unique match is found; any remaining columns of `x` are then ignored. Any nicknames are ignored for the first column but not for subsequent columns.
    - A character vector whose length matches the number of rows of `data`. This will be replaced by `parsename(x)`.

- **data**
  - a character matrix or a `data.frame`. If `surname` and `givenName` are character vectors of names, their length must match the number of rows of `data`. 
### matchName

**Names**

One of the following in which matches for `x` will be sought:

- A character vector or matrix or a `data.frame` for which `NROW(Names) == nrow(data)`.
- Something to select columns of data to produce a character vector or matrix or `data.frame` via `data[, Names]`. In this case, accents will be stripped using `subNonStandardNames`.

**nicknames**

A character matrix with two columns, each row giving a pair of names like "Pete" and "Peter" that should be regarded as equivalent if no exact match(es) is(are) found.

... optional arguments passed to `subNonStandardNames`

**x1**

A character vector of names to match `name`.

NOTE: `matchName` calls `subNonStandardNames`, but `matchName1` does not. Thus, `x1` is assumed to NOT to contain characters not in standard English.

**name**

A character vector or matrix for which `NROW(name) == nrow(data)`.

NOTE: `matchName` calls `subNonStandardNames`, but `matchName1` does not. Thus, `name` is assumed to NOT to contain characters not in standard English.

**namesNotFound**

A character vector passed to `subNonStandardNames` and used to compute any "namesNotFound" attribute of the object returned by `parseName`.

### Details

*** 1. `matchName(x, data, Names, nicknames, ...)`:  
   1.1. if(length(dim(x)<2))x <- `parseName`(x, ...)  
   1.2. x1 <- `matchName1`(x[, 1], cata, Names[1], ...)  
   1.3. For any component `i` of `x1` with multiple rows, let `x1i <- `matchName1`(x[i, 2], x1[[i]], Name[i-1], nicknames=nicknames, ...). If nrow(x1i)>0, x1[[i]] <- x1i; else leave unchanged.  
   1.4. return x1

==============

*** 2. `matchName1(x1, data, name, nicknames, ...)`:  
   2.1. If name indicates a column of data, replace with `data[, name]`.  
   2.2. xsplit <- `strsplit`(x1, ' ')  
   2.3. nx <- length(x1); xlist <- vector(nx, mode='list')  
   2.4. for(j in 1:nx)  
   2.5. xj <- xsplit[[j]]  
   2.6. let jd = the subset of names that match xj or subNonStandardNames(xj) or nicknames of xj; xlist[j] <- jd.  
   2.7. return xlist

### Value

`matchName` returns a list of the same length as `x`, each of whose components is object obtained as a subset of rows of `data` or NULL if no acceptable matches are found. The list may have an attribute "namesNotFound" as determined per the argument of that name.

`matchNames1` returns a list of vectors of integers for subsets of `data` matching `x1`. 
matchName

Author(s)
Spencer Graves

See Also

parseName subNonStandardNames

Examples

```r
## 1. Names to match exercising many possible combinations
## of surname with 0, 1, >1 matches possibly after
## replacing with subNonStandardNames
## combined with possibly multiple givenName combinations
## with 0, 1, >1 matches possibly requiring replacing with
## subNonStandardNames or nicknames
##
## # NOTE: "-" could also be "e" with an accent;
## # not included with this documentation, because
## # non-English characters generate warnings in standard tests.
Names2mtch <- c("Andr_Bruce C_rdenas", "Dolores Ella Feinstein",
                  "George Homer", "Inez Jane Kappa", "Luke Michael Noel",
                  "Oscar Papa", "Quincy Ra_l Stevens",
                  "Thomas U. Vel_zquez", "William X. Young",
                  "Zebra")

## 2. Data = matrix(..., byrow=TRUE) to exercise the combinations
## the combinations from 1
##
## Data1 <- matrix(c("Feld", "Don", "789",
                  "C_rdenas", "Don", "456",
                  "C_rdenas", "Andre B.", "123",
                  "Smith", "George", "aaa",
                  "Young", "Bill", "369"),
                  ncol=3, byrow=TRUE)
Data1. <- subNonStandardNames(Data1)

## 3. matchName1
##
## parceNm1 <- parseName(Names2mtch)
match1.1 <- matchName1(parceNm1[, 'surname'], Data1.)

# check
match1.1s <- vector('list', 10)
match1.1s[[1]] <- 2:3
match1.1s[[9]] <- 5
names(match1.1s) <- parceNm1[, 'surname']

all.equal(match1.1, match1.1s)
```
## 4. matchName1 with name = multiple columns

```r
match1.2 <- matchName1(c('Cardenas', 'Don'), Data1,
  name=Data1[, 1:2])
```

# check
```r
match1.2a <- list(Cardenas=2:3, Don=1:2)
```

all.equal(match1.2, match1.2a)

## 5. matchName

```r
nickNames <- matrix(c("William", "Bill"), 1, byrow=TRUE)
```

```r
match1 <- matchName2mtch(Data1, nicknames=nickNames)
```

# check
```r
match1a <- list("Cardenas, Andre Bruce"=Data1[3, , drop=FALSE ],
  "Feinstein, Dolores Ella"=NULL,
  "Homer, George"=NULL, "Kappa, Inez Jane"=NULL,
  "Noel, Luke Michael"=NULL, "Papa, Oscar"=NULL,
  "Stevens, Quincy Raul"=NULL,
  "Velazquez, Thomas U."=NULL,
  "Young, William X."=Data1[5, , drop=FALSE],
  "Zebra"=NULL)
```

all.equal(match1, match1a)

## 6. namesNotFound

```r
tstNotFound <- matchName('xx_x', Data1)
```

# check
```r
tstNF <- list('xx_x'=NULL)
```

```r
attr(tstNF, 'namesNotFound') <- 'xx_x'
```

all.equal(tstNotFound, tstNF)

## 7. matchName(NULL) to simplify use

```r
mtchNULL <- matchName(NULL, Data1)
```

all.equal(mtchNULL, NULL)

---

matchQuote Match isolated quotes across records
matchQuote

Description

Look for unmatched quotes in a character vector. If found, look for a matching quote starting the
next character string in the vector, possibly after a blank line. If found, merge the two strings and
return the resulting shortened character vector.

Usage

matchQuote(x, Quote='''', sep=' ', maxChars2append=2, ...)

Arguments

  x            a character vector to scan for unmatched Quotes.
  Quote        the Quote character that should appear in pairs
  sep          sep argument passed to paste to combine pairs of successive lines with un-
               matched quotes.
  maxChars2append maximum number of characters in the following string to concatonate two adja-
               cent strings (possibly separated by a blank line) with unmatched Quotes.
  ...          optional arguments for gsub

Details

This function was written to help parse data from the US Department of Health and Human Services
on cyber-security breaches affecting 500 or more individuals. As of 2014-06-03 the csv version of
these data included commas in quotes that are not sep characters, quotes that are not matched, lines
with zero characters, followed by lines with 3 characters being a quote and a comma. This function
was written to drop the blank lines and append the quote-comma line to the preceeding line so it
contained matching quotes.

Value

The input character vector possibly shortened with the following attributes explaining what was
found:

  • unmatchedQuotes indices of the input x with an unmatched Quote.
  • blankLinesDropped indices of the input x that were dropped because they (1) followed an
    unmatched Quote and (2) contained no non-blank characters.
  • quotelinesAppended indices of the input x that were concatonated with a preceeding line be-
    cause the two lines contained unmatched Quote characters, and concatonating them produced
    a line with all Quotes matched.
  • ncharsAppended an integer vector of the same length as quotelinesConcatonated giving
    the number of characters in the second line concatonated onto the previous line.

Author(s)

Spencer Graves
See Also

strsplit1 delimMatch

Examples

```r
chvec <- c('abc', 'de"f", "", "h", matched"quotes", "")
ch. <- matchQuote(chvec)

# check
chv. <- c('abc', 'de"f", "h", matched"quotes", "")
attr(chv., 'unmatchedQuotes') <- c(2, 4, 5)
attr(chv., 'blankLinesDropped') <- 3
attr(chv., 'quoteLinesAppended') <- 4
attr(chv., 'ncharsAppended') <- 2
all.equal(ch., chv.)
```

mergeUShouse.senate

Expand a dataset on some members of the US Congress to the entire membership

Description

Merge a `data.frame` regarding some members of the US Congress with a `data.frame` with general information on all members.

Usage

```r
mergeUShouse.senate(x, UScongress=UShouse.senate(),
    newrows="amount0",
    default=list(member=FALSE, amount=0, vote="notEligible",
    incumbent=TRUE))
```

Arguments

- **x**
  - a `data.frame` to be merged with UScongress
- **UScongress**
  - a `data.frame` to be merged with x.
- **newrows**
  - name of a logical column to add that is TRUE for rows added to x and FALSE otherwise.
- **default**
  - default values for columns of x identified by `regexpr(names(default)[i], tolower(names(x)))`. 
mergeUShouse.senate

Details

1. keyx <- with(x, paste(houseSenate, state, District, sep=":"))
2. keyy <- with(UScongress(houseSenate, state, District, sep=":"))
3. notx <- !is.element(keyy, keyx)
4. Y <- UScongress[notx, ]
5. add default columns to Y
6. if(!newrows is not in names(x))x <- cbind(x, newrows=FALSE)
7. Y[, newrows] <- TRUE
8. xY <- rbind(x, Y[c(names(x))]
9. replace 'Democrat' with 'Democratic' in xY[["Party"]]
10. Look for NAs in "incumbent" who are nevertheless in UScongress; fix. Thus, if x[['incumbent']] is TRUE or FALSE, this value is not checked in UScongress; it's checked only if NA. The check consists of comparing names for a given Office:state:district between strsplit(x[['surname']], ','|[1]|1 and strsplit(UScongress[['surname']], ','|[1]|1] and similarly for givenName. This allows 'Rogers' in x[['surname']] to match 'Rogers (AL)' in UScongress[['surname']], etc. The algorithm is not perfect, but errors should be rare – and could be fixed manually.

Value

a data.frame combining x and UScongress as desired

Author(s)

Spencer Graves

See Also

merge UShouse.senate

Examples

tst <- data.frame(Office=factor(rep(c('House', 'Senate'), c(4, 2))),
   State=factor(c('Missouri', 'Minnesota', 'Tennessee',
      'New York', rep('South Carolina', 2))),
   state=factor(c('MO', 'MN', 'TN', 'NY', 'SC', 'SC'))),
   district=as.character(c(4, 1, 8, 18, 2, 3)),
   surname=c('Hartzler', 'Walz', 'Fincher', 'Maloney',
      'Graham', 'DeMint'),
   givenName=c('Vicky', 'Timothy J.', 'Stephen Lee',
      'Sean Patrick', 'Lindsey', 'Jim'),
   Party=c('Republican', 'Democrat', 'Republican', 'Democrat',
      'Republican', 'Democrat'),
   CommitteeMember=rep(c(TRUE, FALSE), c(4, 2)),
   amount=c(5000, 2000, 29500, 1000, 1000, 11500),
   xvote=c('Y', 'N', 'Y', 'Y', 'notEligible', 'notEligible'),
   incumbent=NA, stringsAsFactors=FALSE )
if(!fda::CRAN()){
  tst2 <- mergeUShouse.senate(tst)

  # A couple of simple tests; don't test too much,
  # because the results of UShouse.senate change,
  # and we don't want this test to fail
  # due to changes that don't affect Ecdat code
  tst3 <- tst2[tst2$amount0, c(1, 4:6, 8:10)]
  row.names(tst) <- row.names(tst3)

  ## Not run:
  all.equal(tst[c(1, 4:6, 8:10)], tst3)

  ## End(Not run)
  # tst3[2] == state = factor with 56 levels,
  # and tst[2] only has 5; compare without this
}

mergeVote

Merge roll call vote record with a data.frame containing other information. The vote records are typically incomplete, so match first on houseSenate and surname. If this match is incomplete, try using givenName. If that fails, try state and district, which may not always be present in vote.

Usage

mergeVote(x, vote, Office="House", vote.x, check.x=TRUE)

Arguments

x
  a data.frame whose columns include Office, surname, and givenName.

vote
  a data.frame with column names which when forced tolower would match surname, givenname, and vote. However, the givenname may not be complete, so use it only if the surname is not sufficient.

Office
  Either "House" or "Senate"; ignored if vote includes a column Office.

vote.x
  name of a column of x containing a vote to be updated with the vote column of the vote data.frame. If missing and x has a column with a name matching "vote", then vote.x is that column. If missing but x has no such column, then append a column to x with the name of the vote column of the vote data.frame.

check.x
  logical: If TRUE, check for rows of x[, vote.x] that are NOT in vote and throw an error if found.
mergeVote

Details

1. Parse `vote.x` to get the name of the column of `x` into which to write the vote column of the vote data.frame.

2. If the vote data.frame contains a column `Office`, ignore the `Office` argument. Otherwise, add the argument `houseSenate` as a column of vote.

3. Create `keyx <- with(x, paste(Office, surname, sep=".")))`, `keyx2 <- paste(keyx, givenName, sep="."))`, `keyx. <- paste(houseSenate, state, district, sep=".")`, and similarly `keyv, keyv2, and keyv.` from vote.

4. Look for `keyx` in `keyx`. When a unique match is found, transfer the vote the vote column of `x`. When no match is found, try for `keyv2` in `keyx2` or `keyv.` in `keyx`. If those fail, print an error message with the information from vote on all failures and ask the user to add state and district information.

5. If `check.x`, check for rows in `x[, vote.x]` that are NOT "notEligible" but are also not in vote: Throw an error if any are found.

Value

a data.frame with the same columns as `x` with its vote column modified per the vote argument.

Author(s)

Spencer Graves

See Also

mergeUShouse.senate

Examples

```r
##
## 1. Test good cases
##

votetest <- data.frame(
  surname=c("Smith", "Jones", "Graves", "Jsn", "Jsn", "Gay"),
  givenName=c("Sam", "", "", "John", "John", ""),
  vote=factor(c("Y", "N", "abstain", "Y", "Y", "Y")),
  state=factor(rep(c("CA", "", "SC", "NY"), c(1, 2, 1, 2))),
  district=factor(rep(c("13", "1", "2", "1"), c(1, 2, 1, 2))),
  stringsAsFactors=FALSE)

x1 <- data.frame(
  Office=factor(rep(c("House", "Senate"), e=8)),
  state=factor(rep(c("NY", "SC", "SD", "CA", "AK", "AR", "NY", "NJ"), 2),
  District=factor(rep(c("2", "2", "At Large", "13", "1", "9", "1", "3"), 2),
  don=1:16, stringsAsFactors=FALSE)
```
x1. <- mergeVote(x1, votetst)

x2 <- cbind(x1, votex=factor(rep(
  c('Y', 'notEligible', 'Y', 'N', 'abstain', 'Y', 'notEligible'),
  c(2,1,1,1,1,1,9) ) ))

allexport(x1., x2)

## 2. Test a case with a vote error in x
##
xla <- cbind(x1, voteerr=rep(
  c('notEligible', 'Y', 'notEligible'), c(7, 1, 8)))
xla. <- try(mergeVote(x1a, votetst))
class(x1a.)=='try-error'

---

**missing0**  
*Missing or length 0*

**Description**  
TRUE if x is missing or if length(x) is 0.

**Usage**  
`missing0(x)`

**Arguments**  
- `x`  
a formal argument as for `missing`

**Details**  
Only makes sense called from within another function

**Value**  
*logical*: TRUE if x is `missing` or if length(x) is 0.
nchar0

Author(s)
Spencer Graves

See Also

missing

Examples

```
tstFn <- function(x) missing0(x)
  # missing
  all.equal(tstFn(), TRUE)

  # length 0
  all.equal(tstFn(logical()), TRUE)

  # supplied
  all.equal(tstFn(1), FALSE)
```

nchar0

Zero characters or NULL

Description

Returns TRUE if (is.null(x) || (length(x) == 0) || (max(nchar(x)) == 0)).

Usage

nchar0(x, ...)

Arguments

x a character vector or something that can be coerced to mode character
...
optional arguments to be passed to nchar

Value

TRUE if x is either NULL or max(nchar(x)) == 0. FALSE otherwise.

Author(s)

Spencer Graves
See Also

nchar

Examples

all.equal(nchar0(NULL), TRUE)

all.equal(nchar0(character(0)), TRUE)

all.equal(nchar0(character(3)), TRUE)

all.equal(nchar0(c('a', 'c')), FALSE)

parseCommas

Convert character string with Dollar signs and commas to numerics

Description

as.numeric of character strings after suppressing commas and dollar signs. This is a generalization of parseDollars.

Usage

parseCommas(x, pattern='\$|,', replacement='', acceptableErrorRate=0, ...)

## Default S3 method:
parseCommas(x, pattern='\$|,', replacement='', acceptableErrorRate=0, ...)

## S3 method for class 'data.frame'
parseCommas(x, pattern='\$|,', replacement='', acceptableErrorRate=0, ...)

Arguments

x vector of character strings to be converted to numerics

pattern regular expression to be replaced by replacement

replacement Character string to substitute for each occurrence of pattern

acceptableErrorRate number indicating the proportion of new NAs to that can be introduced and still assume it's numeric

... optional arguments to pass to gsub
parseDollars

Details

as.numeric(gsub(x, ..., ))

The data.frame method outputs another data.frame with character or factor columns connverted to numerics using parseDollars whenever that can be done without creating NAs.

Value

Numeric vector converted from the character strings in x or a data.frame with columns that are obviously numbers in character format converted to numerics.

Author(s)

Spencer Graves

See Also

gsub asNnumeric parseDollars

Examples

##
## 1.  a character vector
##
X2 <- c('-$2,500', '$5,000.50')
x2 <- parseDollars(X2)

all.equal(x2, c(-2500, 5000.5))

##
## A data.frame
##
chDF <- data.frame(let=letters[1:2], Dol=X2, dol=x2)
numDF <- parseCommas(chDF)

chkDF <- chDF
chkDF$Dol <- x2

all.equal(numDF, chkDF)

<table>
<thead>
<tr>
<th>parseDollars</th>
<th>Convert character string with Dollar signs and commas to numerics</th>
</tr>
</thead>
</table>

Description

as.numeric of character strings after suppressing commas and dollar signs. This is a special case of parseCommas.
parseDollars

Usage

parseDollars(x, pattern='\$|,', replacement=' ', ...)

Arguments

x vector of character strings to be converted to numerics
pattern regular expression to be replaced by replacement
replacement Character string to substitute for each occurrence of pattern
... optional arguments to pass to gsub

Details

as.numeric(gsub(x, ...)). See also parseCommas.

Value

Numeric vector converted from x.

Author(s)

Spencer Graves

See Also

gsub as.numeric parseCommas

Examples

##
## 1. a character vector
##
X2 <- c('-$2,500', '$5,000.50')
x2 <- parseDollars(X2)

all.equal(x2, c(-2500, 5000.5))

##
## A data.frame
##
chDF <- data.frame(let=letters[1:2], Dol=X2, dol=x2)
numDF <- parseCommas(chDF)

chkDF <- chDF
chkDF$Dol <- x2

all.equal(numDF, chkDF)
parseName  

Parse surname and given name

**Description**

Identify the presumed surname in a character string assumed to represent a name and return the result in a character matrix with "surname" followed by "givenName". If only one name is provided (without punctuation), it is assumed to be the givenName; see Wikipedia, "Given name" and "Surname".

**Usage**

```r
parseName(x, surnameFirst=(median(regexpr(',', x))>0),
           suffix=c('Jr.', 'I', 'II', 'III', 'IV', 'Sr.', 'Dr.', 'Jr', 'Sr'),
           fixNonStandard=subNonStandardNames,
           removeSecondLine=TRUE,
           namesNotFound="attr.replacement", ...)
```

**Arguments**

- `x` a character vector
- `surnameFirst` logical: If TRUE, the surname comes first followed by a comma (","), then the given name. If FALSE, parse the surname from a standard Western "John Smith, Jr." format. If missing(surnameFirst), use TRUE if half of the elements of `x` contain a comma.
- `suffix` character vector of strings that are NOT a surname but might appear at the end without a comma that would otherwise identify it as a suffix.
- `fixNonStandard` function to look for and repair nonstandard names such as names containing characters with accent marks that are sometimes mangled by different software. Use `identity` if this is not desired.
- `removeSecondLine` logical: If TRUE, delete anything following "\n" and return it as an attribute "secondLine".
- `namesNotFound` character vector passed to subNonStandardNames and used to compute any "namesNotFound" attribute of the object returned by parseName.
- `...` optional arguments passed to `fixNonStandard`

**Details**

If `surnameFirst` is FALSE:

1. If the last character is ")" and the matching "(" is 3 characters earlier, drop all that stuff. Thus, "John Smith (AL)" becomes "John Smith".
2. Look for commas to identify a suffix like Jr. or III; remove and call the rest `x2`.
3. `split <- strsplit(x2, "\")`
4. Take the last as the surname.

5. If the "Surname" found per 3 is in suffix, save to append it to the givenName and recurse to get the actual surname.

NOTE: This gives the wrong answer with double surnames written without a hyphen in the Spanish tradition, in which, e.g., "Anistasio Somoza Debayle", "Somoza Debayle" give the (first) surnames of Anistasio’s father and mother, respectively: The current algorithm would return "Debayle" as the surname, which is incorrect.

6. Recompose the rest with any suffix as the givenName.

**Value**

a character matrix with two columns: surname and givenName.

This matrix also has a "namesNotFound" attribute if one is returned by subNonStandardNames.

**Author(s)**

Spencer Graves

**See Also**

`strsplit identity subNonStandardNames`

**Examples**

```r
## 1. Parse standard first-last name format

# tstParse <- c('Joe Smith (AL)', 'Teresa Angelica Sanchez de Gomez',
# 'John Brown, Jr.', 'John Brown Jr.',
# 'John W. Brown III', 'John Q. Brown, Jr.',
# 'Linda Rosa Smith-Johnson', 'Anastasio Somoza Debayle',
# 'Raúl Velázquez', 'Sting', 'Colette', '

colnames(tstParse) <- c('Surname', 'GivenName')

taxtParse2 <- matrix(c('Smith', 'Joe', 'Gomez', 'Teresa Angelica Sanchez de',
# 'Brown', 'John, Jr.', 'Brown', 'John, Jr.',
# 'Brown', 'John W., III', 'Brown', 'John Q., Jr.',
# 'Smith-Johnson', 'Linda Rosa', 'Debayle', 'Anastasio Somoza',
# 'Velázquez', 'Raul', '', 'Sting', 'Colette', ''),
# ncol=2, byrow=TRUE)
# NOTE: The 'Anastasio Somoza Debayle' is in the Spanish tradition
# and is handled incorrectly by the current algorithm.
# The correct answer should be "Somoza Debayle", "Anastasio".
# However, fixing that would complicate the algorithm excessively for now.
colnames(tstParse2) <- c('Surname', 'GivenName')

equal(parses, tstParse2)
```
## 2. Parse "surname, given name" format

tst3 <- c('Smith (AL), Joe', 'Sanchez de Gomez, Teresa Angelica',
          'Brown, John, Jr.', 'Brown, John W., III', 'Brown, John Q., I',
          'Smith-Johnson, Linda Rosa', 'Somoza Debayle, Anastasio',
          'Velazquez, Raúl', 'Sting, Colette,')
tst4 <- parseName(tst3)

tst5 <- matrix(c('Smith', 'Joe', 'Sanchez de Gomez', 'Teresa Angelica',
                 'Smith-Johnson', 'Linda Rosa', 'Somoza Debayle', 'Anastasio',
                 'Velazquez', 'Raúl', '', 'Sting', 'Colette', ''),
                 ncol=2, byrow=TRUE)
colnames(tst5) <- c("surname", 'givenName')

equal(tst4, tst5)

## 3. secondLine

L2 <- parseName(c('Adam
2nd line', 'Ed
--Vacancy', 'Frank'))

# check
L2. <- matrix(c('', 'Adam', '', 'Ed', '', 'Frank'),
              ncol=2, byrow=TRUE)
colnames(L2.) <- c('surname', 'givenName')
attr(L2., 'secondLine') <- c('2nd line', 'Vacancy', NA)
equal(L2, L2.)

## 4. Force surnameFirst when in a minority

snf <- c('Sting', 'Madonna', 'Smith, Al')
SNF <- parseName(snf, surnameFirst=TRUE)

# check
SNF2 <- matrix(c('Sting', '', 'Madonna', 'Smith', 'Al'),
               ncol=2, byrow=TRUE)
colnames(SNF2) <- c('surname', 'givenName')
equal(SNF, SNF2)

## 5. nameNotFound

noSub <- parseName('xx_x')
# check
noSub. <- matrix(c('', 'xx_x'), 1)
colnames(noSub.) <- c('surname', 'givenName')
attr(noSub., 'namesNotFound') <- 'xx_x'

all.equal(noSub, noSub.)

---

### Description

***NOTE: THIS IS A PRELIMINARY VERSION OF THIS FUNCTION; ***NOTE: IT MAY BE CHANGED OR REMOVED IN A FUTURE RELEASE.

ping a Uniform resource locator (URL) or Internet Protocol (IP) address.

NOTE: Some Internet Service Providers (ISPs) play games with "ping". That makes the results of Ping unreliable.

### Usage

```r
Ping(url, pingArgs=' ', warn=NA,
    show.output.on.console=FALSE)
```

### Arguments

- **url**
  - a character string of a URL or IP address to ping. If `url` is a vector of length greater than 1, only the first component is used.

- **pingArgs**
  - arguments to pass to the `ping` command of typical operating systems via `pingResult <- system(paste('ping', pingArgs, url), intern=TRUE, ...)`

- **warn**
  - value for `options('warn')` during the call to `system`. `NA` to not change `options('warn')` during this call.

- **show.output.on.console**
  - argument for `system`

### Details

1. `urlSplit0 <- strsplit(url, ':/\')[[1]]`
2. `urlS0 <- urlSplit0[min(2, length(urlSplit0))]`
3. `host <- strsplit(urlS0, '/')[[1]][1]`
4. `pingCmd <- paste('ping', pingArgs, host)`
5. `system(pingCmd, intern=TRUE, ...)"
Value

list with the following components:

- `rawResults` character vector of the raw results from the ping command
- `rawNumbers` numeric vector of the times measured
- `counts` numeric vector of numbers of packets sent, received, and lost
- `p.lost` proportion lost = lost / sent
- `stats` numeric vector of min, avg (mean), max, and mdev (standard deviation) of the measured round trip times

Author(s)

Spencer Graves

See Also

`system`, `options`

Examples

```r
## Some ISPs play games with ping.
## Therefore, the results are not reliable.
##
## Not run:
##
## good
##
## (google <- Ping('http://google.com/ping works on host not pages'))

\dontshow{stopifnot}
with(google, (counts[1]>0) && (counts[3]<1))
\dontshow{}

## ping oops \--- at one time, this failed.
##   However, with some ISPs, it works, so don't test it.
##
## (couldnotfindhost <- Ping('oops'))

\dontshow{stopifnot}
with(couldnotfindhost, 
   length(grep('could not find host', rawResults))>0)
\dontshow{}

## impossible, but not so obvious
##
```
pmatch2 returns a list of the positions of matches or partial matches of \( x \) in \( \text{table} \).
This does sloppy matching to find "Peter" to match "Pete" only if "Pete" is not in \( \text{table} \), and we want "John Peter" if neither "Pete" nor "Peter" are in \( \text{table} \).

**Usage**

\[
\text{pmatch2}(x, \text{table})
\]

**Arguments**

- \( x \)  
  the values to be matched
- \( \text{table} \)  
  the values to be matched against

**Details**

1. \( nx \leftarrow \text{length}(x) \); \( \text{out} \leftarrow \text{vector}(nx, \text{"list"); names(out) \leftarrow x } \)
2. for(i in seq(length=nx)):
3. \( xi \leftarrow \text{which}(x[i]) \text{ in } \text{table} \)
4. if(length(xi)<1) \( xi \leftarrow \text{grep(paste0('^', x[i]), table)} \)
5. if(length(xi)<1)xi <- grep(x[i], table).  
6. out[i]<-xi

**Value**

A list of integer vectors indicating the positions in \( \text{table} \) matching each element of \( x \)

**Author(s)**

Spencer Graves

**See Also**

\text{match pmatch grep matchName}
Examples

```r
##
## 1. common examples
##
x2match <- c('Pete', 'Peter', 'Ma', 'Mo', 'Paul',
              'Cardenas')

tbl <- c('Peter', 'Mary', 'Martha', 'John Paul', 'Peter',
         'Cardenas', 'Cardenas')

x2mtchd <- pmatch2(x2match, tbl)

# answer
x2mtchd. <- list(Pete=c(1, 5), Peter=c(1, 5), Ma=2:3,
                  Mo=integer(0), Paul=4, Cardenas=6:7)

all.equal(x2mtchd, x2mtchd.)

##
## 2. strange cases that caused errors and are now warnings
##
huh <- pmatch2("7", tbl)

# answer
huh. <- list("7"=integer(0))

all.equal(huh, huh.)
```

qqnorm2

Normal Probability Plot with Multiple Symbols

Description

Create a normal probability plot with different symbols for the values of another variable. `qqnorm2` produces an object of class `qqnorm2`, whose plot method produces the plot.

Usage

```r
qqnorm2(y, z, plot.it=TRUE, datax=TRUE, pch=NULL, ...)
## S3 method for class 'qqnorm2'
plot(x, y, ...)
## S3 method for class 'qqnorm2'
lines(x, ...)
## S3 method for class 'qqnorm2'
points(x, ...)
```
Arguments

- **y**: For `qqnorm2`, `y` is a numeric vector for which a normal probability plot is desired. For `plot.qqnorm2`, `y` is ignored; it is included, because the generic `plot` function requires it.

- **z**: A variable to indicate different plotting symbols.

- **plot.it**: logical: Should the result be plotted?

- **datax**: The `datax` argument of `qqnorm`: If TRUE, the data are displayed on the horizontal rather than the vertical axis. (The default value for `datax` is the opposite of that for `qqnorm`.)

- **x**: an object of class `qqnorm2`.

- **pch**: a named vector of the plotting symbols to be used with names corresponding to the levels of `z`.
  - By default, if `z` takes levels `FALSE` and `TRUE` (or 0 and 1), `pch=c(4, 1)` to plot a "x" for `FALSE` and "o" for `TRUE`.
  - If `z` assumes integer values between 0 and 255, by default, the symbols are chosen as described with `points`.
  - Otherwise, by default, `z` is coerced to `character`, and the result is plotted.
  - If `pch` is provided, it must either have names corresponding to levels of `z`, or `z` must be integers between 1 and `length(pch)`.

- **...**: Optional arguments.
  - For `plot.qqnorm2`, they are passed to `plot`.
  - For `qqnorm2`, they are passed to `qqnorm` and to `plot.qqnorm2`.

Details

- For `qqnorm2`:
  1. `q2 <- qqnorm(y, datax=datax, ...)`
  2. `q2["z"] <- z`
  3. `q2["pch"]` gets whatever `pch` decodes to.
  4. Silently return(list(x, y, z, pch)), where "x" and "y" are as returned by `qqnorm` in step 1 above.

- For `plot.qqnorm2`, `plot(x, y, pch=pch[z], ...)`. For `lines.qqnorm2`, `lines(x, y, pch=pch[z], ...)`. For `points.qqnorm2`, `points(x, y, pch=pch[z], ...)`.

Value

`qqnorm2` returns a list with components `x`, `y`, `z`, and `pch`.

Author(s)

Spencer Graves

See Also

`qqnorm`, `qqnorm2s`, `plot`, `points`, `lines`
Examples

```r
##
## a simple test data.frame to illustrate the plot
## but too small to illustrate qqnorm concepts
##
tstDF <- data.frame(y=1:3, z1=1:3, z2=c(TRUE, TRUE, FALSE),
                     z3=c('tell', 'me', 'why'), z4=c(1, 2.4, 3.69))
# plotting symbols circle, triangle, and '+'
qn1 <- with(tstDF, qqnorm2(y, z1))
# plotting symbols "x" and "o"
qn2 <- with(tstDF, qqnorm2(y, z2))
# plotting with "-" and "+
qn. <- with(tstDF, qqnorm2(y, z2, pch=c('FALSE'='-','TRUE'='+')))  
# plotting with "tell", "me", "why"
qn3 <- with(tstDF, qqnorm2(y, z3))
# plotting with the numeric values
qn4 <- with(tstDF, qqnorm2(y, z4))

##
## test plot, lines, points
##
plot(qn4, type='n')  # establish the scales
lines(qn4)         # add a line
points(qn4)        # add points

##
## Check the objects created above
##
# check qn1
qn1. <- qn4
qn1.$xlab <- 'y'
qn1.$ylab <- 'Normal scores'
qn1.$z <- tstDF$z1
qn1.$pch <- 1:3
names(qn1.$pch) <- 1:3
qn1l <- qn1[c(3:4, 1:2, 5:6)]
class(qn1l) <- 'qqnorm2'
all.equal(qn1, qn1l)

# check qn2
qn2. <- qn4
qn2.$xlab <- 'y'
qn2.$ylab <- 'Normal scores'
qn2.$z <- tstDF$z2
qn2.$pch <- c('FALSE'=4, 'TRUE'=1)
qn22 <- qn2[c(3:4, 1:2, 5:6)]
```
### Description

Create a normal probability plot with multiple lines for different variables and different symbols for the values of another variable. `qqnorm2s` produces an object of class `qqnorm2s`, whose plot method produces the plot.
qqnorm2s

Usage

qqnorm2s(y, z, data., plot.it=TRUE, datax=TRUE, outnames=y,
pch=NULL, col=c(1:4, 6), legend.=NULL, ...)
## S3 method for class 'qqnorm2s'
plot(x, y, ...)

Arguments

y For qnorm2s, y is a character vector of names of columns of data, for which normal probability plots are desired. data, is either a data.frame or a list of data.frames of the same length as y, with y[i] being the name of a column of the data.frame data.[[i]]. z is a similar character vector of names of columns of data., which identify symbols for plotting different points in a normal probability plot. The lengths of y, and z must match the number of data.frames in data.; if not, the lengths of the shorter are replicated to the length of the longest before computations begin.
For plot.qnorm2, y is ignored; it is included, because the generic plot function requires it.

z A character vector giving the names of columns of data to indicate different plotting symbols. z should be the same length as y and must equal the number of data.frames in the list data, of data.frames. If not, the shorter are replicated to the length of the longer.

data. a data.frame or a list of data.frames with columns named in y and z.

plot.it logical: Should the result be plotted?

datax The datax argument of qnorm: If TRUE, the data are displayed on the horizontal rather than the vertical axis. (The default value for datax is the opposite of that for qnorm.)

outnames Names for the components of the qnorm2s object returned by the qnorm2s function.

pch a named vector of the plotting symbols to be used with names corresponding to the levels of z.
By default, if z takes levels FALSE and TRUE (or 0 and 1), pch=c(4, 1) to plot a "x" for FALSE and "o" for TRUE.
If z assumes integer values between 0 and 255, by default, the symbols are chosen as described with points.
Otherwise, by default, z is coerced to character, and the result is plotted.
If pch is provided, it must either have names corresponding to levels of z, or z must be integers between 1 and length(pch).

col A vector indicating the colors corresponding to each element of y. Defaults to rep(c(1:4, 6), length=length(y)), with 1:4 and 6 being black, red, green, blue, and pink.

x an object of class qnorm2.

legend. A list with components pch and col providing information for legend to identify the plotting symbols (pch) and colors (col).
By default, pch = list(x='right', legend=names(qq2s[[1]]['pch'])), pch=qq2s[[1]]['pch'])
where qq2s is described below in details.
Similarly, by default, lines = list(x='bottomright', legend=y, lty=1, pch=NA,
col=qq2s[[1]]['col'])

... Optional arguments.
For plot.qqnorm2s, they are passed to plot.
For qqnorm2s, they are passed to qqnorm2 and to plot.qqnorm2s.

Details
For qqnorm2s:
1. Create qq2s = a list of objects of class qqnorm2
2. Add legend. to qq2s.
3. class(qq2s) <- 'qqnorm2s'
4. if(plot.it)plot(qq2s, ...)
5. Silently return(qq2s).
For plot.qqnorm2s, create a plot with one line for each variable named in y.

Value
qqnorm2s returns a named list with components of class qqnorm2 with names = y with each component having an additional component col plus one called "legend."

Author(s)
Spencer Graves

See Also
qqnorm2 plot

Examples

```r
##
## One data.frame
##
tstDF2 <- data.frame(y=1:3, y2=3:5, z2=c(TRUE, TRUE, FALSE),
                    z3=c('tell', 'me', 'why'), z4=c(1, 2.4, 3.69))
# produce the object and plot it
Qn2 <- qqnorm2s(c('y', 'y2'), 'z2', tstDF2)
# plot the object previously created
plot(Qn2)
# Check the object
qy <- with(tstDF2, qqnorm2(y, z2, type='b'))
qy$col <- 1
qy2 <- with(tstDF2, qqnorm2(y2, z2, type='b'))
```
rasterImageAdj

```r
qy2$col <- 2
legend. <- list(pch=list(x='right', legend=c('FALSE', 'TRUE')),
                pch=c('FALSE'=4, 'TRUE'=1),
                col=list(x='bottomright', legend=c('y', 'y2'),
                        lty=1, col=1:2))
Qn2. <- list(y=qy, y2=qy2, legend=legend.)
class(Qn2.) <- 'qqnorm2s'

all.equal(Qn2, Qn2.)

## Two data.frames
##
tstDF2b <- tstDF2
tstDF2b$y <- c(0.1, 0.1, 9)
Qn2b <- qqnorm2s('y', 'z2', list(tstDF2, tstDF2b),
                outnames=c('ok', 'oops'), log='x')
```

---

### Description

Call `rasterImage` to plot image from (xleft, ybottom) to either xright or ytop, shrinking one toward the center to avoid distortion.

angle specifies a rotation around the midpoint ((xleft+xright)/2, (ybottom+ytop)/2). This is different from `rasterImage`, which rotates around (xleft, ybottom).

NOTE: The code may change in the future. The visual image with rotation looks a little off in the examples below, but the code seems correct. If you find an example where this is obviously off, please report to the maintainer – especially if you find a fix for this.

### Usage

```r
rasterImageAdj(image, xleft=par('usr')[1], ybottom=par('usr')[3],
               xright=par('usr')[2], ytop=par('usr')[4], angle = 0,
               interpolate = TRUE, xsub=NULL, ysub=NULL, ...)
```

### Arguments

- **image**: a raster object, or an object that can be coerced to one by `as.raster`.
- **xleft**: a vector (or scalar) of left x positions.
- **ybottom**: a vector (or scalar) of bottom y positions.
- **xright**: a vector (or scalar) of right x positions.
- **ytop**: a vector (or scalar) of top y positions.
angle of rotation in degrees, anti-clockwise about the centroid of image.

NOTE: `rasterImage` rotates around `(xleft, ybottom)`. `rasterImage` rotates around the center `((xleft+xright)/2, (ybottom+ytop)/2)`. See the examples.

`interpolate` a logical vector (or scalar) indicating whether to apply linear interpolation to the image when drawing.

`xsub, ysub` subscripts to subset image

... graphical parameters (see `par`).

Details

1. `imagePixels` = number of (x, y) pixels in image. Do this using `dim(as.raster(image))[2:1]`, because the first dimension of image can be either x or y depending on `class(image)`. For example `link[EBImage]{Image}` returns `dim` with x first then y and an optional third dimension for color. A simple 3-dimensional array is assumed by `rasterImage` to have the y dimension first. `as.raster` puts all these in a standard format with y first, then x.

2. `imageUnits <- c(x=xright-xl, ytop-ybottom)`

3. `xyinches = (x, y) units per inch in the current plot, obtained from `xyinch`.

4. Compute pixel density (pixels per inch) in both x and y dimension: `pixelsPerInch <- imagePixels * xyinches / imageUnits`.

5. Compute `imageUnitsAdj` solving 4 for `imageUnits` and replacing `pixelsPerInch` by the max pixel density: `imageUnitsAdj <- imagePixels * xyinches / max(pixelsPerInch)`.

6. `(dX, dY) = imageUnitsAdj/2 = half of the (width, height) in plotting units.

7. `cntr = (xleft, ybottom) + (dX, dY).` 
   `xleft0 = cntr[1]+sin((angle-90)*pi/180)*dX*sqrt(2); ybottom0 = cntr[2]-cos((angle-90)*pi/180)*dY*sqrt(2);` 
   `(xright0, ytop0) = (upper right without rotation about lower left) xright0 = xleft0+imageUnitsAdj[2] ytop0 = ybottom0+imageUnitsAdj[2]`

8. `rasterImage(image, xleft0, ybottom0, xright0, ytop0, angle, interpolate, ...)`

Value

a named vector giving the values of `xleft, ybottom, xright, and ytop` passed to `rasterImage`. (`rasterImage` returns NULL, at least for some inputs.) This shows the adjustment, shrinking toward the center and rotating as desired.

Author(s)

Spencer Graves

See Also

`rasterImage`
Examples

```r
# something to plot
logo.jpg <- paste(R.home(), "doc", "html", "logo.jpg", 
                   sep = .Platform$file.sep)
if(require(jpeg)){
##
## 1. Shrink as required
##
   Rlogo <- readJPEG(logo.jpg)
   all.equal(dim(Rlogo), c(76, 100, 3))

   plot(1:2)
   # default
   rasterImageAdj(Rlogo)
   plot(1:2, type='n', asp=0.75)
   # Tall and thin
   rasterImage(Rlogo, 1, 1, 1.2, 2)
   # Fix
   rasterImageAdj(Rlogo, 1.2, 1, 1.4, 2)

   # short and wide
   rasterImage(Rlogo, 1.4, 1, 2, 1.2)
   # Fix
   rasterImage(Rlogo, 1.4, 1.2, 2, 1.4)

##
## 2. rotate
##
## 2.1. angle=90: rasterImage left of rasterImageAdj
   plot(0:1, 0:1, type='n', asp=1)
   rasterImageAdj(Rlogo, .5, .5, 1, 1, 90)
   rasterImage(Rlogo, .5, .5, 1, 1, 90)
## 2.2. angle=180: rasterImage left and below
   plot(0:1, 0:1, type='n', asp=1)
   rasterImageAdj(Rlogo, .5, .5, 1, 1, 180)
   rasterImage(Rlogo, .5, .5, 1, 1, 180)
## 2.3. angle=270: rasterImage below
   plot(0:1, 0:1, type='n', asp=1)
   rasterImageAdj(Rlogo, .5, .5, 1, 1, 270)
   rasterImage(Rlogo, .5, .5, 1, 1, 270)
##
## 3. subset
##
   dim(Rlogo)
   # 76 100 3
   Rraster <- as.raster(Rlogo)
   dim(Rraster)
   # 76 100:
   # x=1:100, left to right
   # y=1:76, top to bottom
```

read.testURLs

Read a file produced by testURLs

Description

***NOTE: THIS IS A PRELIMINARY VERSION OF THIS FUNCTION; ***NOTE: IT MAY BE CHANGED OR REMOVED IN A FUTURE RELEASE.

read.table(file.) and return the result as an object of class c('testURLs', 'data.frame').

Usage

read.testURLs(file = 'testURLresults.csv', ...)

Arguments

file. Name of a CSV file to read

... optional arguments for read.csv.

Details

dat <- read.csv(file, ...)
class(dat) <- c('testURLsFile', 'data.frame')

Value

a data.frame from the file written by testURLs, of the same format as the testResults attribute of the testURLs object returned by testURLs.

Author(s)

Spencer Graves

See Also

read.csv
Examples

# Test only 2 web sites, not the default 4,
# and test only twice, not the default 10 times:
tst <- testURLs(c(
  house="http://house.gov/representatives"),
  n=2, maxFail=2)

# The above should have created a file 'testURLresults.csv'
# in the working directory. Read it.

dat <- read.testURLs()

---

read.transpose  Read a data table in transpose form

Description

Read a text (e.g., csv) file, find rows with more than 3 sep characters. Parse the initial contiguous block of those into a matrix. Add attributes headers, footers, and a summary.

The initial application for this function is to read Table 6.16. Income and employment by industry in the National Income and Product Account tables published by the Bureau of Economic Analysis of the United States Department of Commerce.

Usage

read.transpose(file, header=TRUE, sep=',',
  na.strings='---', ...)

Arguments

file        the name of a file from which the data are to be read.
header      Logical: Is the second column of the identified data matrix to be interpreted as variable names?
sep          The field space separator character.
na.strings   character string(s) that translate into NA
...          optional arguments for strsplit

Details

1. txt <- readLines(file)
2. Split into fields.
3. Identify headers, Data, footers.
4. Recombine the second component of each Data row if necessary so all have the same number of fields.
5. Extract variable names
6. Numbers?
7. return the transpose

Value

A matrix of the transpose of the rows with the max number of fields with attributes 'headers', 'footers', 'other', and 'summary'. If this matrix can be coerced to numeric with no NAs, it will be. Otherwise, it will be left as character.

Author(s)

Spencer Graves

References

Table 6.16. Income and employment by industry in the National Income and Product Account tables published by the Bureau of Economic Analysis of the United States Department of Commerce. To get this table from www.bea.gov, under "U.S. Economic Accounts", first select "Corporate Profits" under "National". Then next to "Interactive Tables", select, "National Income and Product Accounts Tables". From there, select "Begin using the data...". Under "Section 6 - income and employment by industry", select each of the tables starting "Table 6.16". As of February 2013, there were 4 such tables available: Table 6.16A, 6.16B, 6.16C and 6.16D. Each of the last three are available in annual and quarterly summaries. The USFinanceIndustry data combined the first 4 rows of the 4 annual summary tables.

See Also

read.table readLines strsplit

Examples

# Find demoFiles/*.csv
demoDir <- system.file('demoFiles', package='Ecdat')
(demoCsv <- dir(demoDir, pattern='csv$', full.names=TRUE))

# Use the fourth example
# to ensure the code will handle commas in a name
# and NAs
nipa6.16D <- read.transpose(demoCsv[4])
str(nipa6.16D)
**readCookPVI**

**Read Cook Partisan Voting Index**

**Description**

Read tables of the Cook Partisan Voting Index and returns a list with components 'House' and 'Senate'. `readCookPVI` returns the tables with the names of the current incumbents per `readUShouse` and `readUSsenate`; `readCookPVI` tables do not include the names of the incumbents.

**Usage**

```r
readCookPVI(url =
  "http://en.wikipedia.org/wiki/Cook_Partisan_Voting_Index")

readCookPVI.(url =
  UShouse=readUShouse(), USsenate=readUSsenate(), ...)
```

**Arguments**

- **url.** Universal resource locator to be read and processed to obtain the desired lists.
- **UShouse, USsenate**
  - data.frames as returned by `readUShouse` and `readUSsenate`, respectively.
- **...** optional arguments passed to `readUShouse` and `readUSsenate`.

**Details**

The primary source for these data is the Cook Political Report web site. However, the current URL we have for these data on that web site includes "2012" in the title. If and when the numbers are updated, we would expect that file name to change.

To avoid that problem the code is currently set to read from the Wikipedia article on "Cook Partisan Voting Index".

The algorithm reads the web site into a list, finds the desired tables on the list, then parses and formats them as desired. Then it merges the results with UShouse and USsenate.

**Value**

A list with components "House" and "Senate". Each contains a data.frame. The "House" data.frame returned by `readCookPVI` includes the following columns:

- **State** name of the state
- **District** District, e.g, 1st, 2nd, At-Large
- **PVI_num** PVI as a number ranging from roughly 50 to 150. 100 means that the vote split in that district was within 0.5 percent of the national average. 101 means that it tilts 1 percent (after rounding) to Republican. 98 means that it tilts 1 percent to Democratic; 99 is not used.
PVIchar

PVI rating in character format. For example, 'D+1' means that the vote tilted 1 percent toward Democratic more than the national average. 'R+1' means that it tilted 1 percent toward Republican.

PartyOfRepresentative

Party of the incumbent, either 'Republican' or 'Democratic'

The 'Senate' data.frame includes the following columns:

State

name of the state

PVINum

PVI numeric, as for 'House'

PVIchar

PVI rating in character format, as for 'House'

PartyOfGovernor

Party of the Governor of the state

PartyInSenate

party of the incumbent senators, either 'Republican', Democratic', or 'Both'.

houseBalanceNum

House balance as a number with 0 = 100 percent Democratic, 99.9 = 100 percent Republican, and 500 for the same number of Republicans as Democrats.

houseBalanceChar

Count by party in the house delegation for that state, e.g., '6R, 1D' for 6 Republicans and 1 Democrat.

readCookPVI. adds to the above the information returned by readUSHouse and readUSSenate.

Author(s)

Spencer Graves

Source

Wikipedia, "Cook Partisan Voting Index" The Cook Political Report

See Also

readUSHouse, readUSSenate

Examples

## Not run:
CookPVI <- readCookPVI()

## End(Not run)

if(!fda::CRAN()){
  CookPVI. <- readCookPVI.()
}
**readFinancialCrisisFiles**

*banking crisis data and function to read financial crisis files*

---

**Description**

Read financial crisis data in files described by an object of class `financialCrisisFiles`. This is designed to read Excel files describing financial crises since 1800 downloaded from [http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/](http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/).

`bankingCrises` is a data.frame created by `readFinancialCrisisFiles()` using 3 files downloaded from [http://www.reinhartandrogoff.com](http://www.reinhartandrogoff.com) and 1 obtained from Prof. Reinhart in January 2013.

**Usage**

```r
readFinancialCrisisFiles(files, crisisType=7, ...)
```

**Arguments**

- `files`: an object of class `financialCrisisFiles`.
- `crisisType`: an integer (vector) between 1 and 8 indicating the type of data to be retrieved: 1=independence year (not a crisis but an indicator), 2=currency, 3=inflation, 4=stock market, 5=domestic sovereign debt crisis, 6=external sovereign debt crisis, 7=banking, 8=tally. ("Type" 1 = year.) These are all 0 or 1 indicating the presence of the event in the given year. Type 8 = sum of types 2 through 7.
- `...`: arguments to pass with file and sheet name to `read.xls` when reading a sheet of an MS Excel file. This is assumed to be the same for all sheets of all files. If this is not the case, the resulting `financialCrisisFiles` object will have to be edited manually before using it to read the data.

**Details**

Reinhart and Rogoff ([http://www.reinhartandrogoff.com](http://www.reinhartandrogoff.com)) provide numerous data sets analyzed in their book, "This Time Is Different: Eight Centuries of Financial Folly". Of interest here are data on financial crises of various types for 70 countries spanning the years 1800 - 2010, downloadable from [http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/](http://www.reinhartandrogoff.com/data/browse-by-topic/topics/7/).

The function `financialCrisisFiles` produces a list of class `financialCrisisFiles` describing different Excel files in very similar formats with one sheet per Country and a few extra descriptor sheets. The data object `financialCrisisFiles` is the default output of that function. `readFinancialCrisisFiles` reads the sheets for the individual countries.

**Value**

- If `length(crisisType) == 1`, a data.frame is returned with the first column being year, and with one other column containing the data for that `crisisType` for each country.
- If `length(crisisType) > 1`, a list is returned containing a data.frame for each country.
readFinancialCrisisFiles

Author(s)
Spencer Graves

Source
http://www.reinhartandrogoff.com

References

See Also
read.xls financialCrisisFiles

Examples

##
## Recreate / update the data object BankingCrises
##
library(Ecdat)

## Not run:
bankingCrises <- readFinancialCrisisFiles(FinancialCrisisFiles)

## End(Not run)

##
## Toy example using local data to check the code
## and illustrate returning all the data not just one crisisType
##
Ecdat.demoFiles <- system.file('demoFiles', package='Ecdat')
Ecdat.xls <- dir(Ecdat.demoFiles, pattern='xls$',
full.names=TRUE)
if(require(gdata)){
tst <- financialCrisisFiles(Ecdat.xls)
bankingCrises.tst <- readFinancialCrisisFiles(tst)
}

# optional tests if not CRAN
if(!fda::CRAN){
allCrises.tst <- readFinancialCrisisFiles(tst, 1:8)
}

# Manually construct tst from allCrises.tst

tst2 <- data.frame(year=1800:1999)
tst2$Algeria <- as.numeric(allCrises.tst$Algeria[-(1:12), 8])
tst2$CentralAfricanRep <- as.numeric(
  allCrises.tst$CentralAfricanRep[-(1:12), 8])
tst2$Taiwan <- as.numeric(allCrises.tst$Taiwan[-(1:11), 8])
tst2$UK <- as.numeric(allCrises.tst$UK[-(1:11), 8])
all.equal(bankingCrises$tst, tst2)

# check
data(bankingCrises)

all.equal(bankingCrises$tst,
  bankingCrises[1:200, c('year', 'Algeria', 'CentralAfricanRep',
                       'Taiwan', 'UK')])

}

readNIPA

---

**Read a National Income and Product Accounts data table**

**Description**

Read multiple files with data in rows using `readNtranspose` and combine the initial columns.

**Usage**

```r
readNIPA(files, sep.footnote='/', ...)  
```

**Arguments**

- `files`          A character vector of names of files from which the data are to be read using `readNtranspose`.
- `sep.footnote`   A single character to identify footnote references in the variable names in some but not all of files.
- `...`            Optional arguments for `readNtranspose`.

**Details**

This is written first and foremost to facilitate updating `USFinanceIndustry` from Table 6.16: Income and employment by industry in the National Income and Product Account tables published by the Bureau of Economic Analysis of the United States Department of Commerce. As of February 2013, this table can be obtained from `http://www.bea.gov`: Under "U.S. Economic Accounts", first select "Corporate Profits" under "National". Then next to "Interactive Tables", select, "National Income and Product Accounts Tables". From there, select "Begin using the data...". Under "Section 6 - income and employment by industry", select each of the tables starting "Table 6.16". As of February 2013, there were 4 such tables available: Table 6.16A, 6.16B, 6.16C and 6.16D. Each of the last three are available in annual and quarterly summaries. The `USFinanceIndustry` data combined the first 4 rows of the 4 annual summary tables.

This is available in 4 separate files, which must be downloaded and combined using `readNIPA`. The first three of these are historical data and are rarely revised. For convenience and for testing, they are provided in the `demoFiles` subdirectory of this Ecdata package.
It has not been tested on other data but should work for annual data with a sufficiently similar structure.

The algorithm proceeds as follows:
1. Data <- lapply(files, read.transpose)
2. Is Data a list of numeric matrices? If no, print an error.
3. cbind common initial variables, averaging overlapping years, reporting percent difference
4. attributes: stats from files and overlap. Stats include the first and last year and the last revision date for each file, plus the number of years overlap with the previous file and the relative change in the common files kept between those two files.

Value

a matrix of the common variables

Author(s)

Spencer Graves

References

United States Department of Commerce Bureau of Economic Analysis National Income and Product Account tables

See Also

read.table readLines strsplit

Examples

# Find demoFiles/*.csv
demoDir <- system.file('demoFiles', package='Ecdat')
(demoCsv <- dir(demoDir, pattern='csv$', full.names=TRUE))
nipa6.16 <- readNIPA(demoCsv)
str(nipa6.16)

readUShouse

Read the list of representatives in the United States House of Representatives

Description

Read the list of representatives in the United States House of Representatives.
Usage

readUShouse(url = "http://www.house.gov/representatives/",
  nonvoting = c('American Samoa', 'District of Columbia',
  'Guam', 'Northern Mariana Islands', 'Puerto Rico',
  'Virgin Islands'),
  fixNonStandard = subNonStandardNames, ...)

Arguments

- **url**: Universal resource locator to be read and processed to obtain the desired list
- **nonvoting**: Character vector of the names of US territories that send a nonvoting delegate to
  the US House.
- **fixNonStandard**: Function to look for and repair nonstandard names such as names containing
  characters with accent marks that are sometimes mangled by different software. Use
  `identity` if this is not desired.
- **...**: Optional arguments passed to `fixNonStandard`

Details

1. `House.gov <- readHTMLTable(url)`. As of April 2013, this is a list of 80 tables. The first 56
   are for the 50 states and 6 territories. The remaining 24 are for the first letter of the last name of
   the representatives.
2. Use `rbind` to collapse these into 2 tables. The first has the district as a number without identifying
   the state (because that was with the names of the first 56 tables in House.gov). The second has
   the state names but with the district numbers in a form not easily parsed.
3. Obtain the state names from the second table to match the names of the representatives in the
   first.
4. Add a `nonvoting` column for those "States" in `nonvoting`
5. Look for and fix surname and `givenName` with nonstandard characters using `fixNonStandard`

Value

`readUShouse` returns a data.frame with the following columns:

- **state**: A factor identifying the state or territory the person represents
- **state**: 2-letter US Postal Service abbreviation for the state or territory
- **district**: The character vector identifying the district each person represents. This is either
  an integer in character format or 0 for "At Large"
- **Name**: A character vector giving the name of each representative (in surname, `givenName`
  format)
- **party**: A factor identifying the party affiliation of each representative ("D" or "R")
- **Room**: Character vector identifying the room number of the office
- **Phone**: Character vector giving the phone number
- **Committees**: Character vector giving the committee assignments of each representative
readUSsenate

surname character vector giving the surname of each representative
givenName given name of each representative (possibly with middle name or initial, a nickname, and a suffix like "Jr.")

Author(s)

Spencer Graves

See Also

getURL readHTMLTable readUSsenate UHouse.senate parseName readUSstateAbbreviations subNonStandardNames readCookPVI

Examples

if(!fda::CRAN(){
  UHouse <- readUHouse()
})

---

readUSsenate Read the list of elected officials in the United States Senate

Description

Read the list of elected officials in the United States Senate.

Usage

readUSsenate(url =
  stateAbbreviations = Ecdat::USstateAbbreviations,
  fixNonStandard = subNonStandardNames, ...)

Arguments

url. Universal resource locator to be read and processed to obtain the desired list

NOTE: On April 26, 2013 the obvious naive use of readHTMLTable worked with the Wikipedia article on the US Senate but did not work with "senate.gov".

stateAbbreviations a data.frame giving names and alternative codes for US states and territories. This must have a column named "Name" giving the names of all 50 states as they appear on the url and another whose name includes "USPS" giving the corresponding 2-letter codes.

fixNonStandard function to look for and repair nonstandard names such as names containing characters with accent marks that are sometimes mangled by different software.

... optional arguments passed to fixNonStandard
Details

1. Senate <- readHTMLTable(url)
2. Use camelParse to remove duplication in Name.
3. Look for and fix surname and givenName with nonstandard characters using fixNonStandard.

Value

readUSsenate returns a data.frame with the following columns:

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>State</td>
<td>A factor identifying the state the person represents</td>
</tr>
<tr>
<td>state</td>
<td>A factor giving the 2-letter USPS code for the state represented</td>
</tr>
<tr>
<td>Class</td>
<td>&quot;1&quot;, &quot;2&quot;, or &quot;3&quot; for election in the 6-year cycle including 2008, 2010, or 2012, respectively.</td>
</tr>
<tr>
<td>Name</td>
<td>A character vector giving the name of each representative (in surname, given name format)</td>
</tr>
<tr>
<td>Party</td>
<td>a factor identifying the party affiliation of each representative (&quot;Democrat&quot;, &quot;Republican&quot;, or &quot;Independent&quot;).</td>
</tr>
<tr>
<td>Experience</td>
<td>character vector highlighting prior experience.</td>
</tr>
<tr>
<td>assumedOffice</td>
<td>character vector giving the date assumed office</td>
</tr>
<tr>
<td>Born</td>
<td>a character vector giving the year of birth</td>
</tr>
<tr>
<td>endOffice</td>
<td>a character vector giving the last day in the present term.</td>
</tr>
<tr>
<td>surname</td>
<td>character vector giving the surname of each representative</td>
</tr>
<tr>
<td>givenName</td>
<td>given name of each representative (possibly with middle name or initial, a nickname, and a suffix like &quot;Jr.&quot;)</td>
</tr>
</tbody>
</table>

Author(s)

Spencer Graves

See Also

getURL readHTMLTable camelParse to remove duplication in Name readUShouse UShouse.senate parseName subNonStandardNames

Examples

```r
if(!fda::CRAN){
  USsenate <- readUSsenate()
}
```
**readUSstateAbbreviations**

*Read a list of abbreviations of states and territories of the United States*

**Description**

Read the list of abbreviations of states and territories of the United states from the relevant Wikipedia article

**Usage**

```r
readUSstateAbbreviations(url =
  clean=TRUE, Names=c('Name', 'Status', 'ISO', 'ANSI.letters',
    'ANSI.digits', 'USPS', 'USCG', 'Old.GPO', 'AP', 'Other'))
```

**Arguments**

- `url` Universal resource locator to be read and processed to obtain the desired list
- `clean` logical: If TRUE, clean the data using `subNonStandardCharacters` and `strsplit(x, "\"\")`
- `Names` names for the columns of the data matrix read; ignored with a warning if the lengths do not match

**Details**

Wrapper for `readHTMLTable`.

NOTE: `readHTMLTable(url)` returns a list of length 7, only one of which is the table we want. Moreover, that table contains some duplicates, which are removed by `readUSstateAbbreviations`. For example, ‘NB’ is an "Obsolete postal code" for Nebraska. If you need this, please consult the Wikipedia article.

**Value**

`readUSstateAbbreviations` returns a data frame from the table in the Wikipedia article on "List of U.S. state abbreviations".

**Author(s)**

Spencer Graves

**See Also**

`getURL` `readHTMLTable` `make.names` `USStateAbbreviations`
Examples

```r
if(!fda::CRAN()){
  abbreviations <- readUSstateAbbreviations()
}
```

__r__

**bivariate recode**

**Description**

Recode x1 and x2 per the lexical codes table.

**Usage**

```r
recode2(x1, x2, codes)
```

**Arguments**

- `x1`, `x2`: vectors of the same length assuming a discrete number of levels
- `codes`: a 2-dimensional matrix indexed by the levels of x1 and x2. If `dimnames(codes)` are not provided, they are assumed to `unique(x1)` (or `unique(x2)`).

**Details**

1. If `length(x1) != length(x2)`, complain.
2. if(is.logical(x1)) l1 <- c(FALSE, TRUE) else l1 <- unique(x1); ditto for x2.
3. If(missing(codes)) codes <- outer(unique(x1), unique(x2))
4. if(is.null(dim(codes))) dim(codes) <- c(length(unique(x1)), length(unique(x2)))
5. If is.null(rownames(codes)), set as follows: If nrow(codes) == length(unique(x1)), rownames(codes) <- unique(x1). Else, if nrow(codes) = max(x1), set rownames(codes) <- seq(1, max(x1)). Else throw an error. Ditto for colnames, ncol, and x2.
6. codes[x1, x2]

**Value**

a vector of the same length as x1 and x2.

**Author(s)**

Spencer Graves

**See Also**

`dim` `rownames` `link{colnames}`
Examples

```r
contrib <- c(-1, 0, 0, 1)
contrib0 <- c(FALSE, FALSE, TRUE, FALSE)

contribCodes <- recode2(contrib>0, contrib0, 
c('returned', 'received', '0', 'ERR')

cC <- c('returned', 'returned', '0', 'received')

all.equal(contribCodes, cC)
```

strsplit1  
Split the first field

Description

Split the first field from `x`, identified as all the characters preceding the first unquoted occurrence of `split`.

Usage

```r
strsplit1(x, split='\', Quote='\', ...)  
```

Arguments

- `x` a character vector to be split
- `split` the split character
- `Quote` a quote character: Occurrences of `split` between pairs of `Quote` are ignored.
- `...` optional arguments for `grep`

Details

This function was written to help parse data from the US Department of Health and Human Services on cybeer-security breaches affecting 500 or more individuals. As of 2014-06-03 the csv version of these data included commas in quotes that are not separator characters. This function was written to split the fields one at a time to allow manual processing to make it easier to correct parsing errors.

Algorithm:

1. `spl1` <- `regexpr(split, x, ...)`
2. `Qt1` <- `regexpr(Quote, x, ...)`
3. For any (`Qt1<spl1`), look for `Qt2` <- `regexpr(Quote, substring(x, Qt1+1))`, then look for `spl1` <- `regexpr(split, substring(x, Qt1+Qt2+1))`
4. `out` <- `list(substr(x, 1, spl1-1), substr(x, spl1+1))`
Value

A list of length 2: The first component of the list contains the character strings found before the first unquoted occurrence of split. The second component contains the character strings remaining after the characters up to the identified split are removed.

Author(s)

Spencer Graves

See Also

strsplit substring grep

Examples

```r
chars2split <- c(qs00='abcdefg', qs01='abc',
     qs10a='abcdefg', qs10b='abcdefg',
     qs1.1='abcdefg', qs20='abcdefg',
     qs2.1='abcdefg', qs21='abcdefg',
     qs22.1='abcdefg')

split <- strsplit1(chars2split)

# answer
split. <- list(c(qs00='abcdefg', qs01='abc', qs10a='abcdefg',
     qs10b='abcdefg', qs1.1='abcdefg', qs20='abcdefg',
     qs2.1='abcdefg', qs21='abcdefg',
     c(qs00='abcdefg', qs01='abcdefg',
     qs10a='abcdefg', qs10b='abcdefg', qs1.1='abcdefg',
     qs20='abcdefg', qs2.1='abcdefg',
     qs21='abcdefg', qs22.1='abcdefg'))

equal(all, split.)
```

Description

First convert to ASCII, stripping standard accents and special characters. Then find the first and last character not in standard characters and replace all between them with replacement. For example, a string like "Ruben" where "e" carries an accent and is mangled by some software would become something like "Rub_n" using the default values for standard characters and replacement.
Usage

```
subNonStandardCharacters(x, standardCharacters=c(letters, LETTERS, ',','.', '?','!', ',', @, 8:9, '/', '*', '$', '%', '\', '^', '+', '&', '_', ';', '(', ')', '[', ']', '\n'), replacement='_', gsubList=list(list(pattern='\\\\\\\\\\\\', replacement='""'), ... ))
```

Arguments

- `x` character vector in which it is desired to find the first and last character not in `standardCharacters` and replace that substring by `replacement`.
- `standardCharacters` a character vector of acceptable characters to keep.
- `replacement` a character to replace the substring starting and ending with characters not in `standardCharacters`.
- `gsubList` list of lists of `pattern` and `replacement` arguments to be called in succession before looking for `nonStandardCharacters`.
- `...` optional arguments passed to `strsplit`.

Details

1. for(il in 1:length(gsubList))x <- gsub( gsubList[[il]]['pattern'], gsubList[[il]]['replacement'], x)
2. x <- stringi::stri_trans_general(x, "Latin-ASCII")
3. nx <- length(x)
4. x. <- strsplit(x, "", ...)
5. for(ix in 1:nx) find the first and last `standardCharacters` in `x.[ix]` and substitute `replacement` for everything in between.

NOTES:

** To find the elements of `x` that have changed, use either `subNonStandardCharacters(x) != x` or `grep(replacement, subNonStandardCharacters(x)), where replacement is the replacement argument = "_" by default.
** On 13 May 2013 Jeff Newmiller at the University of California, Davis, wrote, 'I think it is a fools errand to think that you can automatically "normalize" arbitrary Unicode characters to an ASCII form that everyone will agree on.’ (This was a reply on r-help@r-project.org, subject: "Re: [R] Matching names with non-English characters").
** On 2014-12-15 Ista Zahn suggested `stri_trans_general`. (This was a reply on r-help@r-project.org, subject: "[R] Comparing Latin characters with and without accents?").

Value

a character vector with everything between the first and last character not in `standardCharacters` replaced by `replacement`. 
Author(s)

Spencer Graves with thanks to Jeff Newmiller, who described this as a "fool’s errand", Milan Bouchet-Valat, who directed me to \texttt{iconv}, and Ista Zahn, who suggested \texttt{stri\_trans\_general}.

See Also

\texttt{sub}, \texttt{strsplit}, \texttt{grepNonStandardCharacters}, \texttt{subNonStandardNames} \texttt{encoded\_text\_to\_latex} \texttt{subNonStandardNames} \texttt{iconv} in the base package does some conversion, but is not consistent across platforms, at least using R 3.1.2 on 2015-01-25. \texttt{stri\_trans\_general} seems better.

Examples

```r
## 1. Consider Names = Ruben, Avila and Jose, where "e" and "ä" in
## these examples carry an accent. With the default values
## for standardCharacters and replacement, these might be
## converted to something like Rub_n, _vila, and Jos_, with
## different software possibly mangling the names differently.
## (The standard checks for R packages in an English locale
## complains about non-ASCII characters, because they are
## not portable.)
##
## nonstdNames <- c("Ru\'n", "Ra\'", "\'l", "Torres, Raul",
## "Robert C. \"Bobby\\", NA, '', '\',
## '\$12', '\%')

# confusion in character sets can create
# names like Names[2]
Name2 <- subNonStandardCharacters(nonstdNames)
str(Name2)

# check
Name2. <- c("Ra\'l", "Ra\'", "\'l", nonstdNames[4],
     "Robert C. \"Bobby\", NA, '\', '\',
     '\$12', '\%')
str(Name2.)

all.equal(Name2, Name2.)

## 2. Example from iconv
##
## icx <- c("Ekstr\xf8m", "J\xf6reskog",
## "bi\xdfchen Z\xfcrcher")
icx2 <- subNonStandardCharacters(icx)

# check
icx. <- c("Ekstrom", "Joreskog", "bisschen Zurcher")
```

Description

sub(nonStandardNames[, 1], nonStandardNames[, 2], x)

Accented characters common in non-English languages often get mangled in different ways by different software. For example, the "e" in "Andre" may carry an accent that gets replaced by other characters by different software. This function first converts "Andr*" with "Andr_" for and character "*" not in standard characters. It then looks for "Andr_" in nonstandard names. By default, it will find that and replace it with "Andre".

Usage

subNonStandardNames(x,
    standardCharacters=c(letters, LETTERS, ' ', '.', '?', '!', ', ', 0:9, '/', '*', '$', '%', '\', '¬', '+', '&', '_', ';', '(' , ')', '[', ']', '\n'),
    replacement='_',
    gsubList=list(list(pattern='\\\\\\\\\\\\n', replacement='
'),
        removeSecondLine=TRUE,
        nonStandardNames=Ecdat::nonEnglishNames,
        namesNotFound="attr.replacement", ...)

Arguments

x character vector or matrix or a data.frame of character vectors in which it is desired replace nonStandardNames[, 1] in subNonStandardCharacters(x, ...) with the corresponding element of nonStandardNames[, 2].

standardCharacters, replacement, gsubList, ... arguments passed to subNonStandardCharacters

removeSecondLine

logical: If TRUE, delete anything following "n" and return it as an attribute "secondLine".

nonStandardNames data.frame or character matrix with two columns: Replace any substring of x matching nonStandardNames[, 1] with the corresponding element of nonStandardNames[, 2]
	namesNotFound character vector describing how to treat substitutions not found in nonStandardNames[, 1]:

- "attr.replacement": Return an attribute "namesNotFound" with grep(replacement, subNonStandardCharacters, ...) if any.
subNonStandardNames

- "attr.notFound": Return an attribute "namesNotFound" with x != subNonStandardCharacters(...) if any.
- "print": Print the elements of x "notFound" per either "attr.replacement" or "attr.notFound", as requested.
- "": Do not report any "notFound" elements of x.

NOTE: x = "_" will be identified by "attr.replacement" but not by "attr.notfound" assuming the default value for replacement.

Details

1. removeSecondLine
2. x <- subNonStandardCharacters(x, standardCharacters, replacement, ...)
3. Loop over all rows of nonStandardNames substituting anything matching nonStandardNames[i, 1] with nonStandardNames[i, 2].
4. Eliminate leading and trailing blanks.
5. if(is.matrix(x)) return a matrix; if(is.data.frame(x)) return a data.frame(..., stringsAsFactors=FALSE)

NOTE: On 13 May 2013 Jeff Newmiller at the University of California, Davis, wrote, 'I think it is a fools errand to think that you can automatically "normalize" arbitrary Unicode characters to an ASCII form that everyone will agree on.' (This was a reply on r-help@r-project.org, subject: "Re: [R] Matching names with non-English characters"). Doubtless someone has software to do a better job of this than what this function does, but I've so far been unable to find it in R. If you know of a better solution to this problem, I'd be pleased to hear from you. Spencer Graves

Value

a character vector with all nonStandardCharacters replaced first by replacement and then by the second column of nonStandardNames for any that match the first column. If a secondLine is found on any elements, it is returned as a "secondLine" attribute. If any names with nonStandardCharacters are not found in nonStandardNames[, 1], they are identified in an optional attribute per the namesNotFound argument.

Author(s)

Spencer Graves

See Also

sub nonEnglishNames subNonStandardCharacters

Examples

```r
## 1. Example
##
tstSNSN <- c('Raul', 'Ra\l', 'Torres,Raul', 'Torres, Ra\l',
          "Robert C. \n\Bobby\\\n\n--Vacancy', '"', ' ')

# confusion in character sets can create
```
# names like Names[2]
##
## 2. subNonStandardNames(vector)
##
SNS2 <- subNonStandardNames(tstSNSN)
SNS2

# check
SNS2. <- c("Raul", "Raul", "Torres,Raul", "Torres, Raul",
           "Robert C. "Bobby"", "Ed", "", "")
attr(SNS2., "secondLine") <- c(rep(NA, 5), "--Vacancy", NA, NA)

all.equal(SNS2, SNS2.)

##
## 3. subNonStandardNames(matrix)
##
tstmat <- parseName(tstSNSN, surnameFirst=TRUE)
submat <- subNonStandardNames(tstmat)

# check
SNSmat <- parseName(SNS2., surnameFirst=TRUE)

all.equal(submat, SNSmat)

##
## 4. subNonStandardNames(data.frame)
##
tstdf <- as.data.frame(tstmat)
subdf <- subNonStandardNames(tstdf)

# check
SNSdf <- as.data.frame(SNSmat, stringsAsFactors=FALSE)

all.equal(subdf, SNSdf)

##
## 5. namesNotFound
##
noSub <- subNonStandardNames("xx_x")

# check
noSub. <- "xx_x"
attr(noSub., "namesNotFound") <- "xx_x"

all.equal(noSub, noSub.)
Description

***NOTE: THIS IS A PRELIMINARY VERSION OF THIS FUNCTION; ***NOTE: IT MAY BE CHANGED OR REMOVED IN A FUTURE RELEASE.

try(getURL(...)) to read each element of urls. After each try, write a row to file indicating which of urls was tested, the test time in seconds, and any error message. Repeat any failures up to maxFail times. After testing each element of urls once, repeat n times.

If(ping), proceed each test with "ping url[i]". NOTE: Some Internet Service Providers seem to block some attempts to use "ping" or return fraudulent replies to "ping". It is included in the code, because it seemed like an obvious test. However, it is not executed by default because the results do not necessarily reflect what people might expect from "ping".

Return a list of the last successful version read if any from each element of urls with two attributes: (1) "urls" containing the urls argument. (2) "testResults" being an object of class c('testURLs', 'data.frame') of the test results written to file.

This function was written to diagnose a download problem with a particular Internet Service Provider (ISP). For other tools for testing an ISP, see measurementlab.net or the "Test your ISP" software discussion by the Electronic Frontier Foundation at the URL mentioned in references below.

Usage

testURLs(urls=c(
    wiki="http://en.wikipedia.org",
    house="http://house.gov",
    house.reps="http://house.gov/representatives"),
    file='testURLresults.csv',
    n=10, maxFail=10, warn=-1, tzone='GMT', ping=FALSE, ...)

Arguments

urls a character vector assumed to be universal resource locators to pass to getURL for testing.

The default was selected to provide a 2 x 2 experiment with two different web sites (en.wikipedia.org and house.gov) vs. the landing page and a subordinate page for each site.

file. Name of a CSV file to which to write the results. If the file already exists, new results are appended to it.

n number of times to repeat the cycle testing each member of urls.

maxFail max tests for a continually failing URL. This is designed to make it relatively easy to determine determine dependencies between failures. If the failure rate is constant, the number of consecutive failures will follow a Poisson distribution. Otherwise, it may be possible to evaluate various effects using, e.g., state space
techniques for non-normal time series. This could include daily and weekly cycles possibly with holiday effects and trends as well as drifts suggesting abnormal drifts in web traffic congestion.

**warn**

Argument to pass to `Ping`.

**tzone**

Time zone for `Time`. Defaults to GMT (UTC). `tzone=NULL` will use the current locale.

**ping**

Logical: TRUE to include `Ping`, FALSE otherwise.

**...**

Optional arguments for `Ping`.

---

**Details**

```r
for(i in 1:n):
  1. pingi <- Ping(urls[i], ...)
  2. The time for each call to `getURL` is computed by computing `start.time <- proc.time()` before calling `try(getURL(.))`, then computing the following after:

```
elapsed.time <- max(proc.time() - start.time, na.rm=TRUE)
```

After each of the `urls` is tested, a summary of the results is appended to `file`. This includes the `pingi[['stats']]`, `elapsed.time` and the error message if the download failed.

The Electronic Frontier Foundation provides a table of existing software to "Test your ISP"; see the references below. This table includes a column noting whether the software is "active" (sending test traffic) or "passive" (observing the way the network treats natural traffic). The current `testURLs` function is "active", because it asks for a copy of the code at the indicated URL.

**Value**

An object of class `testURLs`, which in this case is a list of the last successful result returned by `getURL` for each element of `urls` with the following attributes:

- **urls**
  
  The `urls` argument used for this call

- **testURLresults**

  An object of class c('testURLs', 'data.frame') of the data written to `file`. This has the following columns:

  - Time date() for the time a particular test started
  - URL the name in `urls` of the URL tested
  - ping statistics several columns with the count and stats returned by `Ping`.
  - `readTime` time in seconds for the attempt to read the URL `getURL(urls[j])` to complete.
  - `error` character: " if the read attempt was successful; the error message if not.

**Author(s)**

Spencer Graves
trimImage

References

measurementlab.net "Test your ISP" software discussion by the Electronic Frontier Foundation "active" (sending test traffic) or "passive" (observing the way the network treats natural traffic).

See Also

try getURL Ping

Examples

# Test only 2 web sites, not the default 4,
# and test only twice, not the default 10 times:
tst <- testURLs(c(
house="http://house.gov/representatives"),
n=2, maxFail=2)

(class(tst) == 'testURLs') &&
all(names(tst) == c('PVI', 'house')) &&
all(names(attributes(tst)) ==
c('names', 'urls', 'testURLresults', 'class'))

trimImage

Trim zero rows or columns from an object of class Image.

Description

Identify rows or columns of a matrix or 3-dimensional array that are all 0 and remove them.

Usage

trimImage(x, max2trim=.Machine$double.eps, na.rm=TRUE,
returnIndices2Keep=FALSE, ...)

Arguments

x a numeric matrix or 3-dimensional array or an object with subscripting defined so it acts like such.

max2trim a single number indicating the max absolute numeric value to trim.

na.rm logical: If TRUE, NAs will be ignored in determining the max absolute value for the row. If a row or column is all NA, it will be treated as all 0 in deciding whether to trim.
If FALSE, any row or column containing an NA will be retained.
returnIndices2Keep

if TRUE, return a list with 2 integer vectors giving row and column indices to use in selecting the desired subset of x. This allows an array y to be trimmed to match x.

If FALSE, return the desired trimmed version of x.

If this is a list with two two integer vectors, use them to trim x.

Optional arguments; not currently used.

Details

1. Check arguments: 2 <= length(dim(x)) <= 3? is.logical(na.rm)? returnIndices2Keep = logical or list of 2 integer vectors, all the same sign, not exceeding dim(x)?
2. if(is.list(returnIndices2Keep)) check that returnIndices2Keep is a list with 2 integer vectors, all the same sign, not exceeding dim(x). If yes, return x appropriately subsetted.
3. if(!is.logical(returnIndices2Keep)) throw an error message.
5. If(returnIndices2Keep) return (indices2Keep) else return x appropriately susetted.

Value

if(returnIndices2Keep==TRUE) return a list with 2 integer vectors to use as subscripts in trimming objects like x.

Otherwise, return an object like x appropriately trimmed.

Author(s)

Spencer Graves

See Also

trimImage trims raster images, similar to trimImage.
trim trims leading and trailing spaces from character strings and factors. Similar trim functions exist in other packages but without obvious, explicit consideration of factors.

Examples

## 1. trim a simple matrix

```r
tst1 <- matrix(.Machine$double.eps, 3, 3,
    dimnames=list(letters[1:3], LETTERS[1:3]))
tst1[2,2] <- 1
tst1t <- trimImage(tst1)

# check
tst1. <- matrix(1, 1, 1,
    dimnames=list(letters[2], LETTERS[2]))
```
trimImage

all.equal(tst1t, tst1.)

##
## 2. returnIndices2Keep
##
tst2i <- trimImage(tst1, returnIndices2Keep=TRUE)
tst2a <- trimImage(tst1, returnIndices2Keep=tst2i)
tst2i. <- list(index1=2, index2=2)

# check
all.equal(tst2i, tst2i.)

all.equal(tst2a, tst1.)

##
## 3. trim 0's only
##
tst3 <- array(0, dim=3:5)
tst3[2, 2:3, ] <- 0.5*MACHINEdouble.eps
tst3[3,,] <- 1

tst3t <- trimImage(tst3, 0)

# check
tst3t. <- tst3[2:3,, ]

# check
all.equal(tst3t, tst3t.)

##
## 4. trim NAs
##
tst4 <- tst1
tst4[1,1] <- NA
tst4[3,,] <- NA

tst4t <- trimImage(tst4)
# tst4o == tst4
tst4o <- trimImage(tst4, na.rm=FALSE)

# check
all.equal(tst4t, tst1[2, 2, drop=FALSE])
Description
The cumulative distribution function for a truncated distribution is \( P \) for \( x < \text{truncmin} \), \( 1 \) for \( \text{truncmax} < x \), and in between is as follows:
\[
\frac{\text{pdist}(x, \ldots ) - \text{pdist}(	ext{truncmin}, \ldots )}{\text{pdist}(	ext{truncmax}, \ldots ) - \text{pdist}(	ext{truncmin}, \ldots )}
\]
The density, quantile, and random number generation functions are similarly defined from this.

Usage
\[
d\text{truncdist}(x, \ldots , \text{dist}'\text{norm}', \text{truncmin}=-\text{Inf}, \text{truncmax}=\text{Inf})
\]
\[
p\text{truncdist}(q, \ldots , \text{dist}'\text{norm}', \text{truncmin}=-\text{Inf}, \text{truncmax}=\text{Inf})
\]
\[
q\text{truncdist}(p, \ldots , \text{dist}'\text{norm}', \text{truncmin}=-\text{Inf}, \text{truncmax}=\text{Inf})
\]
\[
r\text{truncdist}(n, \ldots , \text{dist}'\text{norm}', \text{truncmin}=-\text{Inf}, \text{truncmax}=\text{Inf})
\]

Arguments
- \( x, q \) numeric vector of quantiles
- \( p \) numeric vector of probabilities
- \( n \) number of observations. If \( \text{length}(n) > 1 \), the length is taken to be the number required.
- \( \ldots \) other arguments to be passed to the corresponding function for the indicated \( \text{dist} \)
- \( \text{dist} \) Standard R name for the family of functions for the desired distribution. By default, this is "norm", so the corresponding function for \( d\text{truncdist} \) is \( \text{dnorm} \), the corresponding function for \( p\text{truncdist} \) is \( \text{pnorm} \), etc.
- \( \text{truncmin}, \text{truncmax} \) lower and upper truncation points, respectively.
Details

NOTE: This is different from "censoring", where it's known that an observation lies between certain limits; it's just not known exactly where it lies between those limits.

For a truncated distribution, observations below truncmin and truncmax are not observed at all. Thus, it's not known how many observations like outside the given range, truncmin to truncmax, if any.

1. Setup
   dots <- list(...)

2. For dtruncdist, return 0 for all x outside truncmin and truncmax. For all others, compute as follows:
   dots$x <- truncmin
dist <- paste0('d', dist) pdist <- paste0('p', dist) p.min <- do.call(pdist, dots)
dots$x <- truncmax
p.max <- do.call(pdist, dots)
dots$x <- x
dx <- do.call(dist, dots)
return(dx / (p.max-p.min))

NOTE: Adjustments must be made if 'log' appears in names(dots)

3. The computations for ptruncdist are similar.

4. The computations for qtruncdist are complementary.

5. For rtruncdist, use qtruncdist(runif(n), ...).

Value

dtruncdist gives the density, ptruncdist gives the distribution function, qtruncdist gives the quantile function, and rtruncdist generates random deviates.

The length of the result is determined by n for rtruncdist and is the maximum of the lengths of the numerical arguments for the other functions.

Author(s)

Spencer Graves

See Also

Distributions Normal

Examples

```r
##
## 1. dtruncdist
##
# 1.1. Normal
dx <- dtruncdist(1:4)

# check
all.equal(dx, dnorm(1:4))
```
# 1.2. Truncated normal between 0 and 1
dx01 <- dtruncdist(seq(-1, 2, .5), truncmin=0, truncmax=1)

# check
dx01. <- c(0, 0, 0, dnorm(.5, 1)/(pnorm(1)-pnorm(0)),
           0, 0)

all.equal(dx01, dx01.)

# 1.3. lognormal mean\(\log(100)\), sd\(\log = 2\), truncmin=500
x10 <- 10^(0:9)
dx10 <- dtruncdist(x10, log(100), 2, dist='lnorm',
                   truncmin=500)

# check
dx10. <- (dtruncdist(log(x10), log(100), 2,
                     truncmin=log(500)) / x10)

all.equal(dx10, dx10.)

# 1.4. log density of the previous example
dx10log <- dtruncdist(x10, log(100), 2, log=TRUE,
                      dist='lnorm', truncmin=500)

all.equal(dx10log, log(dx10))

# 1.5. Poisson without \(\theta\).
dPois0.9 <- dtruncdist(0:9, lambda=1, dist='pois', truncmin=0)

# check
dP0.9 <- c(0, dpois(1:9, lambda=1)/ppois(0, lambda=1, lower.tail=FALSE))

all.equal(dPois0.9, dP0.9)

##
## 2. ptruncdist
##
# 2.1. Normal
px <- ptruncdist(1:4)

# check
all.equal(px, pnorm(1:4))

# 2.2. Truncated normal between 0 and 1
px01 <- ptruncdist(seq(-1, 2, .5), truncmin=0, truncmax=1)
truncdist

# check
px01 <- c(0, 0, (pnorm(c(0, .5, 1)) - pnorm(0))
        /(pnorm(1) - pnorm(0)), 1, 1)

all.equal(px01, px01.)

# 2.3. lognormal meanlog=log(100), sdlog = 2, truncmin=500
x10 <- 10^(0:9)
px10 <- ptruncdist(x10, log(100), 2, dist='lnorm',
                   truncmin=500)

# check
px10. <- (ptruncdist(log(x10), log(100), 2,
                     truncmin=log(500)))

all.equal(px10, px10.)

# 2.4. log of the previous probabilities
px10log <- ptruncdist(x10, log(100), 2, log=TRUE,
                      dist='lnorm', truncmin=500)

all.equal(px10log, log(px10))

##
## 3. qtruncdist
##
## 3.1. Normal
qx <- qtruncdist(seq(0, 1, .2))

# check
all.equal(qx, qnorm(seq(0, 1, .2)))

# 3.2. Normal truncated outside (0, 1)
qx01 <- qtruncdist(seq(0, 1, .2), truncmin=0, truncmax=1)

# check
pxmin <- pnorm(0)
pxmax <- pnorm(1)
unp <- (pxmin + seq(0, 1, .2) * (pxmax - pxmin))
qx01. <- qnorm(unp)

all.equal(qx01, qx01.)

# 3.3. lognormal meanlog=log(100), sdlog=2, truncmin=500
qlx10 <- qtruncdist(seq(0, 1, .2), log(100), 2,
                     dist='lnorm', truncmin=500)
# check
plxmin <- plnorm(500, log(100), 2)
unp. <- (plxmin + seq(0, 1, .2)*(1-plxmin))
qlx10. <- qlnorm(unp., log(100), 2)
all.equal(qlx10, qlx10.)

# 3.4. previous example with log probabilities
qlx10l <- qtruncdist(log(seq(0, 1, .2)), log(100), 2,
                      log.p=TRUE, dist='lnorm', truncmin=500)

# check
all.equal(qlx10, qlx10l)

##
## 4. rtruncdist
##
## 4.1. Normal
set.seed(1)
rx <- rtruncdist(9)

# check
set.seed(1)
all.equal(rx[1], rnorm(1))

# Only the first observation matches; check that.

## 4.2. Normal truncated outside (0, 1)
set.seed(1)
rx01 <- rtruncdist(9, truncmin=0, truncmax=1)

# check
pxmin <- pnorm(0)
pxmax <- pnorm(1)
set.seed(1)
rnp <- (pxmin + runif(9)*(pxmax-pxmin))
rx01. <- qnorm(rnp)
all.equal(rx01, rx01.)

## 4.3. lognormal meanlog=log(100), sdlog=2, truncmin=500
set.seed(1)
rlx10 <- rtruncdist(9, log(100), 2,
                      dist='lnorm', truncmin=500)

# check
plxmin <- plnorm(500, log(100), 2)
Create a list of members of the US House and Senate

**Description**

Combine the output of `readUShouse` and `readUSSenate`.

**Usage**

```r
UShouse.senate(house=readUShouse(), senate=readUSSenate())
```

**Arguments**

- `house`, `senate` - data.frames as returned by the functions `readUShouse` and `readUSSenate`, respectively.

**Details**

Convert the two into a common format and rbind.

**Value**

A data.frame with the following columns:

- **Office**
  A factor identifying "House" vs. "Senate", indicating whether the person is in the US House or Senate
- **state**
  A factor identifying the state using the USPS 2-letter state code (all caps)
- **district**
  "0" or "At-Large" for members of the US House representing an entire state or integers in character format indicating the district. For the Senate, this contains the "class", which codes the year of the next election for that seat is an integer multiple of 6 years after 2012, 2008, or 2010 for class "1", "2", or "3", respectively.
- **Party**
  A factor identifying the party affiliation of each representative, e.g., 'Democratic', 'Republican', 'Democratic-Farmer-Labor', 'Independent'.
- **surname**
  Family name
- **givenname**
  First name with possibly a middle name, nickname, and suffix (e.g., Jr., III).
USsenateClass

Author(s)
Spencer Graves

See Also
readUShouse readUSSenate

Examples

if(!fda::CRAN){
  house <- readUShouse()
  USreps <- UShouse.senate(house)
}

Usage

USSenateClass(x, senate=readUSSenate(),
  Office='Office', state='state',
  surname='surname', district='district', senatePattern='^Senate')

Arguments

x data.frame with character or factor columns Office, state, surname, and district.
senate data.frame as returned by readUSSenate.
Office name of a character or factor variable x in which the members of the US Senate can be identified by grep(senatePattern, x[, Office]).
state Standard 2-letter abbreviation for the state of the US
surname the name of a column of x containing the surname

Description

For all individuals in x with houseSenate == "Senate", look up their state and surname in the reference table senate and return their Class. For individuals not found in senate, return x[[district]]. Senate classes 1, 2 and 3 have their normal elections in 6-year cycles including 2000, 2002, and 2004 (or 2012, 2008, and 2010), respectively. When vacancies occur out of cycle, the vacancy is first filled with appointment by the governor of the state, and an election to fill that seat occurs in the next even-numbered year; the class of that seat does not change.

For example, South Carolina Senator Jim DeMint resigned effective January 1, 2013. South Carolina Governor Nikki Haley appointed Tim Scott to serve until a special election in 2014. This is a Class 3 seat, which means that another election for that seat will occur in 2016.
district       name of a column of x containing the number of the district in the US House. 
               For states with only one representative, this may be 0.

senatePattern a regular expression for identifying the senators from x[,    Office].

Details

The current algorithm may fail if both senators in a state have the same surname.

Value

a data.frame with one row for each row of x and the following columns:

incumbent       logical vector: NA if Office == 'house'. If Office == 'senate', then TRUE if 

    state:surname found in senate and FALSE otherwise.

District        a character vector containing the desired Class for all US Senators found in 

    senate or a guess at the Class for non-incumbents. For members of the House, 

    this returned the previous content of x[[District]].

NOTES:

1. Incumbents can be missed if the spelling of the surname is different between 
   x and senate. This can occur with, for example, Spanish surnames containing 
   an accent.

2. If one but not two incumbents is found, others are currently assigned to the 
   class of an incumbent not found. This could be a mistake, because the person 
   could be a previous incumbent or could have lost to the incumbent in the last 
   election.

Author(s)

Spencer Graves

See Also

readUSsenate

Examples

tst <- data.frame(Office=as.factor(c("House", "Senate", "Senate", 'Senate')),
                   state=as.factor(c('SC', 'SC', 'SC', 'NY')),
                   surname=c("Jones", "DeMint", "Graham", 'Smith'),
                   district=c("9", NA, NA, NA),
                   stringsAsFactors=FALSE)

if(!fda::CRAN()){ 
  tst. <- USSenateClass(tst)

  chk <- data.frame(incumbent=c(NA, FALSE, TRUE, FALSE),
                   district=c("9", "3", "2", "1 or 3"),
                   stringsAsFactors=FALSE)

  all.equal(tst., chk)
## whichAeqB
### Index of a single match

**Description**

Return which(A %in% B) if it has length 1; give an error message otherwise.

**Usage**

```r
whichAeqB(A, B, errNoMatch='no match',
          err2Match='more than one match')
```

**Arguments**

- **A**
  A vector which may have a single match in B.

- **B**
  A vector of possible matches for A.

- **errNoMatch**
  a character string: error message if no match found.

- **err2Match**
  a character string: error message if multiple matches found.

**Value**

a single integer giving the index of the match in A.

**Author(s)**

Spencer Graves

**See Also**

`interpPairs`
Examples

```r
a2b <- whichAeqB(letters, 'b')

all.equal(a2b, 2)
```
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