

Package ‘quickcode’

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Type Package

Title Quick and Essential 'R' Tricks for Better Scripts

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Description The NOT functions, 'R' tricks and a compilation of some simple quick plus of-
ten used 'R' codes to improve your scripts. Improve the quality and reproducibility of 'R' scripts.

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URL <https://quickcode.rpkg.net>

BugReports <https://github.com/oobianom/quickcode/issues>

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add.header *Addin snippet function to add header comment to a current opened file*

Description

Shorthand to add header comment

Usage

```
add.header()
```

Value

Inserts header content for file

Examples

```
if(interactive())
add.header()
```

add.sect.comment *Addin snippet function to custom section comment*

Description

Shorthand to add section comment to current file

Usage

```
add.sect.comment()
```

Value

Inserts section comment content for file

Examples

```
if(interactive())
add.sect.comment()
```

`add.snippet.clear` *Snippet R function to clear console and set directory*

Description

Shorthand to add clear console code to current file

Usage

```
add.snippet.clear()
```

Value

Inserts code to clear console

Examples

```
if(interactive())  
add.snippet.clear()
```

`add_key` *Add index keys to a vector or data frame or list or matrix*

Description

Index a vector or lists and convert to a list of objects

Usage

```
add_key(vector)  
  
indexed(vector, key = key, value = value)
```

Arguments

<code>vector</code>	vector or data frame to transform
<code>key</code>	variable name for keys
<code>value</code>	variable name for values

Details

This function takes a vector and turns it into a list containing 'key' and 'value' for each vector. This allows the output to be used in loops such as for loops or lapply or other functions to track the index of the list content e.g. 1,2,3...

This function also contains a validator to ensure that a vector had not been previously 'keyed', which prevents the user from inadvertently calling the function twice on a vector. Helps especially because the function keys the vector, and sets the new list to the variable name of the original vector.

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Value

a transformed list containing keys along with vector values

Use case

Efficient for loops and for tracking various steps through a vector contents

Note

add_key - resaves the keys and value pairs to original variable

indexed - return the keys and value pairs

Examples

```
# EXAMPLES for add_key()

#ex1 simple conversion of a vector
rti2 <- c("rpkg","obinna", "obianom")
add_key(rti2)
rti2

#ex2 add keys to a vector content for use in downstream processes
ver1 <- c("Test 1","Test 2","Test 3")
add_key(ver1)

#ex3 use keyed ver1 in for loop
for(i in ver1){
  message(sprintf("%s is the key for this %s", i$key, i$value))
}
```

```
}

#ex4 use keyed ver1 in lapply loop
x11 <- lapply(ver1,function(i){
  message(sprintf("lapply - %s is the key for this %s", i$key, i$value))
})

# EXAMPLES for indexed()

#ex1 simple conversion of a vector
rti2 <- c("rpkg","obinna", "obianom")
indexed(rti2)

#ex2 add keys to a vector content for use in downstream processes
ver1 <- c("Test 1","Test 2","Test 3")

#ex3 use keyed ver1 in for loop
for(i in indexed(ver1)){
  message(sprintf("%s is the key for this %s", i$key, i$value))
}

#ex4 use keyed ver1 in for loop
#specify name for key and value
for(i in indexed(ver1,k,v)){
  message(
    sprintf("%s is the new key for this value %s",
            i$k, i$v))
}

#ex5 use keyed ver1 in lapply loop
x11 <- lapply(indexed(ver1),function(i){
  message(sprintf("lapply - %s is the key for this %s", i$key, i$value))
})
```

ai.duplicate

Prompt guided duplication and editing of files

Description

AI like duplication and editing of files

Usage

```
ai.duplicate(file = NULL, new.name = NULL, open = TRUE)
```

Arguments

file	file to duplicate
new.name	OPTIONAL.name of new file
open	open file after duplication

Value

duplicate files with edited texts

Examples

```
if(interactive()){
  file1s <- paste0(tempfile(), ".R")
  writeLines("message(
'Sample items: farm, shinyappstore, rpkg'
)", file1s)
  ai.duplicate(file1s, 'file2.R')
}
```

 archivedPkg

Listing of all CRAN archived R packages

Description

Retrieve a list of all currently archived R packages and their archive date

Usage

```
archivedPkg(
  startsWith = c("all", letters),
  after = NULL,
  inc.date = TRUE,
  as = c("data.frame", "list")
)
```

Arguments

startsWith	one letter that the package name starts with eg. a, e, f
after	packages archived after a specific date eg. 2011-05-10
inc.date	should archive date be included in the result
as	return result as data frame or as list

Value

a data frame or list containing listing of all archived R packages

Use case

This function allows the retrieval of various R packages archived by CRAN along with the respective latest archive date. The packages retrieved include both active and inactive R projects submitted to CRAN. When a new version of an active R package is published, the older versions of the package gets archived. In the same way, when a package is decommissioned from CRAN active projects for one reason or another, it gets archived.

Note

- * The "startsWith" argument should be one letter and should be in lowercase
- * If no argument is provided for "startsWith", all the packages will be retrieved
- * The format of the "after" argument must be YYYY-MM-DD e.g. 2022-04-11

Examples

```
# Task 1: get archived R packages with names beginning with C or All
head(archivedPkg(startsWith = "all"), n= 10) #retrieves all packages
head(archivedPkg(startsWith = "c"), n= 10) #retrieves only packages beginning with a

# Task 2: return the packages from Task 1 without including latest archive date
res.dt2 <- archivedPkg(startsWith = "b", inc.date = FALSE)
res.dt2[1:10,]

# Task 3: return the results from Task 2 as a list
res.dt3 <- archivedPkg(startsWith = "c", inc.date = FALSE, as = "list")
res.dt3$name[1:10]

res.dt3 <- archivedPkg(startsWith = "e", as = "list")
res.dt3$name[1:10]

# Task 4: return the archived packages beginning with Y archived after 2022-08-12
# Note that startsWith should be lowercase

#without archive date
yRPkg <- archivedPkg(startsWith = "y", after= NULL)
nrow(yRPkg) #number of rows returned
head(yRPkg, n = 15) #show first 15 rows

#with archive date
yRPkg2 <- archivedPkg(startsWith = "y", after= "2022-08-12")
nrow(yRPkg2) #number of rows returned
head(yRPkg2, n = 15) #show first 15 rows, notice no archive date before 2022-08-12
```

`as.boolean`*Convert boolean values between formats*

Description

Convert Yes/No to 1/0 or to TRUE/FALSE or vice versa

Usage

```
as.boolean(ds, type = 3)
```

Arguments

<code>ds</code>	item to convert
<code>type</code>	format to convert to, choices 1, 2 or 3

Details

Output various format of booleans into a specified format. Below are the options for the type argument.

type: options are as follows -

- 1 - Yes/No
- 2 - TRUE/FALSE
- 3 - 1/0

Value

output adhering to the format of the type provided

Examples

```
# Task: convert "yes" or "no" to format of TRUE or FALSE
as.boolean("yes",2)
as.boolean("no",2)
as.boolean("YES",2)
as.boolean("NO",2)

# Task: convert "yes" or "no" to format of 1 or 0
as.boolean("yes",3)
as.boolean("no",3)
as.boolean("YES",3)
as.boolean("NO",3)

# Task: convert 1 to format of Yes or No
as.boolean(1,1)
```

```
# Task: convert "T" to format of Yes or No
as.boolean("T",1)

# Task: convert "f" to format of TRUE or FALSE
as.boolean("f",2)

# Task: convert 1 to format of TRUE or FALSE
as.boolean(1,2)

# Task: convert "Y" or "y" to format of Yes or No
as.boolean("Y",1) #uppercase Y
as.boolean("y",1) #lowercase y

# Task: convert TRUE/FALSE to format of 1 or 0
as.boolean(TRUE,3)
as.boolean(FALSE,3)

# Task: convert TRUE/FALSE to format of Yes or No
as.boolean(TRUE,1)
as.boolean(FALSE,1)

# In case of error in argument
# as.boolean("tr",3) #NA
# as.boolean("ye",3) #NA

# vector of mixed boolean to TRUE/FALSE or 1/0
multv <- c(TRUE,"y","n","YES","yes",FALSE,"f","F","T","t")
as.boolean(multv,1) # return vector as Yes/No
as.boolean(multv,2) # return vector as TRUE/FALSE
as.boolean(multv,3) # return vector as 1/0
```

bionic_txt

Generate a bionic text

Description

This function serves as a mechanism enabling the conversion of provided text into a bionic form. Users input the text, and the function, in turn, delivers the text transformed into a bionic format.

Usage

```
bionic_txt(text)
```

Arguments

text input text

Details

A bionic text refers to a transformed version of a given text achieved through a specialized function designed to incorporate elements of advanced technology, enhancing both the form and content of the original input. This function operates by infusing the text with a fusion of various elements, resulting in a synthesis that transcends traditional linguistic boundaries. The function augments the text with dynamic visual representations that adapt to the reader's preferences. The goal is to create a text that not only conveys information but also engages the audience in a more immersive and interactive manner, harnessing the capabilities of modern technology to redefine the traditional concept of textual communication. An example of a bionic text could be a news article that dynamically updates with real-time data, incorporates multimedia elements, and adjusts its presentation style based on the reader's preferences, thereby offering a more enriched and personalized reading experience.

Value

bionic text

References

This idea stems from a blog article published at <https://www.r-bloggers.com/2023/10/little-useless-useful-r-functions-function-for-faster-reading-with-bionic-reading/> and the original source for bionic texts may be found at <https://bionic-reading.com/>

Examples

```
# simple example to show a text
# transformation to bionic text

# text to transform
text1 <- "A tool for nonparametric
estimation and inference
of a non-decreasing
monotone hazard\ratio
from a right censored survival dataset."

# transform text
genbt <- bionic_txt(text1)

# print bionic text as message or cat
message(genbt)
cat(genbt)
```

cat_to_num	<i>Convert Categorical Values to Numeric Improvement to seq_along()</i>
------------	---

Description

Converts categorical values into numeric by sorting unique values alphabetically/numerically and assigning sequential numbers starting from 1

Usage

```
cat_to_num(x, decreasing = FALSE)
cat_to_num2(x, decreasing = FALSE)
```

Arguments

x A vector containing categorical values (character or factor)
decreasing logical. Should the sort be increasing or decreasing?

Value

A numeric vector where each category is mapped to a number based on its sorted position. NA values in the input remain NA in the output.

Examples

```
# Example 1: Basic character vector
fruits <- c("apple", "banana", "apple", "cherry", "banana")
cat_to_num(fruits, decreasing = FALSE) # Returns: 1, 2, 1, 3, 2

# Example 2: Vector with NAs
grades <- c("A", NA, "C", "B", "A", NA)
cat_to_num(grades) # Returns: 1, NA, 3, 2, 1, NA

# Example 3: Factor input
animals <- factor(c("dog", "cat", "bird", "cat", "dog"))
cat_to_num(animals) # Returns: 3, 2, 1, 2, 3

# Example 4: Numeric categories
sizes <- c("XL", "S", "M", "XS", "L")
cat_to_num(sizes) # Returns: 5, 2, 3, 1, 4

# Example 5: Single category
status <- c("active", "active", "active")
cat_to_num(status) # Returns: 1, 1, 1

# Examples for cat_to_num2

# Assign numeric values to a set of categorical values
```

```

# Call the value assignment to assign
# a numeric value to a new categorical value

cat_to_num2(letters[1:7])("b")
cat_to_num2(letters[1:7], decreasing = FALSE)("d")
# the above assigns numeric values to letters a-g
# and then calls the number for "a"

cat_vals_5 <- c("apple", "orange", "guava")
num_equiv <- cat_to_num2(cat_vals_5)

applenum = num_equiv("apple")

```

chain_func

Combine specific functions as store as one function

Description

Use many functions in one call using predeclared calls

Usage

```
chain_func(..., otherargs = list())
```

Arguments

... functions to include. see example for how to use
otherargs other arguments for use in each function

Value

returns a function that combines multiple function

Examples

```

# Example 1 with base functions
comb1 <- chain_func(unique, cumsum, print)
result <- comb1(sample(1:5,10,replace = TRUE))
#or
u = sample(1:5,10,replace = TRUE)
result <- comb1(u)

# Example 2 with base functions with arguments
comb2 <- chain_func(unique, print, otherargs = list(.= c(FALSE),.=c(2)))
result <- comb2(sample(1:3,10,replace = TRUE))

# Example 3 with custom functions

```

```
r = function(a,b,c){
  if(!missing(a))print(a)
  if(!missing(b))print(b)
  if(!missing(c))print(c)
  return(a)
}

r2 = function(a,b,c){
  if(!missing(a))message(a)
  if(!missing(b))message(b)
  if(!missing(c))message(c)
  return(a)
}

comb3 <- chain_func(r,r2, otherargs =list(.=c("apple","cat"),.=c("rice")))
res <- comb3(head(mtcars))
```

clean	<i>Clear environment, clear console, set work directory and load files</i>
-------	--

Description

Shorthand to quickly clear console, clear environment, set working directory, load files

Usage

```
clean(setwd = NULL, source = c(), load = c(), clearPkgs = FALSE)
```

```
clear(setwd = NULL, source = c(), load = c(), clearPkgs = FALSE)
```

Arguments

setwd	OPTIONAL. set working directory
source	OPTIONAL. source in file(s)
load	OPTIONAL. load in Rdata file(s)
clearPkgs	Clear previous loaded packages, TRUE or FALSE

Details

The purpose of this function is provide a one-line code to clear the console, clear the environment, set working directory to a specified path, source in various files into the current file, and load RData files into the current environment. The first process in the sequence of events is to clear the environment. Then the working directory is set, prior to inclusion of various files and RData. With the directory being set first, the path to the sourced in or RData files will not need to be appended to the file name. See examples.

Value

cleared environment and set directory

Examples

```

if(interactive()){
#simply clear environment, clear console and devices
quickcode::clean()

#clear combined with additional arguments
quickcode::clean(
  clearPkgs = FALSE
) #also clear all previously loaded packages if set to true

quickcode::clean(
  setwd = "/home/"
) #clear env and also set working directory

quickcode::clean(
  source = c("/home/file1.R","file2")
) #clear environment and source two files into current document

quickcode::clean(
  setwd = "/home/",
  source = c("file1","file2")
) #clear environment, set working directory and source 2 files into environment

quickcode::clean(
  setwd = "/home/",
  source="file1.R",
  load="obi.RData"
) #clear environment, set working directory, source files and load RData
}

```

cols.rep

Replicate Columns in a Data Frame

Description

This function replicates each column in a data frame a specified number of times.

Usage

```
cols.rep(data, n)
```

Arguments

data	A data frame whose columns are to be replicated.
n	An integer specifying the number of times each column should be replicated.

Value

A data frame with each column replicated ‘n’ times. If ‘n’ is less than or equal to 0, an empty data frame is returned.

Examples

```
# Example with a simple data frame
df <- data.frame(A = c(1, 2), B = c(3, 4))
cols.rep(df, 3)

# Example with no replication (n = 0)
cols.rep(df, 0)
```

 compHist

Compare histograms of two distributions

Description

For comparing histograms of two data distributions. Simply input the two distributions, and it generates a clear and informative histogram that illustrates the differences between the data.

Usage

```
compHist(
  x1,
  x2,
  title,
  col1 = "red",
  col2 = "yellow",
  xlab = "",
  ylab = "Frequency",
  separate = FALSE
)
```

Arguments

x1	NUMERIC. the first distribution
x2	NUMERIC. the second distribution
title	CHARACTER. title of the histogram plot
col1	CHARACTER. color fill for first distribution
col2	CHARACTER. color fill for second distribution
xlab	CHARACTER. label of the x-axis
ylab	CHARACTER. label of the y-axis
separate	LOGICAL. whether to separate the plots

Details

Users have the option to view individual histograms for each distribution before initiating the comparison, allowing for a detailed examination of each dataset's characteristics. This feature ensures a comprehensive understanding of the data and enhances the user's ability to interpret the results of the distribution comparison provided by this function.

Value

return histogram comparison using basic histogram plot

Some recommended color pairs

```
col1 = 'dodgerblue4' (and) col2 = 'darksalmon'
col1 = 'brown' (and) col2 = 'beige'
col1 = 'pink' (and) col2 = 'royalblue4'
col1 = 'red' (and) col2 = 'yellow'
col1 = 'limegreen' (and) col2 = 'blue'
col1 = 'darkred' (and) col2 = 'aquamarine4'
col1 = 'purple' (and) col2 = 'yellow'
```

Note

- Hexadecimal values can also be passed
 in for col1 and col2, see the example section - For best visual results,
 col1 should be a dark color and col2 should be passed as a light color.
 For example, col1 = "black", col2 = "yellow"

Examples

```
# compare two normal distributions with means that differ a lot
# in this case, the overlap will not be observed
set.seed(123)
compHist(
  x1 = rnorm(1000, mean = 3),
  x2 = rnorm(1000, mean = 10),
  title = "Histogram of Distributions With Means 3 & 10",
  col1 = "yellow", col2 = "violet"
)

# compare two normal distributions with means that are close
# in this case, the overlap between the histograms will be observed
set.seed(123)
compHist(
  x1 = rnorm(1000, mean = 0),
  x2 = rnorm(1000, mean = 2),
  title = "Histogram of rnorm Distributions With Means 0 & 2",
  col1 = "lightslateblue", col2 = "salmon"
)
```

```
set.seed(123)
# separate the plots for preview
compHist(
  x1 = rnorm(1000, mean = 0),
  x2 = rnorm(1000, mean = 2),
  title = c("Plot Means 0", "Plot Means 2"),
  col1 = "#F96167", col2 = "#CCF381",
  separate = TRUE
)
```

const

Mathematical constants

Description

List of mathematical functions

Usage

```
const
```

Format

An object of class `list` of length 2.

Details

`const` list contains a list of mathematical constants below

- pi
- e: Euler's number
- golden_ratio
- euler_mascheroni: Euler-Mascheroni constant
- feigenbaum: Feigenbaum constant
- champernowne: Champernowne constant
- apery: Roger Apery's constant
- gelfond_schneider: Gelfond-Schneider constant
- khinchin: Aleksandr Khinchin constant
- ramanujan_soldner: Ramanujan-Soldner constant

Examples

```
# definition of pi
const$defs$pi

# value of pi
const$values$pi
```

create_shape *Create geometric shapes with optional text*

Description

This function creates various geometric shapes using base R graphics. It supports multiple shape types including circle, square, rectangle, triangle, hexagon, star, ellipse, and regular polygons with custom sides.

Usage

```
create_shape(  
  shape = c("circle", "square", "rectangle", "triangle", "hexagon", "star", "ellipse",  
            "polygon"),  
  x_center = 0,  
  y_center = 0,  
  size = 1,  
  width = size,  
  height = size,  
  n_sides = 5,  
  rotation = 0,  
  fill_color = NA,  
  border_color = "black",  
  line_width = 1,  
  title = "Geometric Shape",  
  text = NULL,  
  text_color = "black",  
  text_size = 1,  
  text_font = 1,  
  text_angle = 0,  
  show_axes = FALSE,  
  show_ticks = FALSE,  
  show_axis_labels = FALSE  
)
```

Arguments

shape	Character string specifying the shape to create. Must be one of: "circle", "square", "rectangle", "triangle", "hexagon", "star", "ellipse", "polygon".
x_center	Numeric, x-coordinate of the shape's center. Default is 0.
y_center	Numeric, y-coordinate of the shape's center. Default is 0.
size	Numeric, the primary size parameter (radius for circle, side length for square, etc.). Default is 1.

width	Numeric, width for rectangle and ellipse. Default is size.
height	Numeric, height for rectangle and ellipse. Default is size.
n_sides	Integer, number of sides for polygon. Default is 5.
rotation	Numeric, rotation angle in degrees. Default is 0.
fill_color	Character string or NA, the fill color of the shape. Default is NA (transparent).
border_color	Character string, the border color of the shape. Default is "black".
line_width	Numeric, the width of the border line. Default is 1.
title	Character string, the title of the plot. Default is "Geometric Shape".
text	Character string, optional text to display inside the shape. Default is NULL (no text).
text_color	Character string, color of the text. Default is "black".
text_size	Numeric, size of the text. Default is 5.
text_font	Character string, font family for the text. Default is "".
text_angle	Numeric, rotation angle for the text in degrees. Default is 0.
show_axes	Logical. Whether to display the axes. Default is FALSE.
show_ticks	Logical. Whether to display axis ticks. Default is FALSE.
show_axis_labels	Logical. Whether to display axis labels (X and Y). Default is FALSE.

Value

A list containing:

- plot: The ggplot object
- coordinates: Data frame with x and y coordinates of the shape's vertices
- center: Vector with x and y coordinates of the shape's center
- size: Primary size parameter used
- shape: Shape type
- text: Text displayed inside the shape (if any)

Examples

```
## Not run:
# Example 1: Basic circle
create_shape(shape = "circle", size = 2, fill_color = "lightblue")

# Example 2: Square with rotation and text
create_shape(
  shape = "square",
  size = 3,
  rotation = 45,
  fill_color = "salmon",
  border_color = "darkred",
  line_width = 2,
```

```
    text = "Rotated Square",
    text_color = "darkblue",
    text_size = 3
)

# Example 3: Hexagon with custom title
create_shape(
    shape = "hexagon",
    size = 2.5,
    fill_color = "lightgreen",
    border_color = "darkgreen",
    title = "My Hexagon"
)

# Example 4: Star with custom styling
create_shape(
    shape = "star",
    size = 3,
    rotation = 15,
    fill_color = "gold",
    border_color = "orange",
    line_width = 2,
    title = "Gold Star"
)

# Example 5: Rectangle with custom dimensions
create_shape(
    shape = "rectangle",
    width = 4,
    height = 2,
    fill_color = "lavender",
    text = "Rectangle",
    text_size = 2.5
)

# Example 6: Triangle with rotation and styling
create_shape(
    shape = "triangle",
    size = 3,
    rotation = 180,
    fill_color = "pink",
    border_color = "purple",
    line_width = 2,
    text = "rpkg",
    text_color = "black",
    text_size = 4
)

# Example 7: Ellipse with custom dimensions
create_shape(
    shape = "ellipse",
    width = 5,
    height = 2,
```

```
    rotation = 30,
    fill_color = "lightcyan",
    border_color = "steelblue",
    title = "Rotated Ellipse"
)

# Example 8: Custom polygon (octagon)
create_shape(
  shape = "polygon",
  n_sides = 8,
  size = 2,
  fill_color = "thistle",
  border_color = "purple",
  title = "Octagon",
  text = "8",
  text_color = "darkblue",
  text_size = 3
)

# Example 9: Overlapping shapes (need to call par() between shapes)
create_shape(
  shape = "circle",
  x_center = 0,
  y_center = 0,
  size = 3,
  fill_color = rgb(1, 0, 0, 0.5),
  title = "Overlapping Shapes"
)

par(new = TRUE) # Allow adding to the existing plot
create_shape(
  shape = "circle",
  x_center = 1,
  y_center = 1,
  size = 3,
  fill_color = rgb(0, 0, 1, 0.5),
  title = "" # Empty title to avoid overwriting the previous title
)
# Create a star with text
create_shape("star", size = 2,
            fill_color = "gold",
            text = "*",
            text_size = 8)

## End(Not run)
```

Description

Shorthand to remove elements from a data frame and save as the same name

Usage

```
data_pop(., n = 1, which = c("rows", "cols"), ret = FALSE)
```

Arguments

.	parent data
n	number of elements to remove
which	whether to remove from row or from column
ret	TRUE or FALSE. whether to return value instead of setting it to the parent data

Value

data with elements removed

Examples

```
#basic example: pop off 1 row from sample data
data.frame(ID=1:10,N=number(10)) #before
data_pop(data.frame(ID=1:10,N=number(10))) #after

#using data objects
data.01 <- mtcars[1:7,]

#task: remove 1 element from the end of the data and set it to the data name
data.01 #data.01 data before pop
data_pop(data.01) #does not return anything
data.01 #data.01 data updated after pop

#task: remove 3 columns from the end of the data and set it to the data name
data.01 #data.01 data before pop
data_pop(data.01, n = 3, which = "cols") #does not return anything, but updates data
data.01 #data.01 data updated after pop

#task: remove 5 elements from the end, but do not set it to the data name
data.01 #data.01 data before pop
data_pop(data.01,5, ret = TRUE) #return modified data
data.01 #data.01 data remains the same after pop
```

data_pop_filter	<i>Remove elements from a data matching filter</i>
-----------------	--

Description

Shorthand to remove elements from a data frame based on filter and save as the same name

Usage

```
data_pop_filter(., remove)
```

Arguments

.	data object
remove	expression for filter

Value

data filtered out based on the expression

Examples

```
# this function removes rows matching the filter expression
data.01 <- mtcars
data.02 <- airquality

#task: remove all mpg > 20
data.01 #data.01 data before pop
data_pop_filter(data.01,mpg > 15) #computes and resaves to variable
#note: this is different from subset(data.01,data.01$mpg > 15)
data.01 #modified data after pop based on filter

#task: remove all multiple. remove all elements where Month == 5 or Solar.R > 50
data.02 #data.02 data before pop
data_pop_filter(data.02,Month == 5 | Solar.R > 50) #computes and resaves to variable
data.02 #modified data after pop based on filter
```

data_push	<i>Add data to another data like array_push in PHP</i>
-----------	--

Description

Shorthand to add data to a dataset and save as the same name

Usage

```
data_push(., add, which = c("rows", "cols"))
```

Arguments

.	first data set
add	data set to add
which	where to append the new data e.g. rows or cols

Value

the combined dataset store to a variable with the name of the first

Examples

```
# initialize p1 and p2
init(p1,p2)
p1
p2

# declare p1 and p2 as data frame
p1 <- data.frame(PK=1:10,ID2=1:10)
p2 <- data.frame(PK=11:20,ID2=21:30)

p1
p2

#add p1 to p2 by row, and resave as p1
data_push(p1,p2,"rows")
# p2 # p2 remains the same
p1 #p1 has been updated

# declare a new data frame called p3
p3 <- data.frame(Hindex=number(20),Rindex=number(20,seed=20))

# add p3 to p1 as column, and resave as p1
data_push(p1,p3,"cols")
p1 # p1 has been updated
```

data_rep

Duplicate a data rows or columns X times

Description

Add a data to itself X times by rows or columns

Usage

```
data_rep(., n, which = c("rows", "cols"))
```

Arguments

. data frame variable
 n multiples of duplicate
 which where to append the duplicated data e.g. rows or cols

Value

the duplicated dataset store to a variable with the name of the first

Examples

```
# initialize p1 and p2
init(p1,p2)
p1
p2

# declare p1 and p2 as data frame
p1 <- data.frame(PK=1:10,ID2=1:10)
p2 <- data.frame(PK=11:20,ID2=21:30)

p1
p2

#add p1 twice by row, and resave as p1
data_rep(p1,n=2,"rows")
p1 #p1 has been updated

#add p2 3 times by col, and resave as p2
data_rep(p2,n=3,"cols")
p2 #p2 has been updated
```

 data_shuffle

Shuffle a data frame just like shuffle in PHP

Description

Shorthand to shuffle a data frame and save

Usage

```
data_shuffle(., which = c("rows", "cols"), seed = NULL)
```

Arguments

. data to shuffle as data frame
 which what to shuffle, rows or columns
 seed apply seed if indicated for reproducibility

Value

shuffled data frame of items store to the data frame name

Examples

```
#basic example
data.frame(ID=46:55,PK=c(rep("Treatment",5),rep("Placebo",5))) #before
data_shuffle(
  data.frame(ID=46:55,PK=c(rep("Treatment",5),rep("Placebo",5)))
) #after shuffle row
data_shuffle(
  data.frame(ID=46:55,PK=c(rep("Treatment",5),rep("Placebo",5))),
  which = "cols"
) #after shuffle column

# examples using object
df1<-data.frame(ID=46:55,PK=c(rep("Treatment",5),rep("Placebo",5)))

#illustrate basic functionality
data_shuffle(df1)
df1 #shuffle and resaved to variable

data.f2<-df1
data_shuffle(data.f2)
data.f2 #first output

data.f2<-df1
data_shuffle(data.f2)
data.f2 # different output from first output top

data.f2<-df1
data_shuffle(data.f2,seed = 344L)
data.f2 #second output

data.f2<-df1
data_shuffle(data.f2,seed = 344L)
data.f2 #the same output as second output top
```

date3to1

Combine vector to create Date, or split Date into vector

Description

Combine or split Date into a specified format

Usage

```
date3to1(data, out.format = "%Y-%m-%d", col.YMD = 1:3, as.vector = FALSE)

date1to3(
  data,
  in.format = "%Y-%m-%d",
  date.col = 1,
  out.cols = c("%Y", "%m", "%d")
)
```

Arguments

data	data frame object
out.format	date output format
col.YMD	columns to combine for Year, Month and Day
as.vector	return output as vector, or leave as data frame
in.format	date input format
date.col	numeric value of column within the dataset that contains the dates
out.cols	cols to of date items to split. Make sure to conform to date formats. See "NOTE" section for date formats

Details**NOTE for date3to1**

The three input columns corresponding to "Year Month Day" must be numeric values.

For example, Do not provide the month variable as non-numeric such as "Mar", "Jul", or "Jan".

If the values of the columns are non-numeric, the results will return an "NA" in the output.date column.

Value

date derived from combining values from three columns of a data frame

Note**DATE FORMATS IN R**

Date Specification	Description	Example
%a	Abbreviated weekday	Sun, Thu
%A	Full weekday	Sunday
%b	Abbreviated month	May, Jul
%B	Full month	March, July
%d	Day of the month	27, 07

%j	Day of the year	148, 188
%m	Month	05, 07
%U	Week, with Sunday as first day	22, 27
%w	Weekday, Sunday is 0	0, 4
%W	Week, with Monday as first day	21, 27
%x	Date, locale-specific	
%y	Year without century	84, 05
%Y	Year with century	1984, 2005
%C	Century	19, 20
%D	Date formatted %m/%d/%y	07/17/23
%u	Weekday, Monday is 1	7, 4

References

Adapted from Ecfun R package

Examples

```
# EXAMPLES FOR date3to1

data0 <- data.frame(y=c(NA, -1, 2001:2009),
m=c(1:2, -1, NA, 13, 2, 12, 6:9),
d=c(0, 0:6, NA, -1, 32) )
head(data0)

# combine and convert to date
# return as data frame
date3to1(data0)

# combine and convert to date
# return as vector
date3to1(data0, as.vector = TRUE) #eg. 2004-02-04

# combine and convert to date in the format DD_MM_YYYY
date3to1(data0, out.format = "%d_%m_%Y") #eg. 04_02_1974

# combine and convert to date in the format MM_DD_YY
date3to1(data0, out.format = "%m_%d_%y") #eg. 02_04_74

# combine and convert to date in the various date formats
date3to1(data0, out.format = "%B %d, %y") #eg. February 04, 74
date3to1(data0, out.format = "%a, %b %d, %Y") #eg. Mon, Feb 04, 1974
date3to1(data0, out.format = "%A, %B %d, %Y") #eg. Monday, February 04, 1974
date3to1(data0, out.format = "Day %j in Year %Y") #eg. Day 035 in Year 1974
date3to1(data0, out.format = "Week %U in %Y") #eg. Week 05 in 1974
date3to1(data0, out.format = "Numeric month %m in Year %Y") #eg. Numeric month 02 in Year 1974
```

```

# EXAMPLES FOR date1to3

data1 <- data.frame(Full.Dates =
                    c("2023-02-14", NA, NA,
                      "2002-12-04", "1974-08-04",
                      "2008-11-10"))

head(data1)

# split date with default settings
# return as data frame with columns
# for day(d), month(m) and year(Y)
date1to3(data1)

# split date in the format and only return year in YYYY
date1to3(data1, out.cols = "%Y") #eg. 2002, 2023

# split date in the format and only return month in m
date1to3(data1, out.cols = "%m") #eg. 02, 12, 08

# split date in the format and return multiple date formats columns
date1to3(data1, out.cols = c("%B", "%d") )
date1to3(data1, out.cols = c("%a", "%b", "%y") )
date1to3(data1, out.cols = c("%A", "%B", "%Y", "%y") )
date1to3(data1, out.cols = c("%j", "%Y", "%y", "%m") )
date1to3(data1, out.cols = c("%U", "%Y", "%y", "%x") )
date1to3(data1, out.cols = c("%m", "%Y", "%y", "%C") )

```

detect_outlier

Detect Outliers in a Numeric Vector

Description

This function identifies outliers in a numeric vector using either the interquartile range (IQR) method or the z-score method. The IQR method defines outliers as values below $Q1 - \text{multiplier} * \text{IQR}$ or above $Q3 + \text{multiplier} * \text{IQR}$, where $Q1$ and $Q3$ are the first and third quartiles. The z-score method identifies outliers as values with an absolute z-score exceeding a specified threshold.

Usage

```

detect_outlier(
  x,
  method = "iqr",
  multiplier = 1.5,
  z_threshold = 3,

```

```

    na.rm = TRUE,
    groups = NULL,
    summary = FALSE
)

iqr_outlier(x, multiplier)

zscore_outlier(x, z_threshold)

zscore_outlier2(x, z_threshold)

```

Arguments

<code>x</code>	A numeric vector in which to detect outliers.
<code>method</code>	A character string specifying the outlier detection method. Options are <code>"iqr"</code> (default) for the interquartile range method or <code>"zscore"</code> for the z-score method.
<code>multiplier</code>	A positive numeric value specifying the multiplier for the IQR method. Default is <code>'1.5'</code> , typically used for moderate outliers; <code>'3'</code> is common for extreme outliers. Ignored if <code>'method = "zscore"'</code> .
<code>z_threshold</code>	A positive numeric value specifying the z-score threshold for the <code>'method = "zscore"'</code> option. Default is <code>'3'</code> , meaning values with an absolute z-score greater than 3 are flagged as outliers. Ignored if <code>'method = "iqr"'</code> .
<code>na.rm</code>	A logical value indicating whether to remove <code>'NA'</code> values before computation. Default is <code>'TRUE'</code> . If <code>'FALSE'</code> and <code>'NA'</code> values are present, the function stops with an error.
<code>groups</code>	An optional vector of group names or labels corresponding to each value in <code>'x'</code> . If provided, must be the same length as <code>'x'</code> . Outlier detection is performed separately for each unique group, and results are returned as a nested list. Default is <code>'NULL'</code> (no grouping).
<code>summary</code>	A logical value indicating whether to include a summary in the output. Default is <code>'FALSE'</code> . If <code>'TRUE'</code> , the output list includes a <code>'summary'</code> element with descriptive statistics and outlier counts, either overall or by group if <code>'groups'</code> is provided.

Details

The function returns a list containing the outliers, their indices, detection bounds or thresholds, and a logical vector indicating which elements are outliers. If a grouping vector is provided via `'groups'`, outlier detection is performed separately for each group, and results are returned as a nested list by group. If `'na.rm = TRUE'` (default), missing values (`'NA'`) are removed before computation. If `'na.rm = FALSE'` and `'NA'` values are present, the function stops with an error. The function also stops for non-numeric input, insufficient valid data, or mismatched group lengths.

The function requires at least two non-`'NA'` values per group (if `'groups'` is provided) or overall (if `'groups = NULL'`) to compute meaningful statistics when `'na.rm = TRUE'`. If `'na.rm = FALSE'`, the presence of `'NA'` values triggers an error. If all values in a group are identical or there are insufficient data points, an error is thrown for that group. The IQR method is robust to non-normal data, while the z-score method assumes approximate normality and is sensitive to extreme values.

Value

If `'groups = NULL'` (default), a list with the following components: - `'outliers'`: A numeric vector of the outlier values. - `'indices'`: An integer vector of the indices where outliers occur in the input vector. - `'bounds'` (if `'method = "iqr"'`): A named numeric vector with the `'lower'` and `'upper'` bounds for outliers. - `'threshold'` (if `'method = "zscore"'`): A named numeric vector with the `'lower'` and `'upper'` z-score thresholds. - `'is_outlier'`: A logical vector of the same length as `'x'`, where `'TRUE'` indicates an outlier. - `'summary'` (if `'summary = TRUE'`): A list with summary statistics including the mean, median, standard deviation (for z-score), quartiles (for IQR), and number of outliers.

If `'groups'` is provided, a named list where each element corresponds to a unique group, containing the same components as above but computed separately for that group's values.

Examples

```
# Example 1: Basic IQR method without groups
x <- c(1, 2, 3, 4, 100)
detect_outlier(x)

# IQR method with summary
detect_outlier(x, summary = TRUE)

# Z-score method with custom threshold
y <- c(-10, 1, 2, 3, 4, 5, 20)
detect_outlier(y, method = "zscore", z_threshold = 2.5)

# Handling missing values
z <- c(1, 2, NA, 4, 5, 100)
detect_outlier(z, method = "iqr", multiplier = 3)

# Example 2: IQR method with groups
x2 <- c(1, 2, 3, 100, 5, 6, 7, 200)
groups2 <- c("A", "A", "A", "A", "B", "B", "B", "B")
detect_outlier(x2, groups = groups2)

# Example 3: Z-score method with groups and summary
x3 <- c(-10, 1, 2, 20, 3, 4, 5, 50)
groups3 <- c("X", "X", "X", "X", "Y", "Y", "Y", "Y")
detect_outlier(x3, method = "zscore", z_threshold = 2, groups = groups3, summary = TRUE)

# Example 4: IQR method with groups and NA values
x4 <- c(1, 2, NA, 100, 5, 6, 7, 200, 1000)
groups4 <- c("G1", "G1", "G1", "G1", "G2", "G2", "G2", "G2", "G1")
detect_outlier(x4, groups = groups4)

# Error cases
## Not run:
detect_outlier(c("a", "b")) # Non-numeric input
detect_outlier(c(1), groups = c("A")) # Insufficient data
detect_outlier(c(1, 2), groups = c("A")) # Mismatched group length
detect_outlier(c(1, NA, 3), na.rm = FALSE) # NA with na.rm = FALSE
```

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

detect_outlier2 *Advanced Outlier Detection in Numeric Data with Optional Grouping*

Description

Detects outliers in numeric data using multiple statistical methods and provides comprehensive analysis including visualizations and summary statistics. Supports grouped analysis when group vector is provided. The function supports three detection methods: IQR, Z-score, and Modified Z-score.

Usage

```
detect_outlier2(
  x,
  groups = NULL,
  method = c("iqr", "zscore", "modified_zscore", "all"),
  multiplier = 1.5,
  z_threshold = 3,
  modified_z_threshold = 3.5,
  plot = TRUE,
  plot_groups = TRUE
)
```

Arguments

x	A numeric vector containing the data to analyze
groups	Optional vector of group labels, must be same length as x
method	Character string specifying the outlier detection method. Options are "iqr", "zscore", "modified_zscore", or "all". Default is "iqr"
multiplier	Numeric value specifying the IQR multiplier for outlier detection. Default is 1.5
z_threshold	Numeric value specifying the Z-score threshold. Default is 3
modified_z_threshold	Numeric value specifying the modified Z-score threshold. Default is 3.5
plot	Logical indicating whether to generate visualization plots. Default is TRUE
plot_groups	Logical indicating whether to create separate plots for each group. Only used when groups are provided. Default is TRUE

Value

A list of class "outlier_analysis" containing:

- overall: Overall analysis results
- by_group: Group-specific results (if groups provided)
- plots: List of generated plots

Examples

```

# Example 1: Basic grouped analysis
set.seed(123)
data <- c(rnorm(50), rnorm(50, 2), rnorm(50, 4))
groups <- rep(c("A", "B", "C"), each = 50)
resultA <- detect_outlier2(data) # no groups
resultB <- detect_outlier2(data, groups = groups) # with groups

# Example 2: Custom thresholds by group
test_scores <- c(65, 70, 75, 72, 68, 73, 78, 71, 69, 74,
                90, 85, 92, 88, 95, 87, 91, 89, 86, 93)
class_groups <- rep(c("Morning", "Afternoon"), each = 10)
result <- detect_outlier2(test_scores,
                          groups = class_groups,
                          method = "all",
                          z_threshold = 2)

result$overall
result$by_group
result$plots$overall$boxplot()
result$plots$overall$density()
result$plots$overall$comparison()
result$plots$by_group$Morning$boxplot()
result$plots$by_group$Morning$density()
result$plots$by_group$Afternoon$boxplot()
result$plots$by_group$Afternoon$density()

```

duplicate

Duplicate a file with global text replacement

Description

Shorthand to return a re-sample number of rows in a data frame by unique column

Usage

```
duplicate(file, new.name, pattern = NULL, replacement = NULL, open = TRUE)
```

Arguments

file	data frame to re-sample
new.name	column to uniquely re-sample
pattern	number of rows to return
replacement	unique numeric value for reproducibility
open	description

Value

data frame containing re-sampled rows from an original data frame

Examples

```

if(interactive()){
# example to duplicate a file, and replace text1 within it
# NOTE that, by default, this function will also open the file within RStudio

#create sample file
file1s <- paste0(tempfile(),".R")
writeLines("message(
'Sample items: eggs, coke, fanta, book'
)", file1s)

file2s <- paste0(tempfile(),".R")
file3s <- paste0(tempfile(),".R")

duplicate(
  file = file1s,
  new.name = file2s,
  pattern = 'text1',
  replacement = 'replacement1'
)

# duplicate the file, with multiple replacements
# replace 'book' with 'egg' and 'coke' with 'fanta'
duplicate(
  file1s, file2s,
  pattern = c('book','coke'),
  replacement = c('egg','fanta')
)

# duplicate the file with no replacement
duplicate(file1s,file3s) # this simply performs file.copy, and opens the new file

# duplicate the file but do not open for editing
duplicate(file1s,file3s, open = FALSE) # this does not open file after duplication
}

```

error.out

Error coalescing operator

Description

Alternative return for error statements. Return alternative if the value of expression is erroneous

Usage

```
error.out(test, alternative = "")  
  
test %eo% alternative
```

Arguments

```
test          an object to return  
alternative    alternative object to return
```

Value

value of test if error, else return value of alternative

Examples

```
# The following statement would produce  
# error because 'stat1' does not exist  
  
# stat1 + 1  
  
# To prevent the statement from  
# stopping the process, when can have alternative out  
alt = 0  
error.out(stats1 + 1, alt)
```

extract_comment	<i>Extract all comments or functions from a file</i>
-----------------	--

Description

Vectorize all comments from the file

Usage

```
extract_comment(file)  
  
remove_content_in_quotes(line)  
  
remove_comment(line)  
  
clean_file(file, output_file)  
  
get_func_def(file)
```

Arguments

file file to parse
line string vector to remove contents within quotes or comments
output_file file path to write the output of the new file

Value

vector of all comments within a file

Examples

```
## Not run:  
ex_file1 <- "path/file1.R"  
# get all comments  
cmmts <- extract_comment(ex_file1)  
cmmts  
  
## End(Not run)  
  
## Not run:  
# Ex to clean out comments from file  
file_path <- ".testR"  
output_file_path <- ".cleaned_script.R"  
clean_file(file_path, output_file_path)  
  
## End(Not run)  
  
# example code  
  
## Not run:  
# Ex to get all defined functions  
# within a file  
file_path <- ".testR"  
get_func_def(file_path)  
  
## End(Not run)
```

extract_IP

Extract all IP addresses from a string

Description

Parse a string and vectorize the IP addresses within it

Usage

```
extract_IP(input)
```

Arguments

input input string containing IP

Value

extract_IP returns a vector of IP addresses

Examples

```
# Extract all IP addresses from a string

# Example 1
# This example demonstrates the separate
# extraction of an IP address one per vector element:
str1 = c("Two IP(66.544.33.54) addresses
were discovered in the scan.
One was at 92.234.1.0.",
"The other IP was 62.3.45.255.")
extract_IP(str1)

# Example 2
# This example demonstrates the extraction
# of multiple IP addresses from a single vector element:
str2 = "Two IP addresses were discovered
in the scan. One was at 92.234.1.0.
The other IP was 62.3.45.255."
extract_IP(str2)
```

fAddDate *Append date to filename*

Description

Add today's date to the filename

Usage

```
fAddDate(..., format = "%d-%b-%Y")
```

Arguments

... file name or path to concat
format time format e.g. %d-%b-%Y , refer to [date3to1](#) for date formats

Details

The present function enables users to add the current date to the file name, facilitating the straightforward saving of files with their respective dates. It accepts different file paths and names as arguments, as demonstrated in the example section. This functionality simplifies the process of associating a file's creation date with its name, aiding users in recalling when a file was saved. Moreover, it serves as a preventive measure against unintentional overwriting of files created on different dates.

Value

file name with the current date added as suffix

Examples

```
# Task 1
fAddDate("path1/", "path2/filepre", "filemid", "fileend.png")

# Task 2
fAddDate(c("path1/", "path2/"), "filepre", "filemid", "fileend.png")

# Task 3
fAddDate("one_file_name_fileend.pdf")

# Task 4
fAddDate(c("path1/", "path2/"), "file1-", "filemid", c("fileend.png", ".pdf"))

# Task 5
data("USArrests")
USArrests$fn = paste0(row.names(USArrests), ".txt")
head(fAddDate(USArrests$fn), 10)

# Task 6: format date - month.day.year
fAddDate("sample_file_name.pdf", format = "%B.%d.%YYYY")

# Task 7: format date - year_month
fAddDate("sample_file_name.pdf", format = "%YYYY_%m")
```

find_packages

Fetch R package based on keyword

Description

This function gets R packages based on a keyword in their description.

Usage

```
find_packages(keyword)
```

Arguments

keyword character, the keyword to search package descriptions for.

Value

character vector of matching package names or NULL if no matches.

Note

the function in its current form only searches available.packages()

Examples

```
# find the list of R packages for data or machine learning
matched_pkgs <- suppressWarnings(find_packages("plotting"))
matched_pkgs

matched_pkgs <- suppressWarnings(find_packages("machine learning"))
matched_pkgs
```

from_tensor_slices *Create Tensor-Like Slices from a Data Frame or Matrix*

Description

This function converts a data frame or matrix into row-wise slices, similar to TensorFlow's 'from_tensor_slices()'.

Usage

```
from_tensor_slices(data)
```

Arguments

data A data frame or matrix to be converted into slices.

Value

A list of slices, each corresponding to a row of the input data.

Examples

```
df <- data.frame(A = 1:3, B = c("x", "y", "z"))
from_tensor_slices(df)
```

`geo.cv`*Calculate geometric coefficient of variation, mean, or SD and round*

Description

Calculate the coefficient of variation
Calculate the geometric mean
Calculate the geometric standard deviation

Usage

```
geo.cv(num, round = 2, na.rm = TRUE, neg.rm = TRUE, pct = TRUE)

geo.mean(num, round = 2, na.rm = TRUE, neg.rm = TRUE)

geo.sd(num, round = 2, na.rm = TRUE, neg.rm = TRUE)
```

Arguments

<code>num</code>	vector of numbers
<code>round</code>	round result to decimal place
<code>na.rm</code>	remove NAs from the vector
<code>neg.rm</code>	remove negative values from the vector
<code>pct</code>	TRUE or FALSE. should result be in percent

Value

the geometric cv of a set of numbers
the geometric mean of a set of numbers
the geometric standard deviation of a set of numbers

Examples

```
#simulate numbers using a fixed seed
num1 <- number(n = 1115,max.digits = 4, seed = 10)

#get geometric CV, represent as percent and round to 2 decimal places
geo.cv(num1,round = 2) # result: 60.61%

#or round to 3 decimal places
geo.cv(num1,round = 3) # result: 60.609%

#by default, the above examples return a CV%
#if you do not want the result as percentage, specify "pct"
geo.cv(num1,pct = FALSE) # result: 0.61
```

```

num1 <- sample(300:3000,10)

#get the geometric mean, excluding all negatives and round to 2
geo.mean(num1)

#or
geo.mean(num1)

#get geometric mean, but round the final value to 5 decimal places
geo.mean(num1, round = 5)

num1 <- sample(330:400,20)

#get geometric SD remove negative values and round to 2 decimal places
geo.sd(num1)

#get geometric SD, DON'T remove negative values and round to 2 decimal places
geo.sd(num1,na.rm=FALSE)

#get geometric SD, remove negative values and round to 3 decimal places
geo.sd(num1,round = 3)

```

getDate *Extract all dates from a string*

Description

Extract all dates from a string

Usage

```
getDate(str1, out.format = "%Y-%m-%d")
```

Arguments

str1	string to parse
out.format	date output format

Note

DATE FORMATS IN R

Date Specification	Description	Example
%a	Abbreviated weekday	Sun, Thu
%A	Full weekday	Sunday
%b	Abbreviated month	May, Jul

%B	Full month	March, July
%d	Day of the month	27, 07
%j	Day of the year	148, 188
%m	Month	05, 07
%U	Week, with Sunday as first day	22, 27
%w	Weekday, Sunday is 0	0, 4
%W	Week, with Monday as first day	21, 27
%x	Date, locale-specific	
%y	Year without century	84, 05
%Y	Year with century	1984, 2005
%C	Century	19, 20
%D	Date formatted %m/%d/%y	07/17/23
%u	Weekday, Monday is 1	7, 4

Examples

```
str1 <- "The video was recorded on July 19, 2023."
str2 <- "The video was recorded over a 4 hour period
starting on July 19, 2023."
str3 <- "The first batch aug 12,2024 of aug 12,
2024 reports are due on July 12, 2024; the second
batch on 7/19/24."
str4 <- c("On 3.12.25, Jerry is taking one month
05.13.1895 of leave and is not scheduled to
return until around 4-9-2025.", "The staff
will be out on training on 10/11/24,
Oct 12, 2024, and 10-13-24.")
getDate(str1)
getDate(str2, out.format = "%Y-%m-%d")
getDate(str3, out.format = "%m-%d/%Y")
getDate(str4)
```

```
getGitRepoStart
```

```
Fetch GitHub Repository Creation & Last Updated Date
```

Description

The GitHub REST API is a powerful tool that allows developers to interact with GitHub programmatically. It provides a set of endpoints that allows a user to create integration, retrieve data, and automate workflows related to GitHub repositories. It is a means by which users can interact with GitHub without directly using a web interface.

Usage

```
getGitRepoStart(repo_name, out.format = "%Y-%m-%d")

getGitRepoChange(repo_name, out.format = "%Y-%m-%d")
```

Arguments

repo_name name of the repository
 out.format date output format

Details

The two functions utilize the GitHub REST API to extract important temporal information about a GitHub repository.

- the getGitRepoStart function is used to retrieve the date a GitHub repository was first created.
- the getGitRepoChange function retrieves the date a GitHub repository was last updated.

Value

date of creation of repository as a character
 date of the last update of repository as a character

Examples

```
# Use default date format
getGitRepoStart(repo_name = "oobianom/quickcode")

# Specify date format
getGitRepoStart(repo_name = "oobianom/quickcode", out.format = "%j|%Y")
getGitRepoStart(repo_name = "oobianom/quickcode", out.format = "%D|%j")

getGitRepoChange(repo_name = "oobianom/shinyStorePlus", out.format = "%d-%b-%Y")
getGitRepoChange(repo_name = "oobianom/r2social", out.format = "%Y/%m/%d")
```

 getWeekSeq

Convert Dates into Numeric Week Counts

Description

Convert the range of date to number of weeks

Usage

```
getWeekSeq(start_date, end_date, dates, in.format = "%m/%d/%y")

is.Date(x)
```

```
not.Date(x)
```

```
is.leap(yyyy)
```

Arguments

start_date	A scaler of class Date (if this argument is populated, the date arg must be empty)
end_date	A scaler of class Date; must be later than the start_date (if this argument is populated, the date arg must be empty)
dates	vector of dates that need not be sequential but that reference values of class Date; Note that if this argument is populated, the start_date and end_date arguments must be empty
in.format	date input format
x	date item to check
yyyy	year numeric value eg 1989

Value

data frame of the dates along with their corresponding week

Examples

```
# simple example with start and end date
getWeekSeq(start_date="12/29/25",end_date="1/8/26")
```

```
# enter specific dates instead
# specify format
getWeekSeq(
  dates = c(
    '2025-12-29',
    '2025-12-30',
    '2025-12-31',
    '2026-01-01',
    '2026-01-04',
    '2026-01-05',
    '2026-01-06',
    '2026-01-07',
    '2026-01-08'),
  in.format = "%Y-%m-%d"
)
```

```
getWeekSeq(
  dates = c(
    '12/29/25',
    '12/30/25',
    '12/31/25',
    '01/01/26',
    '01/02/26',
    '01/03/26',
    '01/06/26',
```

```
'01/07/26',  
'01/08/26'  
)  
in.format = "%m/%d/%y"  
)
```

has.error

Check if a call or expression produces errors

Description

Whether a function or series of calls results in error

Usage

```
has.error(...)
```

Arguments

... the expression or function calls

Value

boolean value to indicate if the expression produces errors

Note

More information, check: <https://rpkg.net/package/quickcode>

Examples

```
# this should not produce error  
# so the function result should be FALSE  
has.error({  
  x = 8  
  y = number(10)  
  res = x + y  
})  
  
# this should produce the following error  
# Error in x + y : non-numeric argument to binary operator  
# so the function result should be TRUE  
has.error({  
  x = 8  
  y = "random"  
  res = x + y  
})  
  
# this should result in error because
```

```
# the dataset does not contain a "rpkg.net" column
# the result should be TRUE
df1 = mtcars
has.error(df1[, "rpkg.net"])
```

code header.rmd

Snippet function to add header to a current Rmd opened file

Description

Shorthand to add Rmd header

Usage

```
header.rmd()
```

Value

Inserts header content for Rmd file

Examples

```
if(interactive())
  header.rmd()
```

code in.range

If number falls within a range of values and get closest values

Description

With a defined range of values, the function systematically examines each provided number to determine if it falls within the specified range. It may also provide the values with the range that are closest to a desired number.

Usage

```
in.range(
  value,
  range.min,
  range.max,
  range.vec = NULL,
  closest = FALSE,
  rm.na = FALSE
)
```

Arguments

value	NUMERIC. the vector of numbers to check
range.min	NUMERIC. OPTIONAL. the minimum value of the range
range.max	NUMERIC. OPTIONAL. the maximum value of the range
range.vec	NUMERIC. OPTIONAL. a vector of numbers to use for the range
closest	BOOLEAN. OPTIONAL. return closest value
rm.na	BOOLEAN. OPTIONAL. remove NA values from input

Details

The described function serves the purpose of checking whether a given number or set of numbers falls within a specified range. It operates by taking a range of values as input and then systematically evaluates each provided number to determine if it lies within the defined range. This function proves particularly useful for scenarios where there is a need to assess numeric values against predefined boundaries, ensuring they meet specific criteria or constraints. In the same manner, this function allows the user to also retrieve values within the range that are closest to each provided number.

Value

boolean to indicate if the value or set of values are within the range

Note

The argument `range.vec` is utilized when users opt not to employ the `range.min` or `range.max` arguments. If `range.vec` is specified, `range.min` and `range.max` are disregarded. It's important to note that the use of `range.vec` is optional.

Examples

```
# Task 1: Check if a number is within specified range
in.range(5, range.min = 3, range.max = 10) # TRUE
in.range(25, range.min = 12, range.max = 20) # FALSE

# Task 2: Check if a set of values are within a specified range
in.range(1:5, range.min = 2, range.max = 7) #
in.range(50:60, range.min = 16, range.max = 27) #

# Task 3: Check if a number is within the range of a set of numbers
in.range(5, range.vec = 1:10) # TRUE
in.range(345, range.vec = c(1001,1002,1003,1004,1005,
1006,1007,1008,1009,1010,1011,1012,1013,1014)) # FALSE

# Task 4: Check if a set of values are within the range of a set of numbers
in.range(1:5, range.vec = 4:19) #
in.range(50:60, range.vec = c(55,33,22,56,75,213,120)) #

# Task 5: remove NAs prior to processing
in.range(c(1,3,NA,3,4,NA,8), range.min = 4, range.max = 6, rm.na = FALSE) # do not remove NA
```

```
in.range(c(1,3,NA,3,4,NA,8), range.min = 4, range.max = 6, rm.na = TRUE) # remove NA
#in.range(c(NA), range.min = 4, range.max = 6, rm.na = TRUE) #This will return error

# Task 6: return the closest number to the value
in.range(5:23, range.vec = 7:19, closest = TRUE)
in.range(-5:10, range.vec = -2:19, closest = TRUE)
in.range(c(1:5,NA,6:9), range.vec = 4:19, closest = TRUE)
in.range(c(1:5,NA,6:9), range.vec = 4:19, closest = TRUE, rm.na = TRUE)
```

inc

Increment vector by value

Description

Increment the content of a vector and re-save as the vector

Usage

```
inc(., add = 1L)
```

Arguments

.	vector of number(s)
add	number to add

Details

This function is very useful when writing complex codes involving loops. Apart from the for loop, this can be useful to quickly increment a variable located outside the loop by simply incrementing the variable by 1 or other numbers. Check in the example section for a specific use. Nonetheless, one may also choose to use this function in any other instance, as it's simple purpose is to increase the value of a variable by a number and then re-save the new value to that variable.

Value

a vector incremented by a number

Examples

```
num1 <- sample(330:400,10)
num1#before increment

# increment num1 by 1
inc(num1)
num1 #after increment

# increment num1 by 5
num1 #before increment
```

```

inc(num1, add= 10)
num1 #after increment

#when used in loops

#add and compare directly
rnum = 10
inc(rnum) == 11 #returns TRUE
rnum #the variable was also updated

# use in a for loop
ynum = 1
for( i in c("scientist","dancer","handyman","pharmacist")){
message("This is the item number ")
message(ynum)
message(". For this item, I am a ")
message(i)

#decrement easily at each turn
plus(ynum)
}

#use in a repeat loop
xnum = 1
repeat{ #repeat until xnum is 15
message(xnum)
if(inc(xnum) == 15) break
}

```

init

Initialize new variables and objects

Description

Shorthand to initialize one or more objects

Usage

```
init(..., value = NULL)
```

Arguments

...	variable names to initialize
value	value to initialize them to

Value

initialized objects set to the value specified

Examples

```
init(t,u,v)
message(t) # t = NULL
message(u) # u = NULL
message(v) # v = NULL
init(j,k,m,value = 7)
message(j) # j = 7
message(k) # k = 7
message(m) # m = 7
```

insertInText

Shiny app function to insert string to current file in RStudio

Description

Shorthand to insert content to opened file

Usage

```
insertInText(string)
```

Arguments

string what to insert

Value

Inserts into current position on opened file

Examples

```
if(interactive()){
  insertInText('hello rpkg.net')
  insertInText('hello world')
}
```

is.image	<i>Is file name extension(s) an image</i>
----------	---

Description

Check if one or multiple file name entry is an image

Usage

is.image(x)

Arguments

x vector entry

Details

This current function tests if the extension of the file name provided belongs to any of the image extensions listed below

AI - Adobe Illustrator
BMP - Bitmap Image
CDR - Corel Draw Picture
CGM - Computer Graphics Metafile
CR2 - Canon Raw Version 2
CRW - Canon Raw
CUR - Cursor Image
DNG - Digital Negative
EPS - Encapsulated PostScript
FPX - FlashPix
GIF - Graphics Interchange Format
HEIC - High-Efficiency Image File Format
HEIF - High-Efficiency Image File Format
ICO - Icon Image
IMG - GEM Raster Graphics
JFIF - JPEG File Interchange Format
JPEG - Joint Photographic Experts Group
JPG - Joint Photographic Experts Group
MAC - MacPaint Image
NEF - Nikon Electronic Format
ORF - Olympus Raw Format
PCD - Photo CD
PCX - Paintbrush Bitmap Image
PNG - Portable Network Graphics
PSD - Adobe Photoshop Document
SR2 - Sony Raw Version 2
SVG - Scalable Vector Graphics
TIF - Tagged Image File

TIFF - Tagged Image File Format
WebP - Web Picture Format
WMF - Windows Metafile
WPG - WordPerfect Graphics

Value

a boolean value to indicate if entry is an image

Examples

```
img.1 <- "fjk.jpg"
is.image(img.1)

img.0 <- "fjk.bbVG"
is.image(img.0)

img.2 <- "fjk.bmp"
is.image(img.2)

img.3 <- "fjk.SVG"
is.image(img.3)

# a vector of file names
v <- c("logo.png", "business process.pdf",
      "front_cover.jpg", "intro.docx",
      "financial_future.doc", "2022 buybacks.xlsx")

is.image(v)

# when the file name has no extension
# the function returns NA
v2 <- c("img2.jpg", "northbound.xlsx", "landing", NA)
is.image(v2)
```

is.increasing

Check if numbers in a vector are decreasing or increasing

Description

Test if numeric values of a vector are decreasing or increasing using the radix method

Usage

```
is.increasing(., na.last = TRUE)
```

```
is.decreasing(., na.last = TRUE)
```

Arguments

. a numeric vector
na.last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed.

Value

boolean value to indicate if the values are increasing or decreasing

Examples

```
# example code
doy1 <- rnorm(1e3)
doy2 <- sort(doy1, decreasing = FALSE)
doy3 <- sort(doy1, decreasing = TRUE)

is.increasing(doy1)
is.decreasing(doy1)

is.increasing(doy2)
is.decreasing(doy2)

is.increasing(doy3)
is.decreasing(doy3)
```

is.lognormal

Check if a data fits the distribution

Description

Check whether a vector of data contains values that fit a distribution

Usage

```
is.lognormal(values, alpha = 0.05, method = 1)
is.normal(values, alpha = 0.05, method = 1)
is.uniform(values, alpha = 0.05)
is.poisson(values, alpha = 0.05)
is.gamma(values, alpha = 0.05)
is.logistic(values, alpha = 0.05)
is.weibull(values, alpha = 0.05)
```

```
is.cauchy(values, alpha = 0.05)

setDisAlpha(alpha = 0.05)

unsetDisAlpha()

getDistribution(data, alpha = 0.05)
```

Arguments

values	vector of values
alpha	p-value alpha level
method	method for calculation, where 1 = Shapiro-Wilk test and 2 = Kolmogorov-Smirnov test
data	the data to check against distributions

Details

This function takes a numeric vector as its input. This vector contains the dataset that will be analyzed.

For Normal and LogNormal:

- Method 1: we perform the Shapiro-Wilk test on the (log-transformed) data to test for normality. The null hypothesis of the Shapiro-Wilk test is that the data are normally distributed. If the p-value is greater than the chosen significance level (typically 0.05), we fail to reject the null hypothesis, indicating that the data may follow a log-normal distribution.

- Method 2: we perform the Kolmogorov-Smirnov test on the log-transformed data, comparing it to a normal distribution with the same mean and standard deviation. Again, if the p-value is greater than the chosen significance level, it suggests that the data may follow a log-normal distribution. These tests provide a statistical assessment of whether your data follows a log-normal distribution.

Value

boolean value if lognormal distributed
boolean value if normal distributed
boolean value if uniform distributed
boolean value if poisson distributed
boolean value if gamma distributed
boolean value if logistic distributed
boolean value if logistic distributed
boolean value if cauchy distributed
setDisAlpha sets global significance level for testing of distribution
unsetDisAlpha removes global significance level for testing of distribution

Note

getDistribution() checks if data fits multiple distributions

Examples

```
# Set global alpha for testing significance
setDisAlpha(alpha = 0.05)

# Prepare all data to test
# Set the seed for reproducibility
set.seed(13200323)
lognormal_data <- stats::rlnorm(n = 4000, meanlog = 1, sdlog = 1) #lognormal data
normal_data <- stats::rnorm(n = 4000, mean = 10, sd = 3) #normal data
uniform_data <- stats::runif(4000,min=0,max=10) #uniform data
poisson_data <- stats::rpois(4000, lambda = 5) #poisson data
gamma_data <- stats::rgamma(4000,shape = 5, rate = 2) #gamma data
logis_data <- stats::rlogis(4000, location = 4, scale = 2)#logistic values
weibull_data <- stats::rweibull(4000, shape = 4, scale = 2) #weibull data
cauchy_data <- stats::rcauchy(4000, location = 8, scale = 5) #cauchy data

# EXAMPLE FOR is.lognormal

# Test if the data is lognormal
is.lognormal(lognormal_data)
is.lognormal(normal_data)
is.lognormal(uniform_data)
is.lognormal(poisson_data)
is.lognormal(gamma_data)
is.lognormal(logis_data)
is.lognormal(weibull_data)
is.lognormal(cauchy_data)
is.lognormal(1:4000)

# EXAMPLE FOR is.normal

# Test if the data fits a normal distribution
is.normal(lognormal_data)
is.normal(normal_data)
is.normal(uniform_data)
is.normal(poisson_data)
is.normal(gamma_data)
is.normal(logis_data)
is.normal(weibull_data)
is.normal(cauchy_data)
is.normal(1:4000)

## Not run:
# EXAMPLES for is.uniform

# Test if the data fits a uniform distribution
is.uniform(lognormal_data)
is.uniform(normal_data)
```

```
is.uniform(uniform_data)
is.uniform(poisson_data)
is.uniform(gamma_data)
is.uniform(logis_data)
is.uniform(weibull_data)
is.uniform(cauchy_data)
is.uniform(1:4000)

## End(Not run)
## Not run:
# EXAMPLE for is.poisson

# Test if the data fits a poisson distribution
is.poisson(lognormal_data)
is.poisson(normal_data)
is.poisson(uniform_data)
is.poisson(poisson_data)
is.poisson(gamma_data)
is.poisson(logis_data)
is.poisson(weibull_data)
is.poisson(cauchy_data)
is.poisson(1:4000)

## End(Not run)
## Not run:
# EXAMPLE for is.gamma

# Test if the data fits a gamma distribution
is.gamma(lognormal_data)
is.gamma(normal_data)
is.gamma(uniform_data)
is.gamma(poisson_data)
is.gamma(gamma_data)
is.gamma(logis_data)
is.gamma(weibull_data)
is.gamma(cauchy_data)
is.gamma(1:4000)

## End(Not run)
## Not run:
# EXAMPLE for is.logistic

# Test if the data fits a logistic distribution
is.logistic(lognormal_data)
is.logistic(normal_data)
is.logistic(uniform_data)
is.logistic(poisson_data)
is.logistic(gamma_data)
is.logistic(logis_data)
is.logistic(weibull_data)
is.logistic(cauchy_data)
is.logistic(1:4000)
```

```
## End(Not run)
## Not run:
# Test if the data fits a weibull distribution
is.weibull(lognormal_data)
is.weibull(normal_data)
is.weibull(uniform_data)
is.weibull(poisson_data)
is.weibull(gamma_data)
is.weibull(logis_data)
is.weibull(weibull_data)
is.weibull(cauchy_data)
is.weibull(1:4000)

## End(Not run)
## Not run:
# EXAMPLES for is.cauchy

# Test if the data fits a cauchy distribution
is.cauchy(lognormal_data)
is.cauchy(normal_data)
is.cauchy(uniform_data)
is.cauchy(poisson_data)
is.cauchy(gamma_data)
is.cauchy(logis_data)
is.cauchy(weibull_data)
is.cauchy(cauchy_data)
is.cauchy(1:4000)

## End(Not run)
## Not run:
# set global distribution alpha

# default setting
setDisAlpha()

# set to 0.001
setDisAlpha(alpha = 0.01)

## End(Not run)
## Not run:
# unset global distribution alpha

unsetDisAlpha()

## End(Not run)
```

lastwd

Go back to previous directory

Description

Navigate to previous working directory if the setwd was called by clean() or refresh() function

Usage

```
lastwd()
```

Value

the previous directory

Note

In order to use this function to retrieve the last/previous working directory, you must use the `quickcode::clean()` or `quickcode::refresh()` function to set the working directory.

Examples

```
lastwd()
```

learn_rate_scheduler *Learning Rate Scheduler*

Description

This function adjusts the learning rate at each epoch according to a custom schedule.

Usage

```
learn_rate_scheduler(initial_lr, schedule, epochs)
```

Arguments

<code>initial_lr</code>	A numeric value representing the initial learning rate.
<code>schedule</code>	A function that takes the epoch number as input and returns the adjusted learning rate.
<code>epochs</code>	An integer specifying the total number of epochs.

Value

A numeric vector representing the learning rate for each epoch.

Examples

```
schedule <- function(epoch) { 0.01 * (0.9 ^ epoch) }  
learn_rate_scheduler(0.01, schedule, epochs = 10)
```

libraryAll	<i>Load specific R libraries and clear environment</i>
------------	--

Description

Load specific packages, print a list of the loaded packages along with versions. Only include libraries, don't install if library doesn't exist

Usage

```
libraryAll(  
  ...,  
  lib.loc = NULL,  
  quietly = FALSE,  
  clear = TRUE,  
  clearPkgs = FALSE  
)
```

Arguments

...	multiple library names
lib.loc	OPTIONAL. library store location
quietly	OPTIONAL. attach library quietly
clear	OPTIONAL. clear environment after attach
clearPkgs	Clear previous loaded packages, TRUE or FALSE

Value

loaded libraries and clear environment

Examples

```
# load packages and print their versions to the console  
libraryAll(base) #one package  
  
libraryAll(  
  base,  
  tools,  
  stats  
) #multiple packages  
  
libraryAll("grDevices") #with quotes  
  
libraryAll(  
  stats,  
  utils,  
  quietly = TRUE
```

```
) #load quietly

libraryAll(
  base,
  clear = FALSE) #do not clear console after load

# clear previously loaded packages, then load r2resize and r2social
libraryAll(
  stats,
  utils,
  clearPkgs = TRUE
)
```

list_push

Add elements to a list like array_push in PHP

Description

Shorthand to add elements to a vector and save as the same name

Usage

```
list_push(., add)
```

Arguments

.	first list
add	list to add

Value

vector combining fist and second vector, but have name set to the first

Examples

```
num1 <- list(sample(330:400,10))
num2 <-list("rpkg.net")
list_push(num1, add= num2)
```

list_shuffle	<i>Shuffle a list object just like shuffle in PHP</i>
--------------	---

Description

Shorthand to shuffle a list and save

Usage

```
list_shuffle(., seed = NULL)
```

Arguments

.	list to shuffle
seed	apply seed if indicated for reproducibility

Value

shuffled list of items store to the list name

Examples

```
list001 <- list("a" = 1:5,
              "b" = letters[1:5],
              c = LETTERS[1:10],
              "2" = number(5,5),
              "e" = randString(5,5))
list001 #show initial list

#illustrate basic functionality
list_shuffle(list001)
list001 #shuffle and resaved to variable

list.f2<-list001
list_shuffle(list.f2)
list.f2 #first output

list.f2<-list001
list_shuffle(list.f2)
list.f2 # different output from first output top

list.f2<-list001
list_shuffle(list.f2,seed = 344L)
list.f2 #second output

list.f2<-list001
list_shuffle(list.f2,seed = 344L)
list.f2 #the same output as second output top
```

make_dosing_df	<i>Create Subject-by-Time Dosing Records (Base R) with Optional Subject-Level Covariates</i>
----------------	--

Description

Creates a dosing/event-style dataset by expanding a set of subject IDs ('ID') across a vector of dosing times ('time'). It assigns a dose value to each subject and time combination, and optionally adds additional columns via '...' that can be constant (length 1) or subject-specific (length equal to 'length(ID)').

This utility is useful for quickly generating dosing records for simulation, exploratory analyses, and creating NONMEM-/nlmixr-/Monolix-friendly input structures where each row represents a dosing event (or event-like record) at a given time.

Usage

```
make_dosing_df(ID, time, dose, ...)
```

Arguments

ID	A vector of subject identifiers (numeric, integer, or character). Typically unique.
time	A vector of dosing times. Can be numeric or integer (e.g., hours, days).
dose	Dose value(s). May be: <ul style="list-style-type: none"> • length 1: one common dose applied to all subjects and all times • length equal to 'length(ID)': subject-specific doses mapped by the <i>*position*</i> of each subject in 'ID' (i.e., 'dose[i]' corresponds to 'ID[i]'), then repeated across all times for that subject
...	Named additional columns to include in the output. Each argument in '...' must be either: <ul style="list-style-type: none"> • length 1: a constant value repeated for all rows (e.g., 'STUDY = "CNT01275"') • length equal to 'length(ID)': subject-level values mapped by the <i>*position*</i> of each subject in 'ID' (e.g., 'WT = c(30, 45)' with 'ID = c(1, 2)' assigns 'WT=30' to subject 1 and 'WT=45' to subject 2 across all times) <p>All '...' arguments must be named. Names must not conflict with reserved output column names ('ID', 'time', 'dose').</p>

Details

Mapping rule (important) For any vector argument of length 'length(ID)' (including 'dose' or any '...' column), values are assigned by the **index/position** of 'ID', not by sorted ID or factor level. For example:

```
ID = c(10, 20)
VAR5 = c(4, 6)
```

results in subject '10 -> 4' and subject '20 -> 6' for all time points.

Duplicate IDs If you supply any subject-level vector (i.e., 'dose' length > 1, or any '...' argument length > 1), 'ID' must be unique. Duplicate IDs create ambiguous mappings and will trigger an error.

Output structure The output includes:

- 'ID': subject identifier
- 'time': dosing time
- 'dose': assigned dose
- any additional columns provided via '...'

Rows are sorted by 'ID' then 'time'.

When to use (common scenarios)

1. ****Create dosing records for simulation**** (e.g., one dose level across many subjects, or subject-specific doses).
2. ****Build NONMEM/nlmixr event datasets**** where dosing rows (e.g., 'EVID=1') are generated programmatically; you can add 'EVID', 'CMT', 'RATE', 'STUDY', etc. via '...'.
3. ****Explore exposure metrics**** by generating standardized schedules (e.g., QD times) and attaching covariates like weight, age group, cohort, regimen labels.
4. ****Scenario testing**** for pediatric dosing cutoffs, titration strategies (subject-specific), or bridging comparisons where regimen differs by subgroup.

When not to use Use a different approach if you need:

- time-varying covariates with length equal to 'length(time)' (shared across subjects),
- fully specified matrices of values length 'length(ID) * length(time)',
- multiple event types (dose + observations) in the same call (although you can merge the result with an observation dataset after creation).

Value

A 'data.frame' with 'length(ID) * length(time)' rows and columns 'ID', 'time', 'dose', plus any additional columns passed via '...'.

Examples

```
## Example 1: Simple dataset (10 subjects, dose 40 at times 1:10)
```

```
df1 <- make_dosing_df(ID = 1:10, time = 1:10, dose = 40)
```

```
head(df1)
```

```
## Example 2: Subject-specific dose (one dose per subject)
```

```
df2 <- make_dosing_df(ID = c(1, 2, 3), time = 1:5, dose = c(40, 60, 80))
```

```
head(df2)
```

```
## Example 3: Add constant columns via ...
```

```
df3 <- make_dosing_df(
```

```
  ID = 1:3, time = c(0, 7, 14), dose = 45,
```

```

  STUDY = "CNT01275JPA3001", ROUTE = "SC", CMT = 1
)
df3

## Example 4: Add subject-level covariates via ...
## VAR5 is mapped by position: ID=1 gets 4, ID=2 gets 6
df4 <- make_dosing_df(
  ID = c(1, 2), time = 1:3, dose = 40,
  VAR5 = c(4, 6), WT = c(35.2, 51.0), SEX = c("M", "F")
)
df4

## Example 5: NONMEM-style dosing rows (add EVID and AMT)
df_nm <- make_dosing_df(ID = 1:2, time = c(0, 28, 56), dose = c(45, 45),
  EVID = 1, AMT = NA) # AMT placeholder (optional)
df_nm$AMT <- df_nm$dose
df_nm

```

math.qt

Miscellaneous math computations: Corresponding m-m and quantile for confident intervals

Description

Compute the corresponding quantile given confident interval bounds

Usage

```
math.qt(ci = 0.9)
```

```
math.mm(Vmax, S, Km, V, round = NULL)
```

Arguments

ci	confident interval eg. 0.9 for 90 percent confident intervals
Vmax	The maximum velocity of the enzymatic reaction.
S	The substrate concentration.
Km	The substrate concentration at which the reaction rate is half of Vmax.
V	The current velocity of the enzymatic reaction
round	round result to number of decimal places

Value

vector of two numeric values for the quantile based on the confident interval chosen
 result of calculation of Michaelis-Menten equation

Examples

```
# Get the bounds for 90% confident intervals
math.qt(0.9)

# Get the bounds for 95% confident intervals
# use the bounds to obtain quartile
values = number(100)
values
ci = math.qt(0.95)
getquart = quantile(values, probs = ci)
getquart

math.mm(3,500,0.5)
```

minus	<i>Decrease vector by value</i>
-------	---------------------------------

Description

decrease the content of a vector and re-save as the vector

Usage

```
minus(., minus = 1L)
```

Arguments

.	vector of number(s)
minus	number to minus

Details

Similar to the inc and plus functions, the minus function is very useful when writing complex codes involving loops. Apart from the for loop, minus can be useful to quickly decrease the value of a variable located outside the loop by simply decrement the variable by 1 or other numbers. Check in the example section for a specific use. Given the scope, one may also choose to use this function in any other instances, as it's simple purpose is to decrease the value of a variable by a number and then re-save the new value to that variable.

Value

a vector decreased by a number

Examples

```
num1 <- sample(5:150,10)
num1

# decrease num1 by 1
num1 #before decrease
minus(num1)
num1 #after decrease

# decrease num1 by 5
num1 #before decrease
minus(num1, minus = 5)
num1 #after decrease

#when used in loops

#add and compare directly
rnum = 23
minus(rnum) == 220 #returns FALSE
rnum #the variable was also updated

# use in a for loop
ynum = 100

for( i in c("teacher","student","lawyer","pharmacist")){
message("This is the item number ")
message(ynum)
message(". For this item, I am a ")
message(i)

#decrement easily at each turn
minus(ynum,3)
}

#use in a repeat loop
xnum = 100
repeat{ #repeat until xnum is 85
message(xnum)
if(minus(xnum) == 85) break
}
```

mix.color

Mix or Blend two or more colors

Description

Combine colors to generate a new color

Usage

```
mix.color(color, type = 2, alpha = 1)
```

Arguments

color	CHARACTER. color vector e.g see example
type	NUMERIC. return type of the output
alpha	NUMERIC. alpha or opacity of the resulting color

Value

hex for the combined color

Examples

```
# color vector
colvec <- c("red", "blue", "violet", "green", "#ff0066")

# just one color
mix.color(colvec[1], type = 1, alpha = 1)

# add two colors
mix.color(colvec[1:2], type = 1, alpha = 1)

# add three colors
mix.color(colvec[1:3], type = 1, alpha = 1)

# return type = 2

# just one color
mix.color(colvec[1], type = 2, alpha = 1)

# add two colors
mix.color(colvec[1:2], type = 2, alpha = 1)

# add three colors
mix.color(colvec[1:3], type = 2, alpha = 1)

# opacity or alpha 0.5

# just one color
mix.color(colvec[1], type = 1, alpha = 0.5)

# add two colors
mix.color(colvec[1:2], type = 1, alpha = 0.5)

# add three colors
mix.color(colvec[1:3], type = 1, alpha = 0.5)
```

```
# add all colors
mix.color(colvec, type = 1, alpha = 0.5)
```

mix.cols.btw

Mix or Blend colors between two or more colors

Description

Mix or blend multiple colors between two colors

Usage

```
mix.cols.btw(colors, max = 20, alpha = 1, preview = FALSE)
```

Arguments

colors	the vector of two colors
max	maximum number of colors to blend between
alpha	alpha for the new color blends
preview	LOGICAL. preview all color generated

Value

color hex for all generated colors

Examples

```
# simply mix/blend two colors
mix.cols.btw(c("red", "brown"))

# simply mix/blend two colors, maximum number of colors at the end
mix.cols.btw(c("red", "brown"), max = 8)

# simply mix/blend two colors with alpha=0.2 (opacity=0.2)
mix.cols.btw(c("yellow", "green"), alpha = 0.2)

# also preview after mixing the two colors
mix.cols.btw(c("red", "green"), preview = TRUE)
mix.cols.btw(c("blue", "violet"), alpha = 0.2, preview = TRUE)

mix.cols.btw(c("red", "purple", "yellow", "gray"), preview = TRUE)

mix.cols.btw(c("red", "purple", "yellow", "gray"), alpha = 0.2, preview = TRUE)
```

`mode.calc`*Calculate the Mode of a Numeric or Character Vector*

Description

This function calculates the mode (most frequently occurring value(s)) of a numeric or character vector.

Usage

```
mode.calc(x)
```

Arguments

`x` A numeric or character vector for which the mode is to be calculated.

Value

The mode(s) of the input vector. If multiple values have the same highest frequency, all modes are returned. Returns 'NA' if the input vector is empty.

Examples

```
# Example with a numeric vector
numeric_vector <- c(1, 2, 2, 3, 3, 3, 4, 5)
mode.calc(numeric_vector)

# Example with a character vector
character_vector <- c("apple", "banana", "apple", "orange", "banana", "banana")
mode.calc(character_vector)
```

`multihead_att`*Multi-Head Attention*

Description

This function implements a simplified version of multi-head attention for sequence data.

Usage

```
multihead_att(query, key, value, num_heads)
```

Arguments

query	A numeric matrix representing the query.
key	A numeric matrix representing the key.
value	A numeric matrix representing the value.
num_heads	An integer specifying the number of attention heads.

Value

A numeric matrix representing the attention output.

Examples

```
query <- matrix(rnorm(12), nrow = 3)
key <- matrix(rnorm(12), nrow = 3)
value <- matrix(rnorm(12), nrow = 3)
multihead_att(query, key, value, num_heads = 2)
```

mutate_filter

Mutate only a subset of dataset intact

Description

Extension of the `dplyr::mutate` function that allows the user to mutate only a specific filtered subset of a data, while leaving the other parts of the data intact

Usage

```
mutate_filter(., sub.set, ...)
```

Arguments

.	data object
sub.set	subset of data to modify
...	mutation syntax similar to <code>dplyr::mutate</code>

Value

data frame containing original data, but with a subset mutated

Examples

```
#mutate a subsection filter of mtcars
dt = mtcars
names(dt)
head(dt)
mutate_filter(dt,mpg == 21.0 & cyl == 6, cyl=1000, hp=2000, vs=hp*2)

dt2 = beaver1
names(dt2)
head(dt2)
mutate_filter(dt2, day == 346 & time < 1200, activ = 12, temp = round(temp*10,1))
```

na.cumsum	<i>Cumulative Sum with NA Removal</i>
-----------	---------------------------------------

Description

This function calculates the cumulative sum of a numeric vector, removing any 'NA' values before computation.

Usage

```
na.cumsum(x)
```

Arguments

x A numeric vector for which the cumulative sum is to be calculated.

Value

A numeric vector representing the cumulative sum with 'NA' values removed. Returns an empty vector if the input is empty or contains only 'NA' values.

Examples

```
# Example with numeric vector
numeric_vector <- c(1, 2, NA, 4, 5)
na.cumsum(numeric_vector)

# Example with all NAs
na.cumsum(c(NA, NA))
```

ndecimal	<i>Count the number of decimal places</i>
----------	---

Description

Count the number of decimal places

Usage

```
ndecimal(num)
```

Arguments

num	a numeric value
-----	-----------------

Examples

```
#example 1
ndecimal(9.000322)
```

```
#example 2
ndecimal(34)
```

```
#example 3
ndecimal(45.)
```

newSuperVar	<i>Create and use a super variable with unique capabilities</i>
-------------	---

Description

Create a variable that supersedes other variables and has various functionalities

Usage

```
newSuperVar(variable, value = 0L, lock = FALSE, editn = NULL)
```

Arguments

variable	variable name for super variable
value	value of the variable
lock	lock variable to change
editn	number of times the super variable may be set to a new value using .set(). - Set to NULL to allow unlimited value change - Set to 0 to prevent editing the super variable

Value

no visible return, but variable is created and stored with various functionalities

Note**What you should know about the functionality:**

This function ensures that a variable is created and may not easily be altered. It helps preserve the original variable by providing only limited access to the variable.

Creation of this super variable automatically attached some key functions to it, such that the user is able to call the function like `.set()`, `.rm()`.

Super variable value may be set from any scope using the `.set()` function, which means that it is granted global variable features without being present within the global environment of the current section.

The variable name of the super variable may be overwritten in the local environment, but this would not alter the super variable. It means that once the local variable is removed, the super variable remains available for use.

Use cases:

- Preserve originality of variable within an R session. Avoid inadvertent deletion.
- Widely accessible from any scope e.g functions, lapply, loops, local environment etc
- Restricted mutability of variable using set function e.g `varname.set()`
- Variable with easy function calls by attaching `'.'`
- Variable with un-mutable class when changing its value
- Variable with restricted number of times it can be changed

Examples

```
# Task: create a super variable to
# store dataset that should not be altered
newSuperVar(mtdf, value = austres) # create a super variable
head(mtdf) # view it
mtdf.class # view the store class of the variable, it cannot be changed
# it means that when the super variable is edited, the new value MUST have the same class "ts"

# create and lock super variable by default
# extra security to prevent changing
```

```

newSuperVar(mtdf3, value = beaver1, lock = TRUE)
head(mtdf3) # view
mtdf3.round(1) # round to 1 decimal places
head(mtdf3) # view
mtdf3.signif(2) # round to 2 significant digits
head(mtdf3) # view

# Task: create a new super variable to store numbers
# edit the numbers from various scopes
newSuperVar(edtvec, value = number(5))
edtvec # view content of the vector

# edtvec.set(letters) #ERROR: Cannot set to value with different class than initial value

edtvec.set(number(20)) # set to new numbers
edtvec # view output

for (pu in 1:8) {
  print(edtvec) # view output within loop
  edtvec.set(number(pu)) # set to new numbers within for loop
}

lc <- lapply(1:8, function(pu) {
  print(edtvec) # view output within loop
  edtvec.set(number(pu)) # set to new numbers within lapply loop
})

# see that the above changed the super variable easily.
# local variable will not be altered by the loop
# example
bim <- 198
lc <- lapply(1:8, function(j) {
  print(bim)
  bim <- j # will not alter the value of bim in next round
})

# Task: create and search data.frame
# create a new super variable with value as mtcars
# search if it contains the numeric value 21
newSuperVar(lon2, value = mtcars) # declares lon2
lon2 # view content of lon2
lon2.contains("21.0") # WRONG - since df.col is not specified,
# only the first column is search for the character "21.0"
lon2.contains("21.0", df.col = "mpg") # WRONG - searches mpg column
# for the character "21.0"
lon2.contains(21.0, df.col = "mpg") # CORRECT - search mpg column for the
# numeric value 21.0

# remove lon2 as a super variable
exists("lon2") # before removal
lon2.rm()
exists("lon2") # after removal

```

```
# Task: create and search vector
# create a new super variable with value as 10 random numbers
# search if it contains the numeric value 72
newSuperVar(lon3, value = number(10, seed = 12)) # declares lon3
lon3 # view content of lon3
lon3.contains(72) # should give TRUE or false if the vector contains the value 45
lon3.contains(72, fixed = TRUE) # should give TRUE or false if the vector contains the value 45

# remove lon3 as a super variable
lon3.rm()

#Task: create a super variable that can only be edited 3 times
newSuperVar(man1, value = number(5), editn = 3)
man1 # view value

man1.set(number(10)) # change value first time
man1 # view value

man1.set(number(2)) # change value second time
man1 # view value

man1.set(number(1)) # change value third time
man1 # view value

man1.set(number(5)) # change value forth time,
# should not change because max change times exceeded
man1 # view value
```

normalize.vector	<i>Normalize a Numeric Vector to the Range [0, 1]</i>
------------------	---

Description

This function normalizes a numeric vector so that all values are scaled to the range [0, 1].

Usage

```
normalize.vector(x)
```

Arguments

x A numeric vector to be normalized.

Value

A numeric vector with values scaled to the range [0, 1]. If all values are identical, all values are set to 0.

Examples

```
# Example with a numeric vector
normalize.vector(c(1, 2, 3, 4, 5))

# Example with a vector containing identical values
normalize.vector(c(3, 3, 3))
```

not.data

Not a data

Description

Opposite of `is.data.frame()`. Check if entry is not a data object

Usage

```
not.data(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is a data table

Examples

```
test.dt <- data.frame(ID=1:200,Type="RPKG.net")
test.notenv <- list(t=1)

is.data.frame(test.dt) # TRUE
not.data(test.dt) # FALSE

not.data(test.notenv) # TRUE
if(not.data(test.dt)) message("yes") # NULL
```

not.duplicated	<i>Not duplicated elements</i>
----------------	--------------------------------

Description

Oposite of duplicated(). Checks which elements of a vector or data frame are NOT duplicates of elements with smaller subscripts

Usage

```
not.duplicated(x, incomparables = FALSE, ...)
```

Arguments

x	a vector or a data frame or an array or NULL.
incomparables	a vector of values that cannot be compared. FALSE is a special value, meaning that all values can be compared, and may be the only value accepted for methods other than the default. It will be coerced internally to the same type as x
...	arguments for particular methods.

Value

elements of a vector or data frame that are NOT duplicates

Examples

```
set.seed(08082023)
dtf <- sample(1:10,15, replace = TRUE)
dtf # 3 9 10 3 8 9 6 10 5 1 2 2 2 9 8
dtf[ dtf > 4 & duplicated(dtf) ] # 9 10 9 8
dtf[ dtf > 4 & not.duplicated(dtf) ] # 9 10 8 6 5
```

not.empty	<i>Not empty</i>
-----------	------------------

Description

Check if entry is not empty

Usage

```
not.empty(x)
```

```
is.empty(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is empty

Examples

```
not.empty("empty") # TRUE
not.empty('') # FALSE
not.empty(y<-NULL) # FALSE
if(not.empty('')) message("yes") # NULL
```

not.environment	<i>Not an environment</i>
-----------------	---------------------------

Description

Check if entry is not an environment object

Usage

```
not.environment(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is an environment

Examples

```
test.env <- new.env()
test.notenv <- list(t=1)
not.environment(test.env) # FALSE
not.environment(test.notenv) # TRUE
if(not.environment(test.notenv)) message("yes") # yes
```

not.exists	<i>Not exists</i>
------------	-------------------

Description

Check if object does not exists

Usage

```
not.exists(x)
```

Arguments

x	object
---	--------

Value

a boolean value to indicate if entry does not exists

Examples

```
go = 7
not.exists("exis") # TRUE
not.exists("go") # FALSE
if(not.exists('hallo')) message("yes") # NULL
```

not.image	<i>File name extension(s) is Not an image</i>
-----------	---

Description

Check if one or multiple file name entry is not an image

Usage

```
not.image(x)
```

Arguments

x	vector entry
---	--------------

Details

This current function tests if the extension of the file name provided does NOT belongs to any of the image extensions listed below

- AI - Adobe Illustrator
- BMP - Bitmap Image
- CDR - Corel Draw Picture
- CGM - Computer Graphics Metafile
- CR2 - Canon Raw Version 2
- CRW - Canon Raw
- CUR - Cursor Image
- DNG - Digital Negative
- EPS - Encapsulated PostScript
- FPX - FlashPix
- GIF - Graphics Interchange Format
- HEIC - High-Efficiency Image File Format
- HEIF - High-Efficiency Image File Format
- ICO - Icon Image
- IMG - GEM Raster Graphics
- JFIF - JPEG File Interchange Format
- JPEG - Joint Photographic Experts Group
- JPG - Joint Photographic Experts Group
- MAC - MacPaint Image
- NEF - Nikon Electronic Format
- ORF - Olympus Raw Format
- PCD - Photo CD
- PCX - Paintbrush Bitmap Image
- PNG - Portable Network Graphics
- PSD - Adobe Photoshop Document
- SR2 - Sony Raw Version 2
- SVG - Scalable Vector Graphics
- TIF - Tagged Image File
- TIFF - Tagged Image File Format
- WebP - Web Picture Format
- WMF - Windows Metafile
- WPG - WordPerfect Graphics

Value

a boolean value to indicate if entry is not an image

Examples

```
img.1 <- "fjk.jpg"
not.image(img.1)

img.2 <- "fjk.bmp"
not.image(img.2)

img.3 <- "fjk.SVG"
```

```
not.image(img.3)

# a vector of file names
v <- c("logo.png", "business process.pdf",
"front_cover.jpg", "intro.docx",
"financial_future.doc", "2022 buybacks.xlsx")
not.image(v)

# when the file name has no extension
# the function returns NA
v2 <- c("img2.jpg", NA, "northbound.xlsx", "landing")
not.image(v2)
```

not.inherits	<i>Not inherit from any of the classes specified</i>
--------------	--

Description

Opposite of `base::inherits()`. Indicates whether its first argument inherits from any of the classes specified in the `what` argument. If `which` is `TRUE` then an integer vector of the same length as `what` is returned. Each element indicates the position in the `class(x)` matched by the element of `what`; zero indicates no match. If `which` is `FALSE` then `TRUE` is returned by `inherits` if any of the names in `what` match with any class.

Usage

```
not.inherits(x, what, which = FALSE)
```

Arguments

<code>x</code>	an R object.
<code>what</code>	a character vector naming classes.
<code>which</code>	logical affecting return value: see ‘Details’.

Value

a boolean value to indicate if `!inherits`

Examples

```
keep.cols = "a character"
class(keep.cols) # class is character
not.inherits(keep.cols, "character")

num.var = 1L
class(num.var) # class is integer
not.inherits(num.var, "double")
```

not.integer *Not an integer*

Description

Opposite of is.integer(). Check if entry is not an integer

Usage

```
not.integer(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is an integer

Examples

```
is.integer(78L) #TRUE
not.integer(78L) #FALSE

not.integer(23.43) # TRUE
not.integer(45L) # FALSE
if(not.integer(4L)) message("yes") # NULL
```

not.logical *Not logical*

Description

Opposite of is.logical(). Check if entry is a logical object

Usage

```
not.logical(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is logical

Examples

```
test.env <- TRUE
test.notenv <- 0
not.logical(test.env) # FALSE
not.logical(test.notenv) # TRUE
if(not.logical(test.notenv)) message("yes") # yes
```

not.na	<i>Not NA</i>
--------	---------------

Description

Opposite of is.na(). Check if entry is not NA

Usage

```
not.na(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is NA

Examples

```
not.na(NA) # FALSE
not.na(NULL) # logical(0)
if(not.na(45)) message("something") # TRUE
```

not.null	<i>Not NULL</i>
----------	-----------------

Description

Opposite of is.null(). Check if entry is not NULL

Usage

```
not.null(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is NULL

Examples

```
is.null("") # FALSE
not.null("") # TRUE
not.null(NULL) # FALSE
if(not.null(45)) message("something") # yes
```

not.numeric

Not numeric

Description

Check if entry is not numeric

Usage

```
not.numeric(x)
```

Arguments

x vector entry

Value

a boolean value to indicate if entry is numeric

Examples

```
not.numeric("45") # TRUE
not.numeric(45) # FALSE
if(not.numeric(45)) message("yes") # yes
```

not.vector	<i>Not a vector</i>
------------	---------------------

Description

Opposite of is.vector(). Check if entry is not vector

Usage

```
not.vector(x)
```

Arguments

x	vector entry
---	--------------

Value

a boolean value to indicate if entry is vector

Examples

```
vect1 = list(r=1,t=3:10)
vect2 = LETTERS
is.vector(vect1) # TRUE
not.vector(vect1) # FALSE
not.vector(vect2) # FALSE
if(not.vector(vect1)) message("yes") # NULL
```

number	<i>Generate a random number (integer)</i>
--------	---

Description

Fetch n random integers between 1 and 1,000,000,000

Usage

```
number(n, max.digits = 10, seed = NULL)
```

Arguments

n	how many numbers to generate
max.digits	maximum number of digits in each number
seed	set seed for sampling to maintain reproducibility

Value

random numbers between 1 and 1 billion

Examples

```

number(1)
number(10)
paste0(number(2),LETTERS)

#set maximum number of digits
number(1,max.digits = 5)
number(10,max.digits = 4)

#set seed for reproducibility
#without seed
number(6) #result 1
number(6) #result 2, different from result 1
#with seed
number(6,seed=1)#result 3
number(6,seed=1)#result 4, same as result 3

```

or *Nullish coalescing operator*

Description

Alternative return for empty, null or na statements. Return alternative if the value of expression is empty or NA or NULL

Usage

```

or(test, alternative)

test %or% alternative

```

Arguments

test	an object to return
alternative	alternative object to return

Value

value of test if not null or empty, else return value of alternative

Note

Equivalent to Nullish coalescing operator ?? in javascript or PHP like \$Var = \$operand1 ?? \$operand2;

Examples

```

test1 <- c(4,NA,5,2,0,21)

test2 <- data.frame(ID = 1:10,ED = LETTERS[10:1])

# One may also choose to use

test2[,1] %or% "A"

test2[which(test2$ID == 323),2] %or% "CCCCC"

number(1) %or% "Placeholder"

number(10) %or% "Placeholder"

NA %or% "Random"

NULL %or% "Random"

"" %or% "Random"

or(test1[which(test1==4)],100)

or(test1[which(test1==43)],100)

or(test2[which(test2$ID == 10),2],"BBBBB")

or(test2[which(test2$ID == 323),2],"CCCCC")

```

pairDist

Calculate the distance of points from the center of a cluster

Description

This function operates on multivariate data and calculates the distance of points from the centroid of one or more clusters.

Usage

```
pairDist(data, round)
```

Arguments

data	data frame object or a matrix/array object
round	round result to decimal place

Value

a named vector consisting of a row number and a pair-distance value

Function utility

Used to generate the computations needed to model pair-distance measures in three dimensions

More information about this function

The `pairDist` function is used to quantify how far each data point (row) is from the overall mean across all columns. It's commonly used in multivariate statistics, machine learning, and data analysis to assess the variability or similarity of data points relative to their mean. More specifically, the function is used in outlier detection and cluster analysis to evaluate the dispersion of data. Used in conjunction with other calculations, `pairDist` output can also be used to model data in three dimensions.

References

the current function was adapted from one of the examples in the `svgViewR` package, <https://cran.r-project.org/web/packages/svgViewR/svgViewR.pdf>

Examples

```
data = attenu[,1:2]

# example 1: basic example using data.frame
pairDist(data)

# example 2: basic example using as.matrix
pairDist(as.matrix(data))

# example 3: round results to 2 decimal points
pairDist(data, 2)

# example 4
data = matrix(
  c(1, 5, NA, 4, 5, 6, NA, 8, 9, NA, 12, 23, 6, 0, 3, 3, 8, 15),
  ncol = 3,
  byrow = TRUE)
x = pairDist(data, 3)
data = data.frame(data)
data = cbind(data, x)
```

percent_match	<i>Function to calculate the percentage of matching between two strings</i>
---------------	---

Description

Function to calculate the percentage of matching between two strings

Usage

```
percent_match(  
  string1,  
  string2,  
  case_sensitive = FALSE,  
  ignore_whitespace = TRUE,  
  frag_size = 2  
)  
  
string1 %match% string2  
  
sound_match(string1, string2)
```

Arguments

string1	first string
string2	second string
case_sensitive	if to check case sensitivity
ignore_whitespace	if to ignore whitespace
frag_size	fragment size of string

Details

Case Sensitivity:

The function can optionally consider or ignore case sensitivity based on the `case_sensitive` argument.

Whitespace Handling:

With `ignore_whitespace` set to `TRUE`, the function removes all whitespaces before comparison. This can be useful for matching strings that may have inconsistent spacing.

Exact Character-by-Character Matching:

The function computes the percentage of matching characters in the same positions.

Substring Matching:

The function checks if one string is a substring of the other, awarding a full match if true.

Levenshtein Distance:

The function uses Levenshtein distance to calculate the similarity and integrates this into the overall match percentage.

Fragment Matching:

- A `frag_size` argument is introduced that compares fragments (substrings) of a given size (default is 3) from both strings.
- The function creates unique fragments from each string and compares them to find common fragments.
- The percentage match is calculated based on the ratio of common fragments to the total number of unique fragments.

Combining Metrics:

The overall match percentage is computed as the average of exact match, substring match, Levenshtein match, and fragment match percentages.

Value

numeric value of the match percent

match word sounds

Examples

```
# Example 1: simple match
string1 <- "Hello World"
string2 <- "helo world"

match_percent <- percent_match(string1, string2)
message("Percentage of matching: ", match_percent)
```

```
# Example 2: which date is closest
string0 <- "october 12,1898"
string1 <- "2018-10-12"
string2 <- "1898-10-12"
percent_match(string0, string1)
percent_match(string0, string2)
percent_match(string0, string2, frag_size = 4)
percent_match(string1, string2)
```

```
sound_match("Robert","rupert")
sound_match("rupert","Rubin")
sound_match("book","oops")
```

plus

Increment vector by value

Description

Increment the content of a vector and re-save as the vector

Usage

```
plus(., add = 1L)
```

Arguments

.	vector of number(s)
add	number to add

Details

This function is very useful when writing complex codes involving loops. Apart from the for loop, this can be useful to quickly increment a variable located outside the loop by simply incrementing the variable by 1 or other numbers. Check in the example section for a specific use. Nonetheless, one may also choose to use this function in any other instance, as it's simple purpose is to increase the value of a variable by a number and then re-save the new value to that variable.

Value

a vector incremented by a number

Examples

```
num1 <- sample(330:400,10)
num1#before increment

# increment num1 by 1
inc(num1)
num1 #after increment

# increment num1 by 5
num1 #before increment
inc(num1, add= 10)
num1 #after increment

#when used in loops

#add and compare directly
```

```
rnum = 10
inc(rnum) == 11 #returns TRUE
rnum #the variable was also updated

# use in a for loop
ynum = 1
for( i in c("scientist","dancer","handyman","pharmacist")){
message("This is the item number ")
message(ynum)
message(". For this item, I am a ")
message(i)

#decrement easily at each turn
plus(ynum)
}

#use in a repeat loop
xnum = 1
repeat{ #repeat until xnum is 15
message(xnum)
if(inc(xnum) == 15) break
}
```

randString

Generate a random string

Description

Create a random string of specified length

Usage

```
randString(n, length)
```

Arguments

n	number of strings to create
length	length of string to create

Value

one more random string of specific length

Examples

```
# Task 1: create 1 random string string of length 5
randString(n = 1, length = 5)

# Task 2: create 5 random string string of length 10
randString(n = 5, length = 10)

# Task 3: create 4 random string string of length 16
randString(n = 4, length = 16)
```

rcolorconst	<i>R Color Constant</i>
-------------	-------------------------

Description

This function provides information that describes the color constants that exist in R

Usage

```
rcolorconst(title = "R Color Constants")
```

Arguments

title	title of the output
-------	---------------------

Details

In addition to the color palette in R that can be represented as either color literals or hexadecimal values, numeric values can also be used to add colorization to a plot. Numeric values ranging from 1 to 8 provide 8 basic colors that can be deployed. The rcolorconst function returns both a Named Vector and a color palette plot that connects these numeric values with their corresponding color.

Value

returns color constant

Examples

```
# Without title
ex1 <- rcolorconst()

# With title
ex2 <- rcolorconst("My new color constant")

# More detailed example
set.seed(200)
x = data.frame(
```

```
meas = rnorm(100),
grp = sample(1:8, size = 100,
replace = TRUE))
plot(x, pch = 16, col = x$grp)
colnums = rcolorconst()
```

rDecomPkg

Check whether an R package has been decommissioned in CRAN

Description

Designed to assist users in checking the decommission status of an R package on CRAN. In the context of R language, CRAN stands for the Comprehensive R Archive Network.

Usage

```
rDecomPkg(package)
```

Arguments

package package name to query

Details

CRAN is a network of servers around the world that store R packages and their documentation, providing a centralized repository for the R community. With the current function, users can quickly and easily determine whether a specific R package has been decommissioned on CRAN, ensuring they stay informed about the availability and support status of the packages they rely on for their R programming projects. This tool simplifies the process of package management, helping users maintain up-to-date and reliable dependencies in their R code.

Value

the decommissioned status of a particular package based on the available packages using the `utils` package

Examples

```
## Not run:
# check if cattonum package is decommissioned
# the current package is expected to be decommissioned
rDecomPkg("cattonum")

# check if dplyr is decommissioned
# the current package is expected NOT to be decommissioned
rDecomPkg("dplyr")

# when a package never existed in CRAN
```

```
# the result of the function call should be NA
rDecomPkg("printy")
rDecomPkg("package0002312122312")

## End(Not run)
```

read.csv.print

Read a CSV and preview first X rows and columns

Description

The purpose of this function is combine the functionality of **read.csv** and **print**, which are often used together.

The purpose of this function is to read data from a file into a variable and simultaneously display a preview of the data, showing either the first few rows or columns based on the user's specification. It is important to emphasize that the function expects the user to assign the result of the read operation to a variable in order to achieve its intended purpose. eg. Use **var1 = read.csv.print(file1)** instead of **read.csv.print(file1)**

Usage

```
read.csv.print(
  file,
  header = TRUE,
  sep = ",",
  quote = "\"",
  dec = ".",
  fill = TRUE,
  comment.char = "",
  ...,
  dim = c(10L, 5L)
)
```

Arguments

file the name of the file which the data are to be read from. Each row of the table appears as one line of the file. If it does not contain an *absolute* path, the file name is *relative* to the current working directory, `getwd()`. Tilde-expansion is performed where supported. This can be a compressed file (see [file](#)).

Alternatively, `file` can be a readable text-mode [connection](#) (which will be opened for reading if necessary, and if so `closed` (and hence destroyed) at the end of the function call). (If `stdin()` is used, the prompts for lines may be somewhat confusing. Terminate input with a blank line or an EOF signal, Ctrl-D on Unix and Ctrl-Z on Windows. Any pushback on `stdin()` will be cleared before return.) `file` can also be a complete URL. (For the supported URL schemes, see the 'URLs' section of the help for [url](#).)

header	a logical value indicating whether the file contains the names of the variables as its first line. If missing, the value is determined from the file format: header is set to TRUE if and only if the first row contains one fewer field than the number of columns.
sep	the field separator character. Values on each line of the file are separated by this character. If sep = "" (the default for read.table) the separator is 'white space', that is one or more spaces, tabs, newlines or carriage returns.
quote	the set of quoting characters. To disable quoting altogether, use quote = "". See scan for the behaviour on quotes embedded in quotes. Quoting is only considered for columns read as character, which is all of them unless colClasses is specified.
dec	the character used in the file for decimal points.
fill	logical. If TRUE then in case the rows have unequal length, blank fields are implicitly added. See 'Details'.
comment.char	character: a character vector of length one containing a single character or an empty string. Use "" to turn off the interpretation of comments altogether.
...	Further arguments to be passed to read.table.
dim	dimension of CSV content to show

Details

Read a dataset of type csv and show x rows and y columns with one function call

Value

read csv content and a print out of the data head

Examples

```
## Not run:
# Example: read a csv file and print the first 10 lines
# declare file
new.file <- "test.csv"

# read file and preview default
dth3 <- read.csv.print(file = new.file)

# read file and preview 10 rows and all columns
dth1 <- read.csv.print(file = new.file, dim = 10)

# read file and preview 10 rows and 5 columns
dth2 <- read.csv.print(file = new.file, dim = c(10,5))

## End(Not run)
```

read.table.print	<i>Read in a table and show first X rows and columns</i>
------------------	--

Description

The purpose of this function is combine the functionality of **read.table** and **print**, which are often used together.

The purpose of this function is to read table from a file into a variable and simultaneously display a preview of the data, showing either the first few rows or columns based on the user's specification. It is important to emphasize that the function expects the user to assign the result of the read operation to a variable in order to achieve its intended purpose. eg. Use **var1 = read.table.print(file1)** instead of **read.table.print(file1)**

Usage

```
read.table.print(  
  file,  
  header = FALSE,  
  sep = "",  
  quote = "\"",  
  dec = ".",  
  numerals = c("allow.loss", "warn.loss", "no.loss"),  
  row.names,  
  col.names,  
  as.is = TRUE,  
  na.strings = "NA",  
  colClasses = NA,  
  nrows = -1,  
  skip = 0,  
  check.names = TRUE,  
  fill = NULL,  
  strip.white = FALSE,  
  blank.lines.skip = TRUE,  
  comment.char = "#",  
  allowEscapes = FALSE,  
  flush = FALSE,  
  stringsAsFactors = FALSE,  
  fileEncoding = "",  
  encoding = "unknown",  
  skipNul = FALSE,  
  dim = c(10L, 5L),  
  ...  
)
```

Arguments

file	<p>the name of the file which the data are to be read from. Each row of the table appears as one line of the file. If it does not contain an <i>absolute</i> path, the file name is <i>relative</i> to the current working directory, <code>getwd()</code>. Tilde-expansion is performed where supported. This can be a compressed file (see <code>file</code>).</p> <p>Alternatively, file can be a readable text-mode <code>connection</code> (which will be opened for reading if necessary, and if so <code>closed</code> (and hence destroyed) at the end of the function call). (If <code>stdin()</code> is used, the prompts for lines may be somewhat confusing. Terminate input with a blank line or an EOF signal, <code>Ctrl-D</code> on Unix and <code>Ctrl-Z</code> on Windows. Any pushback on <code>stdin()</code> will be cleared before return.)</p> <p>file can also be a complete URL. (For the supported URL schemes, see the ‘URLs’ section of the help for <code>url</code>.)</p>
header	<p>a logical value indicating whether the file contains the names of the variables as its first line. If missing, the value is determined from the file format: header is set to TRUE if and only if the first row contains one fewer field than the number of columns.</p>
sep	<p>the field separator character. Values on each line of the file are separated by this character. If <code>sep = ""</code> (the default for <code>read.table</code>) the separator is ‘white space’, that is one or more spaces, tabs, newlines or carriage returns.</p>
quote	<p>the set of quoting characters. To disable quoting altogether, use <code>quote = ""</code>. See <code>scan</code> for the behaviour on quotes embedded in quotes. Quoting is only considered for columns read as character, which is all of them unless <code>colClasses</code> is specified.</p>
dec	<p>the character used in the file for decimal points.</p>
numerals	<p>string indicating how to convert numbers whose conversion to double precision would lose accuracy, see <code>type.convert</code>. Can be abbreviated. (Applies also to complex-number inputs.)</p>
row.names	<p>a vector of row names. This can be a vector giving the actual row names, or a single number giving the column of the table which contains the row names, or character string giving the name of the table column containing the row names. If there is a header and the first row contains one fewer field than the number of columns, the first column in the input is used for the row names. Otherwise if <code>row.names</code> is missing, the rows are numbered.</p> <p>Using <code>row.names = NULL</code> forces row numbering. Missing or <code>NULL</code> <code>row.names</code> generate row names that are considered to be ‘automatic’ (and not preserved by <code>as.matrix</code>).</p>
col.names	<p>a vector of optional names for the variables. The default is to use "V" followed by the column number.</p>
as.is	<p>controls conversion of character variables (insofar as they are not converted to logical, numeric or complex) to factors, if not otherwise specified by <code>colClasses</code>. Its value is either a vector of logicals (values are recycled if necessary), or a vector of numeric or character indices which specify which columns should not be converted to factors.</p> <p>Note: to suppress all conversions including those of numeric columns, set <code>colClasses = "character"</code>.</p>

	Note that <code>as.is</code> is specified per column (not per variable) and so includes the column of row names (if any) and any columns to be skipped.
<code>na.strings</code>	a character vector of strings which are to be interpreted as <code>NA</code> values. Blank fields are also considered to be missing values in logical, integer, numeric and complex fields. Note that the test happens <i>after</i> white space is stripped from the input (if enabled), so <code>na.strings</code> values may need their own white space stripped in advance.
<code>colClasses</code>	character. A vector of classes to be assumed for the columns. If unnamed, recycled as necessary. If named, names are matched with unspecified values being taken to be <code>NA</code> . Possible values are <code>NA</code> (the default, when <code>type.convert</code> is used), <code>"NULL"</code> (when the column is skipped), one of the atomic vector classes (logical, integer, numeric, complex, character, raw), or <code>"factor"</code> , <code>"Date"</code> or <code>"POSIXct"</code> . Otherwise there needs to be an <code>as</code> method (from package methods) for conversion from <code>"character"</code> to the specified formal class. Note that <code>colClasses</code> is specified per column (not per variable) and so includes the column of row names (if any).
<code>nrows</code>	integer: the maximum number of rows to read in. Negative and other invalid values are ignored.
<code>skip</code>	integer: the number of lines of the data file to skip before beginning to read data.
<code>check.names</code>	logical. If <code>TRUE</code> then the names of the variables in the data frame are checked to ensure that they are syntactically valid variable names. If necessary they are adjusted (by <code>make.names</code>) so that they are, and also to ensure that there are no duplicates.
<code>fill</code>	logical. If <code>TRUE</code> then in case the rows have unequal length, blank fields are implicitly added. See ‘Details’.
<code>strip.white</code>	logical. Used only when <code>sep</code> has been specified, and allows the stripping of leading and trailing white space from unquoted character fields (numeric fields are always stripped). See <code>scan</code> for further details (including the exact meaning of ‘white space’), remembering that the columns may include the row names.
<code>blank.lines.skip</code>	logical: if <code>TRUE</code> blank lines in the input are ignored.
<code>comment.char</code>	character: a character vector of length one containing a single character or an empty string. Use <code>""</code> to turn off the interpretation of comments altogether.
<code>allowEscapes</code>	logical. Should C-style escapes such as <code>‘\n’</code> be processed or read verbatim (the default)? Note that if not within quotes these could be interpreted as a delimiter (but not as a comment character). For more details see <code>scan</code> .
<code>flush</code>	logical: if <code>TRUE</code> , <code>scan</code> will flush to the end of the line after reading the last of the fields requested. This allows putting comments after the last field.
<code>stringsAsFactors</code>	logical: should character vectors be converted to factors? Note that this is overridden by <code>as.is</code> and <code>colClasses</code> , both of which allow finer control.
<code>fileEncoding</code>	character string: if non-empty declares the encoding used on a file when given as a character string (not on an existing connection) so the character data can be re-encoded. See the ‘Encoding’ section of the help for <code>file</code> , the ‘R Data Import/Export’ manual and ‘Note’.

encoding	encoding to be assumed for input strings. It is used to mark character strings as known to be in Latin-1 or UTF-8 (see Encoding): it is not used to re-encode the input, but allows R to handle encoded strings in their native encoding (if one of those two). See ‘Value’ and ‘Note’.
skipNul	logical: should NULs be skipped?
dim	dimension of table content to show
...	Further arguments to be passed to <code>read.table</code> .

Details

Read a dataset of type table and show x rows and y columns

Value

read table content and a print out of the data head

Examples

```
## Not run:
# Example: read a table file and print the first 10 lines
# declare file
new.file <- "test.csv"

# read file and preview default
dth3 <- read.table.print(file = new.file, sep=",", quote = "\"", dec = ".",
fill = TRUE, comment.char = "", header = TRUE)

# read file and preview 10 rows and all columns
dth1 <- read.table.print(file = new.file, sep=",", quote = "\"", dec = ".",
fill = TRUE, comment.char = "", header = TRUE, dim = 10)

# read file and preview 10 rows and 5 columns
dth2 <- read.table.print(file = new.file, sep=",", quote = "\"", dec = ".",
fill = TRUE, comment.char = "", header = TRUE, dim = c(10,5))

## End(Not run)
```

refresh

Clear environment, clear console, set work directory and load files

Description

Shorthand to quickly clear console, clear environment, set working directory, load files

Usage

```
refresh(setwd = NULL, source = c(), load = c(), clearPkgs = FALSE)
```

Arguments

setwd	OPTIONAL. set working directory
source	OPTIONAL. source in file(s)
load	OPTIONAL. load in Rdata file(s)
clearPkgs	clear previously loaded packages

Details

The purpose of this function is provide a one-line code to clear the console, clear the environment, set working directory to a specified path, source in various files into the current file, and load RData files into the current environment. The first process in the sequence of events is to clear the environment. Then the working directory is set, prior to inclusion of various files and RData. With the directory being set first, the path to the sourced in or RData files will not need to be appended to the file name. See examples.

Value

cleared environment and set directory

Examples

```
if(interactive()){
#exactly like the clean function
#simply clear environment, clear console and devices
quickcode::refresh()

#clear combined with additional arguments
quickcode::refresh(
  clearPkgs = FALSE
) #also clear all previously loaded packages if set to TRUE

quickcode::refresh(
  setwd = "/home/"
) #clear env and also set working directory

quickcode::refresh(
  source = c("/home/file1.R","file2")
) #clear environment and source two files into current document

quickcode::refresh(
  setwd = "/home/",
  source = c("file1","file2")
) #clear environment, set working directory and source 2 files into environment

quickcode::refresh(
  setwd = "/home/",
  source="file1.R",
```

```
  load="obi.RData"  
) #clear environment, set working directory, source files and load RData  
}
```

rows.rep

Replicate Rows in a Data Frame

Description

This function replicates each row in a data frame a specified number of times.

Usage

```
rows.rep(data, n)
```

Arguments

`data` A data frame whose rows are to be replicated.
`n` An integer specifying the number of times each row should be replicated.

Value

A data frame with each row replicated ‘n’ times. If ‘n’ is less than or equal to 0, an empty data frame is returned.

Examples

```
# Example with a simple data frame  
df <- data.frame(A = c(1, 2), B = c("x", "y"))  
rows.rep(df, 3)  
  
# Example with no replication (n = 0)  
rows.rep(df, 0)
```

sample_by_column	<i>Re-sample a dataset by column and return number of entry needed</i>
------------------	--

Description

Shorthand to return a re-sample number of rows in a data frame by unique column

Usage

```
sample_by_column(.dt, col, n, seed = NULL, replace = FALSE)
```

Arguments

.dt	data frame to re-sample
col	column to uniquely re-sample
n	number of rows to return
seed	unique numeric value for reproducibility
replace	should sampling be with replacement

Value

data frame containing re-sampled rows from an original data frame

Examples

```
data1 <- data.frame(ID=1:10,MOT=11:20)
sample_by_column(data1,MOT,3)
sample_by_column(data1,ID,7)
```

setOnce	<i>Set a variable only once</i>
---------	---------------------------------

Description

Facilitates the one-time setting of a variable in R, ensuring its immutability thereafter.

Usage

```
setOnce(., val = 1L, envir = NULL)
```

Arguments

.	variable to set
val	the value to set for the variable
envir	environment where variables resides

Details

With this function, users can establish the change to the initial value of a variable, and it guarantees that any subsequent attempts to modify the variable are ignored. This feature ensures that the variable remains constant and immutable once it has been set, preventing unintentional changes and promoting code stability. This function simplifies the process of managing immutable variables in R, providing a reliable mechanism for enforcing consistency in data throughout the course of a program or script.

Value

the variable set to the new variable, along with a class of once added to the output

Examples

```
# set the value of vector_x1, vector_y1, vector_z1
init(vector_x1, vector_y1, vector_z1, value = 85)

# view the initial values of the variables
vector_x1
vector_y1
vector_z1

# task 1: change the value vector_x1 and prevent further changes
vector_x1 # check value of unchanged
vector_x1 * 0.56 # check value when x 0.56

setOnce(vector_x1, val = 4500) # set vector_x1
vector_x1 # check value
vector_x1 * 0.56 # check value when x 0.56

setOnce(vector_x1, val = 13) # set vector_x1 AGAIN, should not change
vector_x1 # check value
vector_x1 * 0.56 # check value when x 0.56

# task 2: In for loop, change vector_y1 and use later
vector_y1 # check value of unchanged

for(i in 1:20){
  setOnce(vector_y1,as.numeric(Sys.time()))
  # now let's see the difference between vector_y1
  # and the current time as it changes
  message("current vector_y1: ",vector_y1,"; subtraction res: ",as.numeric(Sys.time()) - vector_y1)
}

# task 3: In for lapply, change vector_z1 and use later
vector_z1 # check value of unchanged

invisible(
  lapply(1:20, function(i){
    setOnce(vector_z1,as.numeric(Sys.time()))
    # now let's see the difference between vector_z1
```

```
# and the current time as it changes
message("current vector_z1: ",vector_z1,"; subtraction res: ",as.numeric(Sys.time()) - vector_z1)
})
)

# result of all the tasks
vector_x1
vector_y1
vector_z1
```

sort_length	<i>Sort vector by length or file types of its content</i>
-------------	---

Description

Sort the length or file types of the content of a vector

Takes a vector of file names and sorts them by their file extensions (file type)

Usage

```
sort_length(vec, asc = TRUE)
```

```
sort_file_type(files, asc = TRUE)
```

Arguments

vec a vector

asc A logical value indicating whether to sort in ascending (TRUE) or descending (FALSE) order. Default is TRUE.

files A character vector containing file names to be sorted

Value

vector of items sorted by length

A character vector of sorted file names

Note

This function removes all NAs prior to sorting the vector

Examples

```
# sort by length of content
x = c("acs", "tt", "jdssr", "h", "grab")
sort_length(vec = x) # ascending order of length
sort_length(vec = x, asc = FALSE) # descending order of length

files <- c("doc1.pdf",
  "image.jpg", "house.csv", "notes.txt",
  "patab", "doc2.pdf", "data.csv", "pic.png", "cotab")
sort_file_type(files)
sort_file_type(files, asc = FALSE)
```

strsplit.bool

Split a string of values and return as boolean vector

Description

The purpose of this function is combine the functionality of **strsplit**, **unlist** and **as.logical**, which are often used together.

Usage

```
strsplit.bool(
  x,
  split,
  fixed = FALSE,
  perl = FALSE,
  useBytes = FALSE,
  type = 2
)
```

Arguments

x	character vector, each element of which is to be split. Other inputs, including a factor, will give an error.
split	character vector
fixed	logical. If TRUE match split exactly, otherwise use regular expressions. Has priority over perl.
perl	logical. Should Perl-compatible regexps be used?
useBytes	logical. If TRUE the matching is done byte-by-byte rather than character-by-character, and inputs with marked encodings are not converted.
type	type of return, see the as.boolean function for more info

Details

Given a sting, split by a separator into boolean

Value

boolean values based on split string

Examples

```
# string of numbers
num.01 = "0 1 0 0 1 0 1 T F TRUE FALSE t f"

# split a string of numbers and return as boolean 1/0
strsplit.bool(num.01, split = " ", type = 3)

# split a string of numbers and return as boolean TRUE/FALSE
strsplit.bool(num.01, split = " ", type = 2)

# split a string of numbers and return as boolean Yes/No
strsplit.bool(num.01, split = " ", type = 1)

# string of numbers
num.02 = "0abc1abc0abc0abc1abc0abc1abcTabcFabcTRUEabcFALSEabcf"

# split a string of numbers and return as boolean 1/0
strsplit.bool(num.02, split = "abc", type = 3)

# split a string of numbers and return as boolean TRUE/FALSE
strsplit.bool(num.02, split = "abc", type = 2)

# split a string of numbers and return as boolean Yes/No
strsplit.bool(num.02, split = "abc", type = 1)
```

strsplit.num

Split a string of numbers and return as numeric vector

Description

The purpose of this function is combine the functionality of **strsplit**, **unlist** and **as.numeric**, which are often used together.

Usage

```
strsplit.num(x, split, fixed = FALSE, perl = FALSE, useBytes = FALSE)
```

Arguments

x	character vector, each element of which is to be split. Other inputs, including a factor, will give an error.
split	character vector
fixed	logical. If TRUE match split exactly, otherwise use regular expressions. Has priority over perl.
perl	logical. Should Perl-compatible regexps be used?
useBytes	logical. If TRUE the matching is done byte-by-byte rather than character-by-character, and inputs with marked encodings are not converted.

Details

Given a sting, split by a separator into numbers

Value

numeric values based on split string

Examples

```
# Example 1
# string of numbers with separator " "
num.01 = "5 3 2 3 5 2 33 23 5 32 432 42 23 554"

# split a string of numbers and return as numeric
strsplit.num(num.01, split = " ")

# Example 2
# string of numbers with separator "|||"
num.02 = "0|||1|||4|||43|||6|||8|||00||| 1||| 0 1||| T |||F|||TRUE |||f"

# split a string of numbers and return as numeric
strsplit.num(num.02, split = "[|||]")
```

sub.range

Calculate the Range Difference of a Numeric Vector

Description

This function calculates the difference between the maximum and minimum values of a numeric vector.

Usage

```
sub.range(x)
```

Arguments

x A numeric vector for which the range difference is to be calculated.

Value

A numeric value representing the difference between the maximum and minimum values of the input vector. Returns 'NA' if the input is empty or contains only 'NA' values.

Examples

```
# Example with a numeric vector
numeric_vector <- c(1, 5, 3, 8, 2)
sub.range(numeric_vector)

# Example with missing values
sub.range(c(NA, 4, 7, NA, 10))
```

summarize.envobj *Get all the environment objects and their sizes*

Description

Retrieve the size contribution of all the available objects in the environment

Usage

```
summarize.envobj(envir = parent.frame())
```

Arguments

envir the environment to retrieve objects from

Value

a dataframe of all the variables within the environment

Examples

```
# Get a data frame of all environment objects and their size
summarize.envobj()
```

`switch_cols`*Switch the index of two columns in a data set*

Description

Allows the user to choose precisely which two columns they want to swap places, while optionally preventing some rows within the columns from being altered in the process. Excluded rows within the columns act as anchors that are immune from the switching operation on the selected columns.

Usage

```
switch_cols(data, col1, col2, keep.rows = NULL)
```

Arguments

<code>data</code>	dataset object
<code>col1</code>	numeric or character the first column name or number
<code>col2</code>	numeric or character the second column name or number
<code>keep.rows</code>	numeric. row number to keep

Examples

```
# Example using mtcars
data101 <- mtcars[1:7,]

head(data101) # preview overall data

# task 1: basic result of switching columns 5 and 6
head(switch_cols(data101, 5, 6))

# task 1: basic result of switching columns number 5 and name "gear"
head(switch_cols(data101, 5, "gear"))

# task 1: basic result of switching columns "qsec" and "carb"
head(switch_cols(data101, "qsec", "carb"))

# task 2: switch columns, but retain some rows with the switched columns

# lets exchange some columns, but keep content of row 4, 5 intact
data101[1:6,4:7] # preview the portion that is to be changed
res1 <- switch_cols(data101, col1 = 5, col2 = 6, keep.rows = 4:5) # use column numbers
res1[1:6,4:7] # check result, pay attention to rows 4, 5 of columns 5, 6 as well

data101[1:6,6:11] # preview the portion that is to be changed
res2 <- switch_cols(data101,
  col1 = "qsec",
```

```
col2 = "carb",
keep.rows = c(1,2,3)) # keep 1, 2, 3
res2[1:6,6:11] # check result
```

switch_rows	<i>Switch the index of two rows in a data set</i>
-------------	---

Description

Allows the user to choose precisely which two rows they want to swap places, while optionally preventing some columns from being altered in the process. Excluded columns within the rows act as anchors that are immune from the switching operation on the selected rows.

Usage

```
switch_rows(data, row1, row2, keep.cols = NULL)
```

Arguments

data	dataset object
row1	numeric. the first row number
row2	numeric. the second row number
keep.cols	numeric or character. column number or name to keep

Examples

```
# Example using mtcars
data100 <- mtcars[1:7,]

head(data100) # preview overall data

# task 1: basic result of switching rows 5 and 6
head(switch_rows(data100, 5, 6))

# task 2: switch rows, but retain some columns
data100[5:6,2:10] # preview the portion that is to be changed

# lets switch 2 rows, but keep content of columns 7, 8, 9 10 within the changed rows
res1 <- switch_rows(data100, row1 = 5, row2 = 6, keep.cols = 7:10) # use column numbers
res1[5:6,] # check result, pay attention to columns 9 and 10 as well
res2 <- switch_rows(data100,
row1 = 5,
row2 = 6,
keep.cols = c("disp","cyl")) # use column names
res2[5:6,] # check result, pay attention to columns "disp","cyl" as well
```

`track_func`*Track Function Usage and Performance*

Description

Up count the number of times a particular function is called. This function tracker is a higher-order function or decorator that wraps around other functions to monitor and record their execution patterns, including metrics like call frequency, execution time, argument patterns, and return values. It acts as a transparent layer that doesn't modify the original function's behavior but collects valuable metadata about its usage.

Usage

```
track_func(output.dest = "output_tracking.csv")
```

Arguments

`output.dest` destination of csv file to store outputs

Value

the numeric count of a function usage

Note

The usefulness of function tracking spans several critical areas:

Performance Optimization: By measuring execution times and frequency, developers can identify bottlenecks and frequently called functions that need optimization

Debugging: Tracking argument patterns and function call sequences helps pinpoint issues in complex applications

Usage Analytics: Understanding which features (functions) are most commonly used helps guide development priorities and API design decisions

Resource Management: Monitoring function behavior helps identify memory leaks, resource consumption patterns, and potential optimization opportunities

Testing: Usage patterns can inform test case design and coverage requirements, ensuring critical paths are well-tested

Documentation: Automatically gathering real-world usage examples helps maintain accurate and relevant documentation

Compliance: In regulated environments, function tracking can help maintain audit trails of system behavior

Examples

```
## Not run:  
library(quickcode)  
# Track usage of type2 and type1 functions
```

```
store.usage.file <- tempfile()
type5 <- function(x) type2(x)
type4 <- function(x) type3(x)
type3 <- function(x) type1(x)
type1 <- function(x) {
  mean(x)
  sd(x)
  track_func(store.usage.file)
}
type2 <- function(x) {
  type1(x)
  track_func(store.usage.file)
}

# add usage counts to store.usage.file
type1(number(10))
type2(number(10))
type3(number(10))
type4(number(10))
type5(number(10))

# check the stored function usage file
print(read.csv(store.usage.file))

## End(Not run)
```

trim.file

Remove Empty Lines from a File

Description

This function reads a text file, removes all empty lines, and saves the modified content back to the same file.

Usage

```
trim.file(file_path, resave = TRUE)
```

Arguments

file_path	A character string specifying the path to the text file.
resave	A logical value indicating if the file content should be resaved or returned

Value

NULL This function modifies the file in place and does not return a value.

Examples

```
if(interactive()){
# Remove empty lines from a file
trim.file("path/to/your/file.txt")
}
```

unique_len

Combine unique() and length()

Description

Combine two frequently used function together to return the length of the unique items of a vector

Usage

```
unique_len(.)
```

Arguments

. object such as vector or names(dataframe)

Value

length of the unique items in a vector

Examples

```
frenchnames1 = c("Léa","Obinna","Bastien","Léa","Obinna", "Hugo", "Amélie","Louis")
unique_len(frenchnames1)
```

vector_pop

Remove last n elements or specified elements from a vector like array_pop in PHP

Description

Shorthand to remove elements from a vector and save as the same name

Usage

```
vector_pop(., n = 1, el = NULL, ret = FALSE)
```

Arguments

.	parent vector
n	number of elements to remove
el	vector to remove
ret	TRUE or FALSE. whether to return value instead of setting it to the parent vector

Value

vector with elements removed

Examples

```
# basic example: pop off the last 2 values from vector
c(0,3,"A","Apple", TRUE) #before
vector_pop(c(0,3,"A","Apple", TRUE)) #after 1 pop
vector_pop(c(0,3,"A","Apple", TRUE), n=3) #after 3 pop

# using objects
num1 <- sample(330:400,10)
name1 <- "ObinnaObianomObiObianom"

#task: remove 1 element from the end of the vector and set it to the vector name
num1 #num1 vector before pop
vector_pop(num1) #does not return anything
num1 #num1 vector updated after pop

#task: remove 5 elements from the end, but do not set it to the vector name
num1 #num1 vector before pop
vector_pop(num1,5, ret = TRUE) #return modified vector
num1 #num1 vector remains the same after pop

#task: remove 6 elements from a word, set it back to vector name
name1 #name1 before pop
vector_pop(name1,6) #does not return anything
name1 #name updated after pop

#task: remove 3 elements from a word, Do not set it back to vector name
name1 #name1 before pop
vector_pop(name1,3, ret = TRUE) #returns modified name1
name1 #name1 not updated after pop

#task: remove 4 elements from the end of a vector and return both the removed content and remaining
v_f_num <- paste0(number(20),c("TI")) #simulate 20 numbers and add TI suffix
v_f_num #show simulated numbers
vector_pop(v_f_num, n = 4, ret = TRUE) #get the modified vector
vector_pop(v_f_num, n = 4, ret = "removed") #get the content removed

#task: remove specific items from vector
#note that this aspect of the functionality ignores the 'n' argument
v_f_num_2 <- paste0(number(6, seed = 33),c("AB")) #simulate 6 numbers using seed and add AB suffix
```

```
v_f_num_2 #show numbers
vector_pop(v_f_num_2, e1 = c("403211378AB")) #remove 1 specific entries
v_f_num_2 #show results
vector_pop(v_f_num_2, e1 = c("803690460AB","66592309AB")) #remove 2 specific entries
v_f_num_2 #show results
```

vector_push*Add elements to a vector like array_push in PHP*

Description

Shorthand to add elements to a vector and save as the same name

Usage

```
vector_push(., add, unique = FALSE, rm.na = FALSE, rm.empty = FALSE)
```

Arguments

.	first vector
add	vector to add
unique	remove duplicated entries
rm.na	remove NA values
rm.empty	remove empty values

Details

Note that two vectors are required in order to use this function. Also, note that the final result replaces the content of the first vector. This means that the original content of the 'first vector' will no longer exist after this function executes.

Value

vector combining fist and second vector, but have name set to the first

Use case

This function allows the combination of two vectors in one short line of code. It allows specification of further downstream filtering of the resulting vector such as selecting only unique items, removing NA or empty values. It simplifies a code chunk with many lines of code to concatenate and filter various vectors.

Examples

```
num1 <- number(10, seed = 45)
num2 <- "rpkg.net"

num1
num2

#Task: add num2 to num1 and re-save as num1
vector_push(num1,num2)
num1 #updated with num2
num2 #not updated

#Task: concatenate two vectors and remove duplicates
vector1 = number(4,seed = 5)
vector2 = number(8,seed = 5)
vector3 = number(12,seed = 5)

vector1 #length is 4
vector2 #length is 8
vector3 #length is 12

# with duplicated
vector_push(vector1,vector2, unique = FALSE)
vector1 #return modified vector
length(vector1) #length is 12 because nothing was removed
#duplicates in vector1 is 886905927 100040083 293768998 54080431

# without duplicated
vector_push(vector2,vector3, unique = TRUE)
vector2 #return modified vector
length(vector2) #length is 12 instead of 20
#Total of 8 duplicated numbers were removed

#Task: concatenate two vector and remove NA values
vector1 = number(5)
vector2 = c(4,NA,5,NA)
vector3 = number(5)

# with NA
vector_push(vector1,vector2, rm.na = FALSE)
vector1 #return modified vector

# without NA
vector_push(vector3,vector2, rm.na = TRUE)
vector3 #return modified vector

#Task: concatenate two vector and remove empty values
vector1 = number(5)
vector2 = c(4,'',5,'',NULL,'')
```

```
vector3 = number(5)

# with empty
vector_push(vector1,vector2, rm.empty = FALSE)
vector1 #return modified vector

# without empty
vector_push(vector3,vector2, rm.empty = TRUE)
vector3 #return modified vector
```

vector_shuffle	<i>Shuffle a vector just like shuffle in PHP</i>
----------------	--

Description

Shorthand to shuffle a vector and save

Usage

```
vector_shuffle(., replace = FALSE, prob = NULL, seed = NULL)
```

Arguments

.	vector to shuffle
replace	replace selected value
prob	probability of occurrence
seed	apply seed if indicated for reproducibility

Value

shuffled vector of items store to the vector name

Examples

```
#basic example
vector_shuffle(c(3,45,23,3,2,4,1))

#using objects
v1<-c(3,45,23,3,2,4,1)

#demonstrate vector_shuffle
vector_shuffle(v1)
v1 # show outputs

#demonstrate reproducibility in shuffle with seed
v0<-v1
vector_shuffle(v0)
```

```
v0 #first output

v0<-v1
vector_shuffle(v0)
v0 # different output from first output top

v0<-v1
vector_shuffle(v0,seed = 232L)
v0 #second output

v0<-v1
vector_shuffle(v0,seed = 232L)
v0 #the same output as second output top
```

yesNoBool

Convert Yes/No to Binary or Logical

Description

Seamlessly convert a yes or no to either a binary or logical output

Usage

```
yesNoBool(
  table,
  fldname,
  out = c("change", "append", "vector"),
  type = c("bin", "log")
)
```

Arguments

table	data frame
fldname	field name in the data frame
out	output form, choices - change, append, vector
type	output type, choices - bin, log

Details

type - "bin" for binary, and "log" for logical

Value

converted Yes/No entries into 1/0 or TRUE/FALSE

Examples

```

# Declare data for example
usedata <- data.frame(ID = 1:32)
usedata #view the dataset

usedata$yess = rep(c("yes","n","no","YES","No","NO","yES","Y"),4) #create a new column
usedata #view the modified dataset

# Set all yess field as standardize boolean
# Task: convert the "yess" column content to 1/0 or TRUE/FALSE
# Notice that you have add the column name with or without quotes
yesNoBool(usedata,yess, type="bin") #set all as binary 1/0
yesNoBool(usedata,"yess", type="log") #set all as logical TRUE/FALSE

# Task: By default, the 'out' argument is set to "change"
# means that the original data field will be
# replaced with the results as above

# In this example, set the out variable to
# append data frame with a new column name containing the result

yesNoBool(usedata,yess,"append")
#or yesNoBool(usedata,"yess","append")

# In this example, return as vector
yesNoBool(usedata,yess,"vector")
#or yesNoBool(usedata,"yess","vector")

# Task: Return result as logical
yesNoBool(usedata,"yess",type = "log")

```

zscore

Calculates Z-Scores of a distribution

Description

Calculates Z-Scores based on data

Usage

```
zscore(.data, round, na.rm = TRUE)
```

```
zscoreGrowthCurve(Xi, Mi, Si, Li = !0)
```

Arguments

.data	data object
round	round output to how many decimal place description
na.rm	remove NA values before calculating z-scores
Xi	physical measurement (e.g. weight, length, head circumference, stature or calculated BMI value)
Mi	values from the table (see reference) corresponding to the age in months of the child
Si	values from the table (see reference) corresponding to the age in months of the child
Li	values from the table (see reference) corresponding to the age in months of the child

Value

zscore calculated based on data object or parameters

References

CDC growth chart Z score calculation: <https://www.cdc.gov/growthcharts/cdc-data-files.htm>

Examples

```
# Capture z-score from the following distribution x
x = c(6, 7, 7, 12, 13, 13, 15, 16, 19, 22)
z_scores = zscore(x, round = 2) # limit to 2 decimal place
z_scores = zscore(x) # no decimal place limit

df = data.frame(val = x, zscore = z_scores)
head(df)
#EXAMPLE for zscore based on CDC growth chart

# Calculate the zscore for a patient weighing 50kg
Li=-0.1600954
Mi=9.476500305
Si=0.11218624
Xi=50
zscoreGrowthCurve(Xi,Mi,Si,Li)
```

%nin% *Not in vector or array*

Description

Check if entry is in vector

Usage

```
x %nin% table
```

Arguments

```
x          vector entry
table      table of items to check
```

Value

a boolean value to indicate if entry is present

Examples

```
5 %nin% c(1:10) #FALSE
5 %nin% c(11:20) #TRUE

x = "a"
if(x %nin% letters) x

# let's say we are trying to exclude numbers from a vector
vector_num1 <- number(9, max.digits = 5, seed = 1) #simulate 9 numbers
vector_num1 #values
vector_num1[vector_num1 %nin% c(83615,85229)]#return values not 83615 or 85229
```

%.% *simple function chaining routine*

Description

chain multiple function to a call

Usage

```
obj %.% funcs

chain_sep(sep = "\\.\\"")
```

Arguments

```
obj          data object to apply function
funcs        function chains to apply to the data object
sep          separator for functions argument values
```

Value

the result of applying the chained functions to the data object

Note

chain_sep allows the user to preset the separator for the function chaining

e.g. you can call the function to set sep = "__" before using the

Examples

```
#use default sep ".."
1:3%.%unique..length
sample(1:1000,10,replace=TRUE) %.%unique..length
sample(1:10,10,replace=TRUE) %.%unique..cumsum

# set sep before function chaining
chain_sep("__")
sample(1:10,10,replace=TRUE) %.%unique__cumsum
sample(1:10,10,replace=TRUE) %.%unique__cumsum__length

# set sep before function chaining
chain_sep("X")
sample(1:10,10,replace=TRUE) %.%uniqueXcumsum
```

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