# Package 'sinew'

September 16, 2017
Type Package
Title Create 'roxygen2' Skeleton with Information from Function Script
Version 0.3.2
<b>Date</b> 2017-09-15
Author c(person(``Jonathan", ``Sidi", email = ``yonis@metrumrg.com", role = c(``aut", ``cre")),  person(``Anton", ``Grishin", role = ``ctb"),  person(``Lorenzo", ``Busetto", role = ``ctb"))
Maintainer Jonathan Sidi <yonis@metrumrg.com></yonis@metrumrg.com>
Description
Create 'roxygen2' skeleton populated with information scraped from the within the function scrip Also creates field entries for imports in the 'DESCRIPTION' and import in the 'NAMES-PACE' files.  Can be run from the R console or through the 'RStudio' 'addin' menu.
<b>Depends</b> R (>= 2.3.0)
Imports rstudioapi,miniUI,shiny,utils,yaml
Suggests roxygen2
License GPL-2   GPL-3
<pre>URL https://github.com/metrumresearchgroup/sinew</pre>
<pre>BugReports https://github.com/metrumresearchgroup/sinew/issues</pre>
LazyData true
NeedsCompilation no
RoxygenNote 6.0.1
R topics documented:
interOxyAddIn
ls_param
makeDictionary
makeImport
makeOxyFile
moga
rmOxygen
sinew_opts
tabular

2 ls\_param

Index 11

# Description

Launches an interactive addin for insertion of roxygen2 comments in files. Allows selection of extra parameters for make0xygen

## Usage

```
interOxyAddIn()
```

## **Details**

Open an .R file in Rstudio's source editor. Launch the add-in via Addins -> interactiveOxygen or interOxyAddIn() in the console. Add-in opens in the viewer panel. Select function's/dataset's name in the source editor. If objects cannot be found, the addin prompts to source the file. Choose parameters for makeOxygen. Click Insert. Select next object's name. Rinse.Repeat. Click Quit when done with the file.

## Value

Nothing. Inserts roxygen2 comments in a file opened in the source editor.

# Author(s)

Anton Grishin, Jonathan Sidi

# See Also

```
View(sinew:::oxygenAddin)
```

# **Examples**

```
if(interactive()) interOxyAddIn()
```

ls\_param

Return roxygen2 parameter calls from parameter dictionary

## **Description**

Return roxygen2 parameter calls from the intersection of the parameters listed in the package dictionary and the formals of a function

# Usage

```
ls_param(obj, dictionary = "man-roxygen/Dictionary-1.R", print = TRUE)
```

makeDictionary 3

# **Arguments**

obj function or name of function

dictionary character, path\_to\_dictionary, Default: 'roxygen-man/Dictionary-1.R'

print boolean print output to console, Default: TRUE

## Value

character vector

# **Examples**

```
repo='https://raw.githubusercontent.com/metrumresearchgroup/sinew/master/'
dict_loc=file.path(repo,'man-roxygen/Dictionary-1.R')
ls_param(sinew::make0xygen,dictionary=dict_loc)
```

makeDictionary Parse package R files

Parse package R files to create dictionary of unique parameter defini-

# Description

Given list of R files function returns roxygen2 template consisting of intersecting parameter definitions

# Usage

```
makeDictionary(path, save_path = FALSE)
```

# **Arguments**

path character or character vector of paths to files to parse

save\_path boolean that allows for function to write template to man-roxygen subdirectory,

Default: FALSE

## Value

character/character vector of intersecting parameters

# **Examples**

```
makeDictionary('R')
```

4 makeOxyFile

makeImport	Scrape R script to create namespace calls for R documentation	

# Description

Scrape r script to create namespace calls for roxygen2, namespace or description files

# Usage

```
makeImport(script, cut = NULL, print = TRUE, format = "oxygen")
```

# Arguments

script	character connection to pass to readLines, can be file path, directory path, url path
cut	integer number of functions to write as importFrom until switches to import, Default: $\ensuremath{\text{NULL}}$
print	boolean print output to console, Default: TRUE
format	character the output format must be in c('oxygen', 'namespace', 'description').

Default: 'oxygen'

## **Examples**

```
makeImport('R',format = 'oxygen')
makeImport('R',format = 'namespace')
makeImport('R',format = 'description')
```

makeOxyFile

*Inserts roxygen2 skeletons in file(s).* 

# Description

Applies make0xygen function to all functions/dataframes in supplied file(s)

# Usage

```
makeOxyFile(input = NULL, overwrite = FALSE, verbose = TRUE, ...)
```

# **Arguments**

input	character, vector of path(s) to one or more .R files, a path to directory containing .R files, Default: NULL
overwrite	boolean, If TRUE overwrites file(s), FALSE writes "Oxy"- prefixed files in the same directory, Default: FALSE
verbose	boolean, If TRUE will print output to console and open edited files in the editor viewer, Defulat: TRUE
	additional parameters passed to makeOxygen

makeOxygen 5

#### **Details**

If an object cannot be found it will be sourced into a temporary environment. If the file already contains roxygen2 comments they will be deleted to avoid duplication. Some functions may require attaching additional packages. For instance, if functions were defined with purrr's compose or partial functions, omission of purr:: in definitions will require library(purrr) before proceding with make0xyFile.

#### Value

Nothing. Writes files with roxygen2 comments as a side effect

## Author(s)

Anton Grishin

## See Also

makeOxygen

## **Examples**

```
## Not run:
if(interactive()){
  makeOxyFile() # default args, opens system file selection dialogue
}
makeOxyFile("./myRfunctions/utils.R") # on one R file
makeOxyFile("./myRfunctions/") # on all R files in directory

# adds more fields to defaults, passes "cut" to makeImport

sinew_opts$append(list(add_fields=c("concept", "describeIn")))
makeOxyFile("./myRfunctions/utils.R", cut = 5)

## End(Not run)
```

make0xygen

Creates skeleton roxygen2 with information from within function script

# **Description**

Creates roxygen2 skeleton including title, description, import and other fields

## Usage

```
makeOxygen(obj, add_default = TRUE,
  add_fields = sinew_opts$get("add_fields"), use_dictionary = NULL,
  print = TRUE, ...)
```

6 moga

#### **Arguments**

obj function or name of function

add\_default boolean to add defaults values to the end of the PARAM fields, Default: TRUE

add\_fields character vector to add additional roxygen2 fields, Default: c("details", "examples", "seealso", "rdname

use\_dictionary character, path\_to\_dictionary, Default: NULL
print boolean print output to console, Default: TRUE

... arguments to be passed to makeImport

#### **Details**

add\_fields can include any slot except for the defaults (title,description,param,return). The order in add\_fields determines the order of printout. The roxygen2 fields to add are list below, for more information go to Generating Rd files

Field Skeleton

author AUTHOR [AUTHOR\_2]

backref src/filename.cpp

concept CONCEPT\_TERM\_1 [CONCEPT\_TERM\_2]

describeIn FUNCTION\_NAME DESCRIPTION

details DETAILS

example path/relative/to/packge/root

export

family FAMILY\_TITLE

field FIELD\_IN\_S4\_RefClass DESCRIPTION

format DATA\_STRUCTURE
importClassesFrom importMethodsFrom include DATA\_STRUCTURE

PKG CLASS\_a [CLASS\_b]

PKG METHOD\_a [METHOD\_b]

FILENAME.R [FILENAME\_b.R]

inherit [PKG::]SOURCE\_FUNCTION [FIELD\_a FIELD\_b]

inheritDotParams [PKG::]SOURCE\_FUNCTION

inheritSection [PKG::|SOURCE\_FUNCTION|SECTION\_a SECTION\_b]

keywords KEYWORD\_TERM

name NAME

rdname FUNCTION\_NAME references BIB\_CITATION section SECTION NAME

source \url{http://somewhere.important.com/}

slot SLOTNAME DESCRIPTION

template FILENAME templateVar NAME VALUE

useDynLib PKG [routine\_a routine\_b]

# **Examples**

makeOxygen(stats::lm)

moga Make Oxygen Great Again

rmOxygen 7

# **Description**

Update/append an R file that has roxygen2 headers already with updated information

#### **Usage**

```
moga(path, ..., force.fields = NULL, dry.run = TRUE, overwrite = FALSE)
```

## **Arguments**

path character path to R file

arguments to be passed to new makeOxygen

force.fields character, vector a field names that are in current header that are to be updated

Default: NULL

dry.run boolean, write lines to console the output, Default: TRUE overwrite boolean, overwrite contents of input file, Default: FALSE

#### **Details**

Cross references fields already in the roxygen2 header and adds any new ones from the updated call. To force a change to a field add field name to force.fields.

## Value

character

# **Examples**

 $\verb|moga('https://raw.githubusercontent.com/metrumresearchgroup/ggedit/master/R/aesColour.R')|$ 

rmOxygen

Remove roxygen2 Comments From an .R File

# **Description**

Strips .R files of roxygen2 style comments (#')

# Usage

```
rmOxygen(.file)
```

# Arguments

.file, path to an .R file, character vector of length 1

## Value

Nothing. Overwrites files as a side effect

## Author(s)

Anton Grishin

8 sinew\_opts

#### **Examples**

```
## Not run:
rmOxygen("./myRfunctions/function1.R")
## End(Not run)
```

sinew\_opts

Default and current sinew options

# **Description**

Options for functions in the sinew package. When running R code, the object sinew\_opts (default options) is not modified by chunk headers (local chunk options are merged with default options), whereas sinew\_opts\_current (current options) changes with different chunk headers and it always reflects the options for the current chunk.

## Usage

```
sinew_opts
sinew_opts_current
```

#### **Format**

An object of class list of length 5.

### **Details**

Normally we set up the global options once in the first code chunk in a document using sinew\_opts\$set(), so that all *latter* chunks will use these options. Note the global options set in one chunk will not affect the options in this chunk itself, and that is why we often need to set global options in a separate chunk.

Below is a list of default chunk options, retrieved via sinew\_opts\$get():

List of 28

```
$ add_fields : chr [1:6] "details" "examples" "seealso" "rdname" ...
```

\$ author : chr "Jonathan Sidi" \$ backref : chr "src/filename.cpp"

\$ concept : chr "CONCEPT\_TERM\_1 [CONCEPT\_TERM\_2]"

\$ describeIn : chr "FUNCTION\_NAME DESCRIPTION"

\$ details : chr "DETAILS"

\$ example : chr "path\_to\_file/relative/to/packge/root"

\$ examples : chr "\n#' \\dontrun{\n#' if(interactive()){\n#' #EXAMPLE1\n#' }\n#' }"

\$ export : chr ""

\$ family : chr "FAMILY TITLE"

\$ field : chr "FIELD\_IN\_S4\_RefClass DESCRIPTION"

\$ format : chr "DATA\_STRUCTURE"

\$ importClassesFrom: chr "PKG CLASS\_a [CLASS\_b]"

\$ importMethodsFrom: chr "PKG METHOD\_a [METHOD\_b]"

\$ include : chr "FILENAME.R [FILENAME\_b.R]"

\$ inherit : chr "[PKG::]SOURCE\_FUNCTION [FIELD\_a FIELD\_b]"

\$ inheritDotParams : chr "[PKG::]SOURCE\_FUNCTION"

tabular 9

\$ inheritSection : chr "[PKG::]SOURCE\_FUNCTION [SECTION\_a SECTION\_b]"

\$ keywords : chr "KEYWORD\_TERM"

\$ name : chr "NAME"

\$ rdname : chr "FUNCTION\_NAME"

\$ references : chr "BIB\_CITATION"

\$ section : chr "SECTION\_NAME"

\$ source : chr "\url{http://somewhere.important.com/}"

\$ slot : chr "SLOTNAME DESCRIPTION"

\$ template : chr "FILENAME"
\$ templateVar : chr "NAME VALUE"

\$ useDynLib : chr "PKG [ROUTINE\_a ROUTINE\_b]"

## Note

sinew\_opts\_current is read-only in the sense that it does nothing if you call sinew\_opts\_current\$set();
you can only query the options via sinew\_opts\_current\$get().

## **Examples**

```
sinew_opts$get('add_fields')
```

tabular

Tabular for roxygen2

## **Description**

Convert data.frame to roxygen2 tabular format

## Usage

```
tabular(df, header = TRUE, ...)
```

# Arguments

df data.frame to convert to table

header boolean to control if header is created from names(df), Default: TRUE

.. arguments to pass to format

## Value

character

## Source

roxygen2 formatting

## See Also

format

## **Examples**

```
cat(tabular(mtcars[1:5, 1:5]))
cat(tabular(mtcars[1:5, 1:5],header=FALSE))
```

10 untangle

untangle

Split an R script by functions

# **Description**

Split a R script with multiple functions into multiple single function R files.

## Usage

```
untangle(text = NULL, file = "", dir.out = NULL, keep.body = TRUE)
```

## **Arguments**

text character, vector of R commands, Default: NULL

file character, path to R file, Default: "

dir.out character, path to save new R files, Default: NULL

keep. body boolean, if TRUE all non-funcitons will be saved to body.R, Default: TRUE

# Value

list of seperate functions

## Author(s)

Jonathan Sidi

# **Examples**

```
## Not run:
txt <- "#some comment
#' @import utils
yy <- function(a=4){</pre>
utils::head(runif(10),a)
 # a comment
}
v <- 20
#another comment
#' @import utils
zz <- function(v=10,a=3){</pre>
utils::head(runif(v),pmin(a,v))
}
zz(v)
untangle(txt,dir.out = 'test')
## End(Not run)
```

# **Index**

```
*Topic datasets
sinew_opts, 8

format, 9

interOxyAddIn, 2

ls_param, 2

makeDictionary, 3
makeImport, 4
makeOxyFile, 4
makeOxygen, 5, 5
moga, 6

rmOxygen, 7

sinew_opts, 8
sinew_opts_current (sinew_opts), 8

tabular, 9

untangle, 10
```