

Package ‘itraxR’

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Type Package

Title Itrax Data Analysis Tools

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BugReports <https://github.com/tombishop1/itraxR/issues>

Description Parse, trim, join, visualise and analyse data from Itrax sediment core multi-parameter scanners manufactured by Cox Analytical Systems, Sweden. Functions are provided for parsing XRF-peak area files, line-scan optical images, and radiographic images, alongside accompanying metadata. A variety of data wrangling tasks like trimming, joining and reducing XRF-peak area data are simplified. Principle component analysis (PCA), cluster analysis and associated multivariate methods are implemented with appropriate data transformation.

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Encoding UTF-8

RoxygenNote 7.1.1

LazyData true

Depends dplyr, ggplot2, ggfortify, grid, R (>= 3.5.0)

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R topics documented:

CD166_19_S1	2
itrax_correlation	2

itrax_image	3
itrax_import	4
itrax_join	5
itrax_meta	6
itrax_ordination	6
itrax_radiograph	7
itrax_reduce	8
itrax_restspectra	9
itrax_section	10

Index	11
--------------	-----------

CD166_19_S1	<i>Itrax core scanner data for core CD166_19_S1</i>
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Description

A dataset containing XRF, optical and radiographic images and metadata for ocean core CD166_19_S1.

Usage

CD166_19_S1

Format

A list containing the following:

xrf a tibble of scan variables

rgb a list containing the optical image matrix and associated metadata

rad a list containing the radiographic image matrix and associated metadata

@source Wynn, R. B., and B. T. Cronin. 2005. "RRS "Charles Darwin" Cruise CD166, 29 Oct - 22 Nov 2004. Sedimentary processes and deposits in the Agadir Basin and Gulf of Cadiz." 59. Vol. 44.

itrax_correlation	<i>Calculate a correlation matrix for Itrax result data</i>
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Description

Calculates a correlation matrix for Itrax data results including normalisation and visualisation

Usage

```
itrax_correlation(  
  dataframe,  
  elementsonly = TRUE,  
  zeros = "addone",  
  transform = TRUE,  
  plot = FALSE  
)
```

Arguments

dataframe	pass the name of a dataframe parsed using "itrax_import()" or "itrax_join()"
elementsonly	if TRUE, only chemical elements are included. If FALSE, the data is passed unfiltered, otherwise a character vector of desired variable names can be supplied
zeros	if "addone", adds one to all values. If "limit", replaces zero values with 0.001. Otherwise a function can be supplied to remove zero values.
transform	binary operator that if TRUE will center-log-transform the data, if FALSE will leave the data untransformed. Otherwise, a function can be supplied to transform the data.
plot	set to true if a biplot is required as a side-effect

Value

a correlation matrix object

Examples

```
itrax_correlation(CD166_19_S1$xf, plot = TRUE)
```

itrax_image

Read an Itrax Image File

Description

Reads an Itrax image file and trims it according to the metadata provided.

Usage

```
itrax_image(  
  file = "optical.tif",  
  meta = "document.txt",  
  plot = FALSE,  
  trim = TRUE  
)
```

Arguments

file	defines the name of the datafile to parse
meta	defines the relating metadata
plot	would you like to create a plot as a side-effect?
trim	defines custom trim parameters. The default behaviour uses the limits from the metadata file. Set the false for no trimming, or set the position limits by passing a two element vector.

Value

a matrix of RGB values, and the relevant data from the metadata file relating to the image.

Examples

```
itrax_image(file = system.file("extdata",
                               "CD166_19_S1_optical_lowres.tif",
                               package = "itraxR",
                               mustWork = TRUE),
            meta = system.file("extdata",
                               "CD166_19_S1_xrf_document.txt",
                               package = "itraxR",
                               mustWork = TRUE),
            plot = TRUE)
```

itrax_import

Import Itrax core-scanner result file

Description

Imports and parses data from a results file created by Q-Spec software, part of the Itrax core scanner.

Usage

```
itrax_import(
  filename = "Results.txt",
  depth_top = NA,
  trim_top = 0,
  trim_bottom = 0,
  parameters = "some"
)
```

Arguments

filename	defines the name of the datafile to parse
depth_top	defines the coring in depth of the top of the core, in mm
trim_top	defines the length of any trimming required of data at the top of the core, in mm
trim_bottom	defines the length of any trimming required at the bottom of the core, in mm
parameters	one of all (leave all parameters), some (remove some less useful parameters)

Value

a tibble of the parsed Itrax data

Examples

```
itrax_import(
  filename = system.file("extdata",
                        "CD166_19_S1_Results.txt",
                        package = "itraxR",
                        mustWork = TRUE),
  depth_top = 0)
```

itrax_join	<i>Join two or more Itrax result datasets</i>
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Description

Join two or more Itrax datasets that have been parsed using "itrax_import()"

Usage

```
itrax_join(list)
```

Arguments

list	a list of dataframes that are parsed Itrax result files — this should have been imported using "itrax_import()" and must have a depth variable present. This should take the form "list(core1 = core1, core2 = core2)"
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Value

a tibble of all the input data

Examples

```
itrax_join(list(core1 = CD166_19_S1$xrf, core2 = CD166_19_S1$xrf))
```

itrax_meta	<i>Parse Itrax scan metadata</i>
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Description

Parses the "document.txt files" generated from Itrax core scanners

Usage

```
itrax_meta(datafile = "document.txt")
```

Arguments

datafile a "document.txt files" generated from an Itrax core scanner

Value

a dataframe of all the parsed input data

Examples

```
itrax_meta(system.file("extdata",  
                      "CD166_19_S1_xrf_document.txt",  
                      package = "itraxR",  
                      mustWork = TRUE))
```

itrax_ordination	<i>Principle Component Analysis on Itrax scan data</i>
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Description

Performs and visualises principle component analysis data from Itrax result data

Usage

```
itrax_ordination(  
  dataframe,  
  elementsonly = TRUE,  
  zeros = "addone",  
  transform = TRUE,  
  return = "list",  
  plot = FALSE  
)
```

Arguments

dataframe	pass the name of a dataframe parsed using "itrax_import()" or "itrax_join()"
elementsonly	if TRUE, only chemical elements are included. If FALSE, the data is passed unfiltered, otherwise a character vector of desired variable names can be supplied
zeros	if "addone", adds one to all values. If "limit", replaces zero values with 0.001. Otherwise a function can be supplied to remove zero values.
transform	binary operator that if TRUE will center-log-transform the data, if FALSE will leave the data untransformed. Otherwise, a function can be supplied to transform the data.
return	if "pca" the output of prcomp() is returned, otherwise "list" is a list including the transformed data, sample scores, and the output of prcomp().
plot	set to true if a biplot is required as a side-effect

Value

either an output of prcomp(), or a list including the input data

Examples

```
itrax_ordination(CD166_19_S1$xrf, plot = TRUE)
```

itrax_radiograph	<i>Read an Itrax Radiograph File</i>
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Description

Reads an Itrax radiograph file and trims it according to the metadata provided.

Usage

```
itrax_radiograph(
  file = "radiograph.tif",
  meta = "document.txt",
  plot = FALSE,
  trim = TRUE
)
```

Arguments

file	defines the name of the datafile to parse
meta	defines the relating metadata
plot	would you like to create a plot as a side-effect?
trim	defines positions of the trim if required, input is a vector with min and max positions

Value

a matrix of RGB values, and the relevant data from the metadata file relating to the image. Also computes the aspect ratio of the image.

Examples

```
itrax_radiograph(file = system.file("extdata",
  "CD166_19_S1_radiograph_adj.tif",
  package = "itraxR",
  mustWork = TRUE),
  meta = system.file("extdata",
  "CD166_19_S1_rad_document.txt",
  package = "itraxR",
  mustWork = TRUE),
  plot = TRUE)
```

 itrax_reduce

Reduce Itrax XRF data

Description

Reduces Itrax XRF data into arbitrary chunks using an arbitrary function. This is useful when making direct comparisons between the Itrax XRF data and some other data collected at a lower resolution.

Usage

```
itrax_reduce(
  dataframe,
  names = c(1:length(breaks_lower)),
  breaks_lower,
  breaks_upper,
  fun = mean,
  edges = c(">=", "<"),
  by = NULL
)
```

Arguments

dataframe	defines the name of the XRF data to reduce, usually a itraxR::itrax_import like tibble
names	optional, a vector of the same length as 'breaks'
breaks_lower	a vector of the lower limit of each chunk
breaks_upper	a vector of the upper limit of each chunk
fun	the function to apply in order to reduce the data. Default is mean(), but sd() is also common

edges	a vector of length 2 with the upper and lower bound behaviour; can be any of '<', '<=', '>', '>='
by	if contiguous samples of even sizes are required, 'by' defines the chunk size and will automatically generate 'breaks'

Value

a tibble with the same number of rows as 'breaks' and the same number of columns as 'dataframe'

Examples

```
itrax_reduce(dataframe = CD166_19_S1$xf, by = 10)
```

itrax_restspectra *Make a spectrograph from raw Itrax data spectra files*

Description

Parses a folder full of raw spectra files from an Itrax core scanner and produces a spectral graph of all the data by position

Usage

```
itrax_restspectra(
  foldername = "XRF data",
  datapos = 37,
  plot = TRUE,
  trans = "pseudo_log"
)
```

Arguments

foldername	defines the folder where the spectra "*.spe" files are located
datapos	defines the row at which spectral data begins in the files
plot	TRUE/FALSE, selects whether to create a plot as a side-effect
trans	transformation applied in the plot - see '?ggplot2::scales_colour_gradient()' for options

Value

a dataframe of all the spectral data

Examples

```
## Not run: itrax_restspectra("~/itraxBook/CD166_19_(2020)/CD166_19_S1/CD166_19_S1/XRF data")
```

itrax_section *Cluster analysis and statistical grouping of Itrax data*

Description

Performs a cluster analysis and automatic statistical grouping of parsed Itrax results data to n groups. Also provides information on the most "representative" (central) of each group. These can be used to develop a sub-sampling regime for calibration using another method.

Usage

```
itrax_section(
  dataframe,
  divisions = 30,
  elementsonly = TRUE,
  zeros = "addone",
  transform = TRUE,
  plot = FALSE
)
```

Arguments

dataframe	pass the name of a dataframe parsed using "itrax_import()" or "itrax_join()"
divisions	the number of groups to slice into - also the number of representative samples returned
elementsonly	if TRUE, only chemical elements are included. If FALSE, the data is passed unfiltered, otherwise a character vector of desired variable names can be supplied
zeros	if "addone", adds one to all values. If "limit", replaces zero values with 0.001. Otherwise a function can be supplied to remove zero values.
transform	binary operator that if TRUE will center-log-transform the data, if FALSE will leave the data untransformed. Otherwise, a function can be supplied to transform the data.
plot	set to true if a summary plot is required as a side-effect - the input dataset must have a depth or position variable - depth is used preferentially.

Value

either an output of prcomp(), or a list including the input data

Examples

```
itrax_section(CD166_19_S1$xrf, plot = TRUE)
```

Index

* datasets

CD166_19_S1, [2](#)

CD166_19_S1, [2](#)

itrax_correlation, [2](#)

itrax_image, [3](#)

itrax_import, [4](#)

itrax_join, [5](#)

itrax_meta, [6](#)

itrax_ordination, [6](#)

itrax_radiograph, [7](#)

itrax_reduce, [8](#)

itrax_restspectra, [9](#)

itrax_section, [10](#)