Package 'gamma'

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Title Dose Rate Estimation from in-Situ Gamma-Ray Spectrometry Measurements

Version 1.0.3

Description Process in-situ Gamma-Ray Spectrometry for Luminescence Dating. This package allows to import, inspect and correct the energy shifts of Gamma-ray spectra. It provides methods for estimating the gamma dose rate by the use of a calibration curve as described in Mercier and Falguères (2007). The package only supports Canberra CNF and TKA files.

License GPL-3

URL https://crp2a.github.io/gamma/, https://github.com/crp2a/gamma

BugReports https://github.com/crp2a/gamma/issues

Depends R (>= 3.5)

Imports ggplot2, graphics, IsoplotR, methods, rlang, rxylib, stats, tools, utils

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Collate 'AllClasses.R' 'AllGenerics.R' 'baseline.R'

'baseline_linear.R' 'baseline_rubberband.R' 'baseline_snip.R'

'coerce.R' 'data.R' 'dose_fit.R' 'dose_predict.R'

'energy_calibrate.R' 'gamma-package.R' 'initialize.R'

'mutators.R' 'operators.R' 'peaks_find.R' 'peaks_search.R'

'plot.R' 'read.R' 'show.R' 'signal_integrate.R'

'signal_slice.R' 'signal_split.R' 'signal_stabilize.R'

'smooth.R' 'smooth_rectangular.R' 'smooth_savitzky.R'

'smooth_triangular.R' 'subset.R' 'summarize.R' 'utilities.R'

'validate.R' 'zzz.R'

NeedsCompilation no

2 R topics documented:

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gamma-package	gamma: Dose Rate Estimation from in-Situ Gamma-Ray Spectrometry
	Measurements

Description

Process in-situ Gamma-Ray Spectrometry for Luminescence Dating. This package allows to import, inspect and correct the energy shifts of Gamma-ray spectra. It provides methods for estimating the gamma dose rate by the use of a calibration curve as described in Mercier and Falguères (2007). The package only supports Canberra CNF and TKA files.

Details

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See Also

Useful links:

```
• https://crp2a.github.io/gamma/
```

- https://github.com/crp2a/gamma
- Report bugs at https://github.com/crp2a/gamma/issues

AIX_NaI_1

CEREGE Calibration Curve (NaI)

Description

CEREGE Calibration Curve (NaI)

Usage

```
data(AIX_NaI_1)
```

Format

An object of class CalibrationCurve.

Laboratory CEREGE

Instrument Canberra Inspector 1000

Detector NaI

Authors CEREGE Luminescence Team

See Also

```
Other datasets: BDX_LaBr_1, clermont
```

```
## Load the curve
utils::data(AIX_NaI_1, package = "gamma")
plot(AIX_NaI_1)
```

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baseline

Baseline Estimation and Removal

Description

Baseline Estimation and Removal

Usage

```
signal_baseline(object, ...)
signal_correct(object, ...)
baseline_snip(object, ...)
baseline_rubberband(object, ...)
baseline_linear(object, ...)
## S4 method for signature 'GammaSpectrum'
signal_baseline(object, method = c("SNIP", "rubberband", "linear"), ...)
## S4 method for signature 'GammaSpectra'
signal_baseline(object, method = c("SNIP", "rubberband", "linear"), ...)
## S4 method for signature 'GammaSpectrum'
signal_correct(object, method = c("SNIP", "rubberband", "linear"), ...)
## S4 method for signature 'GammaSpectra'
signal_correct(object, method = c("SNIP", "rubberband", "linear"), ...)
## S4 method for signature 'GammaSpectrum'
baseline_linear(object, from = NULL, to = NULL)
## S4 method for signature 'GammaSpectra'
baseline_linear(object, from = NULL, to = NULL)
## S4 method for signature 'GammaSpectrum'
baseline_rubberband(object, noise = 0, spline = TRUE, ...)
## S4 method for signature 'GammaSpectra'
baseline_rubberband(object, noise = 0, spline = TRUE, ...)
## S4 method for signature 'GammaSpectrum'
baseline_snip(object, LLS = FALSE, decreasing = FALSE, n = 100, ...)
## S4 method for signature 'GammaSpectra'
baseline_snip(object, LLS = FALSE, decreasing = FALSE, n = 100, ...)
```

Arguments

object A GammaSpectrum or GammaSpectra object.

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	Extra parameters to be passed to further methods.
method	A character string specifying the method to be used for baseline estimation (see details). Any unambiguous substring can be given.
from	An integer giving the first channel to be used for linear interpolation. If NULL (the default), channel 1 is used. Only used if method is "linear".
to	An integer giving the last channel to be used for linear interpolation. If NULL (the default), channel max is used. Only used if method is "linear".
noise	A length-one numeric vector giving the noise level. Only used if method is "rubberband".
spline	A logical scalar: should spline interpolation through the support points be used instead of linear interpolation? Only used if method is "rubberband".
LLS	A logical scalar: should the LLS operator be applied on x before employing SNIP algorithm? Only used if method is "SNIP".
decreasing	A logical scalar: should a decreasing clipping window be used? Only used if method is "SNIP".
n	An integer value giving the number of iterations. Only used if method is "SNIP".

Details

The following methods are available for baseline estimation:

SNIP Sensitive Nonlinear Iterative Peak clipping algorithm.

rubberband A convex envelope of the spectrum is determined and the baseline is estimated as the part of the convex envelope lying below the spectrum. Note that the rubber band does not enter the concave regions (if any) of the spectrum.

linear Linear baseline estimation.

Value

- baseline_*() returns a BaseLine object.
- signal_correct() returns a corrected GammaSpectrum or GammaSpectra object (same as object).

Note

baseline_rubberband() is slightly modified from C. Beleites' hyperSpec::spc.rubberband().

Author(s)

N. Frerebeau

References

Liland, K. H. (2015). 4S Peak Filling - baseline estimation by iterative mean suppression. *MethodsX*, 2, 135-140. doi:10.1016/j.mex.2015.02.009.

Morháč, M., Kliman, J., Matoušek, V., Veselský, M. & Turzo, I. (1997). Background elimination methods for multidimensional gamma-ray spectra. *Nuclear Instruments and Methods in Physics Research Section A: Accelerators, Spectrometers, Detectors and Associated Equipment*, 401(1), p. 113-132. doi:10.1016/S01689002(97)010231

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Morháč, M. & Matoušek, V. (2008). Peak Clipping Algorithms for Background Estimation in Spectroscopic Data. *Applied Spectroscopy*, 62(1), p. 91-106. doi:10.1366/000370208783412762

Ryan, C. G., Clayton, E., Griffin, W. L., Sie, S. H. & Cousens, D. R. (1988). SNIP, a statistics-sensitive background treatment for the quantitative analysis of PIXE spectra in geoscience applications. *Nuclear Instruments and Methods in Physics Research Section B: Beam Interactions with Materials and Atoms*, 34(3), p. 396-402. doi:10.1016/0168583X(88)900638

See Also

```
Other signal processing: peaks_find(), peaks_search(), signal_integrate(), signal_slice(), signal_split(), signal_stabilize(), smooth()
```

Examples

```
## Import a CNF file
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")</pre>
spc <- read(spc_file)</pre>
## Remove the first 35 channels
spc <- signal_slice(spc, -c(1:35))</pre>
## Linear baseline
bsl_linear <- baseline_linear(spc, from = 250, to = 750)
plot(spc, bsl_linear)
## SNIP baseline
bsl_snip <- baseline_snip(spc, LLS = FALSE, decreasing = FALSE, n = 100)</pre>
plot(spc, bsl_snip)
## Rubberband baseline
bsl_rubber <- baseline_rubberband(spc)</pre>
plot(spc, bsl_rubber)
## Remove baseline
spc_clean1 <- signal_correct(spc)</pre>
spc_clean2 <- spc - bsl_snip</pre>
all(spc_clean1 == spc_clean2)
plot(spc_clean1)
```

Baseline-class

An S4 Class to Represent a Spectrum Baseline

Description

An S4 Class to Represent a Spectrum Baseline

Note

This class extends the GammaSpectrum class.

Author(s)

N. Frerebeau

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See Also

Other class: CalibrationCurve-class, GammaSpectra-class, GammaSpectrum-class, PeakPosition-class, coerce()

Examples

```
## Import a CNF file
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")</pre>
spc <- read(spc_file)</pre>
## Remove the first 35 channels
spc <- signal_slice(spc, -c(1:35))</pre>
## Linear baseline
bsl_linear <- baseline_linear(spc, from = 250, to = 750)</pre>
plot(spc, bsl_linear)
## SNIP baseline
bsl_snip <- baseline_snip(spc, LLS = FALSE, decreasing = FALSE, n = 100)</pre>
plot(spc, bsl_snip)
## Rubberband baseline
bsl_rubber <- baseline_rubberband(spc)</pre>
plot(spc, bsl_rubber)
## Remove baseline
spc_clean1 <- signal_correct(spc)</pre>
spc_clean2 <- spc - bsl_snip</pre>
all(spc_clean1 == spc_clean2)
plot(spc_clean1)
```

BDX_LaBr_1

CRP2A Calibration Curve (LaBr)

Description

CRP2A Calibration Curve (LaBr)

Usage

```
data(BDX_LaBr_1)
```

Format

An object of class CalibrationCurve.

Laboratory IRAMAT-CRP2A (UMR 5060) **Instrument** Canberra Inspector 1000

Detector LaBr

Authors CRP2A Luminescence Team

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See Also

```
Other datasets: AIX_NaI_1, clermont
```

Examples

```
## Load the curve
utils::data(BDX_LaBr_1, package = "gamma")
plot(BDX_LaBr_1)
```

CalibrationCurve-class

An S4 class to Represent a Dose Rate Calibration Curve

Description

An S4 class to Represent a Dose Rate Calibration Curve

Slots

```
Ni A DoseRateModel object.

NiEi A DoseRateModel object.

details A list of length-one vector giving the curve metadata.

slope A numeric vector.

intercept A numeric vector.

covariance A numeric vector.

MSWD A numeric vector.

df A numeric vector.

p_value A numeric vector.

data A data.frame.

range A numeric vector.

background A numeric vector.
```

Subset

In the code snippets below, x is a CalibrationCurve object.

x[[i]] Extracts information from a slot selected by subscript i. i is a character vector of length one.

Author(s)

N. Frerebeau

See Also

```
Other \ class; \ Baseline-class, Gamma Spectra-class, Gamma Spectrum-class, Peak Position-class, coerce()
```

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clermont

Clermont Reference Data

Description

Clermont Reference Data

Usage

```
data("clermont")
```

Format

TODO

Source

Guérin, G., Mercier, N. & Adamiec, G. (2011). Dose-Rate Conversion Factors: Update. *Ancient TL*, 29(1), p. 5-8.

Miallier, D., Guérin, G., Mercier, N., Pilleyre, T. & Sanzelle, S. (2009). The Clermont Radiometric Reference Rocks: A Convenient Tool for Dosimetric Purposes. *Ancient TL*, 27(2), p. 37-44.

See Also

Other datasets: AIX_NaI_1, BDX_LaBr_1

coerce

Coerce

Description

Coerce

Usage

```
## S3 method for class 'GammaSpectrum'
as.matrix(x, ...)
```

Arguments

x An object to be coerced.

... Currently not used.

Value

A coerced object.

Author(s)

N. Frerebeau

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See Also

 $Other class: \verb|Baseline-class|, CalibrationCurve-class|, GammaSpectra-class|, GammaSpectrum-class|, PeakPosition-class|$

Examples

```
## Import a Canberra CNF file
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")
spc <- read(spc_file)

## Coerce
mtx <- as.matrix(spc)
df <- as.data.frame(spc)
head(df)</pre>
```

decay

Nuclear Decay Data

Description

An internal dataset containing the decay data for the following isotopes: 232-Th, 235-U, 238-U and 40-K.

Format

```
A data. frame with the following columns (absolute errors):
```

```
decay_chain
isotope
occurrence
occurrence_error
post_radon
half_life
half_life_error
energy Energy in keV.
energy_error
gamma_intensity
gamma_intensity_error
counts_chain
counts_chain_error
```

Source

Nuclides And Isotopes: Chart of the Nuclides.

doserate doserate

decay_La

Lanthanum Decay Data

Description

An internal dataset containing the decay data for the Lanthanum isotopes.

Format

```
A data.frame with the following columns (absolute errors):

decay_chain
isotope
occurrence
occurrence_error
post_radon
half_life
half_life_error
energy Energy in keV.
energy_error
gamma_intensity
```

Source

Nuclides And Isotopes: Chart of the Nuclides.

 ${\bf gamma_intensity_error}$

doserate

Dose Rate Estimation

Description

dose_fit builds a calibration curve for gamma dose rate estimation.

Usage

```
dose_fit(object, background, doses, ...)

dose_predict(object, spectrum, ...)

## S4 method for signature 'GammaSpectra,GammaSpectrum,matrix'
dose_fit(
   object,
   background,
   doses,
   range_Ni,
   range_NiEi,
```

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```
details = list(authors = "", date = Sys.time())
)
## S4 method for signature 'GammaSpectra, GammaSpectrum, data.frame'
dose_fit(
  object,
  background,
  doses,
  range_Ni,
  range_NiEi,
  details = list(authors = "", date = Sys.time())
)
## S4 method for signature 'CalibrationCurve, missing'
dose_predict(object, sigma = 1, epsilon = 0)
## S4 method for signature 'CalibrationCurve, GammaSpectrum'
dose_predict(object, spectrum, sigma = 1, epsilon = 0)
## S4 method for signature 'CalibrationCurve, GammaSpectra'
dose_predict(object, spectrum, sigma = 1, epsilon = 0)
```

Arguments

object A GammaSpectra or CalibrationCurve object.

background A GammaSpectrum object of a length-two numeric vector giving the back-

ground noise integration value and error, respectively.

doses A matrix or data.frame TODO.

... Currently not used.

spectrum An optional GammaSpectrum or GammaSpectra object in which to look for

variables with which to predict. If omitted, the fitted values are used.

range_Ni, range_NiEi

A length-two numeric vector giving the energy range to integrate within (in

keV).

details A list of length-one vector specifying additional informations about the instru-

ment for which the curve is built.

sigma A numeric value giving TODO.

epsilon A numeric value giving an extra error term introduced by the calibration of the

energy scale of the spectrum.

Details

dose_predict predicts in situ gamma dose rate.

To estimate the gamma dose rate, one of the calibration curves distributed with this package can be used. These built-in curves are in use in several luminescence dating laboratories and can be used to replicate published results. As these curves are instrument specific, the user may have to build its own curve.

The construction of a calibration curve requires a set of reference spectra for which the gamma dose rate is known and a background noise measurement. First, each reference spectrum is integrated

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over a given interval, then normalized to active time and corrected for background noise. The dose rate is finally modelled by the integrated signal value used as a linear predictor (York *et al.*, 2004). See vignette(doserate) for a reproducible example.

Value

- dose_fit() returns a CalibrationCurve object.
- dose_predict() returns a data.frame with the following columns:

```
name (character) the name of the spectra.
```

- *_signal (numeric) the integrated signal value (according to the value of threshold; see signal_integrate()).
- *_error (numeric) the integrated signal error value (according to the value of threshold; see signal_integrate()).

```
gamma_signal (numeric) the predicted gamma dose rate.
gamma_error (numeric) the predicted gamma dose rate error.
```

Author(s)

N. Frerebeau

References

Mercier, N. & Falguères, C. (2007). Field Gamma Dose-Rate Measurement with a NaI(Tl) Detector: Re-Evaluation of the "Threshold" Technique. *Ancient TL*, 25(1), p. 1-4.

York, D., Evensen, N. M., Martínez, M. L. & De Basabe Delgado, J. (2004). Unified Equations for the Slope, Intercept, and Standard Errors of the Best Straight Line. *American Journal of Physics*, 72(3), p. 367-75. doi:10.1119/1.1632486.

See Also

```
signal_integrate()
```

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```
plot(calib_curve, threshold = "Ni")
## Estimate gamma dose rates
dose_predict(calib_curve, spc)
```

energy

Energy Scale Calibration

Description

Calibrates the energy scale of a gamma spectrum.

Usage

```
energy_calibrate(object, lines, ...)
has_energy(object)

## S4 method for signature 'GammaSpectrum,list'
energy_calibrate(object, lines, ...)

## S4 method for signature 'GammaSpectrum,PeakPosition'
energy_calibrate(object, lines, ...)

## S4 method for signature 'GammaSpectrum'
has_energy(object)

## S4 method for signature 'GammaSpectrum'
has_energy(object)

## S4 method for signature 'GammaSpectrum'
has_calibration(object)

## S4 method for signature 'GammaSpectrum'
has_calibration(object)
```

Arguments

object	A GammaSpectrum or GammaSpectra object.
lines	A PeakPosition object or a list of length two. If a list is provided, each element must be a named numeric vector giving the observed peak position ("channel") and the corresponding expected "energy" value (in keV).
	Currently not used.

Details

The energy calibration of a spectrum is the most tricky part. To do this, the user must specify the position of at least three observed peaks and the corresponding energy value (in keV). A second

order polynomial model is fitted on these energy vs channel values, then used to predict the new energy scale of the spectrum.

The package allows to provide the channel-energy pairs to be use. However, the spectrum can be noisy so it is difficult to properly determine the peak channel. In this case, a better approach may be to pre-process the spectrum (variance-stabilization, smoothing and baseline correction) and perform a peak detection. Once the identified peaks are satisfactory, you can set the corresponding energy values (in keV) and use these lines to calibrate the energy scale of the spectrum.

Regardless of the approach you choose, it is strongly recommended to check the result before proceeding.

Value

- energy_calibrate() returns a GammaSpectrum object.
- has_energy() and has_calibration() return a logical vector.

Author(s)

N. Frerebeau

Examples

```
## Import a CNF file
spc_file <- system.file("extdata/LaBr.TKA", package = "gamma")
(spc <- read(spc_file))

## Set peak positions (channel) and expected energy values
calib_lines <- list(
    channel = c(86, 495, 879),
    energy = c(238, 1461, 2615)
)

## Adjust the energy scale
(spc1 <- energy_calibrate(spc, lines = calib_lines))

## Inspect results
plot(spc1, xaxis = "energy", yaxis = "count") +
    ggplot2::geom_vline(xintercept = c(238, 1461, 2615), linetype = 3)</pre>
```

GammaSpectra-class

An S4 Class to Represent a Collection of Gamma Sectra

Description

Represents a collection of spectra of gamma ray spectrometry measurements.

Details

This class extends the base list and can only contains GammaSpectrum objects.

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Access

```
In the code snippets below, x is a GammaSpectra object.

length(x) Get the number of elements in x.

lengths(x) Get the number of channels in each element of x.

get_names(x), set_names(x) <- value Retrieves or sets the names of x according to value.

get_hash(x) Get the MD5 hash of the raw data files.

get_channels(x) Get the number of channels in each element of x.

get_counts(x) Get the counts of each element of x.

get_energy(x) Get the energy range of each element of x.

get_rates(x) Get the count rates of each element of x.
```

Subset

In the code snippets below, x is a GammaSpectra object.

- x[i] Extracts the elements selected by subscript i. i can be missing or NULL, numeric or character vector or a factor. Returns a new GammaSpectra object.
- x[i, j] Like the above but allows to select a slot thru j (see examples). j is a character vector of length one. Returns a list.
- x[[i]] Extracts the elements selected by subscript i. i can be a numeric or character vector of length one. Returns the corresponding GammaSpectrum object.

Author(s)

N. Frerebeau

See Also

 $Other \ class; Baseline-class, Calibration Curve-class, Gamma Spectrum-class, Peak Position-class, coerce()$

```
## Import all CNF files in a given directory
spc_dir <- system.file("extdata/BDX_LaBr_1/calibration", package = "gamma")
(spc <- read(spc_dir))

## Access
get_hash(spc)
get_names(spc)
get_livetime(spc)
get_realtime(spc)

lengths(spc)
range_energy(spc)

## Subset
spc[] # All spectra
spc[NULL] # All spectra
spc[1] # The first spectrum
spc[-6] # Delete the sixth spectrum</pre>
```

```
spc[1:3] # The first three spectra
spc[c(1, 3)] # The first and third spectra
spc["BRIQUE"] # The spectrum named 'BRIQUE'
spc[c("BRIQUE", "C347")] # The spectra named 'BRIQUE' and 'C347'
spc[1:3, "energy"] # The slot 'energy' of the first three spectra
spc[[1]]
spc[["BRIQUE"]]
```

GammaSpectrum-class

An S4 Class to Represent a Gamma Sectrum

Description

Represents a single spectrum of a gamma ray spectrometry measurement.

Slots

```
hash A character string giving the 32-byte MD5 hash of the imported file.

name A character string the measurement reference.

date A POSIXct element giving the measurement date and time.

instrument A character string giving the instrument name.

file_format A character string giving the format of the imported file.

live_time A numeric value.

real_time A numeric value.

channel A integer vector giving the channel number. Numeric values are coerced to integer as by as.integer() (and hence truncated towards zero).

energy A numeric vector giving the gamma ray's energy (in keV).

count A numeric vector giving the counts number for each channel. Numeric values are coerced to integer as by as.integer() (and hence truncated towards zero).

rate A numeric vector the count rate (1/s) for each channel.

calibration A linear model used for energy scale calibration (see energy_calibrate()).
```

Access

```
In the code snippets below, x is a GammaSpectrum object.

length(x) Get number of channel in x.

get_hash(x) Get the MD5 hash of the raw data file.

get_names(x), set_names(x) <- value Retrieves or sets the name of x according to value.

get_channels(x) Get the number of channels in x.

get_counts(x) Get the counts of x.

get_energy(x) Get the energy range of x.

get_rates(x) Get the count rates of x.
```

Coerce

In the code snippets below, x is a GammaSpectrum object.

```
as.matrix(x) Coerces x to a matrix.
as.data.frame(x) Coerces x to a data.frame.
```

Subset

In the code snippets below, x is a GammaSpectrum object.

x[[i]] Extracts information from a slot selected by subscript i. i is a character vector of length one and will be matched to the name of the slots.

Note

This class retains copy construction.

Author(s)

N. Frerebeau

See Also

 $Other \ class; \ Baseline-class, \ Calibration \ Curve-class, \ Gamma Spectra-class, \ Peak Position-class, \ coerce()$

```
## Import a Canberra CNF file
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")
(spc <- read(spc_file))

## Access
get_hash(spc)
get_names(spc)
get_livetime(spc)
get_realtime(spc)

length(spc)
range_energy(spc)

## Subset
spc[["date"]]
spc[["instrument"]]
spc[["file_format"]]</pre>
```

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mutator

Get or Set Parts of an Object

Description

Getters and setters to extract or replace parts of an object.

Usage

```
get_hash(x)
get_names(x)
set_names(x) \leftarrow value
get_livetime(x)
get_realtime(x)
get_channels(x)
get_counts(x)
get_rates(x)
get_energy(x, ...)
set_energy(x, ...) \leftarrow value
get_method(x)
set_method(x) <- value</pre>
get_residuals(x)
range\_channels(x, ...)
range\_energy(x, ...)
## S4 method for signature 'GammaSpectrum'
length(x)
## S4 method for signature 'GammaSpectrum'
get_hash(x)
## S4 method for signature 'GammaSpectrum'
get_names(x)
## S4 method for signature 'GammaSpectrum'
get_livetime(x)
```

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```
## S4 method for signature 'GammaSpectrum'
get_realtime(x)
## S4 method for signature 'GammaSpectrum'
get_channels(x)
## S4 method for signature 'GammaSpectrum'
get_counts(x)
## S4 method for signature 'GammaSpectrum'
get_rates(x)
## S4 method for signature 'GammaSpectrum'
get_energy(x)
## S4 method for signature 'GammaSpectrum'
range\_energy(x, na.rm = FALSE)
## S4 method for signature 'GammaSpectrum'
range_channels(x, na.rm = FALSE)
## S4 method for signature 'Baseline'
get_method(x)
## S4 method for signature 'GammaSpectra'
get_hash(x)
## S4 method for signature 'GammaSpectra'
get_names(x)
## S4 method for signature 'GammaSpectra'
get_livetime(x)
## S4 method for signature 'GammaSpectra'
get_realtime(x)
## S4 method for signature 'GammaSpectra'
get_channels(x)
## S4 method for signature 'GammaSpectra'
get_counts(x)
## S4 method for signature 'GammaSpectra'
get_rates(x)
## S4 method for signature 'GammaSpectra'
get_energy(x)
## S4 method for signature 'GammaSpectra'
range\_energy(x, na.rm = FALSE)
## S4 method for signature 'GammaSpectra'
```

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```
range_channels(x, na.rm = FALSE)
## S4 method for signature 'DoseRateModel'
get_residuals(x)
## S4 method for signature 'PeakPosition'
get_hash(x)
## S4 method for signature 'PeakPosition'
get_channels(x)
## S4 method for signature 'PeakPosition'
get_energy(x, expected = FALSE)
## S4 replacement method for signature 'GammaSpectrum'
set_names(x) \leftarrow value
## S4 replacement method for signature 'Baseline'
set_method(x) \leftarrow value
## S4 replacement method for signature 'GammaSpectra'
set_names(x) \leftarrow value
## S4 replacement method for signature 'PeakPosition, numeric'
set_energy(x, expected = TRUE) <- value</pre>
```

Arguments

x An object from which to get or set element(s).

value A possible value for the element(s) of x.

... Currently not used.

na.rm A logical scalar: should NA be omitted?

expected TODO.

Value

An object of the same sort as x with the new values assigned.

Author(s)

N. Frerebeau

See Also

Other mutator: subset()

operator 23

operator

Common Operations on GammaSpectrum Objects

Description

Performs common operations on GammaSpectrum objects.

Usage

```
## S4 method for signature 'GammaSpectrum, GammaSpectrum'
Arith(e1, e2)
## S4 method for signature 'GammaSpectrum, numeric'
Arith(e1, e2)
## S4 method for signature 'GammaSpectrum, GammaSpectrum'
Compare(e1, e2)
## S4 method for signature 'GammaSpectrum, numeric'
Compare(e1, e2)
## S4 method for signature 'GammaSpectrum, GammaSpectrum'
Logic(e1, e2)
## S4 method for signature 'GammaSpectrum,numeric'
Logic(e1, e2)
## S4 method for signature 'GammaSpectrum,logical'
Logic(e1, e2)
## S4 method for signature 'GammaSpectrum'
Math(x)
## S4 method for signature 'GammaSpectrum'
Math2(x, digits)
## S4 method for signature 'GammaSpectrum'
Summary(x, ..., na.rm = FALSE)
```

Arguments

x, e1, e2	An object (typically a GammaSpectrum object).
digits	A length-one numeric vector giving the number of digits to be used in round() or signif().
	Further arguments passed to or from methods.
na.rm	A logical scalar: should missing values (including NaN) be omitted from the calculations?

24 PeakPosition-class

Group Generics

GammaSpectrum objects have support for S4 group generic functionality to operate within elements across objects:

Author(s)

N. Frerebeau

Examples

No examples

PeakPosition-class

An S4 Class to Represent a Set of Peaks

Description

An S4 Class to Represent a Set of Peaks

Slots

```
hash A character string giving the 32-byte MD5 hash of the imported spectrum file.

noise_method A character string specifying the method used for peak detection.

noise_threshold A length one numeric vector giving the noise threshold.

window A length one numeric vector giving the half-window size.

channel A integer vector giving the channel number. Numeric values are coerced to integer as by as.integer() (and hence truncated towards zero).

energy_observed A numeric vector giving the observed gamma ray energy (in keV).

energy_expected A numeric vector giving the expected gamma ray energy (in keV).
```

Access

```
In the code snippets below, x is a PeakPosition object.
get_hash(x) Get the MD5 hash of the raw data file.
get_channels(x) Get the channels of x.
get_energy(x), set_energy(x) <- value Retrieves or sets the energy scale of x according to value.</pre>
```

peaks_find 25

Coerce

In the code snippets below, x is a PeakPosition object.

```
as.matrix(x) Coerces x to a matrix.
as.data.frame(x) Coerces x to a data.frame.
```

Subset

In the code snippets below, x is a PeakPosition object.

x[[i]] Extracts information from a slot selected by subscript i. i is a character vector of length one and will be matched to the name of the slots.

Note

This class retains copy construction.

Author(s)

N. Frerebeau

See Also

Other class: Baseline-class, CalibrationCurve-class, GammaSpectra-class, GammaSpectrum-class, coerce()

peaks_find

Find Peaks

Description

Finds local maxima in sequential data.

Usage

```
peaks_find(object, ...)
## S4 method for signature 'GammaSpectrum'
peaks_find(object, method = c("MAD"), SNR = 2, span = NULL, ...)
```

Arguments

object	A GammaSpectrum object.
	Extra parameters to be passed to internal methods.
method	A character string specifying the method to be used for background noise estimation (see below).
SNR	An integer giving the signal-to-noise-ratio for peak detection (see below).
span	An integer giving the half window size (in number of channels). If NULL, 5\ window size.

26 peaks_search

Details

A local maximum has to be the highest one in the given window and has to be higher than $SNR \times noise$ to be recognized as peak.

The following methods are available for noise estimation:

MAD Median Absolute Deviation.

Value

A PeakPosition object.

Author(s)

N. Frerebeau

See Also

```
Other signal processing: baseline, peaks_search(), signal_integrate(), signal_slice(), signal_split(), signal_stabilize(), smooth()
```

Examples

```
## Import a LaBr spectrum
LaBr_file <- system.file("extdata/LaBr.TKA", package = "gamma")
LaBr_spc <- read(LaBr_file)

## Find peaks by channel
(LaBr_pks <- peaks_find(LaBr_spc)) # Ugly
plot(LaBr_spc, LaBr_pks)

## Search peaks by channel
(LaBr_pks <- peaks_search(LaBr_spc, index = c(86L, 207L, 496L), span = 7))
plot(LaBr_spc, LaBr_pks, split = TRUE)

## Import a BEGe spectrum
BEGe_file <- system.file("extdata/BEGe.CNF", package = "gamma")
BEGe_spc <- read(BEGe_file)

## Search peaks by energy
(BEGe_pks <- peaks_search(BEGe_spc, index = c(47, 63, 911, 1460)))
plot(BEGe_spc, BEGe_pks, split = TRUE)</pre>
```

peaks_search

Search Peaks

Description

Search the maxima in sequential data around a given value.

peaks_search 27

Usage

```
peaks_search(object, index, ...)
## S4 method for signature 'GammaSpectrum,integer'
peaks_search(object, index, span = 10, tolerance = 0.025)
## S4 method for signature 'GammaSpectrum,numeric'
peaks_search(object, index, span = 10, tolerance = 0.025)
```

Arguments

object A GammaSpectrum object.

index A vector giving the expected peak position. If index is a numeric vector, peaks

are searched by energy (index is assumed to be expressed in keV). If index is

an integer vector, peaks are searched by channel.

... Currently not used.

span A numeric value giving the half window size for searching. If index is a

numeric vector, span is expressed in keV. If index is an integer vector, span

is expressed in channel.

tolerance A numeric value giving the threshold above which a warning/error is raised.

Value

A PeakPosition object.

Author(s)

N. Frerebeau

See Also

```
Other signal processing: baseline, peaks_find(), signal_integrate(), signal_slice(), signal_split(), signal_stabilize(), smooth()
```

```
## Import a LaBr spectrum
LaBr_file <- system.file("extdata/LaBr.TKA", package = "gamma")
LaBr_spc <- read(LaBr_file)

## Find peaks by channel
(LaBr_pks <- peaks_find(LaBr_spc)) # Ugly
plot(LaBr_spc, LaBr_pks)

## Search peaks by channel
(LaBr_pks <- peaks_search(LaBr_spc, index = c(86L, 207L, 496L), span = 7))
plot(LaBr_spc, LaBr_pks, split = TRUE)

## Import a BEGe spectrum
BEGe_file <- system.file("extdata/BEGe.CNF", package = "gamma")
BEGe_spc <- read(BEGe_file)

## Search peaks by energy</pre>
```

28 plot

```
(BEGe_pks <- peaks_search(BEGe_spc, index = c(47, 63, 911, 1460))) plot(BEGe_spc, BEGe_pks, split = TRUE)
```

plot

Plot

Description

Plot

Usage

```
## S4 method for signature 'GammaSpectrum, missing'
plot(x, xaxis = c("channel", "energy"), yaxis = c("count", "rate"), ...)
## S4 method for signature 'GammaSpectrum, Baseline'
plot(x, y, xaxis = c("channel", "energy"), yaxis = c("count", "rate"), ...)
## S4 method for signature 'GammaSpectra, missing'
plot(
  х,
  xaxis = c("channel", "energy"),
  yaxis = c("count", "rate"),
  select = NULL,
  facet = FALSE,
  nrow = c("fixed", "auto")
## S4 method for signature 'GammaSpectrum, PeakPosition'
plot(x, y, split = FALSE, span = 25)
## S4 method for signature 'CalibrationCurve, missing'
plot(
  х,
  error_ellipse = TRUE,
  error_bar = FALSE,
  energy = FALSE,
  level = 0.95,
  n = 50,
)
```

Arguments

```
    x, y
    Objects to be plotted.
    xaxis, yaxis
    A character string specifying the data to be plotted along each axis. It must be one of "energy" or "channel" (x axis) and "counts" or "rate" (y axis). Any unambiguous substring can be given.
    Currently not used.
    Select
    A numeric or character vector giving the selection of the spectrum that are drawn.
```

plot 29

A logical scalar: should a matrix of panels defined by spectrum be drawn? facet A character string specifying the number of rows. It must be one of "fixed" nrow or "auto". Any unambiguous substring can be given. Only used if facet is TRUE. split A logical scalar: should. An integer giving the half window size (in number of channels). Only used if span split is TRUE. error_ellipse A logical scalar: should error ellipses be plotted? error_bar A logical scalar: should error bars be plotted? A logical scalar: TODO. energy level length-one numeric vector giving the the probability cutoff for the error ellipses.

A length-one numeric vector giving the resolution of the error ellipses.

Value

n

A ggplot2::ggplot object.

Author(s)

N. Frerebeau

See Also

```
IsoplotR::ellipse(), IsoplotR::isochron()
```

```
# Import CNF files
spc_dir <- system.file("extdata/BDX_LaBr_1/calibration", package = "gamma")
spc <- read(spc_dir)

# Plot all spectra
plot(spc, yaxis = "rate", facet = FALSE) +
    ggplot2::theme_bw()

# Plot the spectrum named 'BRIQUE'
plot(spc, xaxis = "energy", yaxis = "count", select = "BRIQUE") +
    ggplot2::theme_bw()

# Plot the first three spectra
plot(spc, xaxis = "channel", yaxis = "rate", select = 1:3, facet = TRUE) +
    ggplot2::theme_bw()</pre>
```

30 read

read

Data Input

Description

Reads a gamma ray spectrum file.

Usage

```
read(file, ...)
## S4 method for signature 'character'
read(file, extensions = c("cnf", "tka"), ...)
```

Arguments

file A character string giving the path of files to be imported.

... Extra parameters to be passed to rxylib::read_xyData().

extensions A character vector specifying the possible file extensions. It must be one or more of "cnf", "tka".

Value

A GammaSpectra object if more than one spectrum are imported at once, else a GammaSpectrum object.

Note

Only supports Canberra CNF and TKA files.

Author(s)

N. Frerebeau

See Also

```
rxylib::read_xyData()
Other IO: summarise()
```

```
## Import a Canberra CNF file
cnf_file <- system.file("extdata/LaBr.CNF", package = "gamma")
(cnf_spc <- read(cnf_file))

## Import a TKA file
tka_file <- system.file("extdata/LaBr.TKA", package = "gamma")
(tka_spc <- read(tka_file))

## Import all files in a given directory
spc_dir <- system.file("extdata/BDX_LaBr_1/calibration", package = "gamma")
(spc <- read(spc_dir))</pre>
```

signal_integrate 31

signal_integrate Signal Integration

Description

Signal Integration

Usage

```
signal_integrate(object, background, ...)
## S4 method for signature 'GammaSpectrum, missing'
signal_integrate(object, range = NULL, energy = FALSE)
## S4 method for signature 'GammaSpectrum, GammaSpectrum'
signal_integrate(object, background, range = NULL, energy = FALSE)
## S4 method for signature 'GammaSpectrum, numeric'
signal_integrate(object, background, range = NULL, energy = FALSE)
## S4 method for signature 'GammaSpectra, missing'
signal_integrate(object, range = NULL, energy = FALSE, simplify = TRUE)
## S4 method for signature 'GammaSpectra, GammaSpectrum'
signal_integrate(
  object,
  background,
  range = NULL,
  energy = FALSE,
  simplify = TRUE
)
## S4 method for signature 'GammaSpectra, numeric'
signal_integrate(
  object,
  background,
  range = NULL,
  energy = FALSE,
  simplify = TRUE
)
```

Arguments

object A GammaSpectrum or GammaSpectra object.

background A GammaSpectrum object.

... Currently not used.

range A length-two numeric vector giving the energy range to integrate within (in

keV).

energy A logical scalar: TODO?

simplify A logical scalar: should the result be simplified to a matrix? The default

value, FALSE, returns a list.

32 signal_slice

Details

It assumes that each spectrum has an energy scale.

Value

If simplify is FALSE (the default) returns a list of numeric vectors (the signal value and its error), else returns a matrix.

Author(s)

N. Frerebeau

References

Guérin, G. & Mercier, M. (2011). Determining Gamma Dose Rates by Field Gamma Spectroscopy in Sedimentary Media: Results of Monte Carlo Simulations. *Radiation Measurements*, 46(2), p. 190-195. doi:10.1016/j.radmeas.2010.10.003.

Mercier, N. & Falguères, C. (2007). Field Gamma Dose-Rate Measurement with a NaI(Tl) Detector: Re-Evaluation of the "Threshold" Technique. *Ancient TL*, 25(1), p. 1-4.

See Also

```
Other signal processing: baseline, peaks_find(), peaks_search(), signal_slice(), signal_split(), signal_stabilize(), smooth()
```

signal_slice

Choose channels by Position

Description

Choose channels by position.

Usage

```
signal_slice(object, ...)
## S4 method for signature 'GammaSpectrum'
signal_slice(object, ...)
## S4 method for signature 'GammaSpectra'
signal_slice(object, ...)
```

Arguments

object A GammaSpectrum or GammaSpectra object.

... integer values giving the channels of the spectrum to be kept/dropped (see below). Numeric values are coerced to integer as by as.integer() (and hence truncated towards zero).

signal_split 33

Details

Either positive values to keep, or negative values to drop, should be provided. The values provided must be either all positive or all negative.

If no value is provided, an attempt is made to define the number of channels to skip at the beginning of the spectrum. This drops all channels before the highest count maximum. This is intended to deal with the artefact produced by the rapid growth of random background noise towards low energies.

Value

A GammaSpectrum or GammaSpectra object.

Author(s)

N. Frerebeau

See Also

```
Other signal processing: baseline, peaks_find(), peaks_search(), signal_integrate(), signal_split(), signal_stabilize(), smooth()
```

Examples

```
## Import CNF files
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")</pre>
spc <- read(spc_file)</pre>
## Plot spectrum
plot(spc)
## Slice
sliced <- signal_slice(spc)</pre>
plot(sliced)
sliced <- signal_slice(spc, -c(1:35))</pre>
plot(sliced)
sliced <- signal_slice(sliced, 450:550)</pre>
plot(sliced)
## Split
g <- rep(c("A", "B", "C"), c(250, 500, 274))
splited <- signal_split(spc, g)</pre>
plot(splited, facet = TRUE)
```

signal_split

Split

Description

Split

34 signal_split

Usage

```
signal_split(object, ...)
## S4 method for signature 'GammaSpectrum'
signal_split(object, groups)
```

Arguments

object A GammaSpectrum object.

... Currently not used.
groups A a factor in the sense that as.factor(groups) defines the grouping (see split).

Value

A GammaSpectra object.

Author(s)

N. Frerebeau

See Also

```
Other signal processing: baseline, peaks_find(), peaks_search(), signal_integrate(), signal_slice(), signal_stabilize(), smooth()
```

```
## Import CNF files
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")</pre>
spc <- read(spc_file)</pre>
## Plot spectrum
plot(spc)
## Slice
sliced <- signal_slice(spc)</pre>
plot(sliced)
sliced <- signal_slice(spc, -c(1:35))</pre>
plot(sliced)
sliced <- signal_slice(sliced, 450:550)</pre>
plot(sliced)
## Split
g <- rep(c("A", "B", "C"), c(250, 500, 274))
splited <- signal_split(spc, g)</pre>
plot(splited, facet = TRUE)
```

signal_stabilize 35

Description

Transform Intensities

Usage

```
signal_stabilize(object, ...)
## S4 method for signature 'GammaSpectrum'
signal_stabilize(object, f, ...)
## S4 method for signature 'GammaSpectra'
signal_stabilize(object, f, ...)
```

Arguments

object	A GammaSpectrum object.
	Extra arguments to be passed to f.
f	A function that takes a numeric vector as argument and returns a numeric vector.

Details

The stabilization step aims at improving the identification of peaks with a low signal-to-noise ratio. This particularly targets higher energy peaks.

Value

A GammaSpectrum or GammaSpectra object with transformed intensities.

Author(s)

N. Frerebeau

See Also

```
Other signal processing: baseline, peaks_find(), peaks_search(), signal_integrate(), signal_slice(), signal_split(), smooth()
```

36 smooth

smooth Smooth

Description

Smoothes intensities.

Usage

```
signal_smooth(object, ...)
smooth_rectangular(object, ...)
smooth_triangular(object, ...)
smooth_savitzky(object, ...)
## S4 method for signature 'GammaSpectrum'
signal_smooth(object, method = c("rectangular", "triangular", "savitzky"), ...)
## S4 method for signature 'GammaSpectra'
signal_smooth(object, method = c("rectangular", "triangular", "savitzky"), ...)
## S4 method for signature 'GammaSpectrum'
smooth_rectangular(object, m = 3, ...)
## S4 method for signature 'GammaSpectra'
smooth_rectangular(object, m = 3, ...)
## S4 method for signature 'GammaSpectrum'
smooth_savitzky(object, m = 3, p = 2, ...)
## S4 method for signature 'GammaSpectra'
smooth_savitzky(object, m = 3, p = 2, ...)
## S4 method for signature 'GammaSpectrum'
smooth_triangular(object, m = 3, ...)
## S4 method for signature 'GammaSpectra'
smooth_triangular(object, m = 3, ...)
```

Arguments

object	A GammaSpectrum or GammaSpectra object.
• • •	Extra parameters to be passed to further methods.
method	A character string specifying the smoothing method to be used. It must be one of "unweighted" (default), "weighted" or "savitzky" (see details). Any unambiguous substring can be given.
m	An odd integer giving the number of adjacent points to be used.
р	An integer giving the polynomial degree. Only used if method is "savitzky".

smooth 37

Details

The following smoothing methods are available:

rectangular Unweighted sliding-average or rectangular smooth. It replaces each point in the signal with the average of m adjacent points.

triangular Weighted sliding-average or triangular smooth. It replaces each point in the signal with the weighted mean of m adjacent points.

savitzky Savitzky-Golay filter. This method is based on the least-squares fitting of polynomials to segments of m adjacent points.

There will be (m-1)/2 points both at the beginning and at the end of the spectrum for which a complete m-width smooth cannot be calculated. To prevent data loss, progressively smaller smooths are used at the ends of the spectrum if method is unweighted or weighted. If the Savitzky-Golay filter is used, the original (m-1)/2 points at the ends of the spectrum are preserved.

Value

A GammaSpectrum or GammaSpectra object.

Author(s)

N. Frerebeau

References

Gorry, P. A. (1990). General Least-Squares Smoothing and Differentiation by the Convolution (Savitzky-Golay) Method. *Analytical Chemistry*, 62(6), p. 570-573. doi:10.1021/ac00205a007.

Savitzky, A. & Golay, M. J. E. (1964). Smoothing and Differentiation of Data by Simplified Least Squares Procedures. *Analytical Chemistry*, 36(8), p. 1627-1639. doi:10.1021/ac60214a047.

See Also

```
Other signal processing: baseline, peaks_find(), peaks_search(), signal_integrate(), signal_slice(), signal_split(), signal_stabilize()
```

```
# Import CNF files
spc_file <- system.file("extdata/LaBr.CNF", package = "gamma")
spc <- read(spc_file)
spc <- signal_slice(spc, -c(1:35))

# Plot raw spectrum
spc_clean <- signal_correct(spc)
plot(spc_clean)

# Rectangular smooth
spc_unweighted <- smooth_rectangular(spc, m = 3)
spc_unweighted_clean <- signal_correct(spc_unweighted)
plot(spc_unweighted_clean)

# Triangular smooth
spc_weighted <- smooth_triangular(spc, m = 5)
spc_weighted_clean <- signal_correct(spc_weighted)</pre>
```

38 subset

```
plot(spc_weighted_clean)

# Savitzky-Golay
spc_savitzky <- smooth_savitzky(spc, m = 21, p = 2)
spc_savitzky_clean <- signal_correct(spc_savitzky)
plot(spc_savitzky_clean)</pre>
```

subset

Extract or Replace Parts of an Object

Description

Operators acting on objects to extract or replace parts.

Usage

```
## S4 method for signature 'GammaSpectrum'
x[[i]]
## S4 method for signature 'GammaSpectra'
x[i, j]
## S4 method for signature 'DoseRateModel'
x[[i]]
## S4 method for signature 'CalibrationCurve'
x[[i]]
## S4 method for signature 'PeakPosition'
x[[i]]
```

Arguments

x An object from which to extract element(s) or in which to replace element(s).

i, j Indices specifying elements to extract or replace. Indices are numeric, integer or character vectors or empty (missing) or NULL. Numeric values are coerced to integer as by as.integer() (and hence truncated towards zero). Character vectors will be matched to the name of the elements. An empty index (a comma separated blank) indicates that all entries in that dimension are selected.

Value

A subsetted object.

Author(s)

N. Frerebeau

See Also

Other mutator: mutator

summarise 39

summarise

Summarize

Description

Summarize

Usage

```
summarise(object, ...)
## S4 method for signature 'GammaSpectrum'
summarise(object)
## S4 method for signature 'GammaSpectra'
summarise(object)
## S4 method for signature 'DoseRateModel'
summarise(object)
## S4 method for signature 'CalibrationCurve'
summarise(object)
```

Arguments

object A GammaSpectrum or GammaSpectra object.
... Currently not used.

Value

A data.frame.

Author(s)

N. Frerebeau

See Also

Other IO: read()

```
## Import a Canberra CNF file
cnf_file <- system.file("extdata/LaBr.CNF", package = "gamma")
spc <- read(cnf_file)
summarise(spc)

## Import all CNF files in a given directory
spc_dir <- system.file("extdata/BDX_LaBr_1/calibration", package = "gamma")
spc <- read(spc_dir)
summarise(spc)</pre>
```

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