Package 'powdR'

August 8, 2022

```
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Maintainer Benjamin Butler <br/> benjamin.butler@hutton.ac.uk>
Description Full pattern summation of X-ray powder diffraction data as
      described in Chipera and Bish (2002) <doi:10.1107/S0021889802017405> and
      Butler and Hillier (2021) <doi:10.1016/j.cageo.2020.104662>.
      Derives quantitative estimates of crystalline and amorphous phase
      concentrations in complex mixtures.
License GPL-2 | file LICENSE
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afps

Automated full pattern summation

Description

afps returns estimates of phase concentrations using automated full pattern summation of X-ray powder diffraction data. It is designed for high-throughput cases involving mineral quantification from large reference libraries.

Usage

```
afps(
  lib,
  smpl,
  harmonise,
  solver,
  obj,
  refs,
  std,
  force,
  std_conc,
  omit\_std,
  closed,
  normalise,
  tth_align,
  align,
  manual_align,
  shift,
  tth_fps,
  lod,
  amorphous,
  amorphous_lod,
  weighting,
)
```

Arguments

1ib A powdRlib object representing the reference library. Created using the powdRlib

constructor function.

smpl A data frame. First column is 2theta, second column is counts

harmonise logical parameter defining whether to harmonise the lib and smpl. Default

= TRUE. Harmonises to the intersecting 2theta range at the coarsest resolution

available using natural splines.

solver The optimisation routine to be used. One of "BFGS", "Nelder-Mead", or "CG".

Default = "BFGS".

obj The objective function to minimise. One of "Delta", "R", "Rwp". Default =

"Rwp". See Chipera and Bish (2002) and page 247 of Bish and Post (1989) for

definitions of these functions.

refs A character string of reference pattern IDs or names from the specified library.

The IDs or names supplied must be present within the lib $phasesphase_id$ or lib $phasesphase_name$ columns. If missing from the function call then all

phases in the reference library will be used.

std The phase ID (e.g. "QUA.1") to be used as internal standard. Must match an ID

provided in the refs parameter.

force An optional string of phase ID's or names specifying which phases should be

forced to remain throughout the automated full pattern summation. The ID's or

names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name

columns.

std_conc The concentration of the internal standard (if known) in weight percent. If un-

known then either omit the argument from the function call of use std_conc = NA, in which case it will be assumed that all phases sum to 100 percent (default).

omit_std A logical parameter to be used when the std_conc argument is defined. When

omit_std = TRUE the phase concentrations are recomputed to account for value

supplied in std_conc. Default = FALSE.

closed A logical parameter to be used when the std_conc argument is defined and

omit_std = TRUE. When closed = TRUE the internal standard concentration is removed and the remaining phase concentrations closed to sum to 100 percent.

Default = FALSE.

normalise deprecated. Please use the omit_std and closed arguments instead.

tth_align A vector defining the minimum and maximum 2theta values to be used during

alignment (e.g. c(5,65)). If not defined, then the full range is used.

align The maximum shift that is allowed during initial 2theta alignment (degrees).

Default = 0.1.

manual_align A logical operator denoting whether to optimise the alignment within the nega-

tive/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. De-

fault = FALSE, i.e. alignment is optimised.

shift A single numeric value denoting the maximum (positive or negative) shift, in

degrees 2theta, that is allowed during the shifting of selected phases. Default =

0.

tth_fps A vector defining the minimum and maximum 2theta values to be used during

automated full pattern summation (e.g. c(5,65)). If not defined, then the full

range is used.

lod Optional parameter used to define the limit of detection (in weight percent) of

the internal standard (i.e. the phase provided in the std argument). The lod value is used to estimate the lod of other phases during the fitting process and hence remove reference patterns that are considered below detection limit. De-

fault = 0.1. If lod = 0 then limits of detection are not computed.

amorphous A character string of any phase IDs that should be treated as amorphous. These

must match phases present in lib\$phases\$phase_id.

amorphous_lod Optional parameter used to exclude amorphous phases if they are below this

specified limit (percent). Must be between 0 and 100. Default = 0.

weighting an optional 2 column data frame specifying the 2theta values in the first column

and a numeric weighting vector in the second column that specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) when minimising the objective function defined in the obj argument. Use this weighting parameter with caution. The default is simply a weighting vector where all values are 1,

which hence has no effect on the computed objective function.

... Other parameters passed to methods e.g. afps.powdRlib

Details

Applies automated full pattern summation to an XRPD measurement to quantify phase concentrations. Requires a powdRlib library of reference patterns with reference intensity ratios in order to derive mineral concentrations. Details provided in Butler and Hillier (2021).

Value

a powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of fitted XRPD pattern

measured a vector of the count intensities of original XRPD measurement (aligned)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped and summed by phase_name

obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

References

Butler, B. M., Hillier, S., 2021.powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Comp. Geo. 147, 104662. doi:10.1016/j.cageo.2020.104662

Chipera, S.J., Bish, D.L., 2013. Fitting Full X-Ray Diffraction Patterns for Quantitative Analysis: A Method for Readily Quantifying Crystalline and Disordered Phases. Adv. Mater. Phys. Chem. 03, 47-53. doi:10.4236/ampc.2013.31A007

Chipera, S.J., Bish, D.L., 2002. FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. J. Appl. Crystallogr. 35, 744-749. doi:10.1107/S0021889802017405

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

```
#Load the minerals library
data(minerals)
# Load the soils data
data(soils)
## Not run:
afps_sand <- afps(lib = minerals,</pre>
                 smpl = soils$sandstone,
                 std = "QUA.2",
                 align = 0.2,
                 lod = 0.2,
                 amorphous = "ORG",
                 amorphous_lod = 1)
afps_lime <- afps(lib = minerals,</pre>
                smpl = soils$limestone,
                std = "QUA.2",
                align = 0.2,
                lod = 0.2,
                 amorphous = "ORG",
                amorphous_lod = 1)
afps_granite <- afps(lib = minerals,</pre>
                   smpl = soils$granite,
                    std = "QUA.2",
                    align = 0.2,
                    lod = 0.2,
                    amorphous = "ORG",
                    amorphous_lod = 1)
#Alternatively run all 3 at once using lapply
afps_soils <- lapply(soils, afps,</pre>
                      lib = minerals,
                      std = "QUA.2",
                      align = 0.2,
                      lod = 0.2,
                      amorphous = "ORG",
                      amorphous_lod = 1)
#Automated quantification using the rockjock library
data(rockjock)
data(rockjock_mixtures)
#This takes a few minutes to run
rockjock_a1 <- afps(lib = rockjock,</pre>
                     smpl = rockjock_mixtures$Mix1,
                     std = "CORUNDUM",
                     align = 0.3,
                     lod = 1)
```

afps.powdRlib

Automated full pattern summation

Description

afps returns estimates of phase concentrations using automated full pattern summation of X-ray powder diffraction data. It is designed for high-throughput cases involving mineral quantification from large reference libraries.

Usage

```
## S3 method for class 'powdRlib'
afps(
  lib,
  smpl,
  harmonise,
  solver,
  obj,
  refs,
  std,
  force,
  std_conc,
  omit_std,
  closed,
  normalise,
  tth_align,
  align,
  manual_align,
  shift,
  tth_fps,
  lod,
  amorphous,
  amorphous_lod,
  weighting,
)
```

Arguments

lib A powdRlib object representing the reference library. Created using the powdRlib

constructor function.

smpl A data frame. First column is 2theta, second column is counts

harmonise logical parameter defining whether to harmonise the lib and smpl. Default

= TRUE. Harmonises to the intersecting 2theta range at the coarsest resolution

available using natural splines.

solver The optimisation routine to be used. One of "BFGS", "Nelder-Mead", or "CG".

Default = "BFGS".

obj The objective function to minimise. One of "Delta", "R", "Rwp". Default =

"Rwp". See Chipera and Bish (2002) and page 247 of Bish and Post (1989) for

definitions of these functions.

refs A character string of reference pattern IDs or names from the specified library.

The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. If missing from the function call then all

phases in the reference library will be used.

std The phase ID (e.g. "QUA.1") to be used as internal standard. Must match an ID

provided in the refs parameter.

force An optional string of phase ID's or names specifying which phases should be

forced to remain throughout the automated full pattern summation. The ID's or

names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name

columns.

known then either omit the argument from the function call of use std_conc = NA, in which case it will be assumed that all phases sum to 100 percent (default).

omit_std A logical parameter to be used when the std_conc argument is defined. When

omit_std = TRUE the phase concentrations are recomputed to account for value

supplied in std_conc. Default = FALSE.

closed A logical parameter to be used when the std_conc argument is defined and

omit_std = TRUE. When closed = TRUE the internal standard concentration is removed and the remaining phase concentrations closed to sum to 100 percent.

Default = FALSE.

normalise deprecated. Please use the omit_std and closed arguments instead.

tth_align A vector defining the minimum and maximum 2theta values to be used during

alignment (e.g. c(5,65)). If not defined, then the full range is used.

align The maximum shift that is allowed during initial 2theta alignment (degrees).

Default = 0.1.

manual_align A logical operator denoting whether to optimise the alignment within the nega-

tive/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. De-

fault = FALSE, i.e. alignment is optimised.

shift A single numeric value denoting the maximum (positive or negative) shift, in

degrees 2theta, that is allowed during the shifting of selected phases. Default =

0.

tth_fps A vector defining the minimum and maximum 2theta values to be used during

automated full pattern summation (e.g. c(5,65)). If not defined, then the full

range is used.

lod Optional parameter used to define the limit of detection (in weight percent) of

the internal standard (i.e. the phase provided in the std argument). The lod value is used to estimate the lod of other phases during the fitting process and hence remove reference patterns that are considered below detection limit. De-

fault = 0.1. If lod = 0 then limits of detection are not computed.

amorphous A character string of any phase IDs that should be treated as amorphous. These

must match phases present in lib\$phases\$phase_id.

amorphous_lod Optional parameter used to exclude amorphous phases if they are below this

specified limit (percent). Must be between 0 and 100. Default = 0.

weighting an optional 2 column data frame specifying the 2theta values in the first column

and a numeric weighting vector in the second column that specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) when minimising the objective function defined in the obj argument. Use this weighting parameter with caution. The default is simply a weighting vector where all values are 1,

which hence has no effect on the computed objective function.

... other arguments

Details

Applies automated full pattern summation to an XRPD measurement to quantify phase concentrations. Requires a powdRlib library of reference patterns with reference intensity ratios in order to derive mineral concentrations. Details provided in Butler and Hillier (2021).

Value

a powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of fitted XRPD pattern

measured a vector of the count intensities of original XRPD measurement (aligned)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped and summed by phase_name

obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

References

Butler, B. M., Hillier, S., 2021.powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Comp. Geo. 147, 104662. doi:10.1016/j.cageo.2020.104662

Bish, D.L., Post, J.E., 1989. Modern powder diffraction. Mineralogical Society of America.

Chipera, S.J., Bish, D.L., 2013. Fitting Full X-Ray Diffraction Patterns for Quantitative Analysis: A Method for Readily Quantifying Crystalline and Disordered Phases. Adv. Mater. Phys. Chem. 03, 47-53. doi:10.4236/ampc.2013.31A007

Chipera, S.J., Bish, D.L., 2002. FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. J. Appl. Crystallogr. 35, 744-749. doi:10.1107/S0021889802017405

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

```
#Load the minerals library
data(minerals)
# Load the soils data
data(soils)
## Not run:
afps_sand <- afps(lib = minerals,</pre>
                 smpl = soils$sandstone,
                  std = "QUA.2",
                  align = 0.2,
                  lod = 0.2,
                  amorphous = "ORG",
                  amorphous_lod = 1)
afps_lime <- afps(lib = minerals,</pre>
                 smpl = soils$limestone,
                 std = "QUA.2",
                 align = 0.2,
                 lod = 0.2,
                 amorphous = "ORG".
                amorphous_lod = 1)
afps_granite <- afps(lib = minerals,</pre>
                    smpl = soils$granite,
                    std = "QUA.2",
                    align = 0.2,
                    lod = 0.2,
                    amorphous = "ORG",
                    amorphous_lod = 1)
#Alternatively run all 3 at once using lapply
afps_soils <- lapply(soils, afps,</pre>
                      lib = minerals,
                      std = "QUA.2",
                      align = 0.2,
                      lod = 0.2,
                      amorphous = "ORG",
                      amorphous_lod = 1)
#Automated quantification using the rockjock library
data(rockjock)
data(rockjock_mixtures)
#This takes a few minutes to run
rockjock_a1 <- afps(lib = rockjock,</pre>
```

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afsis

Africa Soil Information Service (AfSIS) XRPD reference library

Description

A powdR1ib object of 21 pure reference patterns and associated reference intensity ratios for a range of common soil minerals. Data were collected on a Bruker D2 Phaser using Cu K-alpha radiation. All patterns have been normalised to 10,000 counts and reference intensity ratios transformed so that all are relative to that of corundum.

Usage

afsis

Format

A powdRlib object of 3 components

xrd A dataframe of all the count intensities of all reference patterns. Column names denote the unique phase ID of each reference pattern

tth A vector of the 2theta scale for all reference patterns in the library

phases A dataframe the phase IDs, names and reference intensity ratios (RIR)

afsis_codes

Original codes for the afsis reference patterns

Description

A data frame detailing the original codes associated with the afsis reference patterns prior to their addition to powdR.

Usage

afsis_codes

12 align_xy

Format

An 2 column data frame. First column contains the phase IDs from afsis\$phase_id and the second column the original IDs prior to the inclusion in powdR.

afsis_regroup	Regrouping structure for the Africa Soil Information Service (AfSIS)
	XRPD reference library

Description

A data frame containing an example re-grouping structure for the afsis reference library, which results in a slightly coarser description of clay minerals and Fe/Ti-(hydr)oxides in powdRfps or powdRafps objects when used with regroup().

Usage

```
afsis_regroup
```

Format

A data frame with three columns:

```
phase_id the phase IDs present in afsis$phases$phase_id.
phase_name_grouped The phase names that constitute the first regrouping structure.
phase_name_grouped2 The phase names that constitute the second regrouping structure
```

align_xy

Align XRPD data to a given standard

Description

See ?align_xy.XY and align_xy.multiXY for method-specific details.

Usage

```
align_xy(x, std, xmin, xmax, xshift, ...)
```

Arguments

X	an XY or multiXY object.
std	a dataframe of the chosen standard that each sample will be aligned to (column $1=2$ theta, column $2=$ counts)
xmin	the minimum 2theta value used during alignment
xmax	the maximum 2theta value used during alignment
xshift	the maximum (positive and negative) 2theta shift that is allowed during alignment
	other arguments

align_xy

Value

an XY or multiXY object.

```
# Load soils xrd data
data(soils)
#Load minerals library
data(minerals)
## Not run:
#Create a standard quartz pattern to align to
quartz <- data.frame(tth = minerals$tth,</pre>
                     counts = minerals$xrd$QUA.1)
#Plot the main quartz peak prior to alignment
plot(soils, wavelength = "Cu",
     xlim = c(26, 27),
     normalise = TRUE)
#align data
aligned <- align_xy(soils,</pre>
                    std = quartz,
                    xmin = 10,
                     xmax = 60,
                    xshift = 0.2)
#replot data
plot(aligned, wavelength = "Cu",
     xlim = c(26, 27),
     normalise = TRUE)
#Alternatively try with a single XY object
unaligned <- as_multi_xy(list("quartz" = quartz,</pre>
                              "sandstone" = soils$sandstone))
plot(unaligned, wav = "Cu",
     xlim = c(26,27), normalise = TRUE)
sandstone_a <- align_xy(soils$sandstone,</pre>
                         std = quartz,
                         xmin = 10,
                         xmax = 60,
                         xshift = 0.3)
aligned <- as_multi_xy(list("quartz" = quartz,</pre>
                             "sandstone" = sandstone_a))
plot(aligned, wav = "Cu",
     xlim = c(26,27), normalise = TRUE)
## End(Not run)
```

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align_xy.multiXY

Align XRPD data in a multiXY object to a given standard

Description

align_xy.multiXY takes a multiXY object and aligns each of the XY data frames within it to a given standard. An optimisation routine is used that computes a suitable linear shift. After all samples have been aligned, the function harmonises the data to a single 2theta scale.

Usage

```
## S3 method for class 'multiXY'
align_xy(x, std, xmin, xmax, xshift, ...)
```

Arguments

X	a multiXY object.	
std	a dataframe of the chosen standard that each sample will be aligned to (column $1=2$ theta, column $2=$ counts)	
xmin	the minimum 2theta value used during alignment	
xmax	the maximum 2theta value used during alignment	
xshift	the maximum (positive and negative) 2theta shift that is allowed during alignment	
	other arguments	

Value

a multiXY object.

```
# Load soils xrd data
data(soils)
#Load minerals library
data(minerals)
## Not run:
#Create a standard quartz pattern to align to
quartz <- data.frame(tth = minerals$tth,</pre>
                     counts = minerals$xrd$QUA.1)
#Plot the main quartz peak prior to alignment
plot(soils, wavelength = "Cu",
     xlim = c(26, 27),
     normalise = TRUE)
#align data
aligned <- align_xy(soils,</pre>
                     std = quartz,
                     xmin = 10,
```

align_xy.XY

align_xy.XY

Align XRPD data in an XY object to a given standard

Description

align_xy.XY takes an XY object and aligns it to a given standard. An optimisation routine is used that computes a suitable linear shift.

Usage

```
## S3 method for class 'XY'
align_xy(x, std, xmin, xmax, xshift, ...)
```

Arguments

X	an XY object.
std	a dataframe of the chosen standard that each sample is aligned to (column $1 = 2$ theta, column $2 = $ counts)
xmin	the minimum 2theta value used during alignment
xmax	the maximum 2theta value used during alignment
xshift	the maximum (positive and negative) 2theta shift that is allowed during alignment
	other arguments

Value

an XY object.

```
# Load soils xrd data
data(soils)

#Load minerals library
data(minerals)

## Not run:

#Create a standard quartz pattern to align to
quartz <- data.frame(tth = minerals$tth,</pre>
```

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as_multi_xy

Create a multiXY object

Description

as_multi_xy takes a list or data frame of XRPD data and ensures that the data meet various requirements to create a multiXY object. Once a multiXY object has been created, it can easily be plotted using the associated plot.multiXY method.

Usage

```
as_multi_xy(x, ...)
```

Arguments

x a list or data frame of XRPD data
... other arguments

Value

a multiXY object.

```
#EXAMPLE 1
#load soils data
data(soils)
#extract first two samples from the list
```

as_multi_xy.data.frame

as_multi_xy.data.frame

Create a multiXY object from a list of XRPD data

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Description

as_multi_xy.data.frame takes a data frame of XRPD data from multiple samples and ensures that it meets various requirements to create a multiXY object. Once a multiXY object has been created, it can easily be plotted using the associated plot.multiXY method.

Usage

```
## S3 method for class 'data.frame'
as_multi_xy(x, ...)
```

Arguments

x a data frame of XRPD data, with the first column as the 2theta axis and subsequent columns of count intensities.

... other arguments

Value

a multiXY object.

 as_xy

as_multi_xy.list

Create a multiXY object from a list of XRPD data

Description

as_multi_xy.list takes a list of XRPD data and ensures that they meet various requirements to create a multiXY object. These requirements include that each item in the list contains 2 columns of numeric data in a data frame. as_multi_xy.list also checks that all names are unique. Once a multiXY object has been created, it can easily be plotted using the associated plot.multiXY method.

Usage

```
## S3 method for class 'list'
as_multi_xy(x, ...)
```

Arguments

```
x a list of XRPD data frames (column 1 = 2theta, column 2 = counts)
... other arguments
```

Value

```
a multiXY object.
```

Examples

```
#' #load soils data
data(soils)

#extract first two samples from the list
soils <- soils[c(1:2)]

#convert to multiXY
soils <- as_multi_xy(soils)</pre>
```

as_xy

Create an XY object

Description

as_xy takes a data frame of XY XRPD data and ensures that it meets the criteria for an XY object. These requirements include that the data contains 2 columns of numeric data in a dataframe. Once an XY object has been created, it can easily be plotted using the associated plot.XY method.

Usage

```
as_xy(x)
```

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Arguments

```
x a data frame (column 1 = 2theta, column 2 = counts)
```

Value

an XY object.

Examples

```
# Load soils xrd data
data(rockjock_mixtures)

xy <- as_xy(rockjock_mixtures$Mix1)

class(xy)

## Not run:
plot(xy, wavelength = "Cu")
plot(xy, wavelength = "Cu", interactive = TRUE)

## End(Not run)</pre>
```

bkg

Fit a background to XRPD data

Description

bkg fits a background to X-Ray Powder Diffraction data

Usage

```
bkg(xrd, lambda, hwi, it, int)
```

Arguments

xrd an xy data frame of the data to fit a background to. First column is the 2theta

scale, second column is count intensities

lambda second derivative penalty for primary smoothing. Default = 0.5.

hwi Half width of local windows. Default = 25.

it Number of iterations in suppression loop. Default = 50.

int Number of buckets to divide the data into. Default = round(nrow(xrd)/4).

Details

A wrapper for the baseline.fillPeaks in the baseline package.

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Value

a powdRbkg object consisting of of 3 vectors

tth The 2theta axis of the measurement counts The count intensities of the measurement background The count intensities of the fitted background

Examples

```
data(soils)
## Not run:
fit_bkg <- bkg(soils$granite)
plot(bkg)
## End(Not run)</pre>
```

close_quant Close the phase concentration

Close the phase concentration data within a powdRfps or powdRafps

object

Description

close_quant closes the quantitative data within a powdRfps or powdRafps object (derived from fps() and afps(), respectively) by ensuring that the composition sums to 100 percent. See also ?close_quant.powdRfps and ?close_quant.powdRafps.

Usage

```
close_quant(x, ...)
```

Arguments

x A powdRfps or powdRafps object...

... other arguments

Value

a powdRfps or powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (fitted vs measured)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit weighted pure patterns.

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

Examples

```
## Not run:
data(rockjock)
data(rockjock_mixtures)
rockjock_1 <- fps(lib = rockjock,</pre>
                   smpl = rockjock_mixtures$Mix1,
                   refs = c("ORDERED_MICROCLINE",
                             "LABRADORITE",
                             "KAOLINITE_DRY_BRANCH",
                             "MONTMORILLONITE_WYO",
                             "ILLITE_1M_RM30",
                             "{\sf CORUNDUM"}) ,
                  std = "CORUNDUM",
                  align = 0.3,
                  std\_conc = 20)
sum(rockjock_1$phases$phase_percent)
rockjock_1c <- close_quant(rockjock_1)</pre>
sum(rockjock_1c$phases$phase_percent)
rockjock_a1 <- afps(lib = rockjock,</pre>
                     smpl = rockjock_mixtures$Mix1,
                     std = "CORUNDUM",
                     align = 0.3,
                     lod = 1,
                     std\_conc = 20)
sum(rockjock_a1$phases$phase_percent)
rockjock_a1c <- close_quant(rockjock_a1)</pre>
sum(rockjock_a1c$phases$phase_percent)
## End(Not run)
```

 ${\tt close_quant.powdRafps}$ Close the phase concentration data within a ${\tt powdRafps}$ object

Description

close_quant closes the quantitative data within a powdRafps object (derived from afps()) by ensuring that the composition sums to 100 percent.

Usage

```
## S3 method for class 'powdRafps'
close_quant(x, ...)
```

Arguments

x A powdRafps object derived from afps().

... other arguments

Value

a powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (fitted vs measured)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed

obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

close_quant.powdRfps

Description

close_quant closes the quantitative data within a powdRfps object (derived from fps()) by ensuring that the composition sums to 100 percent.

Usage

```
## S3 method for class 'powdRfps'
close_quant(x, ...)
```

Arguments

x A powdRfps object derived from fps().
... other arguments

Value

a powdRfps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (fitted vs measured)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit

 $weighted_pure_patterns$

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

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```
std = "CORUNDUM",
    align = 0.3,
    std_conc = 20)

sum(rockjock_1$phases$phase_percent)

rockjock_1c<- close_quant(rockjock_1)

sum(rockjock_1c$phases$phase_percent)

## End(Not run)</pre>
```

delta

Calculate the Delta value for a fitted pattern

Description

delta computes the absolute difference between a measured and fitted pattern. See equation for Delta in section 2.1 of Butler and Hillier (2021).

Usage

```
delta(measured, fitted, weighting)
```

Arguments

measured a vector of count intensities for a measured pattern fitted a vector of count intensities for a fitted pattern

weighting an optional weighting vector of the same length as those specified in measured

and fitted, which specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) from the calculation. Use with caution. Default is simply a weighting vector where all values are 1, which hence has no effect on the

computed value.

Value

a single numeric value

References

Butler, B.M., Hillier, S., 2021. powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Computers and Geosciences. 147, 104662. doi:10.1016/j.cageo.2020.104662

```
# Load soils xrd data
data(soils)
#Load minerals library
data(minerals)
```

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extract_xy

Import and extract XY data from proprietary files

Description

extract_xy is a wrapper for read_xyData of the rxylib package, which extracts the xy data from various proprietary formats of X-ray powder diffraction data using the xylib C++ library. For more information see ?rxylib and ?rxylib::read_xyData.

Usage

```
extract_xy(files)
```

Arguments

files

path of the file(s) to be imported.

Value

If only one path is supplied then an XY data frame with 2 columns is returned, the first being the 2theta axis and the second being the count intensities. If more than one path is supplied then a multiXY list is returned, with each item in the list being an XY data frame as already described.

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```
## End(Not run)
```

fps

Full pattern summation

Description

fps returns estimates of phase concentrations using full pattern summation of X-ray powder diffraction data.

Usage

```
fps(
  lib,
  smpl,
  harmonise,
  solver,
  obj,
  refs,
  std,
  force,
  std_conc,
  omit\_std,
  normalise,
  closed,
  tth_align,
  align,
  manual_align,
  tth_fps,
  shift,
  remove_trace,
  weighting,
```

Arguments

lib	A powdRlib object representing the reference library. Created using the powdRlib constructor function.	
smpl	A data frame. First column is 2theta, second column is counts	
harmonise	logical parameter defining whether to harmonise the lib and smpl. Default = TRUE. When TRUE the function will harmonise the lib and smpl to the intersecting 2theta range at the coarsest resolution available using natural splines.	
solver	The optimisation routine to be used. One of "BFGS", "Nelder-Mead", "CG", "NNLS". Default = "BFGS".	
obj	The objective function to minimise when "BFGS", "Nelder-Mead", or "CG" are used as the solver argument. One of "Delta", "R", "Rwp". Default = "Rwp". See Chipera and Bish (2002) and page 247 of Bish and Post (1989) for	

definitions of these functions.

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refs A character string of reference pattern IDs or names from the specified library. The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. If missing from the function call then all phases in the reference library will be used. std The phase ID (e.g. "QUA.1") to be used as an internal standard. Must match an ID provided in the refs parameter. force An optional string of phase IDs or names specifying which phases should be forced to remain throughout the automated full pattern summation. The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. std_conc The concentration of the internal standard (if known) in weight percent. If unknown then omit the argument from the function call or use std_conc = NA (default), n which case it will be assumed that all phases sum to 100 percent. A logical parameter to be used when the std_conc argument is defined. When omit_std omit_std = TRUE the phase concentrations are recomputed to account for the value supplied in std_conc. Default = FALSE. normalise deprecated. Please use the omit_std and closed arguments instead. closed A logical parameter to be used when the std_conc argument is defined and omit_std = TRUE. When closed = TRUE the internal standard concentration is removed and the remaining phase concentrations closed to sum to 100 percent. Default = FALSE. tth_align A vector defining the minimum and maximum 2theta values to be used during alignment (e.g. c(5,65)). If not defined, then the full range is used. align The maximum shift that is allowed during initial 2theta alignment (degrees). Default = 0.1. manual_align A logical operator denoting whether to optimise the alignment within the negative/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. Default = FALSE, i.e. alignment is optimised. tth_fps A vector defining the minimum and maximum 2theta values to be used during full pattern summation (e.g. c(5,65)). If not defined, then the full range is used. shift A single numeric value denoting the maximum (positive or negative) shift, in degrees 2theta, that is allowed during the shifting of reference patterns. Default A single numeric value representing the limit for the concentration of trace remove_trace phases to be retained, i.e. any mineral with an estimated concentration below remove_trace will be omitted. Default = 0. weighting an optional 2 column data frame specifying the 2theta values in the first column and a numeric weighting vector in the second column that specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) when minimising the objective function defined in the obj argument. Use this weighting parameter with caution. The default is simply a weighting vector where all values are 1, which hence has no effect on the computed objective function.

Details

Applies full pattern summation (Chipera & Bish, 2002, 2013; Eberl, 2003) to an XRPD measurement to quantify phase concentrations. Requires a powdRlib library of reference patterns with

Other parameters passed to methods e.g. fps.powdRlib

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reference intensity ratios in order to derive mineral concentrations. Details provided in Butler and Hillier (2021).

Value

a powdRfps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

References

Butler, B. M., Hillier, S., 2021.powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Comp. Geo. 147, 104662. doi:10.1016/j.cageo.2020.104662

Chipera, S.J., Bish, D.L., 2013. Fitting Full X-Ray Diffraction Patterns for Quantitative Analysis: A Method for Readily Quantifying Crystalline and Disordered Phases. Adv. Mater. Phys. Chem. 03, 47-53. doi:10.4236/ampc.2013.31A007

Chipera, S.J., Bish, D.L., 2002. FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. J. Appl. Crystallogr. 35, 744-749. doi:10.1107/S0021889802017405

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

```
fps_lime <- fps(lib = minerals,</pre>
                 smpl = soils$limestone,
                refs = minerals$phases$phase_id,
                std = "QUA.1",
                align = 0.2)
fps_granite <- fps(lib = minerals,</pre>
                    smpl = soils$granite,
                    refs = minerals$phases$phase_id,
                    std = "QUA.1",
                    align = 0.2)
#Alternatively run all 3 at once using lapply
fps_soils <- lapply(soils, fps,</pre>
                     lib = minerals,
                     std = "QUA.2",
                     refs = minerals$phases$phase_id,
                     align = 0.2)
#Using the rockjock library:
data(rockjock)
data(rockjock_mixtures)
rockjock_1 <- fps(lib = rockjock,</pre>
                   smpl = rockjock_mixtures$Mix1,
                   refs = c("ORDERED_MICROCLINE",
                            "LABRADORITE",
                            "KAOLINITE_DRY_BRANCH",
                            "MONTMORILLONITE_WYO",
                            "ILLITE_1M_RM30",
                            "CORUNDUM"),
                   std = "CORUNDUM",
                   align = 0.3)
\#Alternatively you can specify the internal standard
#concentration if known:
rockjock_1s <- fps(lib = rockjock,</pre>
                  smpl = rockjock_mixtures$Mix1,
                  refs = c("ORDERED_MICROCLINE",
                           "LABRADORITE",
                           "KAOLINITE_DRY_BRANCH",
                           "MONTMORILLONITE_WYO",
                           "ILLITE_1M_RM30",
                           "CORUNDUM"),
                  std = "CORUNDUM",
                  std\_conc = 20,
                  align = 0.3)
## End(Not run)
```

Description

fps.powdRlib returns estimates of phase concentrations using full pattern summation of X-ray powder diffraction data.

Usage

```
## S3 method for class 'powdRlib'
fps(
  lib,
  smpl,
  harmonise,
  solver,
  obj,
  refs,
  std,
  force,
  std_conc,
  omit_std,
  normalise,
  closed,
  tth_align,
  align,
  manual_align,
  tth_fps,
  shift,
  remove_trace,
  weighting,
)
```

Arguments lib

std

	constructor function.		
smpl	A data frame. First column is 2theta, second column is counts		
harmonise	logical parameter defining whether to harmonise the lib and smpl. Default = TRUE. When TRUE the function will harmonise the lib and smpl to the intersecting 2theta range at the coarsest resolution available using natural splines.		
solver	The optimisation routine to be used. One of "BFGS", "Nelder-Mead", "CG", "NNLS". Default = "BFGS".		
obj	The objective function to minimise when "BFGS", "Nelder-Mead", or "CG" are used as the solver argument. One of "Delta", "R", "Rwp". Default = "Rwp". See Chipera and Bish (2002) and page 247 of Bish and Post (1989) for definitions of these functions.		
refs	A character string of reference pattern IDs or names from the specified library. The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. If missing from the function call then all		

phases in the reference library will be used.

ID provided in the refs parameter.

A powdRlib object representing the reference library. Created using the powdRlib

The phase ID (e.g. "QUA.1") to be used as an internal standard. Must match an

force An optional string of phase IDs or names specifying which phases should be forced to remain throughout the automated full pattern summation. The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. The concentration of the internal standard (if known) in weight percent. If unstd_conc known then omit the argument from the function call or use std_conc = NA (default), n which case it will be assumed that all phases sum to 100 percent. A logical parameter to be used when the std_conc argument is defined. When omit_std omit_std = TRUE the phase concentrations are recomputed to account for the value supplied in std_conc. Default = FALSE. normalise deprecated. Please use the omit_std and closed arguments instead. closed A logical parameter to be used when the std_conc argument is defined and omit_std = TRUE. When closed = TRUE the internal standard concentration is removed and the remaining phase concentrations closed to sum to 100 percent. Default = FALSE. tth_align A vector defining the minimum and maximum 2theta values to be used during alignment (e.g. c(5,65)). If not defined, then the full range is used. The maximum shift that is allowed during initial 2theta alignment (degrees). align Default = 0.1. manual_align A logical operator denoting whether to optimise the alignment within the negative/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. Default = FALSE, i.e. alignment is optimised. A vector defining the minimum and maximum 2theta values to be used during tth_fps full pattern summation (e.g. c(5,65)). If not defined, then the full range is used. shift A single numeric value denoting the maximum (positive or negative) shift, in degrees 2theta, that is allowed during the shifting of reference patterns. Default remove_trace A single numeric value representing the limit for the concentration of trace phases to be retained, i.e. any mineral with an estimated concentration below remove_trace will be omitted. Default = 0. weighting an optional 2 column data frame specifying the 2theta values in the first column and a numeric weighting vector in the second column that specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) when minimising the objective function defined in the obj argument. Use this weighting parameter with caution. The default is simply a weighting vector where all values are 1, which hence has no effect on the computed objective function.

Details

Applies full pattern summation (Chipera & Bish, 2002, 2013; Eberl, 2003) to an XRPD sample to quantify phase concentrations. Requires a powdRlib library of reference patterns with reference intensity ratios in order to derive mineral concentrations. Details provided in Butler and Hillier (2021)

Value

a powdRfps object with components:

other arguments

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

ions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

References

Butler, B. M., Hillier, S., 2021.powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Comp. Geo. 147, 104662. doi:10.1016/j.cageo.2020.104662

Bish, D.L., Post, J.E., 1989. Modern powder diffraction. Mineralogical Society of America.

Chipera, S.J., Bish, D.L., 2013. Fitting Full X-Ray Diffraction Patterns for Quantitative Analysis: A Method for Readily Quantifying Crystalline and Disordered Phases. Adv. Mater. Phys. Chem. 03, 47-53. doi:10.4236/ampc.2013.31A007

Chipera, S.J., Bish, D.L., 2002. FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. J. Appl. Crystallogr. 35, 744-749. doi:10.1107/S0021889802017405

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

```
#Load the minerals library
data(minerals)
# Load the soils data
data(soils)
#Since the reference library is relatively small,
#the whole library can be used at once to get an
#estimate of the phases within each sample.
## Not run:
fps_sand <- fps(lib = minerals,</pre>
                  smpl = soils$sandstone,
                  refs = minerals$phases$phase_id,
                  std = "QUA.1",
                  align = 0.2)
fps_lime <- fps(lib = minerals,</pre>
                 smpl = soils$limestone,
                 refs = minerals$phases$phase_id,
                std = "QUA.1",
                 align = 0.2)
```

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```
fps_granite <- fps(lib = minerals,</pre>
                    smpl = soils$granite,
                    refs = minerals$phases$phase_id,
                    std = "QUA.1",
                    align = 0.2)
#Alternatively run all 3 at once using lapply
fps_soils <- lapply(soils, fps,</pre>
                     lib = minerals,
                     std = "QUA.2",
                     refs = minerals$phases$phase_id,
                     align = 0.2)
#Using the rockjock library:
data(rockjock)
data(rockjock_mixtures)
rockjock_1 <- fps(lib = rockjock,</pre>
                   smpl = rockjock_mixtures$Mix1,
                   refs = c("ORDERED_MICROCLINE",
                             "LABRADORITE",
                            "KAOLINITE_DRY_BRANCH",
                            {\tt "MONTMORILLONITE\_WYO"}\,,
                            "ILLITE_1M_RM30",
                            "CORUNDUM"),
                   std = "CORUNDUM",
                   align = 0.3)
#Alternatively you can specify the internal standard
#concentration if known:
rockjock_1s <- fps(lib = rockjock,</pre>
                  smpl = rockjock_mixtures$Mix1,
                  refs = c("ORDERED_MICROCLINE",
                           "LABRADORITE",
                           "KAOLINITE_DRY_BRANCH",
                           "MONTMORILLONITE_WYO",
                           "ILLITE_1M_RM30",
                           "CORUNDUM"),
                  std = "CORUNDUM",
                  std\_conc = 20,
                  align = 0.3)
## End(Not run)
```

fps_lm

Full pattern summation using linear regression

Description

fps_lm returns a simple fit of a given pattern using linear regression, where coefficients may be either positive or negative. Does not return quantitative data. For quantitative results use fps or afps.

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Usage

```
fps_lm(
    lib,
    smpl,
    harmonise,
    refs,
    std,
    tth_align,
    align,
    manual_align,
    tth_fps,
    shift,
    p,
    ...
)
```

Arguments

р

lib A powdRlib object representing the reference library. Created using the powdRlib

constructor function.

smpl A data frame. First column is 2theta, second column is counts

harmonise logical parameter defining whether to harmonise the lib and smpl. Default =

TRUE. When TRUE the function harmonises the lib and smpl data to the intersecting 2theta range at the coarsest resolution available using natural splines.

refs A character string of reference pattern IDs or names from the specified library.

The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. If missing from the function call then all

phases in the reference library will be used.

std The phase ID (e.g. "QUA.1") to be used as internal standard. Must match an ID

provided in the refs parameter.

tth_align A vector defining the minimum and maximum 2theta values to be used during

alignment (e.g. c(5,65)). If not defined, then the full range is used.

align The maximum shift that is allowed during initial 2theta alignment (degrees).

Default = 0.1.

manual_align A logical operator denoting whether to optimise the alignment within the nega-

tive/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. De-

fault = FALSE, i.e. alignment is optimised.

tth_fps A vector defining the minimum and maximum 2theta values to be used during

full pattern summation (e.g. c(5,65)). If not defined, then the full range is used.

shift A single numeric value denoting the maximum (positive or negative) shift, in

degrees 2theta, that is allowed during the shifting of selected phases. Default = 0

U.

a numeric parameter between 0 and 1 specifying the p-value limit for coeffi-

cients. Any reference patterns with a p-value greater than this value will be omitted from the linear regression and results recomputed. Must be greater than

0.000001 but no greater than 1.

... Other arguments

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Details

Requires a powdRlib library of reference patterns. Mineral concentrations are not quantified and therefore reference intensity ratios are not required.

Value

a powdRlm object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned)

residuals a vector of the residuals (fitted vs measured)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped by phase_name and summed weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

```
data(rockjock)
data(rockjock_mixtures)
#Compute the PCA and loadings
x1 <- xrpd_pca(rockjock_mixtures,</pre>
               mean_center = TRUE,
               bin_size = 1,
               root_transform = 1)
## Not run:
fps_lm_out <- fps_lm(rockjock,</pre>
                      smpl = data.frame("x" = x1$loadings$tth,
                                         y'' = x1$loadings$Dim.1),
                      refs = rockjock$phases$phase_id,
                      std = "QUARTZ",
                      align = 0.3,
                      p = 0.01
plot(fps_lm_out,
     wavelength = "Cu",
     interactive = TRUE,
     group = TRUE)
## End(Not run)
```

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fps_lm.powdRlib

Full pattern summation using linear regression

Description

fps_lm.powdRlib returns a simple fit of a given pattern using linear regression, where coefficients may be either positive or negative. Does not return quantitative data. For quantitative results use fps or afps.

Usage

```
## S3 method for class 'powdRlib'
fps_lm(
    lib,
    smpl,
    harmonise,
    refs,
    std,
    tth_align,
    align,
    manual_align,
    tth_fps,
    shift,
    p,
    ...
)
```

Arguments

lib	A powdRlib object repr	esenting the reference librar	y. Created using the powdRlib
-----	------------------------	-------------------------------	-------------------------------

constructor function.

smpl A data frame. First column is 2theta, second column is counts

harmonise logical parameter defining whether to harmonise the lib and smpl. Default =

TRUE. When TRUE the function harmonises the lib and smpl data to the intersecting 2theta range at the coarsest resolution available using natural splines.

refs A character string of reference pattern IDs or names from the specified library.

The IDs or names supplied must be present within the lib\$phases\$phase_id or lib\$phases\$phase_name columns. If missing from the function call then all

phases in the reference library will be used.

std The phase ID (e.g. "QUA.1") to be used as internal standard. Must match an ID

provided in the refs parameter.

tth_align A vector defining the minimum and maximum 2theta values to be used during

alignment (e.g. c(5,65)). If not defined, then the full range is used.

align The maximum shift that is allowed during initial 2theta alignment (degrees).

Default = 0.1.

manual_align A logical operator denoting whether to optimise the alignment within the nega-

tive/position 2theta range defined in the align argument, or to use the specified value of the align argument for alignment of the sample to the standards. De-

fault = FALSE, i.e. alignment is optimised.

fps_lm.powdRlib 37

A vector defining the minimum and maximum 2theta values to be used during full pattern summation (e.g. c(5,65)). If not defined, then the full range is used.

A single numeric value denoting the maximum (positive or negative) shift, in degrees 2theta, that is allowed during the shifting of selected phases. Default = 0.

p a numeric parameter between 0 and 1 specifying the p-value limit for coefficients. Any reference patterns with a p-value greater than this value will be omitted from the linear regression and results recomputed. Must be greater than 0.000001 but no greater than 1.

... Other arguments

Details

Requires a powdRlib library of reference patterns. Mineral concentrations are not quantified and therefore reference intensity ratios are not required.

Value

a powdRlm object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of the fitted XRPD pattern

measured a vector of the original count intensities of the XRPD measurement (aligned)

residuals a vector of the residuals (fitted vs measured)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

38 interpolate

interpolate

Interpolate an XY, multiXY or powdRlib object to a given 2theta scale.

Description

interpolate takes an XY, multiXY or powdRlib object and interpolates the data onto a new 2theta scale using a natural spline. See additional help via ?interpolate.XY, ?interpolate.multiXY or ?interpolate.powdRlib.

Usage

```
interpolate(x, new_tth, ...)
```

Arguments

```
x an XY or multiXY object.new_tth a numeric vector of the new 2theta scale.... other arguments
```

Value

```
an XY or multiXY object.
```

```
#Define a new 2theta scale:
data(rockjock_mixtures)
tth <- seq(10, 60, 0.04)

#interpolate multiXY object of data onto new scale
i1 <- interpolate(rockjock_mixtures, new_tth = tth)

#interpolate XY object onto new scale
i2 <- interpolate(rockjock_mixtures$Mix1, new_tth = tth)

#interpolate powdRlib object onto new scale
i3 <- interpolate(minerals, new_tth = tth)</pre>
```

interpolate.multiXY 39

interpolate.multiXY Interpolate a multiXY object onto a given 2theta scale.

Description

interpolate takes a multiXY object, which may contain XY data frames with varying 2theta scales, and interpolates all data frames onto the same scale using cubic splines.

Usage

```
## S3 method for class 'multiXY'
interpolate(x, new_tth, ...)
```

Arguments

```
x a multiXY object.new_tth a numeric vector of the new 2theta scale.... other arguments
```

Value

```
a multiXY object.
```

Examples

```
data(rockjock_mixtures)

#Define a new 2theta scale:
tth <- seq(10, 60, 0.04)

#interpolate data onto new scale
i1 <- interpolate(rockjock_mixtures, new_tth = tth)</pre>
```

Description

interpolate takes a powdRlib object and interpolates the data onto a new 2theta scale using a cubic spline.

Usage

```
## S3 method for class 'powdRlib'
interpolate(x, new_tth, ...)
```

Arguments

```
x a powdRlib object.new_tth a numeric vector of the new 2theta scale.... other arguments
```

40 interpolate.XY

Value

```
a powdRlib object.
```

Examples

```
data(minerals)

#Define a new 2theta scale:
tth <- seq(10, 60, 0.04)

#interpolate data onto new scale
i1 <- interpolate(minerals, new_tth = tth)</pre>
```

interpolate.XY

Interpolate an XY object onto a given 2theta scale.

Description

interpolate takes an XY object and interpolates the data onto a new 2theta scale using a cubic spline.

Usage

```
## S3 method for class 'XY'
interpolate(x, new_tth, ...)
```

Arguments

```
x an XY object.new_tth a numeric vector of the new 2theta scale.... other arguments
```

Value

```
an XY object.
```

```
data(rockjock_mixtures)

#Define a new 2theta scale:
tth <- seq(10, 60, 0.04)

#interpolate data onto new scale
i1 <- interpolate(rockjock_mixtures$Mix1, new_tth = tth)</pre>
```

merge.powdRlib 41

merge.powdRlib

Merge two powdRlib objects

Description

merge.powdRlib allows two powdRlib objects (which must have) the same 2theta scale) to be merged into a single powdRlib object.

Usage

```
## S3 method for class 'powdRlib'
merge(x, y, ...)
```

Arguments

```
x a powdRlib object.y a powdRlib object... other arguments
```

Value

a powdRlib object.

Examples

```
#Load the minerals library
data(minerals)

#Load the rockjock library
data(rockjock)

#interpolate minerals library onto same 2theta as rockjock
minerals_i <- interpolate(minerals, new_tth = rockjock$tth)

#merge the libraries
merged_lib <- merge(rockjock, minerals_i)</pre>
```

minerals

An example powdRlib reference library

Description

This powdRlib object, built using the powdRlib constructor function, contains a range of measured XRPD data (collected using Cu K-alpha radiation) along with their reference intensity ratios. The library is designed for simple examples only and can be used with the soils data for relatively fast tests of fps and afps.

Usage

minerals

42 minerals_regroup

Format

A powdRlib object of 3 components

xrd A dataframe of all the count intensities of all reference patterns. Column names denote the unique phase ID of each reference pattern

tth A vector of the 2theta scale for all reference patterns in the library

phases A dataframe the phase IDs, names and reference intensity ratios (RIR)

minerals_phases

Example phases table for a reference library

Description

A data frame of associated phase information for the minerals_xrd data. Together these two data frames can be combined with the powdRlib constructor function to create an example reference library (see ?powdRlib). Use the same layout to create custom reference libraries.

Usage

minerals_phases

Format

A 3 column data frame consisting of:

phase_id A string defining the unique phase IDs that should match those defined as column names of the minerals table (e.g. minerals_xrd).

phase_name A string defining the mineral group that each reference pattern belongs to.

rir A vector defining the reference intensity ratios of each reference pattern.

minerals_regroup

Example regrouping structure for the minerals data

Description

Example regrouping structure for the minerals data

Usage

minerals_regroup

Format

A 2 column data frame.

First column contains the unique phase IDs of all phases in the minerals data. Second column contains the grouping structure for the data (Non-clay, Clay or Amorphous).

minerals_xrd 43

minerals_xrd

Example xrd table for a reference library

Description

A table of 14 reference patterns and their corresponding two theta scale that can be combined with the minerals_phases table to create a powdRlib object using the powdRlib constructor function. Use the same layout to create custom reference libraries.

Usage

```
minerals_xrd
```

Format

A dataframe

The first column defines the two theta scale, and remaining columns are individual reference patterns of pure minerals or amorphous phases. Each column title should be a unique mineral ID

multi_xy_to_df

Convert a multiXY object to a data frame.

Description

multi_xy_to_df converts multiXY objects to a column-wise data frame.

Usage

```
multi_xy_to_df(x, tth, ...)
```

Arguments

x a multiXY object.

tth a logical value denoting whether the 2theta scale is appended as the first column.

Default = TRUE.

... other arguments

Value

```
A data.frame.
```

```
#Load the minerals library
data(soils)

soils_df1 <- multi_xy_to_df(soils, tth = TRUE)
soils_df2 <- multi_xy_to_df(soils, tth = FALSE)</pre>
```

44 omit_std

```
multi_xy_to_df.multiXY
```

Convert a multiXY object to a data frame.

Description

multi_xy_to_df.multiXY converts multiXY objects to a column-wise data frame.

Usage

```
## S3 method for class 'multiXY'
multi_xy_to_df(x, tth, ...)
```

Arguments

```
x a multiXY object.
```

tth a logical value denoting whether the 2theta scale is appended as the first column.

Default = TRUE.

... other arguments

Value

A data.frame.

Examples

```
#Load the minerals library
data(soils)

soils_df1 <- multi_xy_to_df(soils, tth = TRUE)
soils_df2 <- multi_xy_to_df(soils, tth = FALSE)</pre>
```

omit_std

Omit the internal standard from phase concentration data within a powdRfps or powdRafps object

Description

omit_std adjusts phase concentrations in a powdRfps or powdRafps object (derived from fps() and afps(), respectively) by removing the concentrations of the internal standard. Relevant information for the calculation is automatically extracted from x\$inputs\$std and x\$inputs\$std_conc. For more information see ?omit_std.powdRfps and omit_std.powdRafps.

Usage

```
omit_std(x, ...)
```

Arguments

x A powdRfps or powdRafps object..

... other arguments

omit_std 45

Value

a powdRfps or powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed

obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

```
## Not run:
data(rockjock)
data(rockjock_mixtures)
rockjock_1 <- fps(lib = rockjock,</pre>
                   smpl = rockjock_mixtures$Mix1,
                   refs = c("ORDERED_MICROCLINE",
                             "LABRADORITE",
                             "KAOLINITE_DRY_BRANCH",
                             "MONTMORILLONITE_WYO",
                             "ILLITE_1M_RM30",
                             "CORUNDUM"),
                  std = "CORUNDUM",
                  align = 0.3,
                  std\_conc = 20)
rockjock_1o <- omit_std(rockjock_1)</pre>
rockjock_a1 <- afps(lib = rockjock,</pre>
                     smpl = rockjock_mixtures$Mix1,
                     std = "CORUNDUM",
                     align = 0.3,
                     lod = 1,
                     std\_conc = 20)
rockjock_a1o <- omit_std(rockjock_a1)</pre>
## End(Not run)
```

46 omit_std.powdRafps

omit_std.powdRafps

Omit the internal standard from phase concentration data within a powdRafps object

Description

omit_std.powdRafps adjusts phase concentrations in a powdRafps object by removing the concentrations of the internal standard. Relevant information for the calculation is automatically extracted from x\$inputs\$std and x\$inputs\$std_conc.

Usage

```
## S3 method for class 'powdRafps'
omit_std(x, ...)
```

Arguments

x A powdRafps object derived from afps().... other arguments

Value

a powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

omit_std.powdRfps 47

```
std_conc = 20)
rockjock_a1o <- omit_std(rockjock_a1)
## End(Not run)</pre>
```

omit_std.powdRfps

Omit the internal standard from phase concentration data within a powdRfps object

Description

omit_std.powdRfps adjusts phase concentrations in a powdRfps object by removing the concentrations of the internal standard. Relevant information for the calculation is automatically extracted from x\$inputs\$std and x\$inputs\$std_conc.

Usage

```
## S3 method for class 'powdRfps'
omit_std(x, ...)
```

Arguments

x A powdRfps object derived from fps().

... other arguments

Value

a powdRfps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the fitted XRPD pattern

measured a vector of the original XRPD measurement (aligned and harmonised)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern and their concentra-

tions

phases_grouped the phases dataframe grouped by phase_name and concentrations summed obj named vector of the objective parameters summarising the quality of the fit

 $weighted_pure_patterns$

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

48 plot.multiXY

Examples

```
## Not run:
data(rockjock)
data(rockjock_mixtures)
rockjock_1 <- fps(lib = rockjock,</pre>
                   smpl = rockjock_mixtures$Mix1,
                   refs = c("ORDERED_MICROCLINE",
                            "LABRADORITE",
                            "KAOLINITE_DRY_BRANCH",
                            "MONTMORILLONITE_WYO",
                            "ILLITE_1M_RM30",
                            "CORUNDUM"),
                  std = "CORUNDUM",
                  align = 0.3,
                  std\_conc = 20)
rockjock_1o <- omit_std(rockjock_1)</pre>
## End(Not run)
```

plot.multiXY

Plotting a multiXY object

Description

plot.multiXY is designed to provide easy, adaptable plots of multiple XRPD patterns.

Usage

```
## S3 method for class 'multiXY'
plot(x, wavelength, xlim, normalise, interactive, ...)
```

Arguments

X	a multiXY object
wavelength	One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
xlim	A numeric vector providing limits of the x-axis (E.g. c(10, 60)). Defaults to full x-axis unless specified.
normalise	Logical. If TRUE then count intensities will be normalised to a minimum of zero and maximum of 1. Default = FALSE.
interactive	Logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

Plots can be made interactive using the logical interactive argument.

plot.powdRafps 49

Examples

```
# Load the minerals library
data(rockjock_mixtures)
## Not run:
plot(as_multi_xy(rockjock_mixtures), wavelength = "Cu")
plot(as_multi_xy(rockjock_mixtures), wavelength = "Cu", interactive = TRUE)
## End(Not run)
```

plot.powdRafps

Plotting elements of a powdRafps object

Description

plot.powdRafps is designed to provide easy, adaptable plots of full pattern summation outputs produced from afps.

Usage

```
## S3 method for class 'powdRafps'
plot(x, wavelength, mode, group, xlim, show_excluded, interactive, ...)
```

Arguments

x	a powdRafps object
wavelength	One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
mode	One of "fit", "residuals" or "both" defining whether to plot the fitted patterns, the residuals of the fit, or both, respectively. Default = "fit".
group	A logical parameter used to specify whether the plotted data are grouped according to the phase name. Default = FALSE.
xlim	A numeric vector providing limits of the x-axis (E.g. c(10, 60)). Defaults to full x-axis unless specified.
show_excluded	A logical value specifying whether the areas excluded from the fitting are identified in the plot as grey rectangles. Default = TRUE.
interactive	logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

When seeking to inspect the results from full pattern summation, interactive plots are particularly useful and can be specified with the interactive argument.

50 plot.powdRbkg

Examples

plot.powdRbkg

Plotting a powdRbkg object

Description

plot.powdRbkg is designed to provide quick plots to inspect the fitted backgrounds obtained from bkg.

Usage

```
## S3 method for class 'powdRbkg'
plot(x, interactive, ...)
```

Arguments

```
x a powdRbkg object

interactive Logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.

... other arguments
```

Details

The only mandatory argument is x, which must be a powdRbkg object. Plots can be made interactive using the logical interactive argument.

```
# Load the minerals library
data(minerals)
## Not run:
```

plot.powdRfps 51

```
plot(minerals, interactive = TRUE)
## End(Not run)
```

plot.powdRfps

Plotting elements of a powdRfps object

Description

plot.powdRfps is designed to provide easy, adaptable plots of full pattern summation outputs produced from fps.

Usage

```
## S3 method for class 'powdRfps'
plot(x, wavelength, mode, group, xlim, show_excluded, interactive, ...)
```

Arguments

X	a powdRfps object
wavelength	One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
mode	One of "fit", "residuals" or "both" defining whether to plot the fitted patterns, the residuals of the fit, or both, respectively. Default = "fit".
group	A logical parameter used to specify whether the plotted data are grouped according to the phase name. Default = FALSE.
xlim	A numeric vector providing limits of the x-axis (E.g. c(10, 60)). Defaults to full x-axis unless specified.
show_excluded	A logical value specifying whether the areas excluded from the fitting are identified in the plot as grey rectangles. Default = TRUE.
interactive	logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

When seeking to inspect the results from full pattern summation, interactive plots are particularly useful and can be specified with the interactive argument.

```
#Load the minerals library
data(minerals)

# Load the soils data
data(soils)

## Not run:
fps_sand <- fps(lib = minerals,</pre>
```

52 plot.powdRlib

```
smpl = soils$sandstone,
    refs = minerals$phases$phase_id,
    std = "QUA.1",
    align = 0.2)

plot(fps_sand, wavelength = "Cu")
plot(fps_sand, wavelength = "Cu", interactive = TRUE)
## End(Not run)
```

plot.powdRlib

Plotting elements of a powdRlib object

Description

plot.powdRlib is designed to provide easy, adaptable plots of an XRPD reference library built using the powdRlib constructor function.

Usage

```
## S3 method for class 'powdRlib'
plot(x, wavelength, refs, interactive, ...)
```

Arguments

X	a powdRlib object
^	a powdkiło object
wavelength	One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
refs	a character string of reference pattern id's to be plotted
interactive	Logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

Plots can be made interactive using the logical interactive argument.

```
# Load the minerals library
data(minerals)
## Not run:
plot(minerals, wavelength = "Cu", refs = "ALB")
plot(minerals, wavelength = "Cu", refs = "ALB", interactive = TRUE)
## End(Not run)
```

plot.powdRlm 53

|--|

Description

plot.powdRlm is designed to provide easy, adaptable plots of full pattern summation outputs produced from fps_lm.

Usage

```
## S3 method for class 'powdRlm'
plot(x, wavelength, mode, xlim, group, show_excluded, interactive, ...)
```

Arguments

X	a powdRlm object
wavelength	One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
mode	One of "fit", "residuals" or "both" defining whether to plot the fitted patterns, the residuals of the fit, or both, respectively. Default = "fit".
xlim	A numeric vector providing limits of the x-axis (E.g. c(10, 60)). Defaults to full x-axis unless specified.
group	A logical parameter used to specify whether the plotted data are grouped according to the phase name. Default = FALSE.
show_excluded	A logical value specifying whether the areas excluded from the fitting are identified in the plot as grey rectangles. Default = TRUE.
interactive	logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

When seeking to inspect the results from full pattern summation, interactive plots are particularly useful and can be specified with the interactive argument.

54 plot.XY

plot.XY

Plotting an XY object

Description

plot.XY is designed to provide easy, adaptable plots of an XRPD pattern.

Usage

```
## S3 method for class 'XY'
plot(x, wavelength, xlim, normalise, interactive, ...)
```

Arguments

x wavelength	an XY object One of "Cu", "Co" or a custom numeric value defining the wavelength (in Angstroms). Used to compute d-spacings. When "Cu" or "Co" are supplied, wavelengths of 1.54056 or 1.78897 are used, respectively.
xlim	A numeric vector providing limits of the x-axis (E.g. c(10, 60)). Defaults to full x-axis unless specified.
normalise	Logical. If TRUE then count intensities will be normalised to a minimum of zero and maximum of 1. Default = FALSE.
interactive	Logical. If TRUE then the output will be an interactive ggplotly object. If FALSE then the output will be a ggplot object.
	other arguments

Details

Plots can be made interactive using the logical interactive argument.

```
# Load the minerals library
data(rockjock_mixtures)
## Not run:
plot(rockjock_mixtures$Mix1, wavelength = "Cu")
plot(rockjock_mixtures$Mix1, wavelength = "Cu", interactive = TRUE)
## End(Not run)
```

powdR 55

powdR	powdR: Full Pattern Summation of X-Ray Powder Diffraction Data

Description

An implementation of the full pattern summation approach to quantitative mineralogy from X-ray powder diffraction data (Chipera & Bish, 2002, 2013; Eberl, 2003; Butler & Hillier 2021).

Author(s)

Benjamin Butler, The James Hutton Institute, Aberdeen, UK

References

Butler, B. M., Hillier, S., 2021.powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Comp. Geo. 147, 104662. doi:10.1016/j.cageo.2020.104662

Chipera, S.J., Bish, D.L., 2013. Fitting Full X-Ray Diffraction Patterns for Quantitative Analysis: A Method for Readily Quantifying Crystalline and Disordered Phases. Adv. Mater. Phys. Chem. 03, 47-53. doi:10.4236/ampc.2013.31A007

Chipera, S.J., Bish, D.L., 2002. FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. J. Appl. Crystallogr. 35, 744-749. doi:10.1107/S0021889802017405

Eberl, D.D., 2003. User's guide to ROCKJOCK - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

Description

A constructor function for creating a powdRlib object from two tables of data. The resulting powdRlib object is required when using fps or afps.

Usage

```
powdRlib(xrd_table, phases_table, check_names)
```

Arguments

xrd_table	A data frame of the count intensities of the XRPD reference patterns, all scaled to same maximum intensity, with their 2theta axis as the first column.
phases_table	A data frame of the required data (phase ID, phase name, and reference intensity ratio) for each reference pattern.
check_names	A logical argument defining whether the column names in the data supplied in xrd_table are syntactically valid variable names and are not duplicated. Default = TRUE.

56 r

Value

a powdRlib object with components:

a data frame of the count intensities of the reference patterns

tth a vector of the 2theta axis

phases a 3 column data frame of the IDs, names and reference intensity ratios of the

reference pattern

Examples

Calculate the R value for a fitted pattern

r

Description

r computes the difference between a measured and fitted pattern. See equation for R in section 2.1 of Butler and Hillier (2021).

Usage

```
r(measured, fitted, weighting)
```

Arguments

measured a vector of count intensities for a measured pattern fitted a vector of count intensities for a fitted pattern

weighting an optional weighting vector of the same length as those specified in measured

and fitted, which specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) from the calculation. Use with caution. Default is simply a weighting vector where all values are 1, which hence has no effect on the

computed value.

Value

a single numeric value

References

Butler, B.M., Hillier, S., 2021. powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Computers and Geosciences. 147, 104662. doi:10.1016/j.cageo.2020.104662

read_xy 57

Examples

read_xy

Read ASCII XY data

Description

read_xy is a wrapper for read.csv that is designed for space separated XRPD data.

Usage

```
read_xy(files, header, sep)
```

Arguments

files path of the file(s) to be imported.

header a logical value indicating whether the file contains the names of the variables as

its first line. Default = FALSE.

sep the field separator character. Values on each line of the file are separated by this

character. Default = " ", indicating space separated format.

Value

If only one path is supplied then an XY data frame with 2 columns is returned, the first being the 2theta axis and the second being the count intensities. If more than one path is supplied then a multiXY list is returned, with each item in the list being an XY data frame as already described.

58 regroup

Examples

regroup

regroup

Description

regroup allows an alternative grouping structure to be applied to powdRfps and powdRafps objects. For more details see ?regroup.powdRfps or ?regroup.powdRafps.

Usage

```
regroup(x, ...)
```

Arguments

x A powdRfps or powdRafps object

... Other parameters passed to methods e.g. fps.powdRlib

Details

powdRfps and powdRafps objects contain a data frame called phases_grouped that summarises phase concentrations based on defined mineral groups from the powdRlib reference library. regroup allows you to change this grouping structure by supplying new group identities.

Value

a powdRfps or powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of fitted XRPD pattern

measured a vector of the count intensities of original XRPD measurement (aligned)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped and summed by phase_name

obj named vector of the objective parameters summarising the quality of the fit

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```
weighted_pure_patterns
```

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

Examples

```
#Load the minerals library
data(minerals)
#Load the soils data
data(soils)
#Load the regrouping structure
data(minerals_regroup)
## Not run:
fps_sandstone <- fps(lib = minerals,</pre>
                      smpl = soils$sandstone,
                      refs = minerals$phases$phase_id,
                      std = "QUA.1",
                      align = 0.2)
fps_sandstone_regrouped <- regroup(fps_sandstone,</pre>
                                    minerals_regroup)
fps_sandstone_regrouped$phases_grouped
## End(Not run)
```

regroup.powdRafps

regroup

Description

regroup.powdRafps allows an alternative grouping structure to be applied to powdRafps objects.

Usage

```
## S3 method for class 'powdRafps'
regroup(x, y, ...)
```

Arguments

x A	powdRafps	object
-----	-----------	--------

y A data frame. First column contains the phase IDs covering all those present in x\$phases\$phase_id. Second column contains the desired grouping of each phase.

... other arguments

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Details

powdRafps objects contain a data frame called phases_grouped that summarises phase concentrations based on defined mineral groups from the powdRlib reference library. regroup allows you to change this grouping structure by supplying new group identities.

Value

a powdRafps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of fitted XRPD pattern

measured a vector of the count intensities of original XRPD measurement (aligned)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped and summed by phase_name

obj named vector of the objective parameters summarising the quality of the fit

 $weighted_pure_patterns$

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

```
#Load the minerals library
data(minerals)
# Load the soils data
data(soils)
#Load the regrouping structure
data(minerals_regroup)
## Not run:
afps_sandstone <- afps(lib = minerals,</pre>
                        smpl = soils$sandstone,
                        std = "QUA.2",
                        align = 0.2,
                        lod = 0.2,
                        amorphous = "ORG",
                        amorphous_lod = 1)
afps_sandstone_regrouped <- regroup(afps_sandstone,</pre>
                                     minerals_regroup)
afps_sandstone_regrouped$phases_grouped
## End(Not run)
```

regroup.powdRfps 61

Description

regroup.powdRfps allows an alternative grouping structure to be applied to powdRfps objects.

Usage

```
## S3 method for class 'powdRfps'
regroup(x, y, ...)
```

Arguments

x A powdRfps object

y A data frame. First column contains the phase IDs covering all those present

in x\$phases\$phase_id. Second column contains the desired grouping of each

phase.

... other arguments

Details

powdRfps objects contain a data frame called phases_grouped that summarises phase concentrations based on defined mineral groups from the powdRlib reference library. regroup allows you to change this grouping structure by supplying new group identities.

Value

a powdRfps object with components:

tth a vector of the 2theta scale of the fitted data

fitted a vector of the count intensities of fitted XRPD pattern

measured a vector of the count intensities of original XRPD measurement (aligned)

residuals a vector of the residuals (measured minus fitted)

phases a dataframe of the phases used to produce the fitted pattern phases_grouped the phases dataframe grouped and summed by phase_name

obj named vector of the objective parameters summarising the quality of the fit

weighted_pure_patterns

a dataframe of reference patterns used to produce the fitted pattern. All patterns

have been weighted according to the coefficients used in the fit

coefficients a named vector of coefficients used to produce the fitted pattern

inputs a list of input arguments used in the function call

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Examples

```
#Load the minerals library
data(minerals)
#Load the soils data
data(soils)
#Load the regrouping structure
data(minerals_regroup)
## Not run:
fps_sandstone <- fps(lib = minerals,</pre>
                      smpl = soils$sandstone,
                      refs = minerals$phases$phase_id,
                      std = "QUA.1",
                      align = 0.2)
fps_sandstone_regrouped <- regroup(fps_sandstone,</pre>
                                     minerals_regroup)
fps_sandstone_regrouped$phases_grouped
## End(Not run)
```

rockjock

RockJock reference library

Description

A powdRlib object of 168 pure reference patterns from the RockJock library (Cu K-alpha radiation) along with reference intensity ratios. Note that compared to same library supplied with RockJock the powdR patterns have been normalised to 10,000 counts and reference intensity ratios transformed so that all are relative to that of corundum, which has been set to a value of 1.0. Can be used with the fps() and afps() functions for quantitative analysis. Example mixtures for testing the rockjock library with known concentrations are available in the rockjock_mixtures data. See ?rockjock_mixtures.

Usage

rockjock

Format

A powdRlib object of 3 components

xrd A dataframe of all the count intensities of all reference patterns. Column names denote the unique phase ID of each reference pattern

tth A vector of the 2theta scale for all reference patterns in the library

phases A dataframe the phase IDs, names and reference intensity ratios (RIR)

rockjock_mixtures 63

Author(s)

Dennis Eberl

References

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

rockjock_mixtures

RockJock synthetic mixtures

Description

A multiXY list containing 8 XRPD measurements (Cu K-alpha radiation) of synthetic mixtures that can be used to assess accuracy of quantitative analysis from the fps() and afps() functions. The mixtures contain various amounts of quartz (QUARTZ standard of the rockjock library), K-feldspar (ORDERED_MICROCLINE), plagioclase (LABRADORITE), kaolinite (KAOLINITE_DRY_BRANCH), dioctahedral smectite (MONTMORILLIONITE_WYO), illite (ILLITE_1M_RM30) and corundum (CORUNDUM).

Usage

rockjock_mixtures

Format

A multiXY list of 8 components, each comprised of two columns. Column tth specifies the 2theta axis and counts specifies the count intensities. The mixtures have to following compositions that are also tabulated in the rockjock_weights data.

- **Mix1** Contains: 4 % K-feldspar, 20 % plagioclase, 12 % kaolinite, 36 % dioctahedral smectite, 8 % illite and 20 % corundum.
- Mix2 Contains: 4 % quartz, 8 % K-feldspar, 36 % plagioclase, 20 % kaolinite, 12 % illite and 20 % corundum.
- **Mix3** Contains: 8 % quartz, 12 % K-feldspar, 36 % kaolinite, 4 % dioctahedral smectite, 20 % illite and 20 % corundum.
- **Mix4** Contains: 12 % quartz, 20 % K-feldspar, 4 % plagioclase, 8 % dioctahedral smectite, 36 % illite and 20 % corundum.
- **Mix5** Contains: 20 % quartz, 36 % K-feldspar, 8 % plagioclase, 4 % kaolinite, 12 % dioctahedral smectite and 20 % corundum.
- **Mix6** Contains: 36 % quartz, 12 % plagioclase, 8 % kaolinite, 20 % dioctahedral smectite, 4 % illite and 20 % corundum.
- **Mix7** Contains: 8 % K-feldspar, 40 % plagioclase, 4 % kaolinite, 12 % dioctahedral smectite, 16 % illite and 20 % corundum.
- **Mix8** Contains: 8 % quartz, 4 % K-feldspar, 4 % plagioclase, 24 % dioctahedral smectite, 40 % illite and 20 % corundum.

Author(s)

Dennis Eberl

64 rockjock_weights

References

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

rockjock_regroup

Regrouping structure for the rockjock reference library

Description

A data frame containing an example re-grouping structure for the rockjock reference library, which results in a slightly coarser description of clay minerals and Fe/Ti-(hydr)oxides in powdRfps or powdRafps objects when used with regroup().

Usage

rockjock_regroup

Format

A data frame with three columns:

phase_id the phase IDs present in afsis\$phases\$phase_id.
phase_name_grouped The phase names that constitute the first regrouping structure.
phase_name_grouped2 The phase names that constitute the second regrouping structure

rockjock_weights

Mineral concentrations of the rockjock_mixtures *data*

Description

A dataframe summarising the weighed mineral concentrations of the rockjock_mixtures data, all in units of weight percent.

Usage

rockjock_weights

Format

An 8 column dataframe, with each row detailing the composition of a sample.

Author(s)

Dennis Eberl

References

Eberl, D.D., 2003. User's guide to RockJock - A program for determining quantitative mineralogy from powder X-ray diffraction data. Boulder, CA.

run_bkg 65

run_bkg

Run the background fitting shiny app

Description

A wrapper for shiny::runApp to start the powdR background fitting Shiny app.

Usage

```
run_bkg(...)
```

Arguments

... further arguments to pass to shiny::runApp

Examples

```
## Not run:
run_powdR()
## End(Not run)
```

run_powdR

Run the powdR shiny app

Description

A wrapper for shiny::runApp to start the Shiny app for powdR.

Usage

```
run_powdR(...)
```

Arguments

... further arguments to pass to shiny::runApp

```
## Not run:
run_powdR()
## End(Not run)
```

66 rwp

rwp

Calculate the Rwp value for a fitted pattern

Description

rwp computes the difference between a measured and fitted pattern. See equation for Rwp in section 2.1 of Butler and Hillier (2021).

Usage

```
rwp(measured, fitted, weighting)
```

Arguments

measured a vector of count intensities for a measured pattern fitted a vector of count intensities for a fitted pattern

weighting an optional weighting vector of the same length as those specified in measured

and fitted, which specifies areas of the pattern to either emphasise (values > 1) or omit (values = 0) from the calculation. Use with caution. Default is simply a weighting vector where all values are 1, which hence has no effect on the

computed value.

Value

a single numeric value

References

Butler, B.M., Hillier, S., 2021. powdR: An R package for quantitative mineralogy using full pattern summation of X-ray powder diffraction data. Computers and Geosciences. 147, 104662. doi:10.1016/j.cageo.2020.104662

soils 67

```
## End(Not run)
```

soils

Example soil XRPD data

Description

3 soil samples from different parent materials measured by XRPD (Cu K-alpha radiation)

Usage

soils

Format

A multiXY list of 3 XY dataframes (named according to parent material type), with each XY dataframe containing two columns of:

tth The 2theta measurement intervals

counts The count intensities

subset.powdRlib

Subset a powdRlib object

Description

subset.powdRlib is designed to provide an easy way of subsetting a powdRlib object by defining the phase ID's that the user wishes to either keep or remove.

Usage

```
## S3 method for class 'powdRlib'
subset(x, refs, mode, ...)
```

Arguments

x a powdRlib object.

refs a string of the phase IDs or names of reference patterns to be subset. The

ID's or names supplied must be present within the lib\$phases\$phase_id or

lib\$phases\$phase_name columns.

mode denotes whether the phase IDs or names defined in the refs argument are re-

tained ("keep") or removed ("remove").

... other arguments

Value

A powdRlib object.

Examples

 $\begin{array}{ll} {\it summarise_mineralogy} & {\it Summarise_the\ mineralogy\ from\ multiple\ powdRfps\ and\ powdRafps} \\ & outputs \end{array}$

Description

summarise_mineralogy creates a summary table of quantified mineral concentrations across a given dataset using a list of multiple powdRfps or powdRafps derived from fps() and afps(), respectively.

Usage

```
summarise_mineralogy(x, type, order, rwp, r, delta)
```

fitted and measured patterns.

Arguments

a list of powdRfps or powdRafps objects. Х a string specifying whether the table uses all phase ID's, or summarises them type according to the phase name. One of "all" or "grouped". order a logical operator denoting whether the columns of the resulting summary table are ordered in descending order according to the summed abundance of each phase across the dataset. a logical operator denoting whether to include the Rwp value as the final column rwp in the output. This provides an objective measure of the difference between the fitted and measured patterns. a logical operator denoting whether to include the R value as the final column in the output. This provides an objective measure of the difference between the fitted and measured patterns. delta a logical operator denoting whether to include the Delta value as the final column in the output. This provides an objective measure of the difference between the

Value

A data frame

tth_transform 69

Examples

```
data(minerals)
data(soils)
## Not run:
multiple_afps <- lapply(soils, afps,</pre>
                          lib = minerals,
                          std = "QUA.1",
                          align = 0.2,
                          lod = 0.1,
                          amorphous = "ORG",
                          amorphous_lod = 1)
sm1 <- summarise_mineralogy(multiple_afps,</pre>
                              type = "all",
                              order = TRUE)
sm2 <- summarise_mineralogy(multiple_afps,</pre>
                              type = "grouped",
                              order = TRUE)
sm3 <- summarise_mineralogy(multiple_afps,</pre>
                              type = "grouped",
                              order = TRUE,
                              rwp = TRUE)
## End(Not run)
```

tth_transform

Transform a two theta axis between wavelengths

Description

tth_transform converts the two theta axis from one wavelength to another via Bragg's law. Use this function with caution if intending the apply fps() or afps() to wavelength transformed samples or libraries because background signals can vary with wavelength which may therefore affect the quality of the fit.

Usage

```
tth_transform(tth, from, to)
```

Arguments

tth the 2theta vector to be transformed

from numeric value defining the wavelength (Angstroms) to transform from to numeric value defining the wavelength (Angstroms) to transform to

Value

```
a transformed 2theta vector
```

70 xrpd_pca

Examples

```
data(soils)
sandstone2 <- soils$sandstone</pre>
#Convert from Cu (1.54056 Angstroms) to Co (1.78897 Angstroms)
sandstone2$tth <- tth_transform(sandstone2$tth,</pre>
                                 from = 1.54056,
                                 to = 1.78897)
sandstone_list <- as_multi_xy(list("sandstone" = soils$sandstone,</pre>
                                    "sandstone2" = sandstone2))
#plot the change
plot(sandstone_list, wavelength = "Cu")
#Alternatively convert the 2theta axis of a library
data(minerals)
minerals2 <- minerals
minerals2$tth <- tth_transform(minerals2$tth,</pre>
                                 from = 1.54056,
                                 to = 1.78897)
#Plot the difference
plot(x = minerals$tth, y = minerals$xrd$QUA.1,
     type = "1", xlim = c(0, 85))
lines(x = minerals2$tth, y = minerals2$xrd$QUA.1,
      col = "red")
```

xrpd_pca

PCA of XRPD data

Description

xrpd_pca is used to apply principal component analysis to X-ray powder diffraction data.

Usage

```
xrpd_pca(x, mean_center, bin_size, root_transform, components)
```

Arguments

х	A multiXY list containing the XRPD data, where each item in the list is a 2 column XY dataframe defining the x (2theta) and y (counts) axes of each measurement. Each item in the list must have a name corresponding to a unique sample ID.
mean_center	A logical argument defining whether mean centering is applied to the XRPD data (default = $TRUE$).
bin_size	An integer between 1 and 10 defining whether to bin the XRPD data to a lower resolution. This bin_size defines the number of data points used in each bin.
root_transform	An integer between 1 and 8 defining the root transform to apply to the XRPD data $$

xrpd_pca 71

components

An integer defining the number of principal components to include in the output. Must be at least 1 less than the number of XRPD patterns in the dataset (the default).

Details

Applies data pre-treatment and principal components analysis to XRPD data based based on the protocols detailed in Butler et al. (2020).

Value

a list with components:

coords a dataframe containing the sample ID's for each sample and the PCA coordinates

for each dimension

loadings a dataframe containing the 2theta axis and the loading of each dimension

eig a dataframe summarising the variance explained by each dimension

References

Butler, B.M., Sila, A.M., Shepherd, K.D., Nyambura, M., Gilmore, C.J., Kourkoumelis, N., Hillier, S., 2019. Pre-treatment of soil X-ray powder diffraction data for cluster analysis. Geoderma 337, 413-424. doi:10.4236/ampc.2013.31A007

```
data(rockjock_mixtures)
x1 <- xrpd_pca(rockjock_mixtures,</pre>
               mean_center = TRUE,
               bin_size = 1,
               root_transform = 1)
\#Plot the loading of dimension 1
plot(x = x1$loadings$tth,
     y = x1$loadings$Dim.1,
     type = "1")
## Not run:
#Fit loading 1 to the rockjock library
f1 <- fps_lm(rockjock,</pre>
             smpl = data.frame("tth" = x1$loadings$tth,
                                "counts" = x1$loadings$Dim.1),
             refs = rockjock$phases$phase_id,
             std = "QUARTZ",
             align = 0,
             p = 0.05)
plot(f1, wavelength = "Cu", interactive = TRUE)
## End(Not run)
```

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