Package 'handyFunctions'

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checkCols

check the validation and return index of cols given from input in raw-DataFrame

Description

check the validation and return index of cols given from input in rawDataFrame

Usage

```
checkCols(rawDataFrame, cols)
```

Arguments

```
rawDataFrame raw data.frame
cols specific cols given from input
```

Value

return validation (only FALSE if invaild cols input) or index of cols

Examples

```
library(handyFunctions)
data(people)
checkCols(people, c("..name", "..sex"))
# OR
checkCols(people, c(1, 2))
```

checkDtype

Return suggested dtype of vector input

Description

Return suggested dtype of vector input

Usage

```
checkDtype(vector)
```

Arguments

vector

vector/list input

grade 3

Value

Return suggested dtypes of vector

Examples

```
library(handyFunctions)
vector <- c(1, 2, 3, "", NA, " ", "four", "NA", 5)
checkDtype(vector)</pre>
```

grade

Grade records of virtual persons in high school

Description

A dataset containing the personal grade information (chinese, math, english, physics, biology, chemistry) of virtual persons.

Usage

grade

Format

A data frame with 6 rows and 7 variables:

name name, chinese or foreigner, in carats
chinese grade of the chinese, in numbers
math grade of the math, in numbers
english grade of the english, in numbers
physics grade of the physics, in numbers
biology grade of the biology, in numbers
chemistry grade of the chemistry, in numbers ...

Source

"simulated dataset"

4 mergeCustom

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Return the index of source vector matched with query vector

Description

Return the index of source vector matched with query vector

Usage

```
matchIndex(SourceInfo, queryInfo, queryType = TRUE)
```

Arguments

```
SourceInfo the source vector queryInfo the query vector
```

queryType logical If set it to accurate match (default: TRUE)

Value

the index of source vector matched with query vector

Examples

```
library(handyFunctions)
data(grade)
matchIndex(grade[, "name"], c("Ming Li", "Bang Wei"))
```

mergeCustom

merge two data.frame based on xcol and ycol

Description

merge two data.frame based on xcol and ycol

Usage

```
mergeCustom(x, y, xcol, ycol)
```

Arguments

X	the first data.frame
У	the second data.frame
xcol	colnames which you want to merged in first data.frame
ycol	colnames which you want to merged in second data.frame

Value

return the new data.frame merged

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Examples

```
library(handyFunctions)
data(people)
data(grade)
mergeCustom(people, grade, "..name", "name")
```

modifyColNames

Return reformatted data.frame with standard col names

Description

Return reformatted data.frame with standard col names

Usage

```
modifyColNames(rawDataFrame, cols = TRUE, rawSep = "..", sep = "_")
```

Arguments

rawDataFrame	Raw data.frame input
cols	Specific col names or indexes what you want to reform at (default: TRUE, use all cols)
rawSep	Raw odd separation symbol in col names of raw data.frame. Note: it supports regEx (regular expression), so "." means all possible symbols. If you want to use the "." dot notation, please use "[.]".
sep	Separation symbol in col names of modified data.frame

Value

A modified data.frame with col names separated by your given delimitator

Examples

```
library(handyFunctions)
data(people)
modified_people <- modifyColNames(people,rawSep = "[.][.]")</pre>
```

6 modifyRowNames

modifyColTypes	Return s DataFran	 appropriate	dtypes	for	each	column	in	raw-	

Description

Return suggested appropriate dtypes for each column in rawDataFrame

Usage

```
modifyColTypes(rawDataFrame, cols = TRUE, dtype = FALSE, custom = FALSE)
```

Arguments

rawDataFrame	Raw data.frame
cols	Specify cols which you want to change its dtypes when custom is FALSE (default: TRUE, for all cols)
dtype	Specify indexed matched dtypes wheih you want to update when custom is FALSE (default: FALSE, for automatically update)
custom	Option whether set to auto/custom , you can specify your custom dtypes for cols given when setting to TRUE (default: FALSE, for auto)

Value

Return a new data.frame with appropriate dtypes suggested for each cols

Examples

```
library(handyFunctions)
data(people)
modifyColTypes(people)
```

modifyRowNames

Return reformatted data.frame with standard row names

Description

Return reformatted data.frame with standard row names

Usage

```
modifyRowNames(rawDataFrame, rows = TRUE, rawSep = "..", sep = "_")
```

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Arguments

${\tt rawDataFrame}$	Raw data.frame input
rows	Specific row names or indexes what you want to reform at (default: TRUE, use all row) $$
rawSep	Raw odd separation symbol in row names of raw data.frame. Note: it supports regEx (regular expression), so "." means all possible symbols. If you want to use the "." dot notation, please use "[.]".
sep	Separation symbol in row names of modified data.frame

Value

A modified data.frame with row names separated by your given delimitator

Examples

```
library(handyFunctions)
data(people)
modifyRowNames(people)
```

people Basic information of virtual persons	people	Basic information of virtual persons
---	--------	--------------------------------------

Description

A dataset containing the personal basic information (name, sex, age, and death_age) of virtual persons.

Usage

```
people
```

Format

A data frame with 6 rows and 4 variables:

```
..name name, chinese or foreigner, in carats
```

..sex sex of the person, in carats

..age living age in final record, in numbers

..death..age final age when a person is dead, in numbers ...

Source

"simulated dataset"

ShowSNPDensityPlot

queryingInfo	return index of x data.frame with the given vector/list or ycol in				
	data.frame (if set the accurate match or not)				

Description

return index of x data.frame with the given vector/list or yool in data.frame (if set the accurate match or not)

Usage

```
queryingInfo(SourceData, sourceCol, queryCol, queryInfo, queryType = TRUE)
```

Arguments

```
SourceData the source data.frame which you want to query
sourceCol the col names or index of query field in source data.frame
queryCol the col names or index of return field in source data.frame
queryInfo vector/list the query info
queryType logical if set it to accurate match (default: TRUE)
```

Value

a vector in query field matched with query info in source data

Examples

```
library(handyFunctions)
data(grade)
queryingInfo(grade, "name", "chinese", c("Ming Li", "Bang Wei"))
```

ShowSNPDensityPlot Function of showing SNP density at chromosome level

Description

Function of showing SNP density at chromosome level

Usage

```
ShowSNPDensityPlot(
  densityData,
  binSize,
  densityColorBar = c("grey", "darkgreen", "yellow", "red"),
  chromSet = c(1:22),
  withchr = FALSE
)
```

Arguments

densityData the raw density data generated from vcftools binSize the bin size set while generating density data

densityColorBar

vector Specific the color bar for plotting density plot (generally four colors)

chromSet vector Filtered chrom set which you want to plot (it must be matched with the

CHROM column in densityData)

withchr logical If the chromsome labels of density plot is prefixed with "chr". Note: it

cannot work when the filtered chrom set contain other uncommon chrom sym-

bols (e.g. NC0*, etc)

Value

A ggplot2 object for SNP density plot

Examples

```
library(handyFunctions)
data(SNV_1MB_density_data)
ShowSNPDensityPlot(SNV_1MB_density_data, binSize = 1e6, chromSet = c(38:1))
```

```
SNV_1MB_density_data
```

The SNPV number within 1Mb bins at chromosome levels generated from transcriptome dataset of two dog populations (including wild wolf and domesticated dogs).

Description

A dataset containing the SNV number within 1Mb bins called from transcriptome dataset of wild wolf and domesticated dogs.

Usage

```
SNV_1MB_density_data
```

Format

A data frame with 2544 rows and 4 variables:

CHROM chrom id, reference genome of CanFam3.1, in numbers/carats

BIN_START the start genomic coordinate for one bin at relevant chromosome, in numbers **SNP_COUNT** the end genomic coordinate for one bin at relevant chromosome, in numbers

VARIANTS.KB SNV(variants) number within one bin per KB, in numbers ...

Source

"real dataset"

10 unifyDataframe

splitCol	Return	specific-indexed	vector	according	to	given	delimita-
	tor/separator by splitting one co						

Description

Return specific-indexed vector according to given delimitator/separator by splitting one col in data.frame

Usage

```
splitCol(data, col = FALSE, sep, index, fixed = TRUE)
```

Arguments

data	vector or data.frame input
col	the col names or indexes if data.frame input
sep	separation deliminator
index	the index of symbol which you want
fixed	logical. If TRUE match split exactly, otherwise use regular expressions, detailed info can be seen in strsplit.

Value

specific-indexed vector or factor

Examples

```
library(handyFunctions)
data(people)
splitCol(people, col = 1, sep = " ", index = 2)
```

unifyDataframe

Reformat dataframe with the all modifiers simultaneously (colNames, rowNames and dtypes)

Description

 $Reform at data frame \ with \ the \ all \ modifiers \ simultaneously \ (col Names, \ row Names \ and \ dtypes)$

Usage

```
unifyDataframe(
  rawDataFrame,
  rawRowSep = "..",
  rowSep = "_",
  rawColSep = "..",
  colSep = "_",
  changeDtype = TRUE
)
```

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Arguments

rawDataFrame raw data.frame

rawRowSep raw separation deliminator of row names in raw data.frame

rowSep the new separation deliminator of row names

rawColSep raw separation deliminator of col names in raw data.frame

colSep the new separation deliminator of col names

changeDtype if change the dtypes of cols

Value

A modified data.frame with applied to above all modifiers

Examples

```
library(handyFunctions)
data(people)
unifyDataframe(people,rawColSep = "[.][.]")
```

Index

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