Package 'RLumSTARR'

August 25, 2021

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Type Package			
Title Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE)			
Version 0.1.0.9000-85			
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Description Analysing spatially resolved radiofluorescence data using ImageJ. A collection of functions to support early work on the subject.			
License GPL-3			
Depends R (>= 4.0), utils, methods			
Imports Luminescence (>= 0.9.8), cli (>= 3.0.1), rjags (>= 4-8), coda (>= 0.19-1)			
Encoding UTF-8			
Language en-GB			
LazyData true			
RoxygenNote 7.1.1			
R topics documented:			
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RLumSTARR-package

SpaTiAlly Resolved Radiofluorescence

Description

A collection of functions to analyse spatially resolved radiofluorescence data

Details

Funding

• Sebastian Kreutzer received funding from the European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement No 844457 (CREDit).

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) Dirk Mittelstrass (Germany), RLum Developer Team

References

##TODO

create_RFCurveArray

Create Multidimensional Curve Arrays from RF File input

Description

Helper function to create a multidimensional curve array based on RF-file input imported using the function Luminescence::read_RF2R() to prepare the Bayesian modelling process

Usage

```
create_RFCurveArray(files)
```

Arguments

files

list (required): list of .rf files to be imported

Value

Returns a list of class RLumSTARR.RFCurveArrary with two arrays for the RF_nat and the RF_reg curve

Function version

0.1.0

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How to cite

Kreutzer, S., 2021. create_RFCurveArray(): Create Multidimensional Curve Arrays from RF File input. Function version 0.1.0. In: Kreutzer, S., Mittelstrass, D., 2021. RLumSTARR: Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE). R package version 0.1.0.9000-85.

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) , RLum Developer Team

See Also

Luminescence::read_RF2R

Examples

```
## list files using package external data
files <- list.files(system.file("extdata", "", package="RLumSTARR"), full.names=TRUE)
## create curve array
create_RFCurveArray(files = files)</pre>
```

extract_TRUELight

Extract True Light from the Camera Measurements using a Bayesian Approach

Description

A Bayesian modelling approach to extract the true light using the expanding region-of-interest (ROI) approach proposed by Cunningham and Clark-Balzan (2017). The function will return the results for a **single** curve from a **single** ROI

Usage

```
extract_TRUELight(
  data,
  element = c("RF_nat", "RF_reg"),
  ROI = 2,
  stepping = 1,
  method_control = list(),
  verbose = TRUE
)
```

Arguments

data array (required): object created by create_RFCurveArray

element character (with default): element from the input to be analysed, 'supported are

only RF_nat or RF_reg

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ROI numeric (optional): ROI to be analysed, if nothing is given all ROIs are anal-

ysed, however, the first ROIS is discarded!

stepping numeric (with default): stepping parameter that allows you to model only every

xth (the value in stepping). This option can be extremely useful to play with data because it dramatically improves the modelling speed because less data are

considered.

method_control list(optional): parameter to be passed to rjags. Supported are n.chain, n.iter,

thin, variable.names, model, see details for more.

verbose logical (with default): enable/disable terminal feedback

Details

Method control

Supported options to be passed via the parameter method_control, most of them are used internally for the calls to rjags::jags.model and rjags::coda.samples.

PARAMETE	ER TYPE	DESCRIPTION
n.chain	numeric	the number of MCMC chains
n.iter	numeric	number of iterations for the MC runs
thin	numeric	thinning interval used for the monitoring
variable.na	ames character	variable names to monitor, alpha is always monitored
mode1	character	the bugs model

Value

Returns a list with an the following elements:

- ...\$RF_curve: Luminescence::RLum.Data.Curve object (the RF curve with the true light)
- ...\$rjags_output: rjags::coda.samples output for further processing. Note: Regardless the observed variable, the parameter alpha will always be used to create the curve
- ...\$model: the model used to run the Bayesian process, use writeLines to have nicely formatted terminal output

Function version

0.1.0

How to cite

Kreutzer, S., 2021. extract_TRUELight(): Extract True Light from the Camera Measurements using a Bayesian Approach. Function version 0.1.0. In: Kreutzer, S., Mittelstrass, D., 2021. RLum-STARR: Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE). R package version 0.1.0.9000-85.

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) , RLum Developer Team

References

Cunningham, A.C., Clark-Balzan, L., 2017. Overcoming crosstalk in luminescence images of mineral grains. Radiation Measurements 106, 498–505. doi:10.1016/j.radmeas.2017.06.004

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See Also

```
create_RFCurveArray, get_MCMCParameter
```

Examples

```
## list files using package external data
files <- list.files(system.file("extdata", "", package="RLumSTARR"), full.names=TRUE)

## create curve array
dat <- create_RFCurveArray(files = files)
output <-
extract_TRUELight(
data = dat,
ROI = c(4),
stepping = 10,
verbose = FALSE,
method_control = list(
    n.chain = 1,
    n.iter = 50,
    thin = 20))</pre>
```

get_MCMCParameter

Extracts a parameter from an MCMC list

Description

Short cut to extract a parameter from an MCMC list. If more processing is wanted, the 'coda' package can be used

Usage

```
get_MCMCParameter(mcmc, parameter, prob = 0.95, unlist = TRUE)
```

Arguments

mcmc coda::mcmc or coda::mcmc.list (required): input object, if created by extract_TRUELight

the correct object is extracted automatically

parameter character (required): name of the parameter to be extracted

prob numeric (with default): probability for the HPD calculation (cf. coda::HPDinterval) unlist logical (with default): if TRUE the output is a matrix of the means of the lower

and upper intervals of the parameter. If the parameter was estimated based on multiple chains, this chains are also subject to an average calculation. If FALSE

the output is a list as returned by coda::HPDinterval

Value

Returns a matrix with the parameter value or a list

Function version

0.1.0

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How to cite

Kreutzer, S., 2021. get_MCMCParameter(): Extracts a parameter from an MCMC list. Function version 0.1.0. In: Kreutzer, S., Mittelstrass, D., 2021. RLumSTARR: Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE). R package version 0.1.0.9000-85.

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) , RLum Developer Team

See Also

coda::HPDinterval

Examples

##TODO

run_ImageJ

Run ImageJ SR-RF macro

Description

The script runs the SR-RF ImageJ macro in batch mode out of R

Usage

```
run_ImageJ(
 path,
 RF_nat = "default",
 RF_reg = "default",
 bg_rm = "take_from_RF_reg",
  image_group_size = 5,
  image_alignment = TRUE,
 first_slices_rm = FALSE,
 noise_tolorance = 10,
 ROI_size = 10,
 center_x = 0.5,
 center_y = 0.5,
 diameter = 0.9,
 use_predefined_ROIs = FALSE,
 channel_time = 5,
  save_workflow_images = FALSE,
  save_additional_results = FALSE,
  save_signal_decay_videos = FALSE,
 offset_time = 0,
  .ImageJ = "/Applications/Fiji.app/Contents/MacOS/ImageJ-macosx"
```

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Arguments

path character (required): path to files to be analysed character (default): name of the RF_nat file RF_nat RF_reg character (default): name of the RF_reg file bg_rm character (with default): background subtraction options. Allowed are none (no background subtraction), take_from_RF_reg (takes the last 100 channels from the RF_reg signal: dangerous) or <your file name> (this does not work in batch mode) image_group_size numeric (with default): grouping value for running median to remove outliers image_alignment logical (with default): enable/disable image alignment first_slices_rm logical (with default): remove first slice of each curve set noise_tolorance numeric (with default): noise tolerance parameter ROI_size numeric (with default): ROI size in pixel numeric (with default): aliquot ROI centre x-coordinate center_x numeric (with default): aliquot ROI centre y-coordinate center_y diameter numeric (with default): relative diameter aliquot ROI use_predefined_ROIs logical (with default): use pre-defined ROIs imported from a file ROIs.zip found in the same folder as the files channel_time numeric (with default): channel time, this parameter was set the moment the sequence was written save_workflow_images

logical (with default): enable/disable writing of additional workflow images

save_additional_results

logical (with default): enable/disable writing of additional workflow images

 ${\tt save_signal_decay_videos}$

logical (with default): enable/disable writing of additional workflow videos

. ImageJ numeric (with default): Path to ImageJ (the macro is shipped with the package)

Value

This functions returns the path of the analysed data

Function version

0.1.0

How to cite

Kreutzer, S., 2021. run_ImageJ(): Run ImageJ SR-RF macro. Function version 0.1.0. In: Kreutzer, S., Mittelstrass, D., 2021. RLumSTARR: Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE). R package version 0.1.0.9000-85.

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) , RLum Developer Team

Examples

##TODO

run_TRUELightExtraction

Run Automated TRUE-Light Extraction

Description

Runs [extract_TRUELight] automatically in a parallel processing mode using [parallel::mclapply].

Usage

```
run_TRUELightExtraction(
  data,
  ROI,
  stepping = 1,
  mc.cores = max(c(1, parallel::detectCores() - 2)),
  method_control = list(),
  include_jags_output = FALSE,
  verbose = TRUE
)
```

Arguments

data [array] (**required**): object created by [create_RFCurveArray]

ROI [numeric] (*optional*): ROIs to be analysed, if nothing is given all ROIs are

analysed, however, the first ROIS is discarded!

stepping [numeric] (*with default*): the stepping parameter from and to be passed to

[extract_TRUELight]

mc.cores [numeric] (*with default*): number of cores used for the processing, passed to

[parallel::mclapply]

method_control [list] (*optional*): parameters to be passed to [extract_TRUELight]

include_jags_output

[logical] (*with default*): allows to include the output from JAGS as info object in the [Luminescence::RLum.Analysis-class] output objects for further diagnostics. Setting this option to 'TRUE' is not recommended for large datasets, since it will tremendously inflate the size of output and consume a lot of memory. If in-depth diagnostics are required, the function [extract_TRUELight] is recom-

mended instead.

verbose [logical] (*with default*): enable/disable verbose mode. The output of the

MCMC sampling using 'rjags' is **always** silent.

Value

The output is a [list] containing [Luminescence::RLum.Analysis-class] objects with two [Luminescence::RLum.Data.Curclass] objects for 'RF_nat' and 'RF_reg' respectively.

Function version

0.1.0

How to cite

Kreutzer, S., 2021. run_TRUELightExtraction(): Run Automated TRUE-Light Extraction. Function version 0.1.0. In: Kreutzer, S., Mittelstrass, D., 2021. RLumSTARR: Spatially Resolved Radiofluorescence Analysis (EXPERIMENTAL PACKAGE). R package version 0.1.0.9000-85.

Author(s)

Sebastian Kreutzer, Geography & Earth Sciences, Aberystwyth University (United Kingdom) , RLum Developer Team

Examples

```
## Not run:
## list files using package external data
files <- list.files(system.file("extdata", "", package="RLumSTARR"), full.names=TRUE)
## create curve array
dat <- create_RFCurveArray(files = files)
output <- run_TRUELightExtraction(
data = dat,
mc.cores = 1,
ROI = 5,
verbose = TRUE,
method_control = list(
    n.chain = 1,
    n.iter = 50,
    thin = 20))
## End(Not run)</pre>
```

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