Package 'pooledpeaks'

November 20, 2024

```
Title Genetic Analysis of Pooled Samples
Version 1.0.7
Description Analyzing genetic data obtained from pooled samples.
     This package can read in Fragment Analysis output files, process the data,
     and score peaks, as well as facilitate various analyses, including cluster
     analysis, calculation of genetic distances and diversity indices,
     as well as bootstrap resampling for statistical inference. Specifically
     tailored to handle genetic data efficiently, researchers can explore
     population structure, genetic differentiation, and genetic relatedness
     among samples. We updated some functions from Covarrubias-Pazaran et al.
     (2016) <doi:10.1186/s12863-016-0365-6> to allow for the use of new file formats
     and referenced the following to write our genetic analysis functions:
     Long et al. (2022) <doi:10.1038/s41598-022-04776-0>, Jost (2008)
     <doi:10.1111/j.1365-294x.2008.03887.x>, Nei (1973)
     <doi:10.1073/pnas.70.12.3321>, Foulley et al. (2006)
     <doi:10.1016/j.livprodsci.2005.10.021>, Chao et al. (2008)
     <doi:10.1111/j.1541-0420.2008.01010.x>.
URL https://github.com/kmkuesters/pooledpeaks
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AlRich

Calculate Allelic Richness

Description

This function calculates allelic richness based on provided genetic data.

Usage

```
AlRich(datafile = data.frame, n = matrix)
```

Arguments

datafile A data frame containing the data as read in by LoadData

n

A matrix representing the number of markers successfully genotyped like the output of the TypedLoci function.

Value

A vector containing the allelic richness for each locus.

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)

n_alleles <- matrix(c(
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
Alrich(datafile=genetic_data,n=n_alleles)</pre>
```

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associate_dyes

Associate Dye Names in Batch Import Output

Description

This function associates dye info with fragman channel names. It was designed to be performed on any fsa formats after final columns are correctly imported.

Usage

```
associate_dyes(x, y)
```

Arguments

- x The Output list of data frames from fsa_batch_imp.
- y The path to the folder from the current directory where the .fsa files that will be analyzed are stored.

Value

The input dataframe with an added column assigning fluorescent dye colors.

Examples

```
y \leftarrow system.file("extdata", package = "pooledpeaks")
 x \leftarrow fsa\_batch\_imp(y, channels = 5, fourier = FALSE, saturated = FALSE, lets.pullup = FALSE, plotting = FALSE, rawPlot = FALSE, llength = 3000, ulength = 80000) associate_dyes(x,y)
```

BootStrap3

Perform Bootstrap Analysis

Description

This function performs bootstrap analysis on genetic data.

Usage

```
BootStrap3(A = data.frame, Rep = 20, Stat = 1)
```

Arguments

A Data frame containing data as read in by LoadData

Rep Number of bootstrap replicates.

Stat Type of statistic to compute (1 for AlRich, 2 for TwoLevelGST)

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Value

Either a matrix of AlRich statistics or a list containing various statistics computed using TwoLevel-GST

Examples

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)
BootStrap3(A=genetic_data, Rep=10, Stat=1)
BootStrap3(A=genetic_data, Rep=10, Stat=2)</pre>
```

check_fsa_v_batch

Check .fsa Version and Batch Information

Description

This function analyzes .fsa files in a specified folder, providing a summary of their version and batch information.

Usage

```
check_fsa_v_batch(x)
```

Arguments

Х

The path to the folder from the current directory where the .fsa files that will be analyzed are stored.

Value

A written summary of how many .fsa files are in the folder and which version they are.

```
file_path <- system.file("extdata", package = "pooledpeaks")
check_fsa_v_batch(x = file_path)</pre>
```

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clean_scores

Clean Scores Data

Description

This function cleans the score_markers_rev3 data by applying specified patterns and replacements to the ID and filename columns.

Usage

```
clean_scores(
   scores_data,
   pattern1 = NULL,
   replacement1 = NULL,
   pattern2 = NULL,
   replacement2 = NULL,
   pattern3 = NULL,
   replacement3 = NULL
)
```

Arguments

scores_data The list containing the output scores data from the score_markers_rev3.

pattern1 The first pattern to replace in the ID. This is intended to clean up the ID names for

when the machine adds substrings to the names. For example 104.1a_FA060920_2020-

06-09_C05.fsa.1 becomes 104.1a using pattern1="_FA.*" and replacement1=""

replacement 1 Replacement for the first pattern.

pattern2 The second pattern to replace in the ID. See pattern1 for more details.

replacement 2 Replacement for the second pattern.

pattern3 The pattern to replace in the file name. This is intended to clean up the file names

for when the machine adds sub strings to the names. For example 104.1a_FA060920_2020-

06-09_C05.fsa.1 becomes 104.1a_FA060920_2020-06-09_C05.fsa using pattern3=

"\.1*\$" and replacement3= ""

replacement3 Replacement for the file name pattern.

Value

A cleaned long format data frame

```
scores_data <- list(
data.frame(Score = c(90, 85, 70), stringsAsFactors = FALSE),
data.frame(Score = c(80, 75, 60), stringsAsFactors = FALSE)
)
rownames(scores_data[[1]]) <- c("104.1a_FA060920_2020-06-09_C05.fsa_Sa.1",</pre>
```

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cluster

K-means Clustering

Description

K-means Clustering

Usage

```
cluster(RawData = data.frame, K = 2)
```

Arguments

RawData A data frame containing the raw data as read in by LoadData

K An integer specifying the number of clusters.

Value

A list containing the results of the K-means cluster analysis, including cluster assignments and original data.

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)
cluster(RawData=genetic_data, K=2)</pre>
```

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ClusterFromSamples

Cluster From Samples

Description

Perform clustering on samples of loci from a data frame and calculate statistics.

Usage

```
ClusterFromSamples(datafile = data.frame, numloci = 5, reps = 100)
```

Arguments

datafile A data frame containing the input data must be in LoadData style LoadData.

numloci An integer specifying the number of loci to sample.

reps An integer specifying the number of repetitions.

Value

A matrix containing statistics calculated from the clustering results.

Examples

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)
ClusterFromSamples(datafile=genetic_data, numloci=5, reps=10)</pre>
```

data_manipulation

Data Manipulation for Marker Data

Description

This function ensures that at least one peak for each sample is greater than a specified threshold (default: 500) and then formats the data frame for the next steps in the analysis.

Usage

```
data_manipulation(marker, threshold = 500)
```

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Arguments

marker A data frame containing marker data, where each row represents a marker and

each column represents a sample.

threshold The threshold value for peak height. Peaks below this threshold will be replaced

with 0.

Value

A formatted data frame where at least one peak for each sample is greater than the specified threshold.

Examples

```
marker_data <- data.frame(
Sample1 = c(400, 600, 700,0),
Sample2 = c(450, 550, 480,0),
Sample3 = c(300, 200, 400,200),
Sample4 = c(0,0,0,0),
row.names=c(185,188,191,194)
)
data_manipulation(marker_data,threshold=500)</pre>
```

DistCor

Distance Correlation

Description

Calculate the correlation between expected and realized genetic distances and plot them.

Usage

```
DistCor(GD = matrix)
```

Arguments

GD

A matrix containing the genetic distance data.

Value

A plot showing the Expected Genetic Distance versus Realized Genetic Distance

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Examples

```
genetic_distance_matrix <- matrix(c(0.316455, 0.2836333, 0.2760485,
0.2685221, 0.2797302,0.3202661,0.2836333, 0.3106084, 0.2867215, 0.2687472,
0.2596309, 0.2957862,0.2760485,0.2867215, 0.3338663, 0.297918, 0.3057039,
0.3153261,0.2685221, 0.2687472, 0.297918,0.3107094, 0.2753477, 0.3042383,
0.2797302, 0.2596309, 0.3057039, 0.2753477, 0.3761386, 0.3398558,0.3202661,
0.2957862, 0.3153261, 0.3042383, 0.3398558, 0.4402125),
nrow = 6, byrow = TRUE,dimnames = list(c("Sample1", "Sample2", "Sample3",
    "Ind1", "Ind2", "Ind3"),
    c("Sample1", "Sample2", "Sample3", "Ind1", "Ind2", "Ind3")))</pre>
DC<- DistCor(genetic_distance_matrix)
```

EmpiricalSE

Calculate Empirical Standard Error

Description

This function calculates the empirical standard error based on repeated sampling.

Usage

```
EmpiricalSE(datafile = data.frame, NLoci = 10)
```

Arguments

datafile A data frame containing genetic data from the LoadData

NLoci Number of loci to sample in each iteration.

Value

A numeric vector containing the empirical standard error estimates.

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)</pre>
EmpiricalSE(datafile=genetic_data, NLoci=3)
```

fsa_batch_imp

fsa_batch_imp	Batch Import of .fsa files	

Description

This function imports and extracts all of the information out of the .fsa files and combines them into one list type object.fsa_batch_imp is a modification of the original Fragman import script function, storing.inds, This revised script accommodates ABI's .fsa file format up to version 3. It retains Fragman functions for Fourier transformation, saturated peaks, and pull-up correction. Notable adjustments include updating channel parameters, utilizing Dyechannel count from the file directory, and streamlining the script by extracting data only from "DATA" tags. Major changes involve column selection for v3 formats and modifications to the "channel" parameter. Minor changes include allowing relative paths for the data directory, importing only .fsa files, and renaming channels with dye names. This revision ensures successful execution for any format version up to 3.

Usage

```
fsa_batch_imp(
  folder,
  channels = NULL,
  fourier = TRUE,
  saturated = TRUE,
  lets.pullup = FALSE,
  plotting = FALSE,
  rawPlot = FALSE,
  llength = 3000,
  ulength = 80000
)
```

Arguments

folder	The path to the folder from the current directory where the .fsa files that will be analyzed are stored.
channels	The number of dye channels expected, including the ladder.
fourier	True/False Should fourier transformation be applied.
saturated	True/False whether to Check and correct for saturated peaks.
lets.pullup	True/False Applying pull up correction to the samples to decrease noise from channel to channel. The default is FALSE, please do not change this.
plotting	True/False Should plots be drawn of all channels after data cleaning.
rawPlot	True/False indicating whether a plot should be drawn of all vectors.
llength	A numeric value for the minimum number of indexes in each channel.
ulength	A numeric value for the maximum number fo indexes in each channel.

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Value

Output is a LIST where each element of the list is a DATAFRAME with the channels in columns for each FSA file

Examples

```
file_path <- system.file("extdata", package = "pooledpeaks")
fsa_batch_imp(file_path, channels = 5, fourier = FALSE, saturated = FALSE ,
lets.pullup = FALSE,plotting = FALSE, rawPlot = FALSE)</pre>
```

fsa_metadata

Retrieve Metadata

Description

Retrieves basic info from .fsa files about the sample and run,and aggregates multiple samples in a single object.

Usage

```
fsa_metadata(x)
```

Arguments

Χ

The path to the folder from the current directory where the .fsa files that will be analyzed are stored.

Value

A data frame that contains the metadata of the machine and run extracted from the .fsa file. One row for each .fsa file in directory x and the following columns: retrieved_sample_name, batch_container_name, fsa_version, user, run_start_date, run_start_time, machine_type,machineN_serial.

```
file_path <- system.file("extdata", package = "pooledpeaks")
fsa_metadata(x = file_path)</pre>
```

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GeneIdentityMatrix

Gene Identity Matrix

Description

Using the number of typed loci, this function calculates the gene identity between all possible pairwise combinations between individuals for all markers creating a matrix.

Usage

```
GeneIdentityMatrix(RawData = data.frame, LociGenotyped = matrix)
```

Arguments

RawData A data frame containing the input data must be in LoadData style LoadData.

LociGenotyped The Output from the TypedLoci function

Value

The Gene Identity Matrix

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)

n_alleles <- matrix(c(
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
3, 3, 3,
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3, 3, 3, 3,
```

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GeneticDistanceMatrix Genetic Distance Matrix

Description

This function calculates the genetic distance matrix from a given gene identity matrix.

Usage

```
GeneticDistanceMatrix(J = matrix)
```

Arguments

J The Gene Identity Matrix created using GeneIdentityMatrix

Value

The Genetic Distance Matrix

Examples

```
gene_identity_matrix <- matrix(c(
0.3164550, 0.2836333, 0.2760485,
0.2836333, 0.3106084, 0.2867215,
0.2760485, 0.2867215, 0.3338663
), nrow = 3, byrow = TRUE,
dimnames = list(paste0("Sample", 1:3), paste0("Sample", 1:3)))
GeneticDistanceMatrix(gene_identity_matrix)</pre>
```

GST

Nei's GST

Description

This function calculates GST (Nei's standard genetic distance) measure from a gene identity matrix.

Usage

```
GST(J = matrix, pairwise = TRUE)
```

Arguments

J A square matrix representing a gene identity matrix.

pairwise Logical indicating whether to calculate pairwise GST. If set to FALSE, must not

contain any missing data.

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Value

If pairwise = TRUE, returns a matrix of pairwise GST values. If pairwise = FALSE, returns the overall GST value.

Examples

```
gene_identity_matrix <- matrix(c(
0.3164550, 0.2836333, 0.2760485,
0.2836333, 0.3106084, 0.2867215,
0.2760485, 0.2867215, 0.3338663
), nrow = 3, byrow = TRUE,
dimnames = list(paste0("Sample", 1:3), paste0("Sample", 1:3)))

GST(J=gene_identity_matrix, pairwise=TRUE)
GST(J=gene_identity_matrix, pairwise=FALSE)</pre>
```

JostD

Calculate Jost's D

Description

This function calculates Jost's D measure from a gene identity matrix.

Usage

```
JostD(J = matrix, pairwise = TRUE)
```

Arguments

J A gene identity matrix.

pairwise Logical indicating whether to calculate pairwise Jost's D. If pairwise=FALSE,

must not have any missing data.

Value

If pairwise = TRUE, returns a matrix of pairwise Jost's D values. If pairwise = FALSE, returns the overall Jost's D value.

```
gene_identity_matrix <- matrix(c(
0.3164550, 0.2836333, 0.2760485,
0.2836333, 0.3106084, 0.2867215,
0.2760485, 0.2867215, 0.3338663
), nrow = 3, byrow = TRUE,
dimnames = list(paste0("Sample", 1:3), paste0("Sample", 1:3)))

JostD(J=gene_identity_matrix, pairwise=TRUE)
JostD(J=gene_identity_matrix, pairwise=FALSE)</pre>
```

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JostD_KK

Pairwise Jost D between replicates

Description

This function calculates Jost's D between two columns, specifically designed for comparing duplicate samples based on allele frequencies.

Usage

```
JostD_KK(Ni1, Ni2)
```

Arguments

Ni 1 Vector containing the allele frequencies of the first duplicate sample.

Ni2 Vector containing the allele frequencies of the second duplicate sample.

Value

The calculated Jost's D value.

lf_to_tdf

Transform LF to TDF

Description

This function transforms a data frame from LF (long format) to TDF (table format), performing various data manipulation steps such as spreading data across columns, removing NA and/or 0 columns, merging ID allele heights within each replicate, transposing the table, converting from character to numeric class, and replacing empty data with "0".

Usage

```
lf_to_tdf(x)
```

Arguments

Х

A data frame in LF format ideally coming out of the clean_scores function.

Value

A transformed data frame in TDF format.

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Examples

```
scores<- data.frame(ID=c("104.1a","105.2b","106.3c","107.4d","108.5e",
"109.6f"),
filename=c("104.1a_FA060920_2020-06-09_C05.fsa_Sa.1",
"105.2b_FA060920_2020-06-09_C05.fsa_Sa.1",
"106.3c_FA060920_2020-06-09_C05.fsa_Fa.1",
"107.4d_FA060920_2020-06-09_C05.fsa_Sa.1"
"108.5e_FA060920_2020-06-09_C05.fsa_Sa.1"
"109.6f_SA060920_2020-06-09_C05.fsa_Fa.1"),
hei=c(2000,3000,4000,5000,2500, 1000),
pos=c(2000,3000,4000,5000,2500, 1000),
wei=c(290,285,280,275,270,260),
row.names= c("104.1a_FA060920_2020-06-09_C05.fsa_Sa.1",
"105.2b_FA060920_2020-06-09_C05.fsa_Sa.1",
"106.3c_FA060920_2020-06-09_C05.fsa_Fa.1",
"107.4d_FA060920_2020-06-09_C05.fsa_Sa.1" ,
"108.5e_FA060920_2020-06-09_C05.fsa_Sa.1" ,
"109.6f_SA060920_2020-06-09_C05.fsa_Fa.1"))
lf_to_tdf(scores)
```

LoadData

Load Genetic Data

Description

This function imports data for genetic analysis.

Usage

```
LoadData(datafile = NULL)
```

Arguments

datafile

The path to your datafile. The format of your data should be .txt or .csv.

Value

A data frame containing the imported data formatted in the way necessary for downstream population genetic functions.

```
file<-system.file("extdata", "Multiplex_frequencies.txt",
package = "pooledpeaks")
LoadData(file)</pre>
```

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MDSplot

Multi Dimensional Scaling (MDS) Plot

Description

Generate a multidimensional scaling (MDS) plot from genetic distance data.

Usage

```
MDSplot(
  distance = matrix,
  pcs = c(1, 2),
  PF = NULL,
  y = c("dodgerblue", "red", "turquoise3", "purple", "olivedrab3")
)
```

Arguments

distance A matrix containing the genetic distance data.

pcs A numeric vector specifying the principal coordinates to plot.

PF A factor vector specifying population labels.

y A character vector specifying colors for population labels.

Value

The output is the MDS plot for the samples for the specified principal coordinates.

```
genetic_distance_matrix <- matrix(c(
0, 0.2836333, 0.2760485, 0.2685221, 0.2797302,0.3202661,
0.2836333, 0, 0.2867215, 0.2687472, 0.2596309, 0.2957862,
0.2760485,0.2867215, 0, 0.297918, 0.3057039, 0.3153261,
0.2685221, 0.2687472, 0.297918,0, 0.2753477, 0.3042383,
0.2797302, 0.2596309, 0.3057039, 0.2753477, 0,0.3398558,
0.3202661, 0.2957862, 0.3153261, 0.3042383, 0.3398558, 0),
nrow = 6, byrow = TRUE,dimnames = list(c("Sample1", "Sample2", "Sample3", "Ind1", "Ind2", "Ind3"),
c("Sample1", "Sample2", "Sample3", "Ind1", "Ind2", "Ind3")))
MDSplot(distance=genetic_distance_matrix, pcs=c(1,3))</pre>
```

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PCDM

Post-consolidation Data Manipulation

Description

This function manipulates consolidated marker data and egg count data to prepare them for further analysis.

Usage

```
PCDM(consolidated_marker = data.frame, eggcount = data.frame, marker_name)
```

Arguments

consolidated_marker

A data frame containing consolidated marker data.

eggcount A data frame containing egg count data.

marker_name A string specifying the marker name.

Value

A dataframe containing the allele frequencies and eggcounts for each sample.

Examples

```
marker_data <- data.frame(
Sample1 = c(400, 600, 700),
Sample2 = c(450, 550, 480),
Sample3 = c(300, 200, 500),
row.names=c(185,188,191)
)

eggs<-data.frame(
ID=c("Sample1","Sample2","Sample3"),n=c(3000,400,50))

PCDM(consolidated_marker=marker_data, eggcount= eggs,"SMMS2")</pre>
```

preGST

Pre GST Calculation

Description

This function calculates the GST from a gene identity matrix.

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Usage

```
preGST(G = matrix)
```

Arguments

G

The gene identity matrix

Value

The GST value.

preJostD

Calculate Pre-Jost's D

Description

This function calculates the pre-Jost's D measure from a gene identity matrix.

Usage

```
preJostD(G = matrix)
```

Arguments

G

A square matrix representing a gene identity matrix.

Value

The Jost's D value.

Rep_check

Replicate Check for Duplicate Samples

Description

This function checks for duplicate samples in the input data frame and calculates the average peak heights for each sample. If the Jost's D between duplicate samples exceeds 0.05, it flags those samples.

Usage

```
Rep_check(df)
```

Arguments

df

The input data frame containing peak heights for each sample.

RWCDistanceMatrix 21

Value

A data frame containing the average peak heights for each sample, with flagged samples where duplicates have a Jost's D exceeding 0.05.

Examples

```
marker_data <- data.frame(
Sample.1a = c(400, 600, 700),
Sample.1b = c(420, 606, 710),
Sample.2a = c(450, 550, 480),
Sample.2b = c(500, 540, 480),
Sample.3a = c(300, 200, 500),
Sample.3b = c(290, 100, 400),
row.names=c(185,188,191)
)</pre>
Rep_check(marker_data)
```

RWCDistanceMatrix

Random Walk Covariance Distance Matrix

Description

This function calculates the RWC (Random Walk Covariance) distance matrix from a given matrix of genetic distances.

Usage

```
RWCDistanceMatrix(J = matrix)
```

Arguments

J

The Genetic Distance Matrix calculated using GeneticDistanceMatrix

Value

A matrix representing the distance matrix calculated using the Random Walk Covariance method.

```
genetic_distance_matrix <- matrix(c(0.316455, 0.2836333, 0.2760485,
0.2685221, 0.2797302,0.3202661,0.2836333, 0.3106084, 0.2867215, 0.2687472,
0.2596309, 0.2957862,0.2760485,0.2867215, 0.3338663, 0.297918, 0.3057039,
0.3153261,0.2685221, 0.2687472, 0.297918,0.3107094, 0.2753477, 0.3042383,
0.2797302, 0.2596309, 0.3057039, 0.2753477, 0.3761386,0.3398558,0.3202661,
0.2957862, 0.3153261, 0.3042383, 0.3398558, 0.4402125),
nrow = 6, byrow = TRUE, dimnames = list(c("Sample1", "Sample2", "Sample3",
"Ind1", "Ind2", "Ind3"),</pre>
```

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```
c("Sample1", "Sample2", "Sample3", "Ind1", "Ind2", "Ind3")))
RWCDistanceMatrix(genetic_distance_matrix)
```

SampleOfLoci

Sample Of Loci

Description

An internal function that supports ClusterFromSamples. Sample loci from a dataset based on the number of loci specified.

Usage

```
SampleOfLoci(aaax = data.frame, NLoci = max(aaax[, 1]))
```

Arguments

aaax

A data frame containing the input data must be in LoadData style LoadData.

NLoci

An integer specifying the number of loci to sample.

Value

A data frame containing the sampled loci.

score_markers_rev3

Score Markers Wrapper

Description

This is a revision of the Fragman script score.markers, for the original instructions and parameters, run '?score.markers'. This revision designates separate parameters for Left and Right search windows.

Usage

```
score_markers_rev3(
  my.inds,
  channel = 1,
  n.inds = NULL,
  panel = NULL,
  shift = 0.8,
  ladder,
  channel.ladder = NULL,
  ploidy = 2,
  left.cond = c(0.6, 3),
  right.cond = 0.35,
```

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```
warn = FALSE,
windowL = 0.5,
windowR = 0.5,
init.thresh = 200,
ladd.init.thresh = 200,
method = "iter2",
env = parent.frame(),
my.palette = NULL,
plotting = FALSE,
plotdir = "plots_scoring",
pref = 3
```

Arguments

my.inds	The list output from the fsa_batch_imp or storing.inds function that contains the channel information from the individuals that you want to score.
channel	The number of the channel you wish to analyze. Typically 1 is blue, 2 is green, 3 yellow, and 4 red.
n.inds	(optional) A vector specifying which fsa files to score.
panel	A vector containing the expected allele sizes for this marker.
shift	All peaks at that distance from the tallest peak will be ignored and be considered noise.
ladder	A vector containing the expected peaks for your ladder.
channel.ladder	The channel number where your ladder can be found.
ploidy	The name is a relic of the fact that Fragman::score.markers was originally writ-

left.cond

The first part is a percentile (0-1) that corresponds to the height that a peak to the left of the tallest peak must be in order to be considered real. The second argument is a number of base pairs that a peak to the left of the tallest peak must be away to be considered as real.

ten for plants. In the context of pooled egg samples it is used to specify the

right.cond

A percentile (0-1) that corresponds to the height that a peak to the right of the tallest peak must be in order to be real.

warn

TRUE/FAISE Do you want to receive warnings when detecting the ladder?

windowL

the window means that all peaks closer by that distance to the left of the panel

peaks will be accounted as peaks.

number of possible alleles in the marker.

windowR

the window means that all peaks closer by that distance to the right of the panel peaks will be accounted as peaks.

init.thresh

A value that sets a minimum intensity in order for a peak to be called.

ladd.init.thresh

We don't recommend messing with this parameter unless your ladder has special circumstances. See Fragman::score.markers

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method In cases where samples weren't sized using the info.ladder.attach function, this

technique steps in to identify ladder peaks. You have three method options using an argument: "cor" explores all potential peak combinations and thoroughly searches for correlations to identify the correct peaks corresponding to expected DNA weights; "ci" constructs confidence intervals to identify peaks meeting specified conditions from earlier arguments; "iter2" applies an iterative strategy to identify the most likely peaks aligning with your ladder expectations. The

default method is "iter2."

env Please do not change this parameter, it is used to detect the users environment.

my.palette (optional) A character vector specifying which colors to use for the output RFU

plots.

plotting TRUE/FALSE Do you want to create pdf output plots?

plotdir The name of the directory where output pdf plots should be stored.

pref The number of plots to be drawn in the output plot.

Value

The score_markers_rev3 function will return a list containing three variables: \$pos, \$hei, and \$wei. These correspond to the index position for the intensities, the intensity of each peak, and the weight in base pairs based on the ladder respectively. If plotting = TRUE, a pdf file will also have been created in the specified directory. This pdf file allows you to visually inspect how all of the peaks were scored.

```
file_path <- system.file("extdata", package = "pooledpeaks")</pre>
mock_fsa_batch_imp_output<- fsa_batch_imp(file_path, channels = 5,</pre>
fourier = FALSE, saturated = FALSE, lets.pullup = FALSE,
plotting = FALSE, rawPlot = FALSE)
panel <- c(176,179,182,185,188,191,194,197,200,203,206)
ladder <- c( 140, 160, 180, 200, 214, 220,240, 250, 260, 280, 300, 314)
mock_fsa_batch_imp_output <- associate_dyes(mock_fsa_batch_imp_output,</pre>
                              file_path)
score_markers_rev3(my.inds = mock_fsa_batch_imp_output,
                             channel = 1,
                             channel.ladder = 5,
                             panel = "panel",
                             ladder = ladder,
                             init.thresh = 200,
                             ploidy = length(panel),
                             plotting = FALSE)
```

TwoLevelGST 25

TwoLevelGST

Calculate Two-Level GST

Description

This function calculates two-level GST (Nei's standard gene identity) measure from a gene identity matrix.

Usage

```
TwoLevelGST(G = matrix)
```

Arguments

G

A square matrix representing a gene identity matrix.

Value

A list containing the components of two-level GST including within-group gene identity, between-group gene identity, and GST values.

Examples

```
gene_identity_matrix <- matrix(c(
0.3164550, 0.2836333, 0.2760485,
0.2836333, 0.3106084, 0.2867215,
0.2760485, 0.2867215, 0.3338663
), nrow = 3, byrow = TRUE,
dimnames = list(paste0("Sample", 1:3), paste0("Sample", 1:3)))
TwoLevelGST(G=gene_identity_matrix)</pre>
```

TypedLoci

Typed Loci

Description

This function calculates the number of loci successfully genotyped by each individual included in our data set

Usage

```
TypedLoci(datafile = data.frame)
```

Arguments

datafile

A data frame containing the input data must be in LoadData style LoadData.

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Value

A matrix representing processed data.

```
genetic_data <- data.frame(
Locus = c(1, 1, 1, 1, 1, 2, 2, 2, 2, 2),
Locus_allele = c("Marker1", "n", 1, 2, 3, "Marker2", "n", 1, 2, 3),
Sample1 = c(NA, 10, 0.5, 0.5, 0, NA, 10, 0.2, 0.3, 0.5),
Sample2 = c(NA, 20, 0.1, 0.2, 0.7, NA, 20, 0.3, 0.4, 0.3),
Sample3 = c(NA, 30, 0.3, 0.4, 0.3, NA, 30, 0.4, 0.2, 0.4)
)
TypedLoci(datafile=genetic_data)</pre>
```

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```