

Package: sicegar (via r-universe)

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Type Package

Title Analysis of Single-Cell Viral Growth Curves

Version 0.2.4

Description Aims to quantify time intensity data by using sigmoidal and double sigmoidal curves. It fits straight lines, sigmoidal, and double sigmoidal curves on to time vs intensity data. Then all the fits are used to make decision on which model best describes the data. This method was first developed in the context of single-cell viral growth analysis (for details, see Caglar et al. (2018) <[doi:10.7717/peerj.4251](https://doi.org/10.7717/peerj.4251)>), and the package name stands for ``SIngle CEll Growth Analysis in R".

URL <https://github.com/wilkelab/sicegar>

Imports dplyr, minpack.lm, fBasics, ggplot2, stats

License GPL-2 | GPL-3

Suggests covr, cowplot, testthat, knitr, rmarkdown

VignetteBuilder knitr

BugReports <https://github.com/wilkelab/sicegar/issues>

Collate 'categorize.R' 'mainFunctions.R' 'multipleFitFunction.R'
'sigmoidalFitFunctions.R' 'doublesigmoidalFitFunctions.R'
'normalizationFunction.R' 'sicegar.R' 'dataInputCheck.R'
'parameterCalculation.R' 'figureGeneration.R'

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categorize	<i>Categorize input data by comparing the AIC values of the three fitted models.</i>
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Description

Catagorizes the input data using the results of two model fits and chosen thresholds.

Usage

```
categorize(
  parameterVectorSigmoidal,
  parameterVectorDoubleSigmoidal,
  threshold_intensity_range = 0.1,
  threshold_minimum_for_intensity_maximum = 0.3,
  threshold_bonus_sigmoidal_AIC = 0,
  threshold_sm_tmax_IntensityRatio = 0.85,
  threshold_dsm_tmax_IntensityRatio = 0.75,
  threshold_AIC = -10,
  threshold_t0_max_int = 0.05,
  showDetails = FALSE
)
```

Arguments

parameterVectorSigmoidal
Output of the sigmoidalFitFunction.

parameterVectorDoubleSigmoidal
Output of the doublesigmoidalFitFunction.


```

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- sicegar::normalizeData(dataInput,
                                         dataInputName = "sample001")

# Fit sigmoidal model
sigmoidalModel <- sicegar::multipleFitFunction(dataInput = normalizedInput,
                                              model = "sigmoidal",
                                              n_runs_min = 20,
                                              n_runs_max = 500,
                                              showDetails = FALSE)

# Fit double sigmoidal model
doubleSigmoidalModel <- sicegar::multipleFitFunction(dataInput = normalizedInput,
                                                    model = "doublesigmoidal",
                                                    n_runs_min = 20,
                                                    n_runs_max = 500,
                                                    showDetails = FALSE)

# Calculate additional parameters
sigmoidalModel <- sicegar::parameterCalculation(sigmoidalModel)
doubleSigmoidalModel <- sicegar::parameterCalculation(doubleSigmoidalModel)

outputCluster <- sicegar::categorize(parameterVectorSigmoidal = sigmoidalModel,
                                    parameterVectorDoubleSigmoidal = doubleSigmoidalModel)

utils::str(outputCluster)

```

dataCheck

Checks if data is in correct format.

Description

Checks if the input data is appropriate and if it is not, the function converts it into a suitable form. The input data frame should contain two columns named time and intensity related to time variable and intensity variable respectively. If the data frame is in a list its name in the list should be \$timeIntensityData.

Usage

```
dataCheck(data, showDetails = TRUE)
```

Arguments

data the input data. It can be either a list that contains a data frame in `.$timeIntensityData` or can be a data frame by itself.

showDetails logical, if TRUE the function will provide an output "check done" if everything is OK. Default is FALSE

Examples

```
# Example 1

# generate data frame
time <- seq(3, 48, 0.5)
intensity <- runif(length(time), 3.0, 7.5)
dataInput <- data.frame(time, intensity)

# Apply dataCheck function
dataOutputVariable <- dataCheck(dataInput)

# Example 2

# generate data frame
time <- seq(3, 48, 0.5)
intensity <- runif(length(time), 3.0, 7.5)
dataInput <- data.frame(time, intensity)

# Normalize Data
dataOutput <- normalizeData(dataInput)
dataInput2 <- dataOutput

# Apply dataCheck function
dataOutputVariable2 <- dataCheck(dataInput2)
```

doublesigmoidalFitFormula

Double Sigmoidal Formula

Description

Calculates intensities using the double-sigmoidal model fit and the parameters (maximum, final asymptote intensity, slope1Param, midpoint1Param, slope2Param, and mid point distance).

Usage

```
doublesigmoidalFitFormula(
  x,
  finalAsymptoteIntensityRatio,
  maximum,
  slope1Param,
  midPoint1Param,
  slope2Param,
  midPointDistanceParam
)
```

Arguments

<code>x</code>	the "time" (time) column of the dataframe
<code>finalAsymptoteIntensityRatio</code>	This is the ratio between asymptote intensity and maximum intensity of the fitted curve.
<code>maximum</code>	the maximum intensity that the double sigmoidal function reach.
<code>slope1Param</code>	the slope parameter of the sigmoidal function at the steepest point in the exponential phase of the viral production.
<code>midPoint1Param</code>	the x axis value of the steepest point in the function.
<code>slope2Param</code>	the slope parameter of the sigmoidal function at the steepest point in the lysis phase. i.e when the intensity is decreasing.
<code>midPointDistanceParam</code>	the distance between the time of steepest increase and steepest decrease in the intensity data. In other words the distance between the x axis values of arguments of <code>slope1Param</code> and <code>slope2Param</code> .

Value

Returns the predicted intensities for the given time points with the double-sigmoidal fitted parameters for the double sigmoidal fit.

Examples

```
time <- seq(3, 24, 0.1)

#simulate intensity data and add noise
noise_parameter <- 0.2
intensity_noise <- stats::runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- doublesigmoidalFitFormula(time,
                                       finalAsymptoteIntensityRatio = .3,
                                       maximum = 4,
                                       slope1Param = 1,
                                       midPoint1Param = 7,
                                       slope2Param = 1,
                                       midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- normalizeData(dataInput)
parameterVector <- doublesigmoidalFitFunction(normalizedInput, tryCounter = 2)

#Check the results
if(parameterVector$isThisaFit){
  intensityTheoretical <-
    doublesigmoidalFitFormula(
      time,
      finalAsymptoteIntensityRatio = parameterVector$finalAsymptoteIntensityRatio_Estimate,
      maximum = parameterVector$maximum_Estimate,
```

```

      slope1Param = parameterVector$slope1Param_Estimate,
      midPoint1Param = parameterVector$midPoint1Param_Estimate,
      slope2Param = parameterVector$slope2Param_Estimate,
      midPointDistanceParam = parameterVector$midPointDistanceParam_Estimate)

comparisonData <- cbind(dataInput, intensityTheoretical)

require(ggplot2)
ggplot(comparisonData) +
  geom_point(aes(x = time, y = intensity)) +
  geom_line(aes(x = time, y = intensityTheoretical)) +
  expand_limits(x = 0, y = 0)
}

if(!parameterVector$isThisaFit){
  print(parameterVector)
}

```

doublesigmoidalFitFunction

Double sigmoidal fit function.

Description

The function fits a double sigmoidal curve to given data by using likelihood maximization (LM) algorithm and provides the parameters (maximum, final asymptote intensity, slope1Param, mid-point1Param, slope2Param, and midpoint distance) describing the double-sigmoidal fit as output. It also contains information about the goodness of fits such as AIC, BIC, residual sum of squares, and log likelihood.

Usage

```

doublesigmoidalFitFunction(
  dataInput,
  tryCounter,
  startList = list(finalAsymptoteIntensityRatio = 0, maximum = 1, slope1Param = 1,
    midPoint1Param = 0.33, slope2Param = 1, midPointDistanceParam = 0.29),
  lowerBounds = c(finalAsymptoteIntensityRatio = 0, maximum = 0.3, slope1Param = 0.01,
    midPoint1Param = -0.52, slope2Param = 0.01, midPointDistanceParam = 0.04),
  upperBounds = c(finalAsymptoteIntensityRatio = 1, maximum = 1.5, slope1Param = 180,
    midPoint1Param = 1.15, slope2Param = 180, midPointDistanceParam = 0.63),
  min_Factor = 1/2^20,
  n_iterations = 1000
)

```



```

                                midPoint1Param = 7,
                                slope2Param = 1,
                                midPointDistanceParam = 8)
intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- normalizeData(dataInput)
parameterVector <- doublesigmoidalFitFunction(normalizedInput, tryCounter = 2)

#Check the results
if(parameterVector$isThisaFit){
  intensityTheoretical <-
    doublesigmoidalFitFormula(
      time,
      finalAsymptoteIntensityRatio = parameterVector$finalAsymptoteIntensityRatio_Estimate,
      maximum = parameterVector$maximum_Estimate,
      slope1Param = parameterVector$slope1Param_Estimate,
      midPoint1Param = parameterVector$midPoint1Param_Estimate,
      slope2Param = parameterVector$slope2Param_Estimate,
      midPointDistanceParam = parameterVector$midPointDistanceParam_Estimate)

  comparisonData <- cbind(dataInput, intensityTheoretical)

  require(ggplot2)
  ggplot(comparisonData) +
    geom_point(aes(x = time, y = intensity)) +
    geom_line(aes(x = time, y = intensityTheoretical)) +
    expand_limits(x = 0, y = 0)}

if(!parameterVector$isThisaFit) {print(parameterVector)}

```

figureModelCurves *Generate model associated figures.*

Description

Generates figures using ggplot that shows the input data and the fitted curves.

Usage

```

figureModelCurves(
  dataInput,
  sigmoidalFitVector = NULL,
  doubleSigmoidalFitVector = NULL,
  showParameterRelatedLines = FALSE,
  xlabelText = "time",
  ylabelText = "intensity",
  fittedXmin = 0,

```

```

    fittedXmax = NA
  )

```

Arguments

dataInput A data frame or a list containing the dataframe. The data frame should be composed of at least two columns. One represents time, and the other represents intensity. The data should be normalized with the normalize data function `sicegar::normalizeData()` before imported into this function.

sigmoidalFitVector the output of the `sicegar::sigmoidalFitFunction()`, or the augmented version of the output generated by the help of `sicegar::parameterCalculation()`, which contains parameters related with sigmoidal model. Default is NULL.

doubleSigmoidalFitVector the output of the `sicegar::doubleSigmoidalFitFunction()`, or the augmented version of the output generated by the help of `sicegar::parameterCalculation()`, which contains parameters related with double sigmoidal model. Default is NULL.

showParameterRelatedLines if equal to TRUE, figure will show parameter related lines on the curves. Default is FALSE.

xlabelText the x-axis name; with default "time"

ylabelText the y-axis name; with default "intensity"

fittedXmin the minimum of the fitted data that will be plotted (Default 0)

fittedXmax the maximum of the fitted data that will be plotted (Default timeRange)

Value

Returns infection curve figures.

Examples

```

time <- seq(3, 24, 0.1)

#simulate intensity data and add noise
noise_parameter <- 0.2
intensity_noise <- runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- sicegar::doublesigmoidalFitFormula(time,
                                                finalAsymptoteIntensityRatio = .3,
                                                maximum = 4,
                                                slope1Param = 1,
                                                midPoint1Param = 7,
                                                slope2Param = 1,
                                                midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- sicegar::normalizeData(dataInput, dataInputName = "sample001")

```

```

# Do the double sigmoidal fit
doubleSigmoidalModel <- sicegar::multipleFitFunction(dataInput = normalizedInput,
                                                    model = "doublesigmoidal",
                                                    n_runs_min = 20,
                                                    n_runs_max = 500,
                                                    showDetails = FALSE)

doubleSigmoidalModel <- sicegar::parameterCalculation(doubleSigmoidalModel)

fig01 <- sicegar::figureModelCurves(dataInput = normalizedInput,
                                     doubleSigmoidalFitVector = doubleSigmoidalModel,
                                     showParameterRelatedLines = TRUE)

print(fig01)

```

fitAndCategorize	<i>Fit and categorize.</i>
------------------	----------------------------

Description

Fits the sigmoidal and double-sigmoidal models to the data and then categorizes the data according to which model fits best.

Usage

```

fitAndCategorize(
  dataInput,
  dataInputName = NA,
  n_runs_min_sm = 20,
  n_runs_max_sm = 500,
  n_runs_min_dsm = 20,
  n_runs_max_dsm = 500,
  showDetails = FALSE,
  startList_sm = list(maximum = 1, slopeParam = 1, midPoint = 0.33),
  lowerBounds_sm = c(maximum = 0.3, slopeParam = 0.01, midPoint = -0.52),
  upperBounds_sm = c(maximum = 1.5, slopeParam = 180, midPoint = 1.15),
  min_Factor_sm = 1/2^20,
  n_iterations_sm = 1000,
  startList_dsm = list(finalAsymptoteIntensityRatio = 0, maximum = 1, slope1Param = 1,
    midPoint1Param = 0.33, slope2Param = 1, midPointDistanceParam = 0.29),
  lowerBounds_dsm = c(finalAsymptoteIntensityRatio = 0, maximum = 0.3, slope1Param =
    0.01, midPoint1Param = -0.52, slope2Param = 0.01, midPointDistanceParam = 0.04),
  upperBounds_dsm = c(finalAsymptoteIntensityRatio = 1, maximum = 1.5, slope1Param =
    180, midPoint1Param = 1.15, slope2Param = 180, midPointDistanceParam = 0.63),
  min_Factor_dsm = 1/2^20,
  n_iterations_dsm = 1000,
  threshold_intensity_range = 0.1,

```

```

threshold_minimum_for_intensity_maximum = 0.3,
threshold_bonus_sigmoidal_AIC = 0,
threshold_sm_tmax_IntensityRatio = 0.85,
threshold_dsm_tmax_IntensityRatio = 0.75,
threshold_AIC = -10,
threshold_t0_max_int = 0.05,
stepSize = 1e-05,
...
)

```

Arguments

dataInput	Un_normalized input data that will be fitted transferred into related functions
dataInputName	Name of data set (Default is 'NA').
n_runs_min_sm	This number indicates the lower limit of the successful fitting attempts for sigmoidal model. It should be smaller than the upper limit of the fitting attempts (n_runs_max_sm). Default is 20
n_runs_max_sm	This number indicates the upper limit of the fitting attempts for sigmoidal model. Default is 500
n_runs_min_dsm	This number indicates the lower limit of the successful fitting attempts for double sigmoidal model. It should be smaller than the upper limit of the fitting attempts (n_runs_max_dsm). Default is 20
n_runs_max_dsm	This number indicates the upper limit of the fitting attempts for sigmoidal model for double sigmoidal model. Default is 500
showDetails	Logical if TRUE prints details of intermediate steps of individual fits (Default is FALSE).
startList_sm	The initial set of parameters vector that sigmoidal fit algorithm tries for the first fit attempt for the relevant parameters. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 1, slopeParam = 1 and, midPoint = 0.33. The numbers are in normalized time intensity scale.
lowerBounds_sm	The lower bounds for the randomly generated start parameters for the sigmoidal fit. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 0.3, slopeParam = 0.01, and midPoint = -0.52. The numbers are in normalized time intensity scale.
upperBounds_sm	The upper bounds for the randomly generated start parameters for the sigmoidal fit. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 1.5, slopeParam = 180, midPoint = 1.15. The numbers are in normalized time intensity scale.
min_Factor_sm	Defines Defines the minimum step size used by the sigmoidal fit algorithm. Default is $1/2^{20}$.
n_iterations_sm	Defines maximum number of iterations used by the sigmoidal fit algorithm. Default is 1000

- `startList_dsm` The initial set of parameters vector that double sigmoidal fit algorithm tries for the first fit attempt for the relevant parameters. The vector composes of six elements; 'finalAsymptoteIntensityRatio', 'maximum', 'slope1Param', 'midPoint1Param', 'slope2Param', and 'midPointDistanceParam'. Detailed explanations of those parameters can be found in vignettes. Defaults are finalAsymptoteIntensityRatio = 0, maximum = 1, slope1Param = 1, midPoint1Param = 0.33, slope2Param = 1, and midPointDistanceParam=0.29. The numbers are in normalized time intensity scale.
- `lowerBounds_dsm` The lower bounds for the randomly generated start parameters for double sigmoidal fit. The vector composes of six elements; 'finalAsymptoteIntensityRatio', 'maximum', 'slope1Param', 'midPoint1Param', 'slope2Param', and 'midPointDistanceParam'. Detailed explanations of those parameters can be found in vignettes. Defaults are finalAsymptoteIntensityRatio = 0, maximum = 0.3, slope1Param = .01, midPoint1Param = -0.52, slope2Param = .01, and midPointDistanceParam = 0.04. The numbers are in normalized time intensity scale.
- `upperBounds_dsm` The upper bounds for the randomly generated start parameters for double sigmoidal fit. The vector composes of six elements; 'finalAsymptoteIntensityRatio', 'maximum', 'slope1Param', 'midPoint1Param', 'slope2Param', and 'midPointDistanceParam'. Detailed explanations of those parameters can be found in vignettes. Defaults are finalAsymptoteIntensityRatio = 1, maximum = 1.5, slope1Param = 180, midPoint1Param = 1.15, slope2Param = 180, and midPointDistanceParam = 0.63. The numbers are in normalized time intensity scale.
- `min_Factor_dsm` Defines the minimum step size used by the double sigmoidal fit algorithm. Default is $1/2^{20}$.
- `n_iterations_dsm` Define maximum number of iterations used by the double sigmoidal fit algorithm. Default is 1000
- `threshold_intensity_range` Minimum for intensity range, i.e. it is the lower limit for the allowed difference between the maximum and minimum of the intensities (Default is 0.1, and the values are based on actual, not the rescaled data.).
- `threshold_minimum_for_intensity_maximum` Minimum allowed value for intensity maximum. (Default is 0.3, and the values are based on actual, not the rescaled data.).
- `threshold_bonus_sigmoidal_AIC` Bonus AIC points for sigmoidal fit. Negative values help the sigmoidal model to win. Only helps in competition between sigmoidal and double sigmoidal fit at decision step "9", i.e. if none of the models fail in any of the tests and stay as a candidate until the last step (Default is 0).
- `threshold_sm_tmax_IntensityRatio` The threshold for the minimum intensity ratio between the last observed time points intensity and theoretical maximum intensity of the sigmoidal curve. If the value is below the threshold, then the data can not be represented with the sigmoidal model. (Default is 0.85)

threshold_dsm_tmax_IntensityRatio
 The threshold for the minimum intensity ratio between the last observed time points intensity and maximum intensity of the double sigmoidal curve. If the value is above the threshold, then the data can not be represented with the double sigmoidal model. (Default is 0.75)

threshold_AIC Maximum AIC values in order to have a meaningful fit (Default is -10).

threshold_t0_max_int
 Maximum allowed intensity of the fitted curve at time is equal to zero (t=0). (Default is 0.05, and the values are based on actual, not the rescaled data.)

stepSize Step size used by the fitting algorithm. Smaller numbers gave more accurate results than larger numbers, and larger numbers gave the results faster than small numbers. The default value is 0.00001.

... All other arguments that model functions ("sigmoidalFitFunction" and, "double-sigmoidalFitFunction") may need.

Value

Returns the parameters related with the curve fitted to the input data.

Examples

```
# Example 1
time <- seq(3, 24, 0.5)

#simulate intensity data and add noise
noise_parameter <- 0.2
intensity_noise <- stats::runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- sicegar::doublesigmoidalFitFormula(time,
                                                finalAsymptoteIntensityRatio = .3,
                                                maximum = 4,
                                                slope1Param = 1,
                                                midPoint1Param = 7,
                                                slope2Param = 1,
                                                midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)

fitObj <- sicegar::fitAndCategorize(dataInput = dataInput)
```

multipleFitFunction *multiple fit function.*

Description

Calls the fitting algorithms to fit the data multiple times with starting from different randomly generated initial parameters in each run. Multiple attempts at fitting the data are necessary to avoid local minima.


```

n_runs_max = 500)

#Check the results
if(parameterVector$isThisaFit){
  intensityTheoretical <- sigmoidalFitFormula(time,
      maximum = parameterVector$maximum_Estimate,
      slopeParam = parameterVector$slopeParam_Estimate,
      midPoint = parameterVector$midPoint_Estimate)

  comparisonData <- cbind(dataInput, intensityTheoretical)

  print(parameterVector$residual_Sum_of_Squares)

  require(ggplot2)
  ggplot(comparisonData)+
    geom_point(aes(x = time, y = intensity)) +
    geom_line(aes(x = time, y = intensityTheoretical), color = "orange") +
    expand_limits(x = 0, y = 0)
}

if(!parameterVector$isThisaFit){
  print(parameterVector)
}

# Example 2 (doublesigmoidal function with normalization)
time <- seq(3, 24, 0.1)

#simulate intensity data with noise
noise_parameter <- 0.2
intensity_noise <- stats::runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- doublesigmoidalFitFormula(time,
      finalAsymptoteIntensityRatio = .3,
      maximum = 4,
      slope1Param = 1,
      midPoint1Param = 7,
      slope2Param = 1,
      midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- normalizeData(dataInput)
parameterVector <- multipleFitFunction(dataInput = normalizedInput,
      dataInputName="sample001",
      model = "doublesigmoidal",
      n_runs_min = 20,
      n_runs_max = 500,
      showDetails = FALSE)

#Check the results
if(parameterVector$isThisaFit){

```



```

intensityTheoretical <-
  doublesigmoidalFitFormula(
    time,
    finalAsymptoteIntensityRatio = parameterVector$finalAsymptoteIntensityRatio_Estimate,
    maximum = parameterVector$maximum_Estimate,
    slope1Param = parameterVector$slope1Param_Estimate,
    midPoint1Param = parameterVector$midPoint1Param_Estimate,
    slope2Param = parameterVector$slope2Param_Estimate,
    midPointDistanceParam = parameterVector$midPointDistanceParam_Estimate)

comparisonData <- cbind(dataInput, intensityTheoretical)

require(ggplot2)
ggplot(comparisonData) +
  geom_point(aes(x = time, y = intensity)) +
  geom_line(aes(x = time, y = intensityTheoretical), color = "orange") +
  expand_limits(x = 0, y = 0)
}

if(!parameterVector$isThisaFit){
  print(parameterVector)
}

```

normalizeData

Normalization of given data

Description

Maps the given time-intensity data into a rescaled data frame where time is scaled in a way that maximum time point is one and intensity is distributed between [0,1].

Usage

```
normalizeData(dataInput, dataInputName = NA)
```

Arguments

dataInput	A data frame or a list containing the dataframe. The data frame should be composed of at least two columns. One represents time, and the other represents intensity.
dataInputName	experiment name (Default is 'NA').

Value

Function returns a new data frame, scaling factors and scaling constants that connects initial data frame to new one. The new data frame includes 2 columns one is for normalized time and the other is for normalized intensity. The whole time is distributed between 0 and 1 and similarly the whole intensity is distributed between 0 and 1. The time and intensity constants and scaling factors are the parameters to transform data from unnormalized data frame to normalized data frame.


```

                                slope2Param = 1,
                                midPointDistanceParam = 8)
intensity <- intensity+intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- sicegar::normalizeData(dataInput)
parameterVector <- sicegar::multipleFitFunction(dataInput = normalizedInput,
                                                dataInputName = "sample01",
                                                model = "doublesigmoidal",
                                                n_runs_min = 20,
                                                n_runs_max = 500,
                                                showDetails = FALSE)

if(parameterVector$isThisaFit){
  parameterVector <- sicegar::parameterCalculation(parameterVector)
}

print(t(parameterVector))

```

```
preCategorize          Checks for signal in the data.
```

Description

Checks if the signal is present in the data. Often a high percentage of high through-put data does not contain a signal. Checking if data does not contain signal before doing a sigmoidal or double sigmoidal fit can make the analysis of data from high-throughput experiments much faster.

Usage

```
preCategorize(
  normalizedInput,
  threshold_intensity_range = 0.1,
  threshold_minimum_for_intensity_maximum = 0.3
)
```

Arguments

`normalizedInput`
is the output of the `sicegar::normalizeData()` function.

`threshold_intensity_range`
minimum for intensity range, i.e. it is the lower limit for the allowed difference between the maximum and minimum of the intensities (Default is 0.1, and the values are based on actual, not the rescaled data.).

`threshold_minimum_for_intensity_maximum`
minimum allowed value for intensity maximum. (Default is 0.3, and the values are based on actual, not the rescaled data.).

Value

Function returns a brief decision list that includes information about the decision process. Post important part of this information is `decisionList$decision` which might be either "no_signal" or "not_no_signal".

Examples

```
# Example 1 with double sigmoidal data

time=seq(3, 24, 0.1)

#simulate intensity data and add noise
noise_parameter = 0.2
intensity_noise = runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity = sicegar::doublesigmoidalFitFormula(time,
                                               finalAsymptoteIntensityRatio = .3,
                                               maximum = 4,
                                               slope1Param = 1,
                                               midPoint1Param = 7,
                                               slope2Param = 1,
                                               midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- sicegar::normalizeData(dataInput, dataInputName = "sample001")
isThis_nosignal <- sicegar::preCategorize(normalizedInput = normalizedInput)

# Example 2 with no_signal data

time <- seq(3, 24, 0.1)

#simulate intensity data and add noise
noise_parameter <- 0.05
intensity_noise <- runif(n = length(time), min = 0, max = 1) * noise_parameter * 2e-04
intensity <- sicegar::doublesigmoidalFitFormula(time,
                                               finalAsymptoteIntensityRatio = .3,
                                               maximum = 2e-04,
                                               slope1Param = 1,
                                               midPoint1Param = 7,
                                               slope2Param = 1,
                                               midPointDistanceParam = 8)

intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity=intensity, time=time)
normalizedInput <- sicegar::normalizeData(dataInput,dataInputName = "sample001")
isThis_nosignal <- sicegar::preCategorize(normalizedInput = normalizedInput)
```

sameSourceDataCheck *Check is data came from the same source.*

Description

Checks if the provided data and models came from same source by looking to ".dataInputName" columns of the inputs.

Usage

```
sameSourceDataCheck(dataInput, sigmoidalFitVector, doubleSigmoidalFitVector)
```

Arguments

`dataInput` a data frame composed of two columns. One is for time and the other is for intensity. Should be normalized data generated by `normalizeData`.

`sigmoidalFitVector` is the output of `sigmoidalFitFunction`. Default is NULL.

`doubleSigmoidalFitVector` is the output of double sigmoidal fit function. Default is NULL.

Value

Returns TRUE if models can from same source, FALSE otherwise.

`sigmoidalFitFormula` *sigmoidalFitFormula*

Description

Calculates intensities for given time points (x) by using sigmoidal fit model and parameters (maximum, slopeParam, and midPoint).

Usage

```
sigmoidalFitFormula(x, maximum, slopeParam, midPoint)
```

Arguments

`x` the "time" (time) column of the dataframe.

`maximum` the maximum intensity that the sigmoidal function can reach while time approaches infinity.

`slopeParam` the slope parameter of the sigmoidal function at the steepest point.

`midPoint` the x axis value of the steepest point in the function.

Value

Returns the predicted intensities for given time points with the given sigmoidal fit parameters.

Examples

```
time <- seq(3, 24, 0.5)

#simulate intensity data and add noise
noise_parameter <- 0.1
intensity_noise <- stats::runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- sigmoidalFitFormula(time, maximum = 4, slopeParam = 1, midPoint = 8)
intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- normalizeData(dataInput)
parameterVector <- sigmoidalFitFunction(normalizedInput, tryCounter = 2)

#Check the results
if(parameterVector$isThisaFit){
  intensityTheoretical <- sigmoidalFitFormula(time,
                                             maximum = parameterVector$maximum_Estimate,
                                             slopeParam = parameterVector$slopeParam_Estimate,
                                             midPoint = parameterVector$midPoint_Estimate)

  comparisonData <- cbind(dataInput, intensityTheoretical)

  require(ggplot2)
  ggplot(comparisonData) +
    geom_point(aes(x = time, y = intensity)) +
    geom_line(aes(x = time, y = intensityTheoretical)) +
    expand_limits(x = 0, y = 0)
}

if(!parameterVector$isThisaFit){
  print(parameterVector)
}
```

sigmoidalFitFunction *Sigmoidal fit function*

Description

The function fits a sigmoidal curve to given data by using likelihood maximization (LM) algorithm and provides the parameters (maximum, slopeParam and, midPoint) describing the double-sigmoidal fit as output. It also contains information about the goodness of fits such as AIC, BIC, residual sum of squares, and log likelihood.

Usage

```
sigmoidalFitFunction(
  dataInput,
  tryCounter,
  startList = list(maximum = 1, slopeParam = 1, midPoint = 0.33),
  lowerBounds = c(maximum = 0.3, slopeParam = 0.01, midPoint = -0.52),
  upperBounds = c(maximum = 1.5, slopeParam = 180, midPoint = 1.15),
  min_Factor = 1/2^20,
  n_iterations = 1000
)
```

Arguments

dataInput	A data frame or a list containing the dataframe. The data frame should be composed of at least two columns. One represents time, and the other represents intensity. The data should be normalized with the normalize data function <code>sicgar::normalizeData()</code> before imported into this function.
tryCounter	A counter that shows the number of times the data was fit via maximum likelihood function.
startList	The initial set of parameters vector that algorithm tries for the first fit attempt for the relevant parameters. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 1, slopeParam = 1 and, midPoint = 0.33. The numbers are in normalized time intensity scale.
lowerBounds	The lower bounds for the randomly generated start parameters. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 0.3, slopeParam = 0.01, and midPoint = -0.52. The numbers are in normalized time intensity scale.
upperBounds	The upper bounds for the randomly generated start parameters. The vector composes of three elements; 'maximum', 'slopeParam' and, 'midPoint'. Detailed explanations of those parameters can be found in vignettes. Defaults are maximum = 1.5, slopeParam = 180, midPoint = 1.15. The numbers are in normalized time intensity scale.
min_Factor	Defines the minimum step size used by the fitting algorithm. Default is $1/2^{20}$.
n_iterations	Defines maximum number of iterations used by the fitting algorithm. Default is 1000

Value

Returns fitted parameters for the sigmoidal model.

Examples

```
time <- seq(3, 24, 0.5)

#simulate intensity data and add noise
```

```

noise_parameter <- 0.1
intensity_noise <- stats::runif(n = length(time), min = 0, max = 1) * noise_parameter
intensity <- sigmoidalFitFormula(time, maximum = 4, slopeParam = 1, midPoint = 8)
intensity <- intensity + intensity_noise

dataInput <- data.frame(intensity = intensity, time = time)
normalizedInput <- normalizeData(dataInput)
parameterVector <- sigmoidalFitFunction(normalizedInput, tryCounter = 2)

#Check the results
if(parameterVector$isThisaFit){
intensityTheoretical <- sigmoidalFitFormula(time,
                                             maximum = parameterVector$maximum_Estimate,
                                             slopeParam = parameterVector$slopeParam_Estimate,
                                             midPoint = parameterVector$midPoint_Estimate)

comparisonData <- cbind(dataInput, intensityTheoretical)

require(ggplot2)
ggplot(comparisonData) +
  geom_point(aes(x = time, y = intensity)) +
  geom_line(aes(x = time, y = intensityTheoretical)) +
  expand_limits(x = 0, y = 0)
}

if(!parameterVector$isThisaFit){
  print(parameterVector)
}

```

unnormalizeData

Unnormalization of given data

Description

Maps the given time-intensity data into a rescaled frame where time is between [0,1] and similarly intensity is between [0,1].

Usage

```
unnormalizeData(dataInput)
```

Arguments

dataInput a list file composes of two parts First part is the data that will be unnormalized, which is a data frame composed of two columns. One is for time and the other is for intensity Second part is the scaling parameters of the data which is a vector that has three components. The first one of them is related with time and last two of them are related with intensity. The second value represents the min value of the intensity set. First and third values represent the difference between max and min value in the relevant column.

Value

Returns a data frame, scaling factors and scaling constants for time and intensity. The other data frame includes 2 columns one is for normalized time and the other is for normalized intensity. The whole time is distributed between 0 and 1 and similarly the whole intensity is distributed between 0 and 1. The time and intensity constants and scaling factors are the parameters to transform data from given set to scaled set.

Examples

```
# generateRandomData
time <- seq(3, 48, 0.5)
intensity <- runif(length(time), 3.0, 7.5)
dataInput <- data.frame(time, intensity)
# Normalize Data
dataOutput <- normalizeData(dataInput)
dataInput2 <- dataOutput
# Un Normalize it
dataOutput2 <- unnormalizeData(dataInput2)
```

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