Package 'htmrenamer'

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Author Hugo Botelho <hugobotelho@gmail.com></hugobotelho@gmail.com>				
Maintainer Hugo Botelho <hugobotelho@gmail.com></hugobotelho@gmail.com>				
Description Enables the systematic renaming of Zeiss, Olympus and Leica automated microscopes.				
License GPL-3 + file LICENSE				
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echo

Print to renamer GUI

Description

Prints a message to the console and the renamer tool GUI and/or the R console. When printing to the console, just calls the print function.

Usage

```
echo(message, printToGUI = TRUE, printToConsole = TRUE)
```

Arguments

message character, the message to be printed.

printToGUI logical, indicating whether or not to print the message to the gtext widget on the

renamer GUI.

printToConsole logical, indicating whether or not to print the message to the console.

getfiles.XML.LRP

Find Leica MatrixScreener XML and LRP files

Description

Analyzes the Leica MatrixScreener 'AdditionalData' folder and returns the file names of ScanningTemplate files

Usage

```
getfiles.XML.LRP(folder)
```

Arguments

folder character, Leica MatrixScreener export folder.

Value

character vector with file names.

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Note

There must be exactly one copy of these files:

- {ScanningTemplate}templatename.xml
- {ScanningTemplate}templatename.lrp

newinfile.char

Create empty infile (character vector)

Description

Creates an empty microscope infile template, which can be printed to the console and/or saved as a file. The output is structured as a character vector. See the **Note** section for a description of the infile structure.

Usage

```
newinfile.char(numrow = 8, numcol = 12, show = FALSE, saveto = character())
```

Arguments

numrow	integer, the number of rows in the multiwell plate. The default value is 8 (i.e. 96 well plate).
numcol	integer, the number of columns in the multiwell plate. The default value is 12 (i.e. 96 well plate).
show	logical, indicating whether or not to print the result to the console.
saveto	character. If specified, the empty infile template is saved in a file at this location.

Value

Character vector with empty infile template.

Note

The microscope infile is a text file with the following structure:

```
001--A--01--00--00--data1--data2
002--A--02--01--00--data1--data2
003--A--03--02--00--data1--data2
```

In the infile, metadata fields are separated by a double dash (--). The meaning of the metadata fields is the following:

- Well number
- Row label (A, B, ...)
- Column label (01, 02, 03, ...)

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- Column label (00, 01, 02, ...)
- Row label (00, 01, 02, ...)
- Experimental data 1
- Experimental data 2

Wells are numbered in ascending order, line by line. In the case of a 96 well plate, wells are numbered as shown:

The experimental data fields may contain any relevant information describing the well contents.

See Also

Function newinfile.df creates an empty infile in the form of a data frame. Function read.infile.df reads infile text files from disk.

Examples

```
# Infile template for a 96 well plate
newinfile.char()

# Save 384 well plate infile template to the working directory.
# The TXT file extension is recommended.
newinfile.char(numrow = 16, numcol = 24, saveto = "./infile_template.txt")
```

newinfile.df

Create empty infile (data frame)

Description

Creates an empty microscope infile template, which can be printed to the console and/or saved as a file. The output is structured as a data frame. See the **Note** section for a description of the infile structure.

Usage

```
newinfile.df(numrow = 8, numcol = 12, show = FALSE, saveto = character())
```

Arguments

numrow	integer, the number of rows in the multiwell plate. The default value is 8 (i.e. 96 well plate).
numcol	integer, the number of columns in the multiwell plate. The default value is 12 (i.e. 96 well plate).
show	logical, indicating whether or not to print the result to the console.
saveto	character. If specified, the empty infile template is saved in a file at this location.

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Value

Data frame with empty infile template.

Note

The microscope infile is a text file with the following structure:

```
001--A--01--00--00--data1--data2
002--A--02--01--00--data1--data2
003--A--03--02--00--data1--data2
```

In the infile, metadata fields are separated by a double dash (--). The meaning of the metadata fields is the following:

- · Well number
- Row label (A, B, ...)
- Column label (01, 02, 03, ...)
- Column label (00, 01, 02, ...)
- Row label (00, 01, 02, ...)
- Experimental data 1
- Experimental data 2

Wells are numbered in ascending order, line by line. In the case of a 96 well plate, wells are numbered as shown:

The experimental data fields may contain any relevant information describing the well contents.

See Also

Function newinfile.char creates an empty infile in the form of a character vector. Function read.infile.df reads infile text files from disk.

Examples

```
# Infile template for a 96 well plate
newinfile.df()

# Save 384 well plate infile template to the working directory.
# The CSV file extension is recommended.
newinfile.df(numrow = 16, numcol = 24, saveto = "./infile_template.csv")
```

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read.infile.df

Read microscope infiles

Description

Imports a text file representing a microscope infile as a data frame. Skips all lines which do not match the infile structure. See newinfile.df or newinfile.char for a description of the infile structure.

Usage

```
read.infile.df(infilepath)
```

Arguments

infilepath character, the path to the infile.

Value

Data frame with infile.

See Also

Function newinfile.df creates an empty infile in the form of a data frame. Function newinfile.char creates an empty infile in the form of a character vector.

Examples

```
# Create infile in disk
tempfile <- tempfile()
newinfile.char(saveto = tempfile)

# Read infile
myinfile <- read.infile.df(tempfile)
head(myinfile)

# Delete infile file from disk
file.remove(tempfile)</pre>
```

rename_leica

Leica renamer

Description

Renaming of Leica files according to a microscope infile. Auto-detects the software which generated the images to call the apropriate renamer function. See newinfile.df or newinfile.char for a description of the infile structure.

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Usage

```
rename_leica(
   sourcefolder,
   targetfolder,
   infilepath,
   compress = FALSE,
   move = FALSE,
   outputDescriptors = TRUE,
   printMessages = TRUE,
   printFiles = TRUE,
   printToGUI = TRUE
)
```

Arguments

sourcefolder character, the folder with raw images.

targetfolder character, folder where to place renamed files. A subfolder wih the same name

as the infile will be created by the function and must not exist previously.

infilepath character, location of the microscope infile

compress logical, compress raw images?

move logical, move (rather than copy) raw images?

outputDescriptors

logical, save plain text files with experimental metadata?

printMessages,

logical, print messages in the console?

printFiles, logical, print file-by-file progress in the console?
printToGUI logical, print messages to the renamer GUI?

Value

None, this function just calls another function to perform the renaming.

See Also

rename_leica_gui for interacting with this function using a GUI.

rename_leica_gui Leica renamer GUI

Description

Displays the renamer tool GUI for Leica LAS X MatrixsSreener exports. The actual renaming process is performed by function rename_leica().

Usage

```
rename_leica_gui()
```

Value

Null

See Also

rename_leica is the function which implements the actual renaming algorithm. rename_zeiss_gui for the Zeiss renamer GUI.

```
rename_leica_matrixscreener

Leica renamer (MatrixScreener)
```

Description

Renaming of Leica MatrixScreener files according to a microscope infile. See newinfile.df or newinfile.char for a description of the infile structure.

Usage

```
rename_leica_matrixscreener(
   sourcefolder,
   targetfolder,
   infilepath,
   compress = FALSE,
   move = FALSE,
   outputDescriptors = TRUE,
   printMessages = TRUE,
   printFiles = TRUE,
   printToGUI = TRUE
)
```

Arguments

sourcefolder

name structure: demoplate_01--2019_01_01_12_00_00

targetfolder character, folder where to place renamed files. A subfolder wih the same name as the infile will be created by the function and must not exist previously.

infilepath character, location of the microscope infile

compress logical, compress raw images?

character, the folder with raw images. The folder usually has the folllowing

move logical, move (rather than copy) raw images?

outputDescriptors

logical, save plain text files with experimental metadata?

printMessages,

logical, print messages in the console?

printFiles, logical, print file-by-file progress in the console?
printToGUI logical, print messages to the renamer GUI?

Value

Renamed files, metadata and log file

See Also

rename_leica_gui for interacting with this function using a GUI.

```
rename_leica_navigator

Leica renamer (Navigator)
```

Description

Renaming of Leica Navigator files according to a microscope infile. See newinfile.df or newinfile.char for a description of the infile structure.

Usage

```
rename_leica_navigator(
   sourcefolder,
   targetfolder,
   infilepath,
   compress = FALSE,
   move = FALSE,
   outputDescriptors = TRUE,
   printMessages = TRUE,
   printFiles = TRUE,
   printToGUI = TRUE
)
```

Arguments

```
sourcefolder
                  character, the folder with raw images. Images must have been exported as tif
                  files.
                  character, folder where to place renamed files. A subfolder wih the same name
targetfolder
                  as the infile will be created by the function and must not exist previously.
infilepath
                  character, location of the microscope infile
compress
                  logical, compress raw images?
                  logical, move (rather than copy) raw images?
move
outputDescriptors
                  logical, save plain text files with experimental metadata?
printMessages,
                  logical, print messages in the console?
printFiles,
                  logical, print file-by-file progress in the console?
                  logical, print messages to the renamer GUI?
printToGUI
```

Value

Renamed files, metadata and log file

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See Also

rename_leica_gui for interacting with this function using a GUI.

rename_olympus

Olympus renamer

Description

Renaming of Olympus OIB files according to a microscope infile. See newinfile.df or newinfile.char for a description of the infile structure.

Usage

```
rename_olympus(
   sourcefolder,
   targetfolder,
   infilepath,
   numrow,
   numcol,
   REGEXvalidimages = "^.*(\\d{4})_(\\d{2}).oib$",
   printMessages = TRUE,
   printFiles = TRUE,
   printToGUI = FALSE,
   move = FALSE
)
```

Arguments

sourcefolder character, the folder with raw images

targetfolder character, folder where to place renamed files. A subfolder with the same name

as the infile will be created by the function and must not exist previously.

infilepath character, location of the microscope infile
numrow integer, number of rows in the multi well plate
numcol integer, number of columns in the multi well plate

 ${\sf REGEXvalidimages}$

character, regular expression matching all files which should be renamed. Captures (($\1$)) the scene number (i.e. well number) and (($\2$)) the time number in

a time lapse.

printMessages,

logical, print messages in the console?

printFiles, logical, print file-by-file progress in the console?
printToGUI logical, print messages to the renamer GUI?
move logical, move (rather than copy) raw images?

Value

Renamed files, metadata and log file

Null

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See Also

rename_olympus_gui for interacting with this function using a GUI.

rename_olympus_gui

Olympus renamer GUI

Description

Displays the renamer tool GUI for Olympus OIB files. The actual renaming process is performed by function rename_olympus().

Usage

```
rename_olympus_gui()
```

Value

Null

See Also

```
rename_olympus is the function which implements the actual renaming algorithm. rename_leica_gui for the Leica renamer GUI. rename_zeiss_gui for the Zeiss renamer GUI.
```

rename_zeiss

Zeiss renamer

Description

Renaming of Zeiss TIF files exported with Zen Blue according to a microscope infile. See newinfile.df or newinfile.char for a description of the infile structure.

Usage

```
rename_zeiss(
   sourcefolder,
   targetfolder,
   infilepath,
   numrow,
   numcol,
   REGEXvalidimages = "^.*?s.*?t.*?\\.tif$",
   REGEXscenenum = "^.*/.*_s(.*)t.*\\.tif$",
   REGEXtimenum = "^.*/.*_s.*t(.*)\\.tif$",
   printMessages = TRUE,
   printFiles = TRUE,
   printToGUI = FALSE,
   move = FALSE
)
```

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Arguments

sourcefolder character, the folder with raw images.

targetfolder character, folder where to place renamed files. A subfolder with the same name

as the infile will be created by the function and must not exist previously.

infilepath character, location of the microscope infile
numrow integer, number of rows in the multi well plate
numcol integer, number of columns in the multi well plate

REGEXvalidimages

character, regular expression matching all files which should be renamed

REGEXscenenum character, regular expression capturing ((\1)) the scene number (i.e. well num-

ber)

REGEXtimenum character, regular expression capturing ((\1)) the time number in a time lapse

printMessages,

logical, print messages in the console?

printFiles, logical, print file-by-file progress in the console?
printToGUI logical, print messages to the renamer GUI?
move logical, move (rather than copy) raw images?

Value

Renamed files, metadata and log file

Null

See Also

rename_zeiss_gui for interacting with this function using a GUI.

rename_zeiss_gui

Zeiss renamer GUI

Description

Displays the renamer tool GUI for Zeiss Zen Blue exports. The actual renaming process is performed by function rename_zeiss().

Usage

```
rename_zeiss_gui()
```

Value

Null

See Also

```
rename_zeiss is the function which implements the actual renaming algorithm. rename_leica_gui for the Leica renamer GUI.
```

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Description

Determines whether images coming from a Leica microscope were generated with the MatrixScreener or Navigator software.

Usage

whichLeicaSoftware(folder)

Arguments

folder

character, Leica export folder.

Value

character, the name of the software which generated the images.

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