Package 'PCRedux'

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Type Package

Title Quantitative Polymerase Chain Reaction (qPCR) Data Mining and Machine Learning Toolkit

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Description Extracts features from amplification curve data of quantitative Polymerase Chain Reactions (qPCR) (Pabinger S. et al. (2014) <doi:10.1016/j.bdq.2014.08.002>) for machine learning purposes. Helper functions prepare the amplification curve data for processing as functional data (e.g., Hausdorff distance) or enable the plotting of amplification curve classes (negative, ambiguous, positive). The hookreg() and hookregNL() functions (Burdukiewicz M. et al. (2018) <doi:10.1016/j.bdq.2018.08.001>) can be used to predict amplification curves with an hook effect-like curvature. The pcrfit_single() function can be used to extract features from an amplification curve.

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LazyLoad yes

LazyData yes

URL https://CRAN.R-project.org/package=PCRedux

BugReports https://github.com/PCRuniversum/PCRedux/issues

Depends R (>= 3.5.0)

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PCRedux-package

PCRedux - quantitative PCR Data Mining and Machine Learning Toolkit

Description

PCRedux package is a toolbox for the analysis of sigmoid curve (qPCR) data.

Usage

armor

Format

An object of class list of length 11.

Machine learning

In machine learning and statistics, the classification should be used to identify a new unknown observation. This observation is assigned to a number of categories. One basis is training data sets containing observations with known classes. Using the example of sigmoid amplification curves, this could be an assignment to the class "negative", "ambiguous" or "positive". Basically, a number of descriptors (e.g., characteristics of curvature) are required to be able to assign classes. This package contains functions for extracting characteristics. In addition, the package contains data sets of classified amplification curves.

Author(s)

Stefan Roediger, Michal Burdukiewcz, Andrej-Nikolai Spiess, Konstantin A. Blagodatskikh

Examples

```
# Use the mblrr function to analyse amplification curves
library(qpcR)
mblrr(x=boggy[, 1], y=boggy[, 2])
```

armor

armor: fetch errors gently

Description

armor is a helper function that catches errors and creates an output that can be used for further processing.

Usage

 $\operatorname{armor}(f, n = 1)$

Arguments

f	is the function that needs armor.
n	is the number of Zero repeats if a function fails.

Value

gives a numeric value (S3 class) as output for errors

Author(s)

Andrej Nikolai Spiess, Stefan Roediger

See Also

base::suppressMessages() base::inherits()

Examples

```
# Fetch the error from the diffQ function
require(MBmca)
# In the following the approximate derivative of the amplification curve data
# x <- RAS002[, 1] and y <- RAS002[, 2] is calculated by diffQ().</pre>
# This will not give an error.
x <- RAS002[, 1]
y <- RAS002[, 2]
armor_diffQ_passes <- armor(MBmca::diffQ(cbind(x, y), verbose = TRUE)$xy)</pre>
armor_diffQ_passes
#
# In the following the approximate derivative of the sequences x <- 1:40
# and y <- 1:40 is calculated by diffQ(). However, this will fail.</pre>
# This will give the "internal" error
# >
# Error in list.res[[i]][[8]] : subscript out of bounds
# that is resolved to 0.
x <- 1:40
y <- 1:40
armor_diffQ_fails <- armor(MBmca::diffQ(cbind(x, y), verbose = TRUE)$xy)</pre>
armor_diffQ_fails
```

autocorrelation_test A function to test for autocorrelation of amplification curve data from a quantitative PCR experiment

Description

autocorrelation_test is a function for an autocorrelation analysis from a quantitative PCR experiment. The result of the function is a correlation coefficient.

Usage

autocorrelation_test(y, n = 8, sig.level = 0.01)

Arguments

У	is the cycle dependent fluorescence amplitude (y-axis).
n	is the number of lagged cycles (default 12).
sig.level	is the significance level for the correlation test., Default: 0.01

Value

gives a numeric value (S3 class) as output for an autocorrelation

```
autocorrelation_test
```

Author(s)

Stefan Roediger, Michal Burdukiewcz

See Also

as.zoo, lag, cor.test

Examples

```
default.par <- par(no.readonly = TRUE)</pre>
# Test for autocorrelation in amplification curve data
# Load qpcR for the data
library(qpcR)
# Test for autocorrelation in the testdat data set
res_ac <- sapply(2:ncol(testdat), function(i) {</pre>
                    autocorrelation_test(testdat[, i])
                }
         )
# Plot curve data as overview
# Define the colors for the amplification curves
colors <- rainbow(ncol(testdat)-1, alpha=0.3)</pre>
# Names of samplesfile:///home/tux/R_malade
samples <- colnames(testdat)[-1]</pre>
layout(matrix(c(1,2,1,3), 2, 2, byrow = TRUE))
matplot(testdat[, 1], testdat[, -1], xlab="Cycle", ylab="RFU",
        main="testdat data set", type="l", lty=1, col=colors, lwd=2)
legend("topleft", samples, pch=19, col=colors, ncol=2, bty="n")
# Curves rated by a human after analysis of the overview. 1 = positive,
# 0 = negative
human_rating <- c(1,1,0,0,1,1,0,0,
                  1,1,0,0,1,1,0,0,
                  1,1,0,0,1,1,0,0)
# Convert the n.s. (not significant) to 0 and others to 1.
# Combine the results of the aromatic autocorrelation_test as variable "ac",
# the human rated values as variable "hr" in a new data frame (res_ac_hr).
res_ac_hr <- as.matrix(data.frame(ac=ifelse(res_ac=="n.s.", 0, 1),</pre>
                                   hr=human_rating))
res_performeR <- performeR(res_ac_hr[, "ac"], res_ac_hr[, "hr"])</pre>
# Add ratings by human and autocorrelation_test to the plot
par(las=2)
plot(1:nrow(res_ac_hr), res_ac_hr[, "hr"], xlab="Sample", ylab="Decisions",
     xaxt="n", yaxt="n", pch=19)
axis(2, at=c(0,1), labels=c("negative", "positive"), las=2)
axis(1, at=1:nrow(res_ac_hr), labels=colnames(testdat)[-1], las=2)
points(1:nrow(res_ac_hr), res_ac_hr[, "ac"], pch=1, cex=2, col="red")
legend("topleft", c("Human", "autocorrelation_test"), pch=c(19,1),
       bty="n", col=c("black","red"))
```

decision_modus

decision_modus A function to get a decision (modus) from a vector of classes

Description

decision_modus is a function that can be used to find the most frequent (modus) decision. The classes can be defined by the user (e.g., a", "n", "y" -> "ambiguous", "negative", "positive"). This function is useful if large collections of varying decision (e.g., "a", "a", "a", "n", "n") need to be condensed to a single decision $(3 \times "a", 2 \times "n" -> "a")$.

Usage

```
decision_modus(data, variables = c("a", "n", "y"), max_freq = TRUE)
```

Arguments

data	is a table containing the classes.
variables	is the class to look for.
max_freq	is a logical parameter (default == TRUE) delivers either the most occurring class or a summary.

Value

gives a factor (S3 class, type of integer) as output for a decision

Author(s)

Stefan Roediger, Michal Burdukiewcz

Examples

```
# First example
# Enter a string of arbritary of "a","a","y","n"
# Result:
# [1] a
# Levels: a b n y
decision_modus(c("a","a","y","n","b"))
# Second example
# Analyze data from the decision_res_testdat.csv data file
filename <- system.file("decision_res_testdat.csv", package = "PCRedux")</pre>
```

earlyreg

```
my_data <- read.csv(filename)
head(my_data)
dec <- unlist(lapply(1L:nrow(my_data), function(i) {
        decision_modus(my_data[i, 2:4])
}))
names(dec) <- my_data[, 1]
dec</pre>
```

earlyreg

A function to calculate the slope and intercept of an amplification curve data from a quantitative PCR experiment.

Description

earlyreg is a function to calculate the slope and intercept of an amplification curve data from a quantitative PCR experiment. The number of cycles to be analyzed is defined by the user (default 6 cycles). The output contains the Maximal Information Coefficient (MIC), which can be interpreted as a correlation measure with a range of [0,1]. A value of 0 mean statistically independent data and 1 approaches in "probability for noiseless functional relationships" (see original study by Reshef, D. N. et al. Detecting novel associations in large data sets. Science, 334, 1518-1524 (2011)).

Usage

earlyreg(x, y, range = 5, normalize = FALSE)

Arguments

х	is the cycle numbers (x-axis).
У	is the cycle dependent fluorescence amplitude (y-axis).
range	is the number of cycles to be used for the regression.
normalize	is a logical parameter which indicates if the amplification curve data should be normalized to the 99 percent percentile of the amplification curve.

Value

gives a numeric vector (S3 class, type of double) as output for local regression

Author(s)

Stefan Roediger, Michal Burdukiewcz

See Also

lmrob stats::coefficients()

Examples

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```
# Calculate slope and intercept on noise (negative) amplification curve data
# for the cycles 2 to 7 for the C316.amp data set
library(chipPCR)
data(C316.amp)
# Plot the data
plot(C316.amp[, 2], y=C316.amp[, 3], xlab="Cycle", ylab="RFU",
            main="C316.amp data set", lty=1, type="l")
res <- earlyreg(x=C316.amp[, 2], y=C316.amp[, 3], range=5)
res</pre>
```

encu

A function to calculate numerous features from amplification curve data from a quantitative PCR experiment.

Description

encu (ENcode CUrves) is a function to calculate numerous features of a large amplification curve data set. The pcrfit_single is performing the analysis for a single process.

Usage

```
encu(data, detection_chemistry = NA, device = NA)
```

Arguments

data	is the data set containing the cycles and fluorescence amplitudes.	
detection_chemistry		
	contains additional meta information about the detection chemistry (e.g., probes, intercalating dye) that was used.	
device	contains additional meta information about the qPCR system that was used.	

Value

gives a data.frame vector (S3 class, type of list) as output for features

The output of the encu function is identical to the pcrfit_single function.

Author(s)

Stefan Roediger, Michal Burdukiewcz

encu

head2tailratio

Examples

```
# Calculate curve features of an amplification curve data. Note that not all
# available CPU cores are used. If need set "all" to use all available cores.
# In this example the testdat data set from the qpcR package is used.
# The samples F1.1 and F1.2 are positive amplification curves. The samples
# F1.3 and F1.4 are negative.
library(qpcR)
```

res_encu <- encu(testdat[, 1:3])
res_encu</pre>

head2tailratio

A function to calculate to head to tail ratio of amplification curve data from a quantitative PCR experiment

Description

head2tailratio is a function to calculate the ratio of the head and the tail of a quantitative PCR amplification curve. In this test, only the head (first six cycles) and the tail (last six cycles) form the region of interest (ROI).

Usage

```
head2tailratio(y, normalize = FALSE, slope_normalizer = FALSE, verbose = FALSE)
```

Arguments

У	is the cycle dependent fluorescence amplitude (y-axis).	
normalize	is a logical parameter, which indicates if the amplification curve.	
slope_normalizer		
	is a logical parameter, which indicates if the head2tailratio should be normalized to the slope of the ROI.	
verbose	is a logical parameter, which indicates if all the values, parameters and coefficients of the analysis should be shown.	

Value

gives a numeric (S3 class, type of double) as output for the head to tail ratio

Author(s)

Stefan Roediger, Michal Burdukiewcz

Examples

```
# calculate head to tail ratio on amplification curve data
library(qpcR)
res_head2tailratio <- sapply(2:ncol(competimer), function(i) {
    head2tailratio(y=competimer[, i], normalize=TRUE, slope_normalizer=TRUE)
})
res_head2tailratio_cluster <- kmeans(res_head2tailratio, 3)$cluster
matplot(competimer[, 1], competimer[, -1], xlab="Cycle", ylab="RFU",
    main="competimer data set", type="l", lty=1, col=res_head2tailratio_cluster, lwd=2)
```

hookreg	A function to calculate the slope and intercept of an amplification
	curve data from a quantitative PCR experiment at the end of the data
	stream.

Description

hookreg is a function to calculate the slope and intercept of an amplification curve data from a quantitative PCR experiment. The idea is that a strong negative slope at the end of an amplification curve is indicative for a hook effect (see Barratt and Mackay 2002).

Usage

```
hookreg(
    x,
    y,
    normalize = TRUE,
    sig.level = 0.0025,
    CI.level = 0.9975,
    robust = FALSE
)
```

Arguments

х	is the cycle numbers (x-axis).
у	is the cycle dependent fluorescence amplitude (y-axis).
normalize	is a logical parameter indicating if the data should be normalized to the 0.999 quantile
sig.level	defines the significance level to test for a significant regression
CI.level	confidence level required for the slope
robust	is a logical parameter indicating if the data should be analyzed be a robust linear regression (lmrob).

hookregNL

Value

gives a numeric (S3 class, type of double) as output for the detection of a hook

Author(s)

Stefan Roediger, Michal Burdukiewcz

References

K. Barratt, J.F. Mackay, *Improving Real-Time PCR Genotyping Assays by Asymmetric Amplification*, J. Clin. Microbiol. 40 (2002) 1571–1572. doi:10.1128/JCM.40.4.1571-1572.2002.

Examples

```
default.par <- par(no.readonly = TRUE)</pre>
# Calculate slope and intercept on noise (negative) amplification curve data
# for the last eight cycles.
library(qpcR)
res_hook <- data.frame(sample=colnames(boggy)[-1],</pre>
                       t(sapply(2:ncol(boggy), function(i) {
                       hookreg(x=boggy[, 1], y=boggy[, i])}))
res_hook
data_colors <- rainbow(ncol(boggy[, -1]), alpha=0.5)</pre>
cl <- kmeans(na.omit(res_hook[, 2:3]), 2)$cluster</pre>
par(mfrow=c(1,2))
matplot(x=boggy[, 1], y=boggy[, -1], xlab="Cycle", ylab="RFU",
main="boggy Data Set", type="l", lty=1, lwd=2, col=data_colors)
legend("topleft", as.character(res_hook$sample), pch=19,
         col=data_colors, bty="n")
plot(res_hook$intercept, res_hook$slope, pch=19, cex=2, col=data_colors,
xlab="intercept", ylab="Slope",
main="Clusters of Amplification Curves with an Hook Effect-like Curvature\nboggy Data Set")
points(res_hook$intercept, res_hook$slope, col=cl, pch=cl, cex=cl)
legend("topright", c("Strong Hook effect", " Weak Hook effect"), pch=c(1,2), col=c(1,2), bty="n")
 text(res_hook$intercept, res_hook$slope, res_hook$sample)
```

par(default.par)

hookregNL

hookregNL - A function to calculate the slope of amplification curves in the tail region

Description

hookregNL is a function to calculate the slope and intercept of an amplification curve from a quantitative PCR experiment. The idea is that a strong negative slope at the end of an amplification curve is indicative for a hook effect (see Barratt and Mackay 2002). In contrast to hookreg fits this function a sex-parameter model to the amplification curve and extracts the coefficient, which describes the slope.

Usage

hookregNL(x, y, plot = FALSE, level = 0.995, simple = TRUE, manualtrim = 5)

Arguments

Х	is the cycle numbers (x-axis).
У	is the cycle dependent fluorescence amplitude (y-axis).
plot	is a logical parameter indicating if the data should be plotted, Default: FALSE.
level	the confidence level required, Default: 0.99.
simple	is a logical parameter. If TRUE (default) only the slope, confidence interval and decisions are shown as output
manualtrim	is the number of cycles that should be removed from the background. (data.frame). If FALSE, a list including the 6-parameter model is the output.

Value

gives a numeric (S3 class, type of double) as output for the detection of a hook

Author(s)

Andrej-Nikolai Spiess, Stefan Roediger, Michal Burdukiewcz

References

K. Barratt, J.F. Mackay, *Improving Real-Time PCR Genotyping Assays by Asymmetric Amplification*, J. Clin. Microbiol. 40 (2002) 1571–1572. doi:10.1128/JCM.40.4.1571-1572.2002.

See Also

pcrfit confint

Examples

```
# Analyze data from the boggy data set for potential hook effect like
# curvature
library(qpcR)
# has hook
res <- hookregNL(boggy[, 1], boggy[, 2])
res
```

has no hook

humanrater2

```
res <- hookregNL(boggy[, 1], boggy[, 12])
res</pre>
```

humanrater2 Human Rater 2.0

Description

Launches graphical user interface for the manual annotation of large amplification curve data sets, similarly to the humanrater function.

Usage

humanrater2()

Value

No return value, called for side effects

Warning

Any ad-blocking software may cause malfunctions.

mblrr

A function to perform a Local Robust Regression in Ranges defined by Qunantile-filtering

Description

mblrr is a function to perform the Median based Local Robust Regression (mblrr) from a quantitative PCR experiment. In detail, this function attempts to break the amplification curve in two parts (head (~background) and tail (~plateau)). Subsequent, a robust linear regression analysis (1mrob) is preformed individually on both parts. The rational behind this analysis is that the slope and intercept of an amplification curve differ in the background and plateau region.

Usage

mblrr(x, y, sig.level = 0.01, normalize = FALSE)

Arguments

х	is the cycle numbers (x-axis).
У	is the cycle dependent fluorescence amplitude (y-axis).
sig.level	is the significance level for the correlation test.
normalize	is a logical parameter, which indicates if the amplification curve data should be normalized to the 99 percent quantile of the amplification curve.

Details

mblrr_intercept_bg is the intercept of the head region, *mblrr_slope_bg* is the slope of the head region, *mblrr_cor_bg* is the coefficient of correlation of the head region, *mblrr_intercept_pt* is the intercept of the tail region, *mblrr_intercept_pt* is the slope of the tail region, *mblrr_cor_pt* is the coefficient of correlation of the tail region, *mblrr_cor_pt* is the coefficient of correlation of the tail region.

Value

gives a numeric (S3 class, type of double) as output for the regressed regions

Author(s)

Stefan Roediger, Michal Burdukiewcz

Examples

```
# Perform an mblrr analysis on noise (negative) amplification data of qPCR data
# with 35 cycles.
library(qpcR)
mblrr(x=boggy[, 1], y=boggy[, 2], normalize=TRUE)
```

PCRedux_datasets The datasets implemented in PCRedux

Description

A compilation of datasets for method evaluation/comparison.

Usage

```
data_sample
RAS002
RAS002_decisions
kbqPCR
decision_res_kbqPCR
```

Details

data_sample

Setup: Amplification curve data were analyzed with the encu() and the decision_modus() functions. Details:

Data sets: batsch1, boggy, C126EG595, competimer, dil4reps94, guescini1, karlen1, lievens1, reps384, rutledge, testdat, vermeulen1, VIMCFX96_60, stepone_std.rdml, RAS002.rdml, RAS003.rdml, HCU32_aggR.csv, lc96_bACTXY.rdml.

RAS002

Setup: Amplification curve data of the RAS002.rdml data set.

pcrfit_single

Details: Data sets: RAS002.rdml.

RAS002_decisions

Setup: Classes of the amplification curves from the RAS002.rdml data set. Details: Data sets: decision_res_RAS002.csv.

Author(s)

Stefan Roediger

References

Roediger, S., Burdukiewicz, M., Spiess, A.-N. & Blagodatskikh, K. Enabling reproducible real-time quantitative PCR research: the RDML package. *Bioinformatics* (2017). doi:10.1093/bioinformatics/btx528 Roediger, S., Burdukiewicz, M. & Schierack, P. chipPCR: an R package to pre-process raw data of amplification curves. *Bioinformatics* 31, 2900–2902 (2015) Ritz, C. & Spiess, A.-N. qpcR: an R package for sigmoidal model selection in quantitative real-time

polymerase chain reaction analysis. *Bioinformatics* 24, 1549–1551 (2008).

Examples

```
## 'data_sample' dataset.
head(data_sample)
## 'RAS002.rdml' dataset as rda file.
head(RAS002)
```

head(RAS002_decisions)

pcrfit_single pcrfit_single - A function to extract features from an amplification curve

Description

The pcrfit_single is responsible for the extraction of features from amplification curve data. The function can be used for custom functions for a paralleled analysis of amplification curve data. An example is given in the vignette.

Usage

pcrfit_single(x)

Arguments

Х

is the data set containing the fluorescence amplitudes.

Details

Details can be found in the vignette.

Value

Output Description

"cpD1"	maximum of the first derivative curve
"cpD2"	maximum of the second derivative curve
"cpD2_approx"	maximum of the second derivative curve calculated by the approximate derivative
"cpD2_ratio"	a value calculated from the ratio between cpD2 and cpD2_approx
"eff"	qPCR amplification efficiency
"sliwin"	qPCR amplification efficiency according the the 'window-of-linearity' method by Ruijter et al. (200
"cpDdiff"	absolute difference between cpD1 and cpD2
"loglin_slope"	slope determined by a linear model of the data points from the minimum and maximum of the secon
"cpD2_range"	cycle difference between the maximum and the minimum of the second derivative curve
"top"	takeoff point. When no top can be determined, the tob value is set to the first cycle number.
"f.top"	fluorescence at takeoff point. When no f.tdp can be determined, the f.tdp value is set to the RFU val
"tdp"	takes the maximum fluorescence subtracted by reverse values of the fluorescence and calculates the
"f.tdp"	fluorescence at tdp point. When no f.tdp can be determined, the f.tdp value is set to the RFU value a
"bg.stop"	estimates the end (cycle) the amplification curve background based on the bg.max function and nor
"amp.stop"	estimates the end (cycle) of the amplification curve based in the bg.max function and normalizes it t
"b_slope"	Is the slope of the seven parameter model
"b_model_param"	Is the b model parameter of the model optimally fitted according to the AIC
"c_model_param"	Is the c model parameter of the model optimally fitted according to the AIC
"d_model_param"	Is the d model parameter of the model optimally fitted according to the AIC
"e_model_param"	Is the e model parameter of the model optimally fitted according to the AIC
"f_model_param"	Is the f model parameter of the model optimally fitted according to the AIC
"f_intercept"	Is the intercept of the seven parameter model
"convInfo_iteratons"	Number of iterations needed to fit the 7 parameter model
"qPCRmodel"	non-linear model determined for the analysis
"qPCRmodelRF"	non-linear model determined for the analysis of the reversed amplification curve
"minRFU"	minimum of fluorescence amplitude
"maxRFU"	maximum of fluorescence amplitude
"init2"	initial template fluorescence from an exponential model
"fluo"	raw fluorescence value at the point defined by cpD2
"slope_bg"	slope of the first cycles
"k1_model_param"	Is the k1 model parameter of the seven parameter model
"k2_model_param"	Is the k2 model parameter of the seven parameter model
"intercept_bg"	intercept of the first cycles
"sigma_bg"	sigma of background
"sd_bg"	standard deviation of the background (ground phase) region (start to takeoff point)
"head2tail_ratio"	ratio between the signal of the background and tail region
"mblrr_intercept_bg"	the value of the intercept in the estimated background region of the amplification curve
"mblrr_slope_bg"	the value of the slope in the estimated background region of the amplification curve
"mblrr_cor_bg"	the value of the linear correlation coefficient in the estimated background region of the amplification
"mblrr_intercept_pt"	the value of the intercept in the estimated plateau phase of the amplification curve
"mblrr_slope_pt"	the value of the slope in the estimated plateau phase of the amplification curve

the value of the linear correlation coefficient in the estimated plateau phase of the amplification curv "mblrr_cor_pt" "polyarea" area of a polygon given by the vertices in the vectors cycles and fluorescence "peaks_ratio" Takes the estimate approximate local minimums and maximums is a value of autocorrelation of a gain curve from a quantitative PCR experiment "autocorrelation" "cp_e.agglo" agglomerative hierarchical estimate for multiple change points "cp_bcp" change point by Bayesian analysis methods "amptester_shapiro" tests based on the Shapiro-Wilk normality test if the amplification curve is just noise "amptester lrt" performs a cycle dependent linear regression and determines if the coefficients of determination dev Resids growth test (RGt) tests if fluorescence values in a linear phase are stable "amptester_rgt" "amptester_tht" Threshold test (THt) takes the first 20 percent and the last 15 percent of any input data set and perfo "amptester_slt" Signal level test compares 1. the signals by a robust "sigma" rule by median + 2 * mad and 2. by co pco test (pco) determines if the points in an amplification curve (like a polygon, in particular non-cc "amptester_polygon" "amptester_slope.ratio" SIR uses the inder function to find the approximated first derivative maximum, second derivative mi "hookreg_hook" estimate of hook effect like curvature "hookreg_hook_slope" estimate of slope of the hook effect like curvature "hookreg_hook_delta" Estimated value for the number of cycles from the qPCR cycle where the hook effect was determine "central_angle" shows the central angle calculated from the maximum and minimum of the second derivatives, with "sd_bg" shows the standard deviation of the fluorescence in the ground phase "number_of_cycles" Number of cylces test if the maximum of the first derivative is positive or negative "direction" "range" outputs the difference of fluorescence between 0.99 and 0.01 percentile. The value thus corresponds "polyarea_trapz" calculates trapezoidal integration. The calculation stops when the difference from one step to the ne "cor" is the value of the correlation coefficient from a linear correlation analysis according to Pearson between the second seco "res_coef_pcrfit.b" is the parameter from the adjustment with a nonlinear (sigmoid) four-parametric model which descr "res_coef_pcrfit.c" is the parameter from the adjustment with a nonlinear (sigmoid) four-parametric model which descr "res_coef_pcrfit.d" is the parameter from the adjustment with a nonlinear (sigmoid) four-parametric model which descr is the parameter from the adjustment with a nonlinear (sigmoid) four-parametric model, which desc "res_coef_pcrfit.e" "fitAIC" is the value of the Akaike's second-order corrects Information Criterion, which was determined on a "fitIter" Number of iterations needed to fit the 4 parameter model "segment_x" Adjusts a regression model with segmented (linear) relationships between fluorescence and PCR cy "segment_U1.x" Adjusts a regression model with segmented (linear) relationships between fluorescence and PCR cy "segment_U2.x" Adjusts a regression model with segmented (linear) relationships between fluorescence and PCR cy "segment_psi1.x" Adjusts a regression model with segmented (linear) relationships between fluorescence and PCR cy "segment_psi2.x" Adjusts a regression model with segmented (linear) relationships between fluorescence and PCR cy-"sumdiff" describes proportion of cycles x in which the fluorescence signal of x is smaller than in x+1is a value of a third-order polynomial $a + b^*x + c^*x^2 + d^*x^3$ is fitted to the curve data, where the "poly_1" "poly_2" is a value of a third-order polynomial $a + b^*x + c^*x^2 + d^*x^3$ is fitted to the curve data, where the "poly_3" is a value of a third-order polynomial $a + b^*x + c^*x^2 + d^*x^3$ is fitted to the curve data, where the "poly 4" is a value of a third-order polynomial $a + b^*x + c^*x^2 + d^*x^3$ is fitted to the curve data, where the The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_1" "window_Win_2" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window Win 3" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_4" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_5" "window Win 6" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_7" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_8" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s "window_Win_9" The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s

pcrfit_single

"window_Win_10"The complete curve trajectory is segmented into 10 equidistant windows by fitting an interpolating s"sd_plateau"describes the standard deviation in the late phase of an amplification curve (last five cycles). With id

gives a data.frame (S3 class, type of list) as output for the curve features

Author(s)

Stefan Roediger, Michal Burdukiewcz

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S. Roediger, M. Burdukiewicz, P. Schierack, *chipPCR: an R package to pre-process raw data of amplification curves, Bioinformatics.* 31 (2015) 2900–2902. doi:10.1093/bioinformatics/btv205.

See Also

bcp bg.max,amptester,smoother e.agglo diffQ,mcaPeaks,diffQ2 head2tailratio,earlyreg,hookreg,hookregNL,mblr polyarea pcrfit,takeoff,sliwin,efficiency diff quantile segmented

Examples

```
# Load the chipPCR package and analyze from the C126EG685 the first qPCR run
# "A01" (column 2).
library(chipPCR)
res <- pcrfit_single(C126EG685[, 2])</pre>
```

performeR

Description

This function performs an analysis sensitivity and specificity to asses the performance of a binary classification test. For further reading the studies by Brenner and Gefeller 1997, James 2013 by Kuhn 2008 are a good starting point.

Usage

performeR(sample, reference)

Arguments

sample	is a vector with logical decisions $(0, 1)$ of the test system.
reference	is a vector with logical decisions $(0, 1)$ of the reference system.

Details

TP, true positive; FP, false positive; TN, true negative; FN, false negative Sensitivity - TPR, true positive rate TPR = TP / (TP + FN) Specificity - SPC, true negative rate SPC = TN / (TN + FP) Precision - PPV, positive predictive value PPV = TP / (TP + FP) Negative predictive value - NPV NPV = TN / (TN + FN) Fall-out, FPR, false positive rate FPR = FP / (FP + TN) = 1 - SPC False negative rate - FNR FNR = FN / (TN + FN) = 1 - TPR False discovery rate - FDR FDR = FP / (TP + FP) = 1 - PPV Accuracy - ACC ACC = (TP + TN) / (TP + FP + FN + TN) F1 score F1 = 2TP / (2TP + FP + FN) Likelihood ratio positive - LRp LRp = TPR/(1-SPC) Matthews correlation coefficient (MCC) MCC = (TP*TN - FP*FN) / sqrt(TN + FP) * sqrt(TN+FN)) Cohen's kappa (binary classification) kappa=(p0-pc)/(1-p0)

r (reference) is the trusted label and s (sample) is the predicted value

	r=1	r=0
s=1	а	b
s=0	с	d

$$n = a + b + c + d$$

```
pc=((a+b)/n)((a+c)/n)+((c+d)/n)((b+d)/n)
po=(a+d)/n
```

Value

gives a data.frame (S3 class, type of list) as output for the performance

Author(s)

Stefan Roediger, Michal Burdukiewcz

References

H. Brenner, O. Gefeller, others, Variation of sensitivity, specificity, likelihood ratios and predictive values with disease prevalence, *Statistics in Medicine*. 16 (1997) 981–991.

M. Kuhn, Building Predictive Models in R Using the caret Package, *Journal of Statistical Software*. 28 (2008). doi:10.18637/jss.v028.i05.

G. James, D. Witten, T. Hastie, R. Tibshirani, An Introduction to Statistical Learning, *Springer New York, New York, NY*, (2013). doi:10.1007/978-1-4614-7138-7.

Examples

```
# Do the statistical analysis with the performeR function
performeR(sample=test_data, reference=reference_data)
```

qPCR2fdata

A helper function to convert amplification curve data to the fdata format.

Description

qPCR2fdata is a helper function to convert qPCR data to the functional fdata class as proposed by Febrero-Bande & de la Fuente (2012). This function prepares the data for further analysis with the fda.usc package, which includes utilities for functional data analysis (e.g., Hausdorff distance).

qPCR2fdata

Usage

qPCR2fdata(data, preprocess = FALSE)

Arguments

data	is a data set containing the amplification cycles (1. column) and the fluorescence (subsequent columns).
preprocess	is a logical parameter (default FALSE). If TRUE, the CPP function from the chipPCR package (Roediger et al. 2015) is used to pre-process the data (e.g., imputation of missing values). and the fluorescence (subsequent columns).

Value

gives an fdata object (S3 class, type of list) as output for a converted amplification curve.

Author(s)

Stefan Roediger, Michal Burdukiewcz

References

M. Febrero-Bande, M.O. de la Fuente, others, *Statistical computing in functional data analysis: The R package fda.usc*, Journal of Statistical Software. 51 (2012) 1–28. http://www.jstatsoft.org/v51/i04/

S. Roediger, M. Burdukiewicz, P. Schierack, *chipPCR: an R package to pre-process raw data of amplification curves*, Bioinformatics. 31 (2015) 2900–2902. doi:10.1093/bioinformatics/btv205.

Examples

```
default.par <- par(no.readonly = TRUE)</pre>
# Calculate slope and intercept on noise (negative) amplification curve data
# for the last eight cycles.
library(qpcR)
library(fda.usc)
# Convert the gPCR data set to the fdata format
res_fdata <- qPCR2fdata(testdat)</pre>
# Extract column names and create rainbow color to label the data
res_fdata_colnames <- colnames(testdat[-1])</pre>
data_colors <- rainbow(length(res_fdata_colnames), alpha=0.5)</pre>
# Plot the converted gPCR data
par(mfrow=c(1,2))
plot(res_fdata, xlab="cycles", ylab="RFU", main="testdat", type="l",
                   lty=1, lwd=2, col=data_colors)
legend("topleft", as.character(res_fdata_colnames), pch=19,
         col=data_colors, bty="n", ncol=2)
# Calculate the Hausdorff distance (fda.usc) package and plot the distances
```

tReem

PCRedux app

run_PCRedux

Description

A graphical user interface for computing the properties of amplification curves. Take a look at the vignette to learn more about the different ways to start the app.

Usage

run_PCRedux()

Value

null.

No return value, called for side effects

Note

Any ad-blocking software may cause malfunctions.

See Also

encu, runApp.

tReem

A function to Group Amplification Curves According to their Shape

Description

tReem is a function to group amplification curves from a quantitative PCR experiment according to their shape. Either the Pearson correlation coefficient or the Hausdorff distance is used as measure. In most cases the grouping based on the Pearson correlation coefficient is sufficient. The grouping based on the Hausdorff distance can be very slow for large data sets.

Usage

tReem(data, cor = TRUE, k = 2)

winklR

Arguments

data	is the cycle dependent fluorescence amplitude (y-axis).
cor	is a logical parameter. If set true, the Pearson correlation is used as distance measure. If set FALSE the Hausdorff distance will be used.
k	an integer scalar or vector with the desired number of groups.

Value

gives a data.frame (S3 class, type of list) as output for the manual analyzed data

Author(s)

Stefan Roediger, Andrej-Nikolai Spiess

See Also

metric.hausdorff, cutree, qPCR2fdata, hclust, cor

Examples

Classify amplification curve data by Hausdorff distance

library(qpcR)
tReem(testdat[, 1:5])

winklR

winklR: A function to calculate the angle based on the first and the second derivative of an amplification curve data from a quantitative *PCR* experiment.

Description

winklR is a function to calculate the in the trajectory of the first and the second derivatives maxima and minima of an amplification curve data from a quantitative PCR experiment. For the determination of the angle (central angle), the origin is the maximum of the first derivative. On this basis, the vectors to the minimum and maximum of the second derivative are determined. This means that systematic off-sets, such as those caused by background, are taken into account. The output contains the angle.

Usage

winklR(x, y, normalize = FALSE, preprocess = TRUE)

Arguments

х	is the cycle numbers (x-axis). By default the first ten cycles are removed.
У	is the cycle dependent fluorescence amplitude (y-axis).
normalize	is a logical parameter, which indicates if the amplification curve data should be normalized to the 99 percent percentile of the amplification curve.
preprocess	is a logical parameter, which indicates if the amplification curve data should be smoothed (moving average filter, useful for noisy, jagged data).

Value

gives a list (S3 class, type of list) as output for the angles from an amplification curve.

Author(s)

Stefan Roediger

See Also

acos diffQ2

Examples

Calculate the angles for amplification curve data from the RAS002 data set data(RAS002)

```
# Plot the data
plot(RAS002[, 1],
  y = RAS002[, 2], xlab = "Cycle", ylab = "RFU",
 main = "RAS002 data set", lty = 1, type = "l"
)
res <- winklR(x = RAS002[, 1], y = RAS002[, 2])</pre>
res
plot(rbind(res$origin, res$p1, res$p2), col = c("black", "green", "blue"))
plot(RAS002[, 1],
 y = RAS002[, 7], xlab = "Cycle", ylab = "RFU",
  main = "RAS002 data set", lty = 1, type = "1"
)
res <- winklR(x = RAS002[, 1], y = RAS002[, 7])</pre>
res
plot(rbind(res$origin, res$p1, res$p2), col = c("black", "green", "blue"))
res_angles <- unlist(lapply(2:21, function(i) {</pre>
  winklR(RAS002[, 1], RAS002[, i])$angle
}))
cdplot(RAS002_decisions[1L:20] ~ res_angles, xlab = "angle", ylab = "decision")
```

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