# Package 'HGNChelper'

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| <b>Depends</b> R (>= 2.10)  |
| Author Levi Waldron and Markus Riester  |
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| License GPL (>=2.0)   |
| <b>Title</b> Handy Functions for Working With HGNC Gene Symbols and Affymetrix Probeset Identifiers.  |
| Description Contains functions for identifying and correcting HGNC gene symbols which have been converted to date format by Excel, for reversibly converting between HGNC symbols and valid R names, identifying invalid HGNC symbols and correcting synonyms and outdated symbols which can be mapped to an official symbol. |
| <pre>URL https://bitbucket.org/lwaldron/hgnchelper</pre>  |
| BugReports https://bitbucket.org/lwaldron/hgnchelper/issues   |
| R topics documented:  |
| HGNChelper-package affyToR checkGeneSymbols findExcelGeneSymbols hgnc.table rToAffy rToSymbol symbolToR   |
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HGNChelper-package Handy functions for working with HGNC gene symbols and Affymetrix probeset identifiers.

**Description** 

Contains functions for identifying and correcting HGNC gene symbols which have been converted to date format by Excel, for reversibly converting between HGNC symbols and valid R names, identifying invalid HGNC symbols and correcting synonyms and outdated symbols which can be mapped to an official symbol.

#### **Details**

Package: HGNChelper

Maintainer: Levi Waldron < lwaldron@hsph.harvard.edu>

Depends: R (>= 2.10)

Author: Levi Waldron and Markus Riester

Version: 0.3.1 Date: 2014-12-03 License: GPL (>2.0)

Title: Handy functions for working with HGNC gene symbols and Affymetrix probeset identifiers.

URL: https://bitbucket.org/lwaldron/hgnchelper BugReports: https://bitbucket.org/lwaldron/hgnchelper

#### Author(s)

Levi Waldron and Markus Riester

affyToR function to convert Affymetrix probeset identifiers to valid R names

#### **Description**

This function simply prepends "affy." to the probeset IDs to create valid R names. Reverse operation is done by the rToAffy function.

#### **Usage**

affyToR(x)

#### **Arguments**

vector of Affymetrix probeset identifiers, or any identifier which may with a digit.

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#### Value

a character vector that is simply x with "affy." prepended to each value.

#### Author(s)

Levi Waldron and Markus Riester

checkGeneSymbols

function to identify outdated or Excel-mogrified gene symbols

#### **Description**

This function identifies gene symbols which are outdated or may have been mogrified by Excel or other spreadsheet programs. If output is assigned to a variable, it returns a data frame of the same number of rows as the input, with a second column indicating whether the symbols are valid and a third column with a corrected gene list.

#### Usage

checkGeneSymbols(x, unmapped.as.na=TRUE, hgnc.table=NULL)

#### **Arguments**

x Vector of gene symbols to check for mogrified or outdated values

unmapped.as.na If TRUE, unmapped symbols will appear as NA in the Suggested.Symbol col-

umn. If FALSE, the original unmapped symbol will be kept when no correction

can be found.

hgnc.table If hgnc.table is a data.frame with colnames(hgnc.table) identical to c("Symbol",

"Approved.Symbol"), it is used to correct gene symbols in x. Otherwise, the default table data("hgnc.table", package="HGNChelper") is used. The function used for creating this the hgnc.table dataframe can be found in the inst/hgncLookup.R file, in the source of this package. By default the hgnc.table dataframe shipped

with this package is used (see ?hgnc.table)

#### Value

The function will return a data.frame of the same number of rows as the input, with corrections possible from hgnc.table.

#### Author(s)

Levi Waldron and Markus Riester

#### See Also

hgnc.table

#### **Examples**

findExcelGeneSymbols function to identify Excel-mogrified gene symbols

#### **Description**

This function identifies gene symbols which may have been mogrified by Excel or other spreadsheet programs. If output is assigned to a variable, it returns a vector of the same length where symbols which could be mapped have been mapped. Note that this function is superceded by checkGeneSymbols, which corrects Excel-mogrified gene symbols as well as aliases and outdated symbols.

#### Usage

#### **Arguments**

х

Vector of gene symbols to check for mogrified values

mog.map

Map of known mogrifications. A default map is available with this package by data(mog.map), but any map may be used. This should be a dataframe with two columns: original and mogrified, containing the correct and incorrect symbols, respectively.

regex

Regular expression, recognized by the base::grep function which is called with ignore.case=TRUE, to identify mogrified symbols. It is not necessary for all matches to have a corresponding entry in mog.map\$mogrified; values in x which are matched by this regex but are not found in mog.map\$mogrified simply will not be corrected. This regex is based that provided by Zeeberg et al., Mistaken Identifiers: Gene name errors can be introduced inadvertently when using Excel in bioinformatics. BMC Bioinformatics 2004, 5:80.

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#### Value

if the return value of the function is assigned to a variable, the function will return a vector of the same length as the input, with corrections possible from mog.map made.

#### Author(s)

Levi Waldron and Markus Riester

#### See Also

checkGeneSymbols

hgnc.table

All current and withdrawn HGNC symbols.

#### **Description**

All current and withdrawn symbols from genenames.org.

#### **Format**

A dataframe with the first column providing a gene symbol or known alias (including withdrawn symbols), second column providing the approved HGNC symbol.

#### Source

Extracted from genenames.org prior to package build.

#### **Examples**

```
data("hgnc.table", package="HGNChelper", envir=environment())
head(hgnc.table)
```

rToAffy

function to convert the output of affyToR back to the original Affymetrix probeset identifiers.

#### **Description**

This function simply strips the "affy." added by the affyToR function.

#### Usage

rToAffy(x)

#### **Arguments**

Х

the character vector returned by the affyToR function.

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#### Value

a character vector of Affymetrix probeset identifiers.

#### Author(s)

Levi Waldron and Markus Riester

rToSymbol

function to reverse the conversion made by symbolToR

#### Description

This function reverses the actions of the symbolToR function.

#### Usage

```
rToSymbol(x)
```

#### **Arguments**

х

the character vector returned by the symbolToR function.

#### Value

a character vector of HGNC gene symbols, which are not in general valid R names.

#### Author(s)

Levi Waldron and Markus Riester

#### **Examples**

```
library(HGNChelper)

data("hgnc.table", envir=environment())
hgnc.symbols <- as.character(na.omit(unique(hgnc.table[ ,2])))
if( !identical(all.equal(hgnc.symbols, rToSymbol(make.names(symbolToR(hgnc.symbols)))), TRUE))
    stop("HGNC mapping was not reversible.")</pre>
```

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| names. | symbolToR | function to *reversibly* convert HGNC gene symbols to valid R names. |
|--------|-----------|--|
|--------|-----------|--|

#### Description

This function reversibly converts HGNC gene symbols to valid R names by prepending "symbol.", and making the following substitutions: "-" to "hyphen", "@" to "ampersand", and "/" to "forward-slash".

#### Usage

```
symbolToR(x)
```

#### **Arguments**

Χ

vector of HGNC symbols

#### Value

a vector of valid R names, of the same length as x, which can be converted to the same HGNC symbols using the rToSymbol function.

#### Author(s)

Levi Waldron and Markus Riester

#### **Examples**

```
library(HGNChelper)

data("hgnc.table", envir=environment())

hgnc.symbols <- as.character(na.omit(unique(hgnc.table[ ,2])))
if( !identical(all.equal(hgnc.symbols, rToSymbol(make.names(symbolToR(hgnc.symbols)))), TRUE))
    stop("HGNC mapping was not reversible.")</pre>
```

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