## Package ‘diffractometry’

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**Type** Package

**Title** Baseline identification and peak decomposition for x-ray diffractograms

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**Description** Residual-based baseline identification and peak decomposition for x-ray diffractograms as introduced in Davies/Gather/Mergel/Meise/Mildenberger (2008).

**License** GPL (>= 2)

**NeedsCompilation** yes

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Baseline identification and peak decomposition for x-ray diffractograms

Description

Residual-based baseline identification and peak decomposition for x-ray diffractograms as introduced in Davies et al. (2008).

Details

| Package: | diffractometry |
| Type:    | Package        |
| Version: | 0.1-02         |
| Date:    | 2010-03-05     |
| License: | GPL (>= 2)     |

The package diffractometry contains an implementation of the automatic procedure for analysing x-ray diffractograms of thin films introduced in Davies et al. (2008). The function diffractogram can be used for a complete analysis, while baselinefit and pkdecomp perform baseline estimation and peak decomposition separately. The dataset indiumoxide is the diffractogram used as an example in the article.

Author(s)


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References


See Also

diffractogram, baselinefit, pkdecomp
Baseline and peak intervals for diffractometry data

Description

Calculates Taut String and Weighted Smoothing Spline approximations of a diffractogram and identifies the baseline and peak intervals.

Usage

```r
baselinefit(data, tau=2.5, gam=1, scl.factor=1.2, maxwdth=5)
```

Arguments

- `data`: A diffractogram given as a matrix, where the first column gives the angles of diffraction in degrees $2\theta$ and the second column gives the corresponding photon counts. Angles of diffraction are taken to be equidistant.
- `tau`: Value of Parameter tau used in the residual criterion.
- `gam`: Factor which is used in the separation between peaks and baseline.
- `scl.factor`: Factor which is used in the heteroscedastic residual criterion.
- `maxwdth`: Maximum width of the peaks. By default set to 5 degrees.

Value

Returns a list with components:

- `pmg`: Output (list) of the taut string procedure `fnpreg()`. Contains the estimated fit `pmg$fn`.
- `spl`: Output (list) of the weighted smoothing spline procedure `wsspoiss()`. Contains the estimated fit `spl$reg`.
- `baseline`: Output (list) of the baseline fit procedure `basiserg()`. Contains the baseline fit `baseline$basisl`.
- `npks`: Number of peaks.
- `indlsep`: Left indices of the peak intervals.
- `indrsep`: Right indices of the peak intervals.
- `indextr`: Indices of the extreme values.
- `bs`: Output of the weighted smoothing spline procedure for the data with removed peaks. Contains the baseline for non-peak intervals `bs$reg`.
- `pks`: A vector which contains the data for the peak intervals and zero otherwise.
- `exb`: Output (list) of the routine `exber.maxwdth()` which computes the peak intervals. Contains again `indlsep` and `indrsep`.
- `x`: x values of the data.
- `y`: y values of the data.
Author(s)

M. Meise and P.L. Davies

References


See Also
diffractogram, pkdecomp

Examples

## baseline fit and peak interval identification for a small segment of the indiumoxide data
## baseline plotted in red and data plotted in peak intervals in blue

data(indiumoxide)
index<-indiumoxide[1901:2400,]
plot(index,xlab="",ylab="")
base<-baselinefit(index)
lines(index[,1],base$baseline$basisl,col="red")
points(index[base$indextr[,],col="blue"]

---

**diffractogram**

*Complete analysis of diffractograms as described in Davies et al. (2008)*

**Description**

Performs a complete analysis of x-ray diffractogram, i.e. calculation of the baseline and the peak intervals as well as decomposition of the peaks.

**Usage**

diffractogram(data, tau=2.5,gam=1, scl.factor=1.2, maxwidth=5, intnum=0, alpha=0.1, maxiter1=500, maxiter=10000, hmax=5, maxsolutions=3, heterosk=TRUE, baselim=c(0.05,5), dispers=1)
diffractogram

Arguments

data A diffractogram given as a matrix, where the first column gives the angles of diffraction in $2\theta$ and the second column gives the corresponding photon counts. Angles of diffraction are taken to be equidistant.

tau Value of Parameter tau used in the residual criterion.

gam Factor which is used in the separation between peaks and baseline.
scl.factor Factor which is used in the heteroscedastic residual criterion.

maxwidth Maximum width of the peaks. By default set to 5 degrees.

intnum Vector of numbers of intervals. If intnum = 0, all intervals are used

alpha Test level for residual criterion

maxiter1 Number of attempts to fit a model with 1 component

maxiter Number of attempts to fit a model with $k > 1$ components

hmax Maximum number of components

maxsolutions Number of solutions with $k$ components

heterosk If TRUE, the estimate of noise level given in baslfit is used (default); otherwise noise level is taken to be proportional to signal height

baselim Limits for changes in the baseline estimate; first component is given in percent of the baseline height, second in counts per $2\theta$

dispers Additional dispersion factor; not used if heterosk==T

Details

diffractogram first calls baselinefit with the specified parameters and then pkdecomp for the output of baselinefit. For further details, see the corresponding help pages.

Value

Returns a LIST with components

basl Output of baselinefit

pks Output of pkdecomp

Author(s)

P.L. Davies, M. Meise, T. Mildenberger

References

See Also

baselinefit, pkdecomp

Examples

```r
## Complete Analysis of a segment from indiumoxide data:
## Identification of baseline and peak as well as decomposition of the peaks
## Plot shows baseline and decomposition of first found peak into two components

set.seed(0)
par(mfrow=c(3,1))
data(indiumoxide)
indox<-indiumoxide[1901:2400,]
sol<-diffractogram(indox,maxsolutions=1,maxiter1=50,maxiter=250)
ind<-c(sol$basis$indepsol[1],sol$basis$indrsep[1])
plot(indox,xlab="",ylab="")
lines(indox[,1],sol$basis$basis1,ylab="red")
points(indox[ind[1]:ind[2]],ylab="blue")

plot(indox[ind[1]:ind[2]],sol$bas$basel$peaks[ind[1]:ind[2]],xlab="",ylab="")
lines(indox[ind[1]:ind[2]],sol$bas$basel$peaks[ind[2]]$fit,ylab="red")
plot(indox[ind[1]:ind[2]],sol$bas$basel$peaks[ind[2]]$fitpk[1],ylab="1",ylab="blue")
lines(indox[ind[1]:ind[2]],sol$bas$basel$peaks[ind[2]]$fitpk[2])
```

---

**indiumoxide**  
*Diffactogram of Indium oxide doped with tin*

Description

Example of a diffractogram used in Davies et al. (2008)

Usage

```r
data(indiumoxide)
```
The diffractogram was taken on a thin film of $In_2O_3:Sn$, i.e. indium oxide doped with tin. This material is usually called ITO (indium-tin-oxide). It has a good electrical conductivity and is transparent in the visible wave length region. The first column of the matrix contains the angles of diffraction, the second column contains the corresponding photon counts.

Author(s)
D. Mergel

References

**pkdecomp**

*Decomposition of peaks for the whole data set*

**Description**
Calculates decompositions of peaks for the whole diffractogram

**Usage**

```
pkdecomp(baslfit,intnum=0, alpha=0.1, maxiter1=500, maxiter=10000, hmax=5, maxsolutions=3,heterosk=TRUE, baselim=c(0.05,5), dispers=1)
```

**Arguments**

- **baslfit**: Output of `baselinefit`
- **intnum**: Vector of numbers of intervals. If intnum = 0, all intervals are used
- **alpha**: Test level for residual criterion
- **maxiter1**: Number of attempts to fit a model with 1 component
- **maxiter**: Number of attempts to fit a model with k > 1 components
- **hmax**: Maximum number of components
- **maxsolutions**: Number of solutions with k components
- **heterosk**: If TRUE, the estimate of noise level given in baslfit is used (default); otherwise noise level is taken to be proportional to signal height
- **baselim**: Limits for changes in the baseline estimate; first component is given in percent of the baseline height, second in counts/2theta
- **dispers**: Additional dispersion factor; not used if heterosk==T
Details

Calls `pkdecompint` to decompose the peaks found by `baselinefit` into Pearson Type VII kernels. For every interval, first one kernel is tried. The number of kernels is increased until either a solution accepted by the residual criterion is found or the maximum number of kernels $h_{max}$ is reached. After a solution is accepted, for $max_{solutions}$ greater than 1, further decompositions with the same number of kernels can be produced.

Value

A vector of lists as given by `pkdecompint`

Author(s)

T. Mildenberger; Algorithm for residual criterion by T. Bernholt and T. Hofmeister

References


See Also

diffractogram, baselinefit, pkdecompint

Examples

```r
## Decomposition of data in peak interval into two components

set.seed(0)

par(mfrow=c(2,1))

data(indiumoxide)
indox<-indiumoxide[1901:2400,]
base<-baselinefit(indox)
ind<-c(base$indlsep[1],base$indrsep[1])
plot(indox[ind[1]:ind[2],], base$baseline$peaks[ind[1]:ind[2]],xlab="",ylab="")

pks<-pkdecomp(base,intnum=1,maxsolutions=1,maxiter=50,maxiter=258)

lines(indox[ind[1]:ind[2],],pks[[2]]$fit,col="red")
plot(indox[ind[1]:ind[2],],pks[[2]]$fittpk[1],ylim=c(0,1800),type="l",xlab="",ylab="")
```
De decomposition of peaks in an interval of the diffractogram

Usage

pkdecompint(baslfit, intnum, k, thresh=0, alpha=0.1, heterosk=TRUE, maxiter=10000, dispers=1, baselim=c(0.05,5))

Arguments

- baslfit: Output of baslfit
- intnum: Number of interval
- k: Number of peak components to fit
- thresh: Threshold for residual criterion
- alpha: Test level for residual criterion
- heterosk: If TRUE, the estimate of noise level given in baslfit is used (default); otherwise noise level is taken to be proportional to signal height
- maxiter: Number of attempts to fit a model with k components
- dispers: Additional dispersion factor; not used if heterosk==T
- baselim: Limits for changes in the baseline estimate; first component is given in percent of the baseline height, second in counts/2theta

Value

Returns a LIST with components

- intnr: Number of interval
- x: values of 2theta
- y: the diffractogram with baseline removed
- fit: the resulting fit, evaluated at all points of x
- fitpk: a matrix with num. ker rows that contain fits of the individual peak components
- basl: the baseline estimate as given in baslfit
- baslchg: change of baseline estimate
- rss: residual sum of squares, standardized by noise level estimate
- num.ker: number of peak components
par     parameter vector as given in section 8 of Davies et al. (2008)
parbl   intercept and slope of the baseline change
parpks  physical characteristics of the peaks
accept  is the fit accepted by the residual criterion
alpha   test level for residual criterion
thresh  threshold used in residual criterion

Author(s)
T. Mildenberger; Algorithm for residual criterion by T. Bernholt and T. Hofmeister

References


See Also
diffractogram, baselinefit, pkdecomp

Examples
## Decomposition of data in peak interval into two components

set.seed(0)
par(mfrow=c(2,1))
data(indiumoxide)
indox<-indiumoxide[1901:2400,]
base<-baselinefit(indox)
ind<-c(base$indlsep[1],base$indrsep[1])
plot(indox[ind[1]:ind[2]],base$baseline$peaks[ind[1]:ind[2]],xlab="",ylab="")
pks<-pkdecompint(base,intnum=1,k=2)
lines(indox[ind[1]:ind[2]],pks$fitpk[1],col="red")
plot(indox[ind[1]:ind[2]],pks$fitpk[1],ylim=c(0,1800),type="l",xlab="",ylab="")
lines(indox[ind[1]:ind[2]],pks$fitpk[2,])
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