Package ‘RpeakChrom’

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Type Package

Title Tools for Chromatographic Column Characterization and Modelling

Chromatographic Peak

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Author Manuel David Peris Diaz, Maria Isabel Alcoriza Balaguer

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Maintainer Manuel David Peris Diaz <madape@alumni.uv.es>

Description The quantitative measurement and detection of molecules in HPLC should be carried out by an accurate description of chromatographic peaks. In this package non-linear fitting using a modified Gaussian model with a parabolic variance (PVMG) has been implemented to obtain the retention time and height at the peak maximum. This package also includes the traditional Van Deemter approach and two alternatives approaches to characterize chromatographic column.

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LazyData TRUE

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**parameters_col_metoxi**

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<tr>
<th>col</th>
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<tbody>
<tr>
<td>Parameters data frame for columnar measurements.</td>
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**Description**

Data frame containing the parameters calculated with processPeak function for columnar measurements of several compounds.

**Usage**

data("col")

**Format**

A data frame with 50 observations on the following 9 variables.

- compound: a factor with levels sulfi cloro dimeti Mera
- flow: a numeric vector
- tr: a numeric vector
- Hmax: a numeric vector
- A60: a numeric vector
- B60: a numeric vector
- A10: a numeric vector
- B10: a numeric vector
- area: a numeric vector

**parameters_col_metoxi**

<table>
<thead>
<tr>
<th>parameters_col_metoxi</th>
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</thead>
<tbody>
<tr>
<td>Parameters data frame for sulphadimetroxine columnar measurements.</td>
</tr>
</tbody>
</table>

**Description**

Data frame containing the parameters calculated with processPeak function for columnar measurements of sulphadimetroxine.

**Usage**

data("parameters_col_metoxi")
**parameters_dead**

**Format**

A data frame with 12 observations on the following 9 variables.

- `compound`  a factor with levels `metoxi`
- `flow`  a numeric vector
- `tr`  a numeric vector
- `Hmax`  a numeric vector
- `A60`  a numeric vector
- `B60`  a numeric vector
- `A10`  a numeric vector
- `B10`  a numeric vector
- `area`  a numeric vector

**Parameters data frame for dead marker measurements.**

**Description**

Data frame containing the parameters calculated with `processPeak` function for dead marker measurements.

**Usage**

```r
data("parameters_dead")
```

**Format**

A data frame with 13 observations on the following 9 variables.

- `compound`  a factor with levels `Kbr`
- `flow`  a numeric vector
- `tr`  a numeric vector
- `Hmax`  a numeric vector
- `A60`  a numeric vector
- `B60`  a numeric vector
- `A10`  a numeric vector
- `B10`  a numeric vector
- `area`  a numeric vector
parameters_ext  Parameters data frame for Kbr extracolumnar measurements.

Description
Data frame containing the parameters calculated with processPeak function for extracolumnar measurements of kbr.

Usage
data("parameters_ext")

Format
A data frame with 13 observations on the following 9 variables.

compound  a factor with levels Kbre
flow  a numeric vector
tr  a numeric vector
Hmax  a numeric vector
A60  a numeric vector
B60  a numeric vector
A10  a numeric vector
B10  a numeric vector
area  a numeric vector

peak  Peak read using readChrom function.

Description
Data frame with 2 columns (x, y) corresponding to a chromatographic peak that has been read and extracted using readChrom function.

Usage
data("peak")

Format
A data frame with 12000 observations on the following 2 variables.

V1  a numeric vector
V2  a numeric vector
processPeak

Estimating the main parameters of a chromatographic peak.

Description

Function processPeak uses peak data to calculate the main parameters of the peak: retention time, maximum high, A60, B60, A10, B10 and area.

Usage

processPeak(peak, baseline=FALSE, method, flow=FALSE, compound=FALSE, area=FALSE)

Arguments

peak data frame. The input is a peak selected by the readChrom function.
baseline if TRUE, the function estimates a baseline using asymmetric least squares and subtracts it from the data. By default, FALSE.
flow numeric. This value will be written in the output data frame.
method string indicating the method used to process the peak. "pvmg": the upper part of the peak is filtered and fitted by the PVMG model by non-linear fitting obtaining the time and height at the peak maximum. Then cubic splines is applied obtaining the half-widths measured at 60.65% and 10% peak height. "splines": interpolation by cubic splines to obtain both half-widths left and right either at 60.65% and 10% peak height from the signal of the peak. "direct": Interpolation of the signal of the peak to estimate the half-widths.
compound string. The name of the compound will be written in the output data frame.
area if TRUE, the peak area is estimated using by trapezoidal numerical integration (pracma package). By default, FALSE.

Details

The PVMG model peak (Parabolic Variance Modified Gaussian) is a simplification of the PLMG model (Parabolic Lorentzian Modified Gaussian) removing the Lorentzian function in the PLMG model and using a Gaussian model with a variance showing only a parabolic change with time. This simplified model gives good performance in relatively narrow ranges along the peak elution. It fits accurately the upper region of the peak. The estimation of the time and height at the peak maximum is carried out by non-linear fittings of the chromatographic data to the PVMG equation.

The difference between "pvmg" and "splines" is that the height and time at the peak maximum are calculated either by the PVGM model or by interpolation using the natural signal without being modelled. Then interpolation by cubic splines is applied for obtaining the half-widths at 60.65% and 10% of peak height. The method "direct" obtain the half-widths directly from the natural signal by interpolation.

If an error "singular gradient matrix at initial parameter estimates" occurs, the retention times used to subset the peak by the readChrom function need to be fitted again. See examples.
Value

This function returns a data frame with the following items: compound (if supplied), flow (if supplied), retention time (tr), maximum high (Hmax), A60, B60, A10, B10, area, RSE (square root of the estimated variance of the random error, MeanError (prediction error) and correlation (R).

Note

To get compounds and flows in the output data frame, they need to be written by the user when using this function.

Author(s)

Manuel David Peris, Maria Isabel Alcoriza Balaguer

References


See Also

readChrom, vanDeemterAlternative, vanDeemter

Examples

```r
# peak <- readChrom("file.csv", do.plot = T,
# t1=28, t2=29.5)
# parameters <- processPeak(peak, baseline=FALSE, flow=0.1,
# method="pvmg", compound="alanine", area=TRUE)
```
## Description

Function `readChrom` allows to read a chromatogram from a csv or txt file with two columns (x, y) and subset the peak of interest if the user gives the limiting retention times. Also, this function can draw the peak to check if the given retention times are correct.

## Usage

```r
readChrom(filepath, do.plot=TRUE, t1=0, t2=0)
```

## Arguments

- **filepath**: string indicating the path to the file. The first column represent the retention times and the second one the intensities.
- **do.plot**: if TRUE, the function prints the chromatogram in the interval selected by t1 and t2. By default, it is TRUE.
- **t1**: numeric. Filter for measurements with retention time >= t1.
- **t2**: numeric. Filter for measurements with retention time <= t2.

## Details

Setting t1 and t2 allows you to filter your peak in the chromatogram. To insurance the correct selection of the peak, the argument "do.plot" should be used.

## Value

The return value is a data frame containing two columns, the retention time and intensity.

## Author(s)

Manuel David Peris, Maria Isabel Alcoriza Balaguer
References


See Also

processPeak, vanDeemterAlternative, vanDeemter

Examples

# Substitute the file path argument for a csv or txt file

# To see the whole chromatogram
# peak <- readChrom("example_file.csv", do.plot = TRUE)

# To subset the peak make use of t1 and t2 arguments, for example:
# peak <- readChrom("example_file.csv", do.plot = TRUE, t1 = 2, t2 = 2.5)
vanDeemter

Usage

vanDeemter(col, ext, dead, length, A, B, C, Foley=FALSE, GG=FALSE, do.plot=TRUE)

Arguments

col data frame of the columnar measurements obtained using processPeak function.

ext data frame of the extracolumnar measurements obtained using processPeak function.

dead data frame of the dead marker measurements obtained using processPeak function.

length numeric value indicating the column length in mm.

A numeric value indicating the initial value of the parameter A from the van Deemter equation.

B numeric value indicating the initial value of the parameter B from the van Deemter equation.

C numeric value indicating the initial value of the parameter C from the Van Deemter equation.

Foley if TRUE, Foley and Dorsey approach is used to estimate the variance from the half-widths measured at 10% peak height.

GG if TRUE the variance and retention time are calculated by using the Grushka and Giddings approach.

do.plot logical

Details

The Van Deemter approach has been widely used in column performance in HPLC from the information obtained in the elution of probe compounds at different flow rates, which relate the column plate height to the linear mobile phase velocity given solute, column and mobile phase composition. In this function the approaches for obtaining the retention time and variance are based on Grushka and Giddings, or Foley and Dorsey. The Grushka and Giddings approach make use of the half-widths measured at 60.65% peak height whereas Foley and Dorsey approach is based on the measurements at 10% peak height where the peak asymmetry is higher. The theoretical plate height (H) is determined according to the Martin and Synge plate model taking into account the measurement of the extra-column contribution.

Value


Author(s)

Manuel David Peris, Maria Isabel Alcoriza Balaguer
References


See Also

readChrom, processPeak, vanDeemterAlternative

Examples

```r
ggmetoxi <- vanDeemter(col = parameters_col_metoxi, ext = parameters_ext, 
dead = parameters_dead, length = 150, A = 6, B = 200, C = 0.04, 
GG = TRUE, Foley = FALSE, do.plot = TRUE)

foleymetoxi <- vanDeemter(col = parameters_col_metoxi, ext = parameters_ext, 
dead = parameters_dead, length = 150, A = 6, B = 200, C = 0.04, 
GG = FALSE, Foley = TRUE, do.plot = TRUE)
```

vanDeemterAlternative  Characterization of chromatographic columns using a new approximation to vanDeemter equations.

Description

Characterization of chromatographic columns using a new approximation to vanDeemter equations.
vanDeemterAlternative

Usage

vanDeemterAlternative(col, ext, dead, length, approachI=FALSE, A, B, C, approachII=FALSE)

Arguments

col data frame of the columnar measurements obtained using processPeak function.

ext data frame of the extracolumnar measurements obtained using processPeak function.

dead data frame of the dead marker measurements obtained using processPeak function.

length numeric value indicating the column length in mm.

approachI If TRUE approach I is performed.

A numeric value indicating the initial value of the parameter A from the van Deemter equation.

B numeric value indicating the initial value of the parameter B from the van Deemter equation.

C numeric value indicating the initial value of the parameter C from the van Deemter equation.

approachII If TRUE approachII is performed.

Details

In the ApproachI the parameters A, B and C from the Van Deemter equation are obtained in two steps. First the variance in volume for a set of compounds eluted a several flows are linearly fitted versus the retention volume to obtain the plate height. In the second step, the obtained slopes at several flow rates are non-linearly correlated with the linear mobile phase velocity. In the ApproachII in the first step the parabolic behavior for the variance in volume units for each compound in the set is fitted against the flow rate. In the second step, the A, B and C coefficients for the different compounds are linearly correlated with their retention volume. The slopes in the straight-lines are the model parameters A, B and C in the Van Deemter equation.

Value

For Approach I: list containing 6 items. Table: a summary of slope estimated values at several flows. Coefficients: A, B and C coefficients already fitted. Step I: coefficient of the linear fitting for the first step (R2). Correlation: R for the second step of the graphic approach (H vs u, non-linear fitting). Mean error: for the graphic (H vs u). RSE: square root of the estimated variance of the random error for the nls graphic.

For Approach II: list containing 8 items Table: a summary of slope estimated values at several flows. Coefficients: A, B and C coefficients already fitted. r2A: R2 for the coefficient A. r2B: R2 for the coefficient B. r2C: R2 for the coefficient C. MREA: mean relative prediction error for the coefficient A. MREB: mean relative prediction error for the coefficient B. MREC: mean relative prediction error for the coefficient C.
Author(s)
Manuel David Peris, Maria Isabel Alcoriza Balaguer

References

See Also
readChrom, processPeak, vanDeemter

Examples
coeff1 <- vanDeemterAlternative(col = col, ext = parameters_ext,
dead = parameters_dead, length = 150, approachI = TRUE, A = 6, B = 200,
C = 0.04, approachII = FALSE)
coeff2 <- vanDeemterAlternative(col = col, ext = parameters_ext,
dead = parameters_dead, length = 150, approachI = FALSE, A = 6, B = 200,
C = 0.04, approachII = TRUE)
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