Package ‘Scale4C’

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Type     Package
Title     Scale4C: an R/Bioconductor package for scale-space
transformation of 4C-seq data
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           SummarizedExperiment
Suggests

Description Scale4C is an R/Bioconductor package for scale-space
transformation and visualization of 4C-seq data. The
scale-space transformation is a multi-scale visualization
technique to transform a 2D signal (e.g. 4C-seq reads on a
genomic interval of choice) into a tesselation in the scale
space (2D, genomic position x scale factor) by applying
different smoothing kernels (Gauss, with increasing sigma).
This transformation allows for explorative analysis and
comparisons of the data's structure with other samples.

License  LGPL-3

biocViews Visualization, QualityControl, DataImport, Sequencing,
               Coverage

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Repository Bioconductor 3.18

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Description

The function `addPointsOfInterest` adds marker points to a Scale4C object, which are subsequently used to mark points of interest in created plots.

Usage

```r
addPointsOfInterest(data, poi)
```

Arguments

- `data` 
  Scale4C object the points are to be added to
- `poi` 
  Points of interest data, in a GRanges object. Important: column names must be specified and include 'colour' and 'name' for each point of interest with appropriate values

Details

The function `addPointsOfInterest` adds predefined points of interest to a Scale4C object. Each point of interest is defined by 'chr', 'start', 'end', 'colour', and 'name'. A bed file or text file can be used to store the information, however, column names have to be added before import. Other additional columns are ignored by the function. The function then converts the information to a GRanges object.

Value

A data frame that contains the data for all points of interest
Author(s)

Carolin Walter

Examples

# import provided point of interest example, and check if import was
# successful
data(liverData)
poiFile <- system.file("extdata", "vp.txt", package="Scale4C")
pointsOfInterest(liverData) <- addPointsOfInterest(liverData,
  read.csv(poiFile, sep = "\t", stringsAsFactor = FALSE))
head(pointsOfInterest(liverData))

```
calculateFingerprintMap

Calculate scale-space fingerprint map for given 4C-seq data
```

Description

calculateFingerprintMap uses the scale space map to calculate the inflection points of the smoothed signals.

Usage

calculateFingerprintMap(data, maxSQSigma = 5000, epsilon = 0.0000001)

Arguments

data Scale-space object for the 4C-seq data
maxSQSigma Maximum square sigma used to calculate the fingerprint map
epsilon Small numeric value (can also be zero); used to test for inflection points

Details

Scale4C uses Gauss kernels of increasing (square) sigma to smooth the original 4C-seq signal. The resulting inflection points for a chosen sigma are stored in the corresponding line of the fingerprint map, i.e. a 2D matrix (position x sigma).

Value

A Scale4C object containing the fingerprint map for a Scale4C object, i.e. a matrix with quite a lot of zeros and the occasional 2 or -1 as symbols for inflection points. The fingerprint map is included as second assay of the Scale4C object’s scaleSpace SummarizedExperiment slot.

Author(s)

Carolin Walter
Examples

```r
# read prepared example data
data(liverData)
# use small maxSQSigma for a fast example
liverData <- calculateFingerprintMap(liverData, maxSQSigma = 50)
head(t(assay(scaleSpace(liverData), 2))[1:10, 1:20])
```

---

**Description**

`Scale4C` uses Gauss kernels of increasing (square) sigma to smooth the original 4C-seq signal. The resulting data is stored in a 2D matrix (position x sigma).

**Usage**

```r
calculateScaleSpace(data, maxSQSigma = 5000)
```

**Arguments**

- `data` Scale-space object for the 4C-seq data
- `maxSQSigma` Maximum square sigma used to calculate the scale space

**Details**

The central idea of the scale-space transformation is to smooth the original signal with increasing strength, identify inflection points, track those inflection points throughout the different smoothing layers, and find singularities in those inflection point 'lines'. In case of 4C-seq data, this corresponds to smoothing the signal gradually, while making notes when features such as 'peaks' or 'valleys' disappear by merging with other features. `calculateScaleSpace` smoothes the original signal up to a provided smoothing factor square sigma (Gauss kernel).

**Value**

A `SummarizedExperiment` that contains in its first assay the scale space representation for a `Scale4C` object.

**Author(s)**

Carolin Walter

**Examples**

```r
# read prepared example data
data(liverData)
# use small maxSQSigma for a fast example
scaleSpace(liverData) <- calculateScaleSpace(liverData, maxSQSigma = 10)
head(t(assay(scaleSpace(liverData), 1))[1:10, 1:20])
```
findSingularities  

Find singularities in a provided scale-space fingerprint map

Description

This function allows to identify singular points in a scale-space fingerprint map.

Usage

```r
findSingularities(data, minSQSigma = 5, outputTrackingInfo = FALSE, 
guessViewpoint = FALSE, useIndex = TRUE)
```

Arguments

- **data**: Scale-space object for the 4C-seq data
- **minSQSigma**: Minimum square sigma used to calculate singularities; for a square sigma of 1, the data can be quite chaotic and identified singularities are less prone to error when a minSQSigma of 2 or higher is used
- **outputTrackingInfo**: If TRUE, notify the user that a certain position / singularity causes problems during tracking in the fingerprint map
- **guessViewpoint**: If TRUE, add another 'peak' at the coordinates of the viewpoint, if provided. Extra singularities can also be added manually. The idea is to decrease running speed significantly by not actually calculating the largest singularity for a typical 4C-seq experiment, i.e. the main viewpoint peak. Its inflection point contours should easily be visible in the fingerprint map, provided that the viewpoint position is actually included in the raw data, but calculating the full contours requires a very high sigma that should usually not be needed to identify other singularities in the area. Cave: Viewpoint contours don’t have to start directly next to the viewpoint coordinates.
- **useIndex**: If TRUE, use fragment index instead of genomic position data

Details

`findSingularities` identifies possible singular points in the fingerprint map's contours, i.e. points where a line of '2' and '-1' in the matrix meet. Starting from those points in scale-space, the contours are traced back down. This 'localization step' ensures that the coordinates for a feature ('peak' or 'valley') corresponding to a given singular point are as accurate as possible: Smoothing with a high-sigma Gauss kernel distorts the original signal somewhat, so that the inflection points identifying the start and the end of a certain feature 'move outwards'.

Value

A data frame that lists the position where a singular point occurs (genomic position and scale-space sigma), plus the size of the feature as given by its minimal / left and maximal / right position.
importBasic4CseqData

Author(s)
Carolin Walter

Examples

# read prepared example data
data(liverData)
singularities(liverData) = findSingularities(liverData, 5, useIndex = TRUE)
singularities(liverData)

importBasic4CseqData  Import raw data from a provided Basic4Cseq output file

Description
A convenience function to easily include Basic4Cseq output data into Scale4C, importBasic4CseqData extracts valid fragments or valid fragment ends from a typical Basic4Cseq output table.

Usage
importBasic4CseqData(rawFile, viewpoint, viewpointChromosome, distance, useFragEnds = TRUE)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>rawFile</td>
<td>Name for the raw file</td>
</tr>
<tr>
<td>viewpoint</td>
<td>Viewpoint position: only fragments around a certain point of interest are imported (doesn’t have to be the actual viewpoint of the experiment, though)</td>
</tr>
<tr>
<td>viewpointChromosome</td>
<td>Viewpoint chromosome of the experiment</td>
</tr>
<tr>
<td>distance</td>
<td>Distance from the viewpoint: only fragments within a certain distance of the viewpoint are imported</td>
</tr>
<tr>
<td>useFragEnds</td>
<td>If TRUE, use full fragment end data, if FALSE, merge fragmentStart and fragmentEnd to a single item per fragment</td>
</tr>
</tbody>
</table>

Details

importBasic4CseqData is a convenience function to import data from Basic4Cseq. It can be ignored altogether if raw experimental data is imported from another source or with another function into R.

Value
A GRanges object that includes the experiment’s raw data for further processing
Author(s)
Carolin Walter

Examples

```r
csvFile <- system.file("extdata", "liverData.csv", package="Scale4C")
liverReads <- importBasic4CseqData(csvFile, viewpoint = 21160072,
                                    viewpointChromosome = "chr10", distance = 1000000)
head(liverReads)
```

---

**liverData**  
*Example 4C-seq data set of fetal liver data*

**Description**
This data set contains an instance of a Scale4C object.
The 4C-seq data was taken from Stadhouders et al’s fetal liver data set.

**Usage**
```
data("liverData")
```

**Format**
Formal class 'Scale4C'

**Value**
A pre-computed instance of a Scale4C object with fingerprint map and singularities. Scale-space image is reduced to save space.

**Source**
Shortened version of Stadhouders et al’s fetal liver data:

**Examples**
```
data("liverData")
liverData
```
Example 4C-seq data set of fetal liver data, with added VP

Description

This data set contains an instance of a Scale4C object.
The 4C-seq data was taken from Stadhouders et al’s fetal liver data set. It contains a manually added viewpoint peak and a peak with a manual coordinate correction.

Usage

data("liverDataVP")

Format

Formal class 'Scale4C'

Value

A pre-computed instance of a Scale4C object with fingerprint map and singularities

Source

Shortened version of Stadhouders et al’s fetal liver data:

Examples

data("liverDataVP")
liverDataVP

Output list of all features for a given scale-space map

Description

This function provides a list of features for a given fingerprint map in scale-space, with position and range of sigma for which the feature in question exists.

Usage

outputScaleSpaceTree(data, outputPeaks = TRUE, useLog = TRUE, useIndex = TRUE)
plotInflectionPoints

Arguments

data            a Scale4C object with singularity data
outputPeaks     If TRUE, output GRanges peak list only, if FALSE, also output valley data in a larger table
useLog          If TRUE, use a log2 transformation on the square sigma values (fewer changes and fewer singularities for high sigma, in contrast to low sigma)
useIndex        If TRUE, use fragment position

Details

Similar to plotTesselation, outputScaleSpaceTree analyzes a list of singular points and calculates corresponding features, i.e. 'peaks' and 'valleys'. Each singular point marks the disappearance (or occurrence, depending on the view) of a feature in scale space: With increasing square sigma as smoothing parameter for the Gauss kernel, smaller features are merged into larger features. In case of Gauss smoothing, one feature is always surrounded by two features of the opposite type, e.g. a 'peak' is surrounded by two 'valleys'. If a 'peak' is smoothed out, it is replaced by a new valley formed of the former peak's adjacent valleys. The singularity list contains only direct information on those 'central' features; outputScaleSpaceTree adds data on the direct neighbours / adjacent features and also provides the sigma ranges for the features as a measure of their stability throughout the smoothing process. Mean read counts for the identified features are also provided ("signal"). If outputPeaks is true, a reduced list of peaks is printed, while omitting valleys or the central-left-right structural information.

Value

A GRanges object that includes all features as identified through singular points, plus 'neighbour features' at each side (each 'peak' is surrounded by two 'valleys' and vice versa for Gauss kernel smoothing), with positions and range of sigma for which the feature in question remains stable.

Author(s)

Carolin Walter

Examples

# read prepared example data
data(liverDataVP)
output = outputScaleSpaceTree(liverDataVP, useLog = FALSE)
head(output)

plotInflectionPoints  Draw a smoothed near-cis profile with marked inflection points for a 4C-seq signal
Description

plotInflectionPoints plots the inflection points for a given square sigma (i.e. a row of the fingerprint map) onto a corresponding smoothed near-cis plot for the 4C-seq signal. This allows to check problematic parts of the fingerprint map in more detail (e.g. unclear tracking areas with close contours), and to improve possible corrections in the singularity list. Plotting the smoothed signal for a given square sigma before calculation of the fingerprint map is also possible.

Usage

plotInflectionPoints(data, sqsigma, fileName = "inflectionPlot.pdf", width = 9, height = 5, maxVis = 5000, useIndex = TRUE, plotIP = TRUE)

Arguments

data : Scale4C object with experimental 4C-seq data to be smoothed and processed
sqsigma : Chosen square sigma, i.e. row of the fingerprint map to pick the inflection points from
fileName : Optional name for export file (pdf)
width : Width of the plot
height : Height of the plot
maxVis : Maximum y-axis value (read number, not sigma!) for visualization
useIndex : If TRUE, use fragment index for x-axis
plotIP : If TRUE, then mark chosen inflection points, if FALSE, simply plot smoothed data

Value

A near-cis plot of the smoothed data with (optional) marked inflection points in darker or lighter grey, depending on their direction

Note

PDF export is supported. If no plot file name is provided, the result is plotted on screen.

Author(s)

Carolin Walter

Examples

data(liverData)
plotInflectionPoints(liverData, 50)
plotTesselation

**Description**

This method draws the final scale space tesselation, as specified by the list of singularities identified for a Scale4C object. Features are marked with different colours; for the default colour scheme, brown corresponds to 'peaks' and blue to 'valleys', while slightly darker colours mark features originating from singularities ('central' features in a set of three features, e.g. 'valley-peak-valley' or 'peak-valley-peak') and lighter colours the two adjacent features. Different colours for 'central' and 'adjacent' features allow for optical quality control of the tesselation: a 'central' / dark feature's direct predecessor or successor (y-axis) can't be of the same colour (i.e. a 'peak' that passes through a singularity is smoothed out into a 'valley'), and neighbouring intervals have to be of the opposing (but lighter) colour (i.e. each 'peak' is surrounded by two 'valleys' for Gauss kernel smoothing). The same is not necessarily true for an 'adjacent' / light feature, however.

**Usage**

```r
plotTesselation(data, minSQSigma = 5, maxSQSigma = -1, maxVis = -1,
                 fileName = "tesselationPlot.pdf", width = 5, height = 5, xInterval = 100,
                 yInterval = 50, chosenColour = c("grey50", "moccasin", "lightskyblue1",
                                            "beige", "azure"), useIndex = TRUE)
```

**Arguments**

- `data` (Scale4C object with singularity data)
- `minSQSigma` (Minimum square sigma to consider)
- `maxSQSigma` (Maximum square sigma to consider; if -1 then the number of rows in the fingerprint map is used)
- `maxVis` (Maximum y value for visualization (doesn't have to be maxSQSigma); if -1 also defaults to number of rows in the fingerprint map)
- `fileName` (Optional name for export file (pdf))
- `width` (Width of the plot)
- `height` (Height of the plot)
- `xInterval` (Interval length for x-axis)
- `yInterval` (Interval length for y-axis)
- `chosenColour` (Chosen colours for the tesselation plot, five in total. Colour 1 is used for the actual lines of the plot, colour 2 for 'central peaks', colour 3 for 'central valleys', colour 4 for 'adjacent peaks', and colour 5 for 'adjacent valleys')
- `useIndex` (If TRUE, use fragment index for x-axis)

**Value**

A tesselation plot, showing different features of the scale space with their range of existance (square sigma) and position
Note
PDF export is supported. If no plot file name is provided, the result is plotted on screen.

Author(s)
Carolin Walter

Examples

```r
if(interactive()) {
  data(liverData)
  plotTesselation(liverData)
}
```

`plotTraceback`  
*Draw the traceback results for a list of singular points on a fingerprint map*

Description
This method plots the traceback results together with fingerprint data, allowing to check for possible errors during tracking. Problems during tracking can occur if contours are very close, have holes, or if the singularity in question is not recognized at all due to holes at the meeting point of both contours that form a singular point. Each singular point is marked with a grey triangle, and the traced left and right end of the corresponding feature are connected with grey lines. If a contour’s end doesn’t match the traceback line, manual correction is possible in the singularity list.

Usage

```r
plotTraceback(data, maxSQSigma = -1, fileName = "tracebackPlot.pdf", width = 15, height = 15, useIndex = TRUE)
```

Arguments

- `data`  Scale4C object with singularity data
- `maxSQSigma`  Maximum square sigma (i.e. maximum y value) to be drawn; if -1 then all available rows in the fingerprint map are used
- `fileName`  Optional name for export file (pdf)
- `width`  Width of the plot
- `height`  Height of the plot
- `useIndex`  If TRUE, use fragment index for x-axis

Value

A traceback plot, showing the traced singular points with their points of origin throughout different smoothing layers
Note

PDF export is supported. If no plot file name is provided, the result is plotted on screen.

Author(s)

Carolin Walter

Examples

```r
if(interactive()) {
  data(liverData)
  plotTraceback(liverData)
}
```

Description

This function creates a Scale4C object. Data on the 4C-seq experiment, i.e. read counts per fragment and viewpoint coordinates, are stored and checked for plausibility.

Usage

```
Scale4C(viewpoint, viewpointChromosome, rawData)
```

Arguments

- **viewpoint**: The experiment’s viewpoint (start, single coordinate)
- **viewpointChromosome**: The experiment’s viewpoint Chromosome
- **rawData**: Reads of the 4C-seq experiment per fragment on an interval of interest (GRanges object with position and read data)

Details

A Scale4C object contains the basic information on a 4C-seq experiment for a certain interval of interest, i.e. read counts at given positions. See `Scale4C-class` for details. Scale-space features such as fingerprint maps or tesselation are calculated during further steps of the analysis by the appropriate functions.

Scale4C expects the raw data to be in a simple data frame consisting of ‘position’ and ‘reads’. `importBasic4CseqData` allows to import fragment data from Basic4Cseq for convenience, however, preparing and importing a simple table with two columns into R is sufficient.

Value

An instance of the Scale4C class.
Author(s)

Carolin Walter

See Also

Scale4C-class

Examples

```r
# create a Scale4C object from a Basic4Cseq export table with added
# viewpoint data
csvFile <- system.file("extdata", "liverData.csv", package="Scale4C")
liverReads <- importBasic4CseqData(csvFile, viewpoint = 21160072,
  viewpointChromosome = "chr10", distance = 1000000)
liverData = Scale4C(rawData = liverReads, viewpoint = 21160072,
  viewpointChromosome = "chr10")
liverData
```

Scale4C-class

Class "Scale4C"

Description

This class is a container for information on a specific 4C-seq scale-space transformation. Stored information includes raw read data, the experiment’s viewpoint location (optional), possible points of interest, the scale-space fingerprint map, and a list of identified singularities in scale-space.

Objects from the class

Objects can be created by calls of the form `new("Scale4C", ...)`.

Slots

- **viewpoint**: Object of class "numeric" representing the viewpoint's location
- **viewpointChromosome**: Object of class "character" representing the viewpoint's chromosome
- **pointsOfInterest**: Object of class "GRanges" representing any points of interest to be marked in the visualizations (usually near-cis based, i.e. close to the viewpoint)
- **rawData**: Object of class "GRanges" representing the 4C-seq reads (or signal strength) of the experiment at given genomic positions
- **scaleSpace**: Object of class "SummarizedExperiment" representing the gradually smoothed 4C-seq signal ('scale space') in its first assay and the corresponding fingerprint map in its second assay.
- **singularities**: Object of class "GRanges" representing singularities in the fingerprint map for the given 4C-seq signal
Methods

**viewpoint**<- signature(object = "Scale4C", value = "numeric"): Setter-method for the viewpoint slot.

**viewpoint** signature(object = "Scale4C"): Getter-method for the viewpoint slot.

**viewpointChromosome**<- signature(object = "Scale4C", value = "character"): Setter-method for the viewpointChromosome slot.

**viewpointChromosome** signature(object = "Scale4C"): Getter-method for the viewpointChromosome slot.

**pointsOfInterest**<- signature(object = "Scale4C", value = "GRanges"): Setter-method for the pointsOfInterest slot.

**pointsOfInterest** signature(object = "Scale4C"): Getter-method for the pointsOfInterest slot.

**rawData**<- signature(object = "Scale4C", value = "GRanges"): Setter-method for the rawData slot.

**rawData** signature(object = "Scale4C"): Getter-method for the rawData slot.

**scaleSpace**<- signature(object = "Scale4C", value = "matrix"): Setter-method for the scaleSpace slot.

**scaleSpace** signature(object = "Scale4C"): Getter-method for the scaleSpace slot.

**singularities**<- signature(object = "Scale4C", value = "GRanges"): Setter-method for the singularities slot.

**singularities** signature(object = "Scale4C"): Getter-method for the singularities slot.

Author(s)

Carolin Walter

Examples

```r
showClass("Scale4C")
```
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