

Package ‘mQTL.NMR’

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Type Package

Title Metabolomic Quantitative Trait Locus Mapping for 1H NMR data

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Description mQTL.NMR provides a complete mQTL analysis pipeline for 1H NMR data. Distinctive features include normalisation using most-used approaches, peak alignment using RSPA approach, dimensionality reduction using SRV and binning approaches, and mQTL analysis for animal and human cohorts.

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URL <http://www.ican-institute.org/tools/>

LazyLoad yes

LazyData yes

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biocViews Cheminformatics

Depends R (>= 2.15.0)

Imports qtl, GenABEL, MASS, outliers, graphics, stats, utils

Suggests BiocStyle

R topics documented:

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mQTL.NMR-package	<i>Metabolomic Quantitative Trait Locus mapping for 1H NMR data</i>
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Description

mQTL.NMR provides a complete mQTL analysis pipeline for 1H NMR data. Distinctive features include normalisation using most-used approaches, peak alignment using RSPA approach, dimensionality reduction using SRV and binning approaches, and mQTL analysis for animal and human cohorts.

Details

Package: mQTL.NMR
 Type: Package
 Version: 0.99.2
 Link: <http://www.ican-institute.org/tools>
 Date: 2014-05-19
 License: Artistic-2.0

Main functions:

- `format_mQTL`: generates the proper format of animal crosses data
- `format_mGWA`: generates the proper format of human data
- `align_mQTL`: peak alignment
- `normalise_mQTL`: normalisation of metabolomic data using different approaches (Probabilistic quotient, constant sum,...)
- `pre_mQTL`: dimension reduction by statistical recoupling of variables or binning
- `process_mQTL`: computes LODs using extended Haley-Knott method for animal crosses
- `process_mGWA`: computes p-values using a standard linear regression approach for human
- `post_mQTL`: plots the results of a given run
- `summary_mQTL`: provides the results as a table
- `simple.plot`: Plots a region of NMR profile
- `SRV.plot`: Plots the regions identified by SRV in NMR profiles
- `ppersp`: Plot 3-D profile of LODs as function of genomic position and chemical shift
- `pplot`: Plot a color scale layer
- `Top_SRV.plot`: Plot top SRV clusters for structural assignment
- `circle_mQTL`: Plot a circular genome-metabolome plot

Author(s)

Lyamine Hedjazi and Jean-Baptiste Cazier

Maintainer: Lyamine Hedjazi <<mqtl@ican-institute.org>>

Examples

```
# Download data files

load_datafiles()

# Format data

format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

# Constant Sum normalisation
nmeth<-CS
normalise_mQTL(cleandat,CSnorm,nmeth)

# Alignment
align_mQTL(CSnorm,aligdat)

# Dimensionality reduction
met="rectangle" # choose the statistical summarizing measure ("max","sum","trapez",...)
RedMet="SRV" # reduction method ("SRV" or "bin")
```

```

pre_mQTL(aligdat, reducedF, RedMet="SRV",met, corrT=0.9)

# mQTL mapping
results<- list() # a list to stock the mQTL mapping results
nperm<- 0 # number of permutations if required
results<-process_mQTL(reducedF, cleangen, nperm)

## Post-Process
post_mQTL(results)

## Summarize
redfile<-"rectangle_SRV.ppm"
summary_mQTL(results,redfile,T=8)

#plot circular genome
circle_mQTL(results, Th=8,spacing=0)

## visualisation and metabolite identification
#plot NMR profile
simple.plot(file=cleandat,lo=3.02,hi=3.08,k=1:20,title="NMR profile")

#plot SRV regions
SRV.plot(file1=cleandat,file2=rectangle_SRV,lo=3.02,hi=3.08,k=1:20,title="Cluster plot")

#plot lod for the region of interest
SRV_lod.plot(results,rectangle_SRV,Th=1)

#plot top lod SRV regions
Top_SRV.plot(file1=cleandat,file2=rectangle_SRV,results=results,met=met,intMeth="mean")

```

alignSp

Base function for Spectrum Alignment

Description

Alignment of spectrum segment to the spectrum of interest

Usage

```
alignSp(refSp, refSegments, intSp, intSegments, recursion, MAX_DIST_FACTOR, MIN_RC)
```

Arguments

refSp	a vector specifying the reference spectrum
refSegments	a list characterizing the reference segments (start, end, peaks, ...)
intSp	a vector specifying the spectrum of interest
intSegments	a list characterizing the segment of interest (start, end, peaks, ...)

recursion	A list defining default values of the parameters of recursive alignment(minimal segment width, recursion step, resamblance, acceptance, ...)
MAX_DIST_FACTOR	distance matching parameter (0.5*peak width)
MIN_RC	minimum resamblance coefficient

Value

alignedSpectrum
aligned spectrum as a vector

Author(s)

Lyamine Hedjazi

See Also

[align_mQTL](#)

Examples

```
## Data
load_datafiles()
Sp<-t(read.table(phenofile))
ppm<-as.numeric(colnames(Sp))

## Normalization
normSp<-normalise(abs(Sp),CS)

##Segmentation and matching parameters
setupRSPA(ppm)

##reference spectrum selection
attach(normSp)
index<-selectRefSp(Sp,recursion$step)
refSp<-Sp[index,]

##segmentate a reference spectrum
refSegments<- segmentateSp(refSp, peakParam)

##segmentate a test spectrum
testSegments<- segmentateSp(Sp[1,], peakParam)

##attach test and reference segments
attachedSegs<-attachSegments(refSegments, testSegments)

##Match test and reference segments
attach(attachedSegs)
Segs<-matchSegments(refSp, Sp[1,], testSegmentsNew, refSegmentsNew, MAX_DIST_FACTOR, MIN_RC)

##Align test spectrum
```

```
attach(Segs)  
SpAlg<- alignSp(refSp,refSegs,Sp[1,],testSegs,recursion,MAX_DIST_FACTOR,MIN_RC)
```

align_mQTL

Peak alignment and normalisation of metabolomic data

Description

Recursive Segment-Wise Peak Alignment (RSPA) for accounting peak position variation across metabolomic data

Usage

```
align_mQTL(datafile, outdat,idx)
```

Arguments

datafile	The main input file of raw spectra in the csvs format
outdat	The output file of aligned spectra in the csvs format
idx	index of reference spectrum

Details

The algorithm is based on the following workflow:

1. Automatic selection of a reference spectrum (if required).
2. Segmentate a reference spectrum.
3. Then for each test spectrum:
 - segmentate a test spectrum.
 - match test and reference segments.
 - align a test spectrum.

Value

It returns a file with aligned data in the csvs format.

Author(s)

Lyamine Hedjazi

References

Veselkov,K. et al (2009) Recursive Segment-Wise Peak Alignment of Biological ¹H NMR Spectra for Improved Metabolic Biomarker Recovery, *Anal. Chem.*, 81(1), 56-66.

See Also

[alignSp](#), [attachSegments](#), [matchSegments](#), [segmentateSp](#), [format_mQTL](#), [format_mQTL](#)

Examples

```
# Download data files

load_datafiles()

# Format data

format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

# Constant Sum normalisation
nmeth<-CS
normalise_mQTL(cleandat,CSnorm,nmeth)

# Alignment
align_mQTL(CSnorm,aligdat)
```

attachSegments	<i>Concatenation of test and reference segments</i>
----------------	---

Description

Concatenation of test and reference segments to ensure one-to-one correspondence.

Usage

```
attachSegments(refSegments, testSegments)
```

Arguments

refSegments	a list characterizing the segments of the reference spectrum (start, end, peaks, center)
testSegments	a list characterizing the segments of the test spectrum (start,end, peaks, center)

Details

The algorithm:

1. For each reference segment within segment boundaries, i.e. between initial and final positions, find all centre (middle) positions of test segments and merge those segments, if more than one centre position is found
2. Apply the same procedure for each test segment

Value

A list:

segments\$start a vector specifying the starting of each concatenated test segment
 segments\$PeakLeftBoundary a list defining the peak left boundary of each concatenated test segment
 segments\$PeakRightBoundary a list defining the peak right boundary of each concatenated test segment
 segments\$Peaks a list specifying the peaks information of each concatenated test segment (max position, start position, end position,...)
 segments\$end a vector specifying the end of each concatenated test segment
 segments\$end a vector specifying the center of each concatenated test segment

Author(s)

Lyamine Hedjazi

References

Veselkov, K. et al (2009) Recursive Segment-Wise Peak Alignment of Biological ¹H NMR Spectra for Improved Metabolic Biomarker Recovery, *Anal. Chem.*, 81(1), 56-66.

See Also

[matchSegments](#)

Examples

```
## Data
load_datafiles()
Sp<-t(read.table(phenofile))
ppm<-as.numeric(colnames(Sp))

## Normalization
normSp<-normalise(abs(Sp),CS)

##Segmentation and matching parameters
setupRSPA(ppm)

##reference spectrum selection
attach(normSp)
index<-selectRefSp(Sp,recursion$step)
refSp<-Sp[index,]

##segmentate a reference spectrum
refSegments<- segmentateSp(refSp, peakParam) # segmentate reference spectrum

##segmentate a test spectrum
testSegments<- segmentateSp(Sp[1,], peakParam) # segmentate test spectrum (1st sample)
```



```
##attach test and reference segments
attachedSegs<-attachSegments(refSegments, testSegments)
```

`circle_mQTL`*Circular genome-metabolome plot for mQTL.NMR*

Description

shows mQTL locations and relations with the metabolome on a central chemical axis

Usage

```
circle_mQTL(results, Th = 0, chr = 9, spacing = 25)
```

Arguments

<code>results</code>	a list containing mQTL mapping results generated by mQTL.NMR package
<code>Th</code>	a numerical parameter specifying LOD threshold
<code>chr</code>	a numerical value defining the chromosomes to show if necessary
<code>spacing</code>	a numerical parameter specifying the spacing between chromosomes on the circular genome

Value

A circular plot where the central horizontal line corresponds to the NMR chemical axis, the circle represents the chromosomal positions, and the colored lines significant association between a shift and genomic location.

Author(s)

Lyamine Hedjazi

See Also

[pplot](#)

Examples

```
load_datafiles()
load(results)

circle_mQTL(results, Th=8,spacing=0)
```

configureRSPA	<i>segmentaion and recursive alignment parameters</i>
---------------	---

Description

The routine used to change and improve the RSPA algorithm performance

Usage

```
configureRSPA(ppm)
```

Arguments

ppm a numerical vector defining the chemical shift scale

Author(s)

Jean-Baptiste Cazier

See Also

[setupRSPA](#)

Examples

```
load_datafiles()

load(results)
ppm<-results$ppm
configureRSPA(ppm)
```

format_mGWA	<i>Routine to reformat the data into the required format to perform mG-WAS</i>
-------------	--

Description

This function enables to reformat data into the proper format. The user should provides in input metabolomic file, Genotype file, map file and a file containing sex, age and individual IDs.

Usage

```
format_mGWA(datafile, genofile1, genofile2, covarfile, outdat, outgeno)
```

Arguments

datafile	metabolomic data file
genofile1	genotype file in the "ped" format
genofile2	map file containing more information on SNP marker (position, ...)
covarfile	a text file contains covariates such as age or sex
outdat	output data file with formatted phenotype data in csvs format
outgeno	output data file with formatted genotype data in csvs format

Value

formatted phenotype and genotype data files (in format csvs) are written to the user working directory (it is therefore preferable that the user create a new directory to be used throughout the study)

Author(s)

Lyamine Hedjazi

See Also

[format_mQTL](#), [process_mGWA](#)

Examples

```
load_datafiles()
format_mGWA(human.pheno, human.geno, humanMap, covarFile, cleandat, cleangen)
```

format_mQTL	<i>Routine to reformat the data of animal crosses into the required format to perform mQTL mapping</i>
-------------	--

Description

This function enables to reformat data into the proper format. The user should provides in input metabolomic file, Genotype file and a file containing sex and pgm (parental grandmother).

Usage

```
format_mQTL(datafile, genofile, physdat, outdat, outgeno)
```

Arguments

datafile	metabolomic data file in text format
genofile	genotype data file in text format
physdat	a file containing sex and pgm in text format
outdat	Output data file with formatted phenotype data (metabolomic data + sex + pgm) in the format csvs
outgeno	Output data file with formatted genotype data in the csvs format

Value

formatted phenotype and genotype data files (in format csvs) are written to the user working directory (it is therefore preferable that the user create a new directory to be used throughout the study)

Author(s)

Lyamine Hedjazi

See Also

[align_mQTL](#),

Examples

```
# Download data files
load_datafiles()

# Format data

format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)
```

load_datafiles

Load data files for examples

Description

Data files are downloaded from the extdata directory to the user's working directory.

Usage

```
load_datafiles()
```

Value

Loaded data files concern four datasets: raw metabolomic data ('phenofile.txt'), genomic data ('genofile.txt'), additional data ('physiodat.txt').

Author(s)

Lyamine Hedjazi

See Also

[format_mQTL](#)

Examples

```
# Load data files
load_datafiles()
```

load_demo_data	<i>Load demo data files</i>
----------------	-----------------------------

Description

Data files are downloaded from the sourceforge.net website to the user's working directory.

Usage

```
load_demo_data()
```

Value

Loaded data files concern four datasets: raw metabolomic data (Metabofile.txt), genomic data (Genofile.txt), additional data (physiodat.txt), formatted metabolomic data (met.clean.txt) and formatted genomic data (gen.clean.txt). Data files specifying additional information and results are also provided such as: result of SRV clustering (ur.rectangle.alig.txt), aligned data (ur.alig.txt), normalized data by CS and PQN methods (cs.norm.txt and pqn.norm.txt) and SRV clusters parameters (rectangle_SRV.txt)

Author(s)

Lyamine Hedjazi

See Also

[format_mQTL](#)

Examples

```
## Not run:

# Load demo data files
load_demo_data()

## End(Not run)
```

matchSegments *Matching the segment of interest to the corresponding reference*

Description

The algorithm makes use of a fuzzy logic approach to match the segment of interest to the corresponding reference

Usage

```
matchSegments(refSp, intSp, intSegments, refSegments, MAX_DIST_FACTOR, MIN_RC)
```

Arguments

refSp	a vector specifying the spectrum of reference
intSp	a vector specifying the spectrum of interest (test spectrum)
intSegments	a list characterizing the segments of spectrum of interest
refSegments	a list characterizing the segments of the reference spectrum (start, end, peaks, center)
MAX_DIST_FACTOR	distance matching parameter ($0.5 * \text{peak_width}$)
MIN_RC	minimum resamblance coefficient

Details

Algorithm:

1. pick-up segment of interest
2. pick-up reference segments
3. calculate relative distance between them
4. calculate relative resamblance between them
5. find min value of relative distance and resamblance
6. use it as representative of similiarity between target and reference segments
7. find the segment that has the highest value of both relative distance and resamblance

Value

A list:

testSegs	a list characterizing the matched test segments
refSegs	a list characterizing the matched reference segments

Author(s)

Lyamine Hedjazi

References

Veselkov, K. et al (2009) Recursive Segment-Wise Peak Alignment of Biological ¹H NMR Spectra for Improved Metabolic Biomarker Recovery, *Anal. Chem.*, 81(1), 56-66.

See Also

[attachSegments](#)

Examples

```
## Data
load_datafiles()
Sp<-t(read.table(phenofile))
ppm<-as.numeric(colnames(Sp))

## Normalization
normSp<-normalise(abs(Sp),CS)

##Segmentation and matching parameters
setupRSPA(ppm)

##reference spectrum selection
attach(normSp)
index<-selectRefSp(Sp,recursion$step)
refSp<-Sp[index,]

##segmentate a reference spectrum
refSegments<- segmentateSp(refSp, peakParam) # segmentate reference spectrum

##segmentate a test spectrum
testSegments<- segmentateSp(Sp[1,], peakParam) # segmentate test spectrum (1st sample)

##attach test and reference segments
attachedSegs<-attachSegments(refSegments, testSegments)

##Match test and reference segments
attach(attachedSegs)
Segs<-matchSegments(refSp, Sp[1,], testSegmentsNew, refSegmentsNew, MAX_DIST_FACTOR, MIN_RC)
```

normalise

Base function of normalisation

Description

Removing dilutions between biofluid samples (normalisation of spectra)

Usage

```
normalise(X, method, refIdx, noiseInt)
```

Arguments

X	A matrix specifying metabolomic data
method	A character defining the normalization method. Constant sum normalisation (method<-'CS'), Constant noise normalisation (method<-'CN'), Quotient probabilistic method (method<-'PQN'), Linear baseline normalisation (method<-'LBN'), Auto scaling (method<-'AS'), Pareto scaling (method<-'PS').
refIdx	index of reference individual (set by the user if necessary)
noiseInt	noise region on the resonance axis as an interval (ex. [11,12] ppm)

Value

A matrix defining normalised spectrum

Author(s)

Lyamine Hedjazi

References

- Probabilistic quotient normalisation: Dieterle, F., Ross, A., Schlotterbeck, G., & Senn, H. (2006). Probabilistic quotient normalization as robust method to account for dilution of complex biological mixtures. Application to 1H NMR metabolomics. *Analytical Chemistry*, 78, 4281-4290.
- Constant sum (total area) normalisation: Craig, A., Cloarec, O., Holmes, E., Nicholson, J. K., Lindon, J. C., Scaling and normalization effects in NMR spectroscopic metabonomic data sets. *Anal Chem* 2006, 78, (7), 2262-2267.
- Linear baseline normalisation: Bolstad, B. M., Irizarry, R. A., Astrand, M., & Speed, T. P. (2003). A comparison of normalization methods for high density oligonucleotide array data based on variance and bias. *Bioinformatics*, 19, 185-193.
- Auto-scaling: Jackson, J. E. (2003). *A user's guide to principal components*. Hoboken, NJ: Wiley-Interscience.
- Pareto scaling: Eriksson, L., Antti, H., Gottfries, J., Holmes, E., Johansson, E., Lindgren, F., et al. (2004). Using chemometrics for navigating in the large data sets of genomics, proteomics, and metabonomics (gpm). *Analytical and Bioanalytical Chemistry*, 380, 419-429.

See Also

[normalise_mQTL](#)

Examples

```
## Data
Sp=matrix(rnorm(10*5000,mean=0,sd=1), nrow=10,ncol=5000)

## Quotient probabilistic normalisation
NormDat<-normalise(abs(Sp),PQN)
```

normalise_mQTL	<i>Normalisation of metabolomic data</i>
----------------	--

Description

Takes use of the base function [normalise](#) to provide a normalised metabolomic data file.

Usage

```
normalise_mQTL(infile,outfile,method,refIdx=1, noiseInt=c(11,12))
```

Arguments

infile	a text file with non-normalised spectra profiles
outfile	a text file with normalised spectra profiles
method	a character defining the normalization method: - Constant sum normalisation (method<-'CS') - Constant noise normalisation (method<-'CN') - Quotient probabilistic method (method<-'PQN') - Linear baseline normalisation (method<-'LBN') - Auto-scaling (method<-'AS') - Pareto scaling (method<-'PS')
refIdx	index of reference individual (set by the user)
noiseInt	noise region on the resonance axis as an interval (ex. [11,12] ppm)

Value

a file containing normalised spectra profiles

Author(s)

Lyamine Hedjazi

See Also

[normalise](#)

Examples

```
# Download data files
load_datafiles()

# Format data

format_mQTL(phenofile, genofile, physiodat, cleandat, cleangen)

# Constant Sum normalisation
nmeth<-CS
normalise_mQTL(cleandat, CSnorm, nmeth)
```

peakPeaks

Peak picking algorithm

Description

Identification of peaks in metabolomic data based on the calculation of smoothed derivatives using Savitzky-Golay filter. The peak is identified if derivative crosses zero, i.e. $\text{sign}(X'(i)) > \text{sign}(X'(i+1))$.

Usage

```
peakPeaks(SpSmooth, dpDerivs, Sp)
```

Arguments

SpSmooth	a vector specifying smoothed spectrum
dpDerivs	a vector specifying smoothed derivative of the spectrum
Sp	a vector specifying the spectrum of interest

Value

identified peaks

Author(s)

Lyamine Hedjazi

References

Veselkov, K. et al (2009) Recursive Segment-Wise Peak Alignment of Biological ¹H NMR Spectra for Improved Metabolic Biomarker Recovery, *Anal. Chem.*, 81(1), 56-66.

See Also

[sgolayDeriv](#)

Examples

```
load_datafiles()
Sp<-t(read.table(phenofile))

## Peak picking
Spectrum<-Sp[1,]
iOrder <- 3
iFrameLen<- 11

SpDerivs<-sgolayDeriv(Spectrum,iOrder,iFrameLen,2)
SpSmooth<-sgolayDeriv(Spectrum,iOrder,iFrameLen,1)
peaks<-peakPeaks(SpSmooth,SpDerivs,Spectrum)
```

post_mQTL	<i>Plot top LOD results</i>
-----------	-----------------------------

Description

plot the results of a given run

Usage

```
post_mQTL(results, probs = c(0.95, 0.99, 0.999, 0.9999))
```

Arguments

results	a list containing the results of mQTL analysis.
probs	a numerical vector of probabilities with values in [0,1]. (Values up to 2e-14 outside that range are accepted and moved to the nearby endpoint).

Details

This function plots different results corresponding to top LOD marker

Value

It returns one window gathering all figures of the mQTL analysis. Each figure is also saved separately in the user's working space.

Author(s)

Hedjazi Lyamine

See Also

[pre_mQTL](#)

Examples

```
# Download data files
load_datafiles()

# mQTL mapping results
load(results)

# Plot mQTL mapping results
post_mQTL(results)
```

ppersp

Plot a 3-D profile of LODs

Description

Plot 3-D profile of LODs as function of genomic position and chemical shift

Usage

```
ppersp(z, ppm, title, theta=-15, phi=15, r=50)
```

Arguments

z	a matrix specifying metabolome genome-wide mQTL mapping results
ppm	a vector of chemical shift
title	plot title
theta	angle defining the viewing direction (azimuthal direction)
phi	angle defining the viewing direction (colatitude direction)
r	the distance of the eyepoint from the centre of the plotting box.

Value

plot 2D-profile

Author(s)

Jean-Baptiste Cazier

See Also

[pplot](#)

Examples

```
# Download data files
load_datafiles()

# mQTL mapping results
load(results)

## Plot 3D profile
dev.new(width=5,height=5,pointsize=5)
ppersp(results$res, results$ppm, title="Example plot")
```

pplot

Plot a color scale layer

Description

Plot the results with a color scale y layer over 3 in 2D

Usage

```
pplot(z, title, ppm, res, LT = c(5,10,15,20))
```

Arguments

z	a matrix specifying metabolome genome-wide mQTL mapping results
title	figure title
ppm	a vector of chemical shift
res	mQTL results to be plotted (scanone object)
LT	quantil(res,probs), res: matrix of mQTL mapping results and probs: vector of probabilities

Value

plot of 2-D profile

Author(s)

Jean-Baptiste Cazier

See Also

[ppersp](#)

Examples

```
# Download data files
load_datafiles()

# mQTL mapping results
load(results)

## Plot 3D profile

dev.new(width=5,height=5,pointsize=5)

probs=c(0.95,0.99,0.999,0.9999) ## probabilities

pplot(results$res,"Full 2D Profile", results$ppm, results$best, quantile(results$res,probs=probs))
```

pre_mQTL

*Statistical Recoupling of variables for mQTL analysis***Description**

Makes use of SRV to preprocess metabolomic data for dimensionality reduction by statistical recoupling of variables

Usage

```
pre_mQTL(infile, outfile, RedMet="SRV", met="sum", corrT = 0.9, BinWidth=0.01)
```

Arguments

infile	metabolomic datafile in csvs format
outfile	reduced metabolomic datafile in csvs format
met	a character specifying the used statistical summary
RedMet	a character indicating the used dimensionality reduction method: Redmet="SRV" for statistical recoupling of variables and Redmet="bin" to apply the binning approach
corrT	a numerical parameter indicating correlation threshold
BinWidth	a numerical parameter indicating the binning width

Details

mQTL-NMR package implements two dimensionality reduction methods. The first one concerns the SRV algorithm which forms clusters of variables using a measure of a local spectral dependency. The second one concerns the classical binning method which divides the spectra into evenly spaced windows (bins) whose width commonly ranges between 0.001 and 0.05 ppm.

Value

variables are associated into a series of clusters (or bins). This function provides in output the paramaters of the clusters (min and max borders, mean,...)

Author(s)

Lyamine Hedjazi

References

- Blaise,B. et al (2009) Statistical recoupling prior to significance testing in nuclear magnetic resonance based metabonomics, Anal. Chem., 81(15), 6242-6251. - S praul, M.; Neidig, P.; Klauck, U.; Kessler, P.; Holmes, E.; Nicholson, J. K.; Sweatman, B.C.; Salman, S.R.; Farrant, R.D.; Rahr, E.; et al. J.Pharm. Biomed. Anal. 1994, 12, 1215-1225.

See Also

[SRV,post_mQTL](#)

Examples

```
# Download data files

load_datafiles()

# Format data

format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

# Constant Sum normlisation
nmeth<-CS
normalise_mQTL(cleandat,CSnorm,nmeth)

# Alignment
align_mQTL(CSnorm,aligdat)

# Dimensionality reduction
met="rectangle" # choose the statistical summarizing measure ("max","sum","trapez",...)
RedMet="SRV" # reduction method ("SRV" or "bin")

pre_mQTL(aligdat, reducedF, RedMet="SRV",met, corrT=0.9)
```

process_mGWA	<i>Metabolomic Genome-Wide Association analysis for a set of independent individuals</i>
--------------	--

Description

Test for association between a trait and genetic polymorphism

Usage

```
process_mGWA(phenofile = phenofile, genofile = genofile, nperm = 0, gtmodel = "overdominant", covarList
```

Arguments

phenofile	a text file with phenotype data
genofile	a text file with genotype data
nperm	number of permutations
gtmodel	genetic model ("additive","recessive","dominant","overdominant")
covarList	covariate variables ("sex" and/or "age")

Details

This function makes use of metabolomic and genotype data to perform genome-wide association analysis using a standard regression method based on the GenABEL package.

Value

2D score tables (-log₁₀(p-value))

Author(s)

Lyamine Hedjazi

References

Aulchenko, Y.S.; Ripke, S.; Isaacs, A.; van Duijn, C.M. *Bioinformatics* 2007, 23, 1294-1296.

See Also

[format_mGWA](#)

Examples

```
load_datafiles()
format_mGWA(human.pheno, human.geno, humanMap, covarFile,hcleandat, hcleangen)

# mQTL mapping
results<- list() # a list to stock the mQTL mapping results
nperm<- 0 # number of permutations if required

results<-process_mGWA(phenofile=hreducedF, genofile=hcleangen,nperm=0, gtmodel="additive")
```

process_mQTL	<i>mQTL mapping</i>
--------------	---------------------

Description

Function to process the tissue extract of the individuals for QTL analysis

Usage

```
process_mQTL(datfile, genfile, nperm = 0)
```

Arguments

datfile	a text file with phenotype data
genfile	a text file with genotype data
nperm	nperm

Details

This function makes use of metabolomic and genotype data to perform QTL analysis based on the R/QTL package, for mapping quantitative trait loci. In particular, it makes use of the extended Haley-Knott method to optimize the LOD score evaluation and avoid problems with missing genotypes.

Value

2D LOD score table

Author(s)

Jean-Baptiste Cazier and Hedjazi Lyamine

References

Broman,K., et al (2006) R/qtl: QTL mapping in experimental crosses, *Bioinformatics*, 19(7), 889-890.

See Also[post_mQTL](#)**Examples**

```
# Download data files

load_datafiles()

# mQTL mapping
results<- list() # a list to stock the mQTL mapping results
nperm<- 0 # number of permutations if required
results<-process_mQTL(reducedF, cleangen, nperm)
```

segmentateSp

*Segmentation of a spectrum of interest***Description**

Determination of highly intensive peaks in the spectrum of interest and subsequent concatenation of closely located peaks into larger segments

Usage

```
segmentateSp(Sp, peakParam)
```

Arguments

Sp	a vector defining the spectrum
peakParam	a list: <ul style="list-style-type: none"> • ampThr: amplitude threshold [default 2*median(peaksMaxValues)] • iFrameLen: Savitzky-Golay frame length • iOrder: polynomial order of Savitzky - Golay filter • iFrameLen: Savitzky-Golay frame length • minPeakWidth: min peak size • ppmDist: distance to concatenate adjacent peaks

Value

A list:

testSegmentsNew

a list specifying the new test segments

refSegmentsNew

a list specifying the new reference segments

Author(s)

Lyamine Hedjazi

References

Veselkov, K. et al (2009) Recursive Segment-Wise Peak Alignment of Biological ¹H NMR Spectra for Improved Metabolic Biomarker Recovery, *Anal. Chem.*, 81(1), 56-66.

See Also[attachSegments](#), [matchSegments](#)**Examples**

```
## Data
load_datafiles()
Sp<-t(read.table(phenofile))
ppm<-as.numeric(colnames(Sp))

## Normalization
normSp<-normalise(abs(Sp),CS)

##Segmentation and matching parameters
setupRSPA(ppm)

##reference spectrum selection
attach(normSp)
index<-selectRefSp(Sp,recursion$step)
refSp<-Sp[index,]

##segmentate a reference spectrum
refSegments<- segmentateSp(refSp, peakParam) # segmentate reference spectrum
```

`selectRefSp`*Automated selection of a reference spectrum*

Description

The selection of reference spectrum among all spectrums is based on the highest similarity to all other spectra

Usage`selectRefSp(X, step)`

Arguments

X matrix of spectra
step a numerical parameter used to scale spectral regions down to specific bin size

Value

returns the index of selected spectrum

Author(s)

Lyamine Hedjazi

See Also

[alignSp](#)

Examples

```
# Data  
  
Sp=matrix(rnorm(10*5000,mean=0,sd=1), nrow=10,ncol=5000)  
  
# Reference spectrum selection  
  
step=0.02 # Recursion step (default 0.02)  
index<-selectRefSp(Sp,step)
```

setupRSPA *setup of alignment parameters*

Description

Configuration of the RSPA algorithm invariant parameters

Usage

```
setupRSPA(ppm)
```

Arguments

ppm a vector defining chemical shift scale

Author(s)

Jean-Baptiste Cazier

See Also[configureRSPA](#)**Examples**

```
load_datafiles()

load(results)
ppm<-results$ppm
setupRSPA(ppm)
```

sgolay*Find the matrix of differentiation filters*

Description

designs a Savitzky-Golay (polynomial) FIR smoothing filter. The polynomial order must be less than the frame size which must be odd.

Usage

```
sgolay(k,F,W)
```

Arguments

k	a numerical value of polynomial order
F	a numerical value of frame size
W	weighting matrix

Value

matrix of differentiators

Author(s)

Lyamine Hedjazi

References

Sophocles J. Orfanidis, INTRODUCTION TO SIGNAL PROCESSING, Prentice-Hall, 1995, Chapter 8

See Also[sgolayDeriv](#)

Examples

```
k <- 3
F <- 11

Sg=sgolay(k,F)
```

`sgolayDeriv`*Calculate smoothed derivatives*

Description

Calculate smoothed derivatives using Savitzky-Golay filter

Usage

```
sgolayDeriv(dpSpectr, iOrder, iFrameLen, j)
```

Arguments

<code>dpSpectr</code>	a vector specifying the input spectrum
<code>iOrder</code>	polynomial order of Savitzky - Golay filter
<code>iFrameLen</code>	Savitzky-Golay frame length in ppm scale
<code>j</code>	order of derivative

Value

`j`th derivative of the spectrum

Author(s)

Lyamine Hedjazi

See Also

[sgolay](#)

Examples

```
## Data

Sp=matrix(rnorm(10*13454,mean=0,sd=1), nrow=10,ncol=13454)

## Peak picking
Spectrum<-Sp[10,]
iOrder <- 3
iFrameLen<- 11
j<-2
```

```
SpDerivs<-sgolayDeriv(Spectrum,iOrder,iFrameLen,j)
```

`simple.plot`*Plot NMR profile plus SRV regions*

Description

Plot NMR profile plus SRV regions and consensus across the various statistics

Usage

```
simple.plot(file,lo,hi,k,title)
```

Arguments

<code>file</code>	a text file containing NMR data
<code>lo</code>	starting point on the chemical axis
<code>hi</code>	ending point on the chemical axis
<code>k</code>	number of samples
<code>title</code>	title of the plot

Value

NMR profile and SRV region plot with peak calling consensus

Author(s)

Jean-Baptiste Cazier

See Also

[SRV.plot](#)

Examples

```
# Load data files
load_datafiles()

# Format data
format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

# Plot NMR profile
simple.plot(file=cleandat,lo=3.02,hi=3.08,k=1:20,title="NMR profile")
```

SRV*Statistical Recoupling of Variables*

Description

Base function for dimensionality reduction by statistical recoupling of variables

Usage

```
SRV(X, minsize, correl, clustf = median)
```

Arguments

X	matrix of metabolomic data
minsize	a numerical value defining the singlet size
correl	a numerical value defining the bucketting resolution
clustf	a numerical value defining the correlation threshold

Value

A list:

indicesdebf	a vector indicating the starting border of superclusters
indicesfinf	a vector indicating the ending border of superclusters
Xcluster	matrix of reduced data

Author(s)

Jean-Baptiste Cazier

References

Blaise,B. et al (2009) Statistical recoupling prior to significance testing in nuclear magnetic resonance based metabonomics, *Anal. Chem.*, 81(15), 6242-6251.

See Also

[pre_mQTL](#)

Examples

```
# Load data files

load_datafiles()

Sp<-read.table(phenofile, as.is=TRUE, header=TRUE, sep=\t)

# Perform the SRV analysis to reduce the number of dimension of Spectra #data (Sp)

corrT=0.9 # correlation threshold
minsize=10 # singlet size
met="rectangle" # summary measure

SRV<-SRV(t(Sp), minsize, corrT,clustf=met)
```

SRV.plot

Plot SRV clusters

Description

Plot arrows defined by SRV on data

Usage

```
SRV.plot(file1,file2,lo,hi,k,title)
```

Arguments

file1	a text file with NMR data
file2	a text file with SRV results
lo	starting point on chemical shift
hi	ending point on chemical shift
k	number of samples
title	title of the plot

Author(s)

Lyamine Hedjazi

See Also

[simple.plot](#)

Examples

```
# Load data files
load_datafiles()

# Format data
format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

## Plot SRV profile
SRV.plot(file1=cleandat,file2=rectangle_SRV,lo=3.02,hi=3.08,k=1:168,title="Cluster plot")
```

SRV_lod.plot

Plot top lod SRV clusters

Description

Plot all SRV clusters associated with the top lod locus

Usage

```
SRV_lod.plot(results, file, Th)
```

Arguments

results	a list specifying the results of mQTL mapping
file	a text file contains resulting clusters
Th	a numerical value of LOD threshold

Author(s)

Lyamine Hedjazi

See Also

[SRV.plot](#)

Examples

```
load_datafiles()
load(results)

## Plot LOD profile
SRV_lod.plot(results,rectangle_SRV,T=1)
```

summary_mQTL	<i>Function to summarize the mQTL mapping results of all the runs and their differences</i>
--------------	---

Description

This function generates a table containing the genetic markers and their associated metabolomic variables and estimated LOD score.

Usage

```
summary_mQTL(results, redfile,Th = 5)
```

Arguments

results	a list specifying the mQTL mapping results
redfile	a text file containing the parameters of identified clusters(.PPM file)
Th	a numerical parameter indicating the threshold of top accepted score (LOD or $-\log_{10}(\text{p-value})$)

Details

Generates a text file containing a table of summary of mQTL mapping results

Value

returns Summaries

Author(s)

Jean-Baptiste Cazier and Lyamine Hedjazi

See Also

[pre_mQTL](#)

Examples

```
load_datafiles()
load(results)

Th<-10 ## LOD threshold
summary_mQTL(results,rectangle_SRV,Th)## summarizes mQTL results in a table
```

Top_SRV.plot	<i>Plot top SRV clusters</i>
--------------	------------------------------

Description

Plot lines defined by SRV on top SRV clusters

Usage

```
Top_SRV.plot(file1,file2,results,met,intMeth,clustidx)
```

Arguments

file1	a text file with NMR data
file2	a text file with SRV clusters
results	a list containinig results of mQTL mapping
met	a character specifying the summarizing statistical measure of peaks
intMeth	a charcter specifying summarizing method across samples ("mean" or "max")
clustidx	index specifying the SRV cluser of interest (optinal)

Author(s)

Lyamine Hedjazi

See Also

[SRV.plot](#)

Examples

```
load_datafiles()

load(results)

# Format data

format_mQTL(phenofile,genofile,physiodat,cleandat,cleangen)

## Plot SRV profile
Top_SRV.plot(file1=cleandat,file2=rectangle_SRV,results=results,met=met,intMeth="mean")
```

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