# Package 'MetaboAnalystR'

January 14, 2020

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Title An R Package for Comprehensive Analysis of Metabolomics Data

Version 2.0.2

Author Jianguo Xia [aut, cre], Jasmine Chong [aut]

Maintainer Jasmine Chong <jasmine.chong@mail.mcgill.ca>

BugReports https://github.com/xia-lab/MetaboAnalystR/issues
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Description This package contains the R functions and libraries underlying the popular MetaboAnalyst web server, including 500 functions for data processing, normalization, statistical analysis, metabolite set enrichment analysis, metabolic pathway analysis, and biomarker analysis. The package is synchronized with the web server. After installing and loading the package, users will be able to reproduce the same results from their local computers using the corresponding R command history downloaded from MetaboAnalyst, to achieve maximum flexibility and reproducibility.

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Depends R (>= 3.5.2), lattice, methods, pls, data.table
License GPL-3
Encoding UTF-8
LazyData true
RoxygenNote 7.0.2
Imports Rserve,
     ellipse,
     scatterplot3d,
     Cairo,
     randomForest,
     caTools,
     e1071,
     som,
     impute,
     pcaMethods,
     RJSONIO,
     ROCR,
     globaltest,
     GlobalAncova,
     Rgraphviz,
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2 R topics documented:

genefilter, pheatmap, SSPA, SSPA, SVA, Repp. pROC, limma, car, fitdistrplus, lars, Hmise, magrittr, xtable, caret, igraph, gplots, KEGGgraph, reshape, RColorBrewer, tibble, RSQLite, spls. siggenes, ggplot2, BiocParallel, metap, scales Suggests knitr, rmarkdown, devtools, testhat, plotly, reshape2, MSnbase, xcms, CAMERA  VignetteBuilder knitr  R topics documented:read.metaboanalyst.libreadDataTable AddErrMsg analyze.lipids ANOVA.Anal		preprocesscore,
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Repp, pROC, limma, car, fitdistrplus, lars, lars, Hmisc, magrittr, xtable, caret, igraph, gplots, KEGGgraph, reshape, RColorBrewer, tibble, RSQLite, spls, siggenes, ggplot2, BiocParallel, metap, scales Suggests knitr, rmarkdown, devtools, testthat, plotty, reshape2, MSnbase, xcms, cAMERA VignetteBuilder knitr  R topics documented:  .read.metaboanalyst.libreadDataTable		sva.
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read.metaboanalyst.lib	O	
read.metaboanalyst.lib		
read.metaboanalyst.lib	R to	opics documented:
.readDataTable       13         AddErrMsg       14         analyze.lipids       14         ANOVA.Anal       15		•
.readDataTable       13         AddErrMsg       14         analyze.lipids       14         ANOVA.Anal       15		.read.metaboanalyst.lib
AddErrMsg       14         analyze.lipids       14         ANOVA.Anal       15		·
analyze.lipids		
ANOVA. Anal		
		•
ANUVAZ.Anal		
		ANOVAZ.Anal

aof	16
aov.between	16
aov.repeated	17
aov.within	17
ASCAfun.res	18
ASCAfun1	18
ASCAfun2	19
	19
	20
	20
	21
••	21
•	2.2.
	22
	23
	23
	23 24
	24 24
	24 25
	25
	26
6	26
	27
	27
1	27
I	28
	28
6	29
CreateAnalNullMsg	29
CreateANOVAdoc	30
CreateAOV2doc	30
CreateASCAdoc	31
CreateBiomarkerInputDoc	31
	32
	32
	32
	33
<u>i</u>	33
	34
	34
	35
	35
1	36
	36
	30 37
	37
	38
CreateEnrichRnwReport	38

CreateEnrichSSPdoc		39
CreateFooter		39
CreateGraph		40
CreateGSEAAnalTable		40
CreateHCdoc		41
CreateHeatmap2doc		41
CreateIntegPathwayAnalysisRnwReport		42
CreateIntegratedPathwayAnalInputDoc		42
CreateIntegratedPathwayAnalIntr		43
CreateIntegratedPathwayDoc		43
CreateIntegratedPathwayGeneMapTable		44
CreateIntegratedPathwayNameMapTable		44
CreateIntegratedPathwayResultsTable		45
CreateiPCAdoc		45
		45
CreateKMdoc		
CreateLadder		46
CreateLibFromKEGG		47
CreateMappingResultTable		47
CreateMBdoc		47
CreateMetaAnalTable		48
CreateMetaAnalysisDEdoc		48
CreateMetaAnalysisInputDoc		49
CreateMetaAnalysisIntr		49
CreateMetaAnalysisNORMdoc		50
CreateMetaAnalysisOutput		50
CreateMetaAnalysisOverview		51
CreateMetaAnalysisRnwReport		51
CreateMetaTable		52
CreateModelBiomarkersDoc		52
CreateMultiBiomarkersDoc		53
CreateMummichogAnalTable		53
CreateMummichogAnalysisDoc		54
CreateMummichogInputDoc		54
CreateMummichogIntro		55
		55 55
CreateMummichogLibs		
CreateMummichogOverview		56
CreateMummichogRnwReport		56
CreateNetworkExplorerDoc		57
CreateNetworkExplorerInputDoc		57
CreateNetworkExplorerIntr		58
CreateNetworkExplorerOverview		58
CreateNetworkExplorerRnwReport		59
CreateNetworkGeneMapTable		59
CreateNetworkNameMapTable		60
CreateNORMdoc		60
CreateOPLSDAdoc	'	61
CreatePathAnalDoc		61
CreatePathInputDoc		62

CreatePathIntr
CreatePathProcessDoc
CreatePathResultDoc
CreatePathRnwReport
CreatePCAdoc
CreatePLSdoc
CreatePowerAnalDoc
CreatePowerInputDoc
CreatePowerIntr
CreatePowerOverview
CreatePowerParametersDoc
CreatePowerRnwReport
CreateRatioTable
CreateRFdoc
CreateRHistAppendix
CreateROCLabelsTable
CreateSAMdoc
CreateSemiTransColors
CreateSOMdoc
CreateSPLSDAdoc
CreateStatIntr
CreateStatIOdoc
CreateStatRnwReport
CreateSummaryTable
CreateSVMdoc
CreateTimeSeriesAnalNullMsg
CreateTimeSeriesIOdoc
CreateTimeSeriesRnwReport
CreateUnivarBiomarkersDoc
CreateUNIVdoc
CreateUnivROCTable
Create VennMetaTable
CrossReferencing
CVTest.LRmodel
descendMin
doCompoundMapping
doGeneIDMapping
doKEGG2NameMapping
doKOFiltering
EBAM.Init
ExtractMS2data
FC.Anal.paired
FC.Anal.unpaired
FeatureCorrelation
fgsea2
fillpathways
FilterVariable
findEqualGreaterM

FisherLSD	
FormatPeakList	
genLogisticRegMdl	87
Get.asca.tss	88
Get.bwss	88
Get.ConcRef	89
Get.Leverage	89
Get.pAUC	90
Get.pred	90
Get.rpart.summary	91
Get.VIP	91
GetAbundanceLabel	
GetAccuracyInfo	92
GetAllDataNames	92
GetAllKMClusterMembers	93
GetAllSOMClusterMembers	
GetCandidateList	
GetCircleInfo	
GetCIs	
GetCMD	
GetCompoundDetails	
GetConvertFullPath	
getDataFromTextArea	97
GetExtendRange	
GetFC	
GetFeatureNumbers	
GetFinalNameMap	
GetFisherPvalue	
GetHTMLMetSet	
GetHTMLPathSet	
GetImpFeatureMat	
GetKEGGNodeInfo	
GetKMClusterMembers	
GetLassoFreqs	
GetLimmaResTable	
GetMapTable	
GetMaxPCAComp	103
GetMeanROC	
GetMetaResultMatrix	104
GetMetaSigHitsTable	105
GetMetSetName	
GetMsetLibCheckMsg	
GetMsetLibSearchResult	
GetMsetNames	
GetMummichogPathSetDetails	
GetNetworkGeneMappingResultTable	
GetNewSampleNames	
GetORA.pathNames	

GetORA.smpdbIDs	
GetORATable	
GetQEA.keggIDs	
GetQEA.pathNames	
GetQEATable	
GetRCommandHistory	
GetRFConf.Table	. 111
GetRFConfMat	. 112
GetRFOOB	. 112
GetRFSigMat	. 113
GetROC.coords	
GetROCLassoFreq	. 114
GetROCTtestP	. 114
GetSampleSizeLadder	
GetSelectedDataNames	. 115
GetSelectedDataNumber	. 115
GetSigTable	
GetSigTable.Anova	. 116
GetSigTable.Aov2	
GetSigTable.ASCA	
GetSigTable.Corr	
GetSigTable.EBAM	
GetSigTable.FC	
GetSigTable.MB	
GetSigTable.RF	
GetSigTable.SAM	
GetSigTable.SVM	
GetSigTable.TT	
GetSigTable.Volcano	
GetSOMClusterMembers	
GetSSPTable	
GetSuggestedSAMDelta	
GetSVMSigMat	
GetTopInx	
GetTrainTestSplitMat	
GetTtestRes	
GetTTSigMat	
GetUnivReport	
GetVariableLabel	
Get Venn Gene Names	
GetXYCluster	
GroupPeakList	
heckbert	
HMDBID2KEGGID	
HMDBID2Name	
ImportRawMSData	
•	
ImportRawMSDataList	
ImputeVar	. 131

8

InitDataObjects	131
InitPowerAnal	132
InitStatAnalMode	132
InitTimeSeriesAnal	
iPCA.Anal	
isEmptyMatrix	
IsSmallSmplSize	
IsSpectraProcessingOK	
KEGGID2HMDBID	
KEGGID2Name	
KEGGPATHID2SMPDBIDs	
Kmeans.Anal	
kwtest	
LoadKEGGKO_lib	
LoadKEGGLib	
LoadSmpLib	
LogNorm	
LSD.test	
make_cpdlib	
make_cpdlist	
map	
MapCmpd2KEGGNodes	
MapKO2KEGGEdges	
Match.Pattern	
MergeDatasets	
MergeDuplicates	
MetaboliteMappingExact	
MSspec.fillPeaks	
MSspec.rtCorrection	
multi.stat	
Normalization	
OPLSDA.Permut	
OPLSR.Anal	
parseFisher	
parseTukey	
PCA.Anal	
PCA.Flip	149
PCA.GENES	150
Perform.ASCA	150
Perform.ASCA.permute	151
Perform.Permut	151
Perform.permutation	152
Perform.UnivROC	152
PerformAdductMapping	153
PerformApproxMatch	154
PerformBatchCorrection	
PerformCurrencyMapping	
PerformCV.explore	

PerformCV.test
PerformDetailMatch
PerformEachDEAnal
PerformIndNormalization
PerformIntegCmpdMapping
PerformIntegGeneMapping
PerformIntegPathwayAnalysis
PerformKOEnrichAnalysis_KO01100
PerformKOEnrichAnalysis_List
PerformLimmaDE
PerformMapping
performMB
PerformMetaMerge
PerformMultiMatch
PerformPeakAnnotation
PerformPeakProfiling
PerformPowerProfiling
PerformPSEA
PerformPvalCombination
PerformVoteCounting
Plot.Permutation
PlotAccuracy
PlotANOVA
PlotANOVA2
PlotASCA.Permutation
PlotAscaImpVar
PlotASCAModel
PlotBoxPlot
PlotCmpdSummary
PlotCmpdView
PlotConcRange
PlotCorr
PlotCorrHeatMap
PlotDetailROC
PlotEBAM.Cmpd
PlotEIC
PlotEnrichNet Overview 177
PlotFC
PlotHCTree
PlotHeatMap
PlotHeatMap2
PlotImpVar
PlotImpVars
PlotInmexGraph
PlotInmexPath
PlotIntegPaths
PlotInteraction
PlotKFGGPath 186

PlotKmeans	
PlotLoadingCmpd	187
PlotMBTimeProfile	188
PlotMetaVenn	188
PlotMetpaPath	189
PlotModelScree	189
PlotMS.RT	190
PlotMS2Spectra	
PlotMSEA.Overview	191
PlotMSPeaksPerm	
PlotNormSummary	
PlotOPLS.MDL	193
PlotOPLS.Permutation	194
PlotOPLS.Splot	
PlotOPLS2DScore	
PlotORA	
PlotPathSummary	
PlotPathwayMZHits	
PlotPCA.overview	
PlotPCA2DScore	
PlotPCA3DScore	
PlotPCA3DScoreImg	
PlotPCABiplot	
PlotPCALoading	
PlotPCAPairSummary	
PlotPCAScree	
PlotPeaks2Paths	
PlotPLS.Classification	
PlotPLS.Imp	
PlotPLS.Permutation	
PlotPLS2DScore	
PlotPLS3DScore	
PlotPLS3DScoreImg	
PlotPLSLoading	
PlotPLSPairSummary	
PlotPowerProfile	
PlotPowerStat	
PlotProbView	
plotProfile	
PlotQEA.MetSet	
PlotQEA.Overview	
PlotRF.Classify	
PlotRF.Outlier	
PlotRF.VIP	
PlotROC	
PlotROC.LRmodel	
PlotROCTest	
PlotRSVM.Classification	

PlotRSVM.Cmpd	
PlotSAM.Cmpd	
PlotSAM.FDR	
PlotSampleNormSummary	
PlotSelectedFeature	
PlotSigVar	
PlotSOM	
PlotSPLS2DScore	
PlotSPLS3DScore	. 227
PlotSPLS3DScoreImg	. 228
PlotSPLSDA.Classification	. 229
PlotSPLSLoading	. 230
PlotSPLSPairSummary	. 231
PlotSubHeatMap	. 231
PlotTestAccuracy	. 233
PlotTT	. 233
PlotVolcano	
PLSDA.CV	. 235
PLSDA.Permut	
PLSR.Anal	
Predict.class	
PrepareIntegData	
PrepareNetworkData	
PreparePDFReport	
PreparePermResult	
PreparePrenormData	
PrepareQueryJson	
PrepareROCData	
PrepareROCDetails	
PrepareVennData	
RankFeatures	
rda2list	
Read.BatchCSVdata	
Read.MSspec	
Read.PeakList	
Read.PeakListData	
Read TextData	
ReadIndData	. 211
ReadPairFile	
RecordRCommand	
rectUnique	
RegisterData	
e e e e e e e e e e e e e e e e e e e	
RemoveCmpd	
RemoveData	
RemoveDuplicates	
RemoveFile	
RemoveFolder	
RemoveGene	. 250

12

RemoveMissingPercent	
ReplaceMin	
RerenderMetPAGraph	251
RF.Anal	252
ROCPredSamplesTable	252
RSVM	253
RSVM.Anal	253
SAM.Anal	254
SanityCheckData	254
SanityCheckIndData	255
SanityCheckMummichogData	255
SaveTransformedData	256
SearchByCompound	
SearchByName	
SearchMsetLibraries	
SearchNetDB	
SelectMultiData	
SetAnalysisMode	
SetAnnotationParam	
SetCachexiaSetUsed	
SetCandidate	
SetClass	
SetCurrentGroups	
SetCurrentMsetLib	
SetCustomData	
SetDesignType	
SetKEGG.PathLib	
SetMetabolomeFilter	
SetMummichogPval	
SetMummichogPvalFromPercent	
SetOrganism	
SetPeakEnrichMethod	
SetPeakFormat	
SetPeakList.GroupValues	
SetPeakParam	
SetSMPDB.PathLib	
Setup.AdductData	
Setup.BiofluidType	
Setup.ConcData	
Setup.HMDBReferenceMetabolome	
Setup.KEGGReferenceMetabolome	
Setup.MapData	
Setup.UserMsetLibData	
SetupKEGGLinks	
SetupMSdataMatrix	
SetupSMPDBLinks	
SOM.Anal	
sparse.mint.block_iteration	
	213

.read	.metaboa	analv	st.lib

4	1
	- 4

Index		286
	XSet2MSet	. 285
	Volcano.Anal	
	usr2png	. 284
	UpdatePLS.Loading	. 283
	UpdatePCA.Loading	. 283
	UpdateOPLS.Splot	. 282
	UpdateIntegPathwayAnalysis	. 281
	UpdateInstrumentParameters	. 281
	UpdateGraphSettings	. 280
	UpdateData	. 280
	UnzipUploadedFile	. 279
	Ttests.Anal	. 279
	template.match	. 278
	SumNorm	. 278
	SPLSR.Anal	. 277
	splsda	. 276

.read.metaboanalyst.lib

Read RDS files from the internet

# Description

Function downloads the required file and reads it only if not already in working directory. Need to specify the file URL and the destfile.

### Usage

.read.metaboanalyst.lib(filenm)

### **Arguments**

filenm

Input the name of the file to download

.readDataTable

Read data table

### **Description**

Function to read in a data table. First, it will try to use fread, however, it has issues with some windows 10 files. In such case, use the slower read.table method.

# Usage

.readDataTable(fileName)

14 analyze.lipids

### **Arguments**

fileName

Input filename

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

AddErrMsg

Adds an error message

# Description

The error message will be printed in all cases. Used in higher functions.

### Usage

AddErrMsg(msg)

# Arguments

msg

Error message to print

analyze.lipids

Lipid analysis pipeliner

# Description

Lipid analysis pipeliner

### Usage

```
analyze.lipids(inFile, iso = "y")
```

# Arguments

inFile

Input the file to read in

iso

Default is set to "y"

# Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

ANOVA.Anal

	ANOVA.Anal	Perform ANOVA analysis
--	------------	------------------------

# Description

ANOVA analysis

# Usage

```
ANOVA.Anal(mSetObj=NA, nonpar=F, thresh=0.05, post.hoc="fisher")
```

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
nonpar	Logical, use a non-parametric test (T) or not (F)
thresh	Numeric, from 0 to 1, indicate the p-value threshold
post.hoc	Input the name of the post-hoc test, "fisher" or "tukey"
all_results	Logical, if TRUE, it will output the ANOVA results for all compounds with no post-hoc tests performed.

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

ANOVA2.Anal	Perform Two-way ANOVA

# **Description**

Perform Two-way ANOVA

# Usage

```
ANOVA2.Anal(mSetObj=NA, thresh=0.05, p.cor="fdr", type="time0", aov.type=1, use.interact=1)
```

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
thresh	Input the p-value threshold
p.cor	Select method for p-value correction, bonferroni, holm or fdr
type	Select b to perform between-subjects ANOVA, and w for within-subjects ANOVA
aov.type	Specify 1 for ANOVA type 1, or 3 for ANOVA type 3
use.interact	Numeric, whether to consider interaction in two-way repeated ANOVA (1) or not (0).

16 aov.between

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

aof

**ANOVA** 

### **Description**

Perform anova and only return p values and MSres (for Fisher's LSD)

### Usage

```
aof(x, cls)
```

# Arguments

x Input the data to perform ANOVA

cls Input class labels

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

aov.between

Perform Two-way ANOVA

# Description

Perform Two-way ANOVA Perform between-subjects anova

### Usage

```
aov.between(x)
```

# Arguments

x Input data to perform 2-way ANOVA

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

aov.repeated 17

aov.repeated

Perform Two-way ANOVA

# Description

Perform Two-way ANOVA Perform repeated measure one-way anova

### Usage

```
aov.repeated(x, time.fac)
```

### **Arguments**

x Input the data

time.fac Input the time factor

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

aov.within

Perform Two-way ANOVA

# Description

Perform Two-way ANOVA Perform within-subjects anova

### Usage

```
aov.within(x, time.fac)
```

# Arguments

x Input the data

time.fac Input the time factor

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

18 ASCAfun1

ASCAfun.res

Function to perform ASCA

### **Description**

Perform ASCA

# Usage

```
ASCAfun.res(X, Fac)
```

# Arguments

X Input list of compounds

Fac Numeric McGill University, Canada License: GNU GPL (>= 2)

### Author(s)

Jeff Xia < jeff.xia@mcgill.ca>

ASCAfun1

Function to perform ASCA

# Description

Perform ASCA

# Usage

```
ASCAfun1(X, Design, Fac)
```

# Arguments

X Numeric, number of compoundsDesign Number of levels in the factor

Fac Numeric, the factor

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

ASCAfun2

ASCAfun2	Function to perform ASCA
----------	--------------------------

### **Description**

Perform ASCA

### Usage

```
ASCAfun2(X, Desa, Desb, Fac)
```

### Arguments

Χ	Numeric, number of compounds
Desa	Number of levels in the factor TIME
Desb	Number of levels in the other factor

Fac Numeric, the factor

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

calculateConcISO Calculate Concentration ISO

# Description

Assuming independent random distribution of FA, the most probable frequency will be the product of the each component. Note: the data is concentration, we need to get frequencies - percentage w.r.t the total nmol. the result is the saved as separate files for each lipid class data for each FA class, first col is sample name

### Usage

```
calculateConcISO(dat, cls.name, cls.num, min.file, prob.file)
```

### **Arguments**

dat	Input the data
cls.name	Input the class names
cls.num	Input the number of classes
min.file	Input the min file
prob.file	Input the prob file

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CalculateGlobalTestScore

CalculateFeatureRanking

Calculates feature importance

# **Description**

Perform calculation of feature importance (AUC, p value, fold change)

### Usage

```
CalculateFeatureRanking(mSetObj=NA, clust.num=5)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)
clust.num Numeric, input the number of clusters for cluster-analysis

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt CalculateGlobalTestScore}$ 

Quantitative enrichment analysis with globaltest

### **Description**

Various enrichment analysis algorithms

### Usage

```
CalculateGlobalTestScore(mSetObj = NA)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CalculateHyperScore 21

CalculateHyperScore

Over-representation analysis using hypergeometric tests

#### Description

Over-representation analysis using hypergeometric tests The probability is calculated from obtaining equal or higher number of hits using 1-phyper. Since phyper is a cumulative probability, to get P(X>=hit.num) => P(X>(hit.num-1))

### Usage

```
CalculateHyperScore(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CalculateImpVarCutoff Calculate the Important Variable Cutoff

### Description

This function calculates the all important features based on a specfic cutoff.

### Usage

```
CalculateImpVarCutoff(mSetObj, spe.thresh, lev.thresh)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

spe.thresh alpha threshold, less is better, default less than 5 percentile based chi-square

note: spe and leverage are vectors, not a single value, but a list to store the result note: the last model is Model.res, no spe Calculate leverage cutoff based on permutation Calculate the reference distribution of leverages note: leverage.perm

is a list with each member in a 3 column matrix

lev. thresh leverage threshold, the higher better, default more than 95 percentile of permuted

leverage

```
Jeff Xia < jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

22 CalculatePairwiseDiff

CalculateOraScore

Calculate ORA score

# Description

Calculate the over representation analysis score

### Usage

CalculateOraScore(mSetObj=NA, nodeImp, method)

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

nodeImp Indicate the pathway topology analysis, "rbc" for relative-betweeness centrality,

and "dgr" for out-degree centrality.

method is "fisher" or "hyperg"

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CalculatePairwiseDiff Calculate Pairwise Differences

### **Description**

Mat are log normalized, diff will be ratio. Used in higher functions.

### Usage

CalculatePairwiseDiff(mat)

### **Arguments**

mat

Input matrix of data to calculate pair-wise differences.

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CalculateQeaScore 23

CalculateQeaScore Calculate quantitative enrichment score
---

# Description

Calculate quantitative enrichment score

### Usage

```
CalculateQeaScore(mSetObj=NA, nodeImp, method)
```

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

nodeImp Indicate the pathway topology analysis, "rbc" for relative-betweeness centrality,

and "dgr" for out-degree centrality.

method Indicate the pathway enrichment analysis, global test is "gt" and global ancova

is "ga".

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CalculateSSP

Single sample profiling to compare with

# Description

reference concentrations stored in the library

#### Usage

```
CalculateSSP(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

24 CleanData

CheckMetaDataConsistency

Check if data are ready for meta-analysis

### **Description**

This function determines if all annotated data are ready for meta-analysis

# Usage

```
CheckMetaDataConsistency(mSetObj = NA, combat = TRUE)
```

### Arguments

mSetObj Input name of the created mSet Object

combat Adjust for batch effects, logical variable: TRUE = adjust for batch effects us-

ing an empirical Bayes framework (R package sva), FALSE = no batch effect

adjustment.

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CleanData

Perform data cleaning

### **Description**

Cleans data and removes -Inf, Inf, NA, negative and 0s.

### Usage

```
CleanData(bdata, removeNA = T, removeNeg = T, removeConst = T)
```

### **Arguments**

bdata Input data to clean

 $\label{eq:logical} \textbf{Logical}, \textbf{T} \ \textbf{to} \ \textbf{remove} \ \textbf{NAs}, \textbf{F} \ \textbf{to} \ \textbf{not}.$ 

removeNeg Logical, T to remove negative numbers, F to not.

removeConst Logical, T to remove samples/features with 0s, F to not.

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CleanDataMatrix 25

CleanDataMatrix

Clean the data matrix

# Description

Function used in higher functinos to clean data matrix

# Usage

CleanDataMatrix(ndata)

# Arguments

ndata

Input the data to be cleaned

CleanNumber

Replace infinite numbers

# Description

Replace -Inf, Inf to 99999 and -99999

# Usage

CleanNumber(bdata)

# Arguments

bdata

Input matrix to clean numbers

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

26 ClearStrings

ClearNegatives

Data processing: Dealing with negative values

# Description

Operates on dataSet\$proc after dealing with missing values

### Usage

```
ClearNegatives(mSetObj = NA, method = "abs")
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

method Input the method to clear negatives

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

ClearStrings

Remove spaces

# Description

Remove from, within, leading and trailing spaces

# Usage

```
ClearStrings(query)
```

# Arguments

query

Input the query to clear

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

ClearUserDir 27

ClearUserDir

Clear folder and memory

# Description

Clear the current folder and objects in memory

### Usage

```
ClearUserDir(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt Compound\_function\_mzlist}$ 

Makes adducts

# Description

Makes adducts

### Usage

```
Compound_function_mzlist(ms_mode, mw)
```

 ${\tt ComputeAverageCurve}$ 

Compute average ROC curve

# Description

Compute the average ROC curve

# Usage

ComputeAverageCurve(perf, avg.method)

28 ComputeHighLow

### **Arguments**

perf Input the average

avg. method Input the name of the method to compute the average curve

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

computeConc

Lipid analysis

### **Description**

The upper limit for each combination is considered to be the minimal of the fatty acid concentration (nmol fatty acid/gram of sample) X is the lopomics data obtained above the result is the saved as separate files for each lipid class

### Usage

```
computeConc(X, iso = "y")
```

### **Arguments**

X Input the data

iso Default is set to "y"

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

ComputeHighLow

Compute the 95 percent interval for threshold ROC

# Description

Computes the 95 percent interval only for the y-axis. Utility function, called upon by higher functions

### Usage

```
ComputeHighLow(perf)
```

### **Arguments**

perf Input the performance

Convert2Mummichog 29

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Convert2Mummichog

Convert mSetObj to proper format for MS Peaks to Pathways module

# Description

Following t-test analysis, this functions converts the results from the mSetObj to the proper format for mummichog analysis

### Usage

```
Convert2Mummichog(mSet0bj = NA, rt = FALSE)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj.

### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateAnalNullMsg

Create null message for analysis Creates a message for the Sweave report

### **Description**

Creates a message stating that no analyses were performed on your data.

# Usage

CreateAnalNullMsg()

30 CreateAOV2doc

CreateANOVAdoc

Create report of analyses

### **Description**

Report generation using Sweave Create ANOVA document

### Usage

```
CreateANOVAdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateAOV2doc

Create report of analyses

# Description

Report generation using Sweave ANOVA

### Usage

```
CreateAOV2doc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateASCAdoc 31

CreateASCAdoc

Create report of analyses

# Description

Report generation using Sweave Random Forest ASCA

### Usage

```
CreateASCAdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt CreateBiomarkerInputDoc}$ 

Create biomarker analysis report: Data Input

### **Description**

Report generation using Sweave Power analysis report, data input documentation.

# Usage

```
CreateBiomarkerInputDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateBiomarkerIntr

Create biomarker analysis report: Introduction

### **Description**

Report generation using Sweave Biomarker analysis report introduction

### Usage

CreateBiomarkerIntr()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateBiomarkerOverview

Create biomarker analysis report: Overview

### **Description**

Report generation using Sweave Power analysis report overview

### Usage

CreateBiomarkerOverview()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateBiomarkerRatioOverview

Create biomarker analysis report: Normalization, ratio

# Description

Report generation using Sweave Biomarker analysis, ratio option

### Usage

CreateBiomarkerRatioOverview(mSetObj = NA)

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateBiomarkerRnwReport

Create report of analyses (Biomarker)

### **Description**

Report generation using Sweave Puts together the analysis report

### Usage

CreateBiomarkerRnwReport(mSetObj, usrName)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateCorrDoc

Create report of analyses

### **Description**

Report generation using Sweave Create correlation document

### Usage

```
CreateCorrDoc(mSetObj = NA)
```

### **Arguments**

mSetObj Input the name

Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

34 CreateEBAMdoc

createCVset

Separate data set using k-fold cross validation (CV)

### **Description**

Separate data set with k-fold CV, used in higher function

# Usage

```
createCVset(groupN, kfold, rseed)
```

### Arguments

groupN Input the size of the group

kfold Input the number of cross-validations

rseed Input the random seed

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateEBAMdoc

Create report of analyses

### **Description**

Report generation using Sweave Create EBAM document Note: the search for delta (SAM) and a0 (EBAM) will not be plotted it is only exploration, and may cause potential inconsistentcies.

# Usage

```
CreateEBAMdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichAnalDoc 35

 ${\tt CreateEnrichAnalDoc}$ 

Create report of analyses (Met Enrichment)

# **Description**

Report generation using Sweave Metabolite enrichment analysis report, analysis

### Usage

CreateEnrichAnalDoc()

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateEnrichInputDoc Create report of analyses (Met Enrichment)

# Description

Report generation using Sweave Metabolite enrichment analysis report data input

### Usage

```
CreateEnrichInputDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

36 CreateEnrichORAdoc

CreateEnrichIntr

Create report of analyses (Met Enrichment)

### **Description**

Report generation using Sweave Metabolite enrichment analysis report introduction

# Usage

```
CreateEnrichIntr()
```

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichORAdoc

Create report of analyses (Met Enrichment)

### **Description**

Report generation using Sweave Metabolite enrichment analysis report, over representation analysis (ORA)

# Usage

```
CreateEnrichORAdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichOverview 37

CreateEnrichOverview Create report of analyses (Met Enrichment)

# Description

Report generation using Sweave Metabolite enrichment analysis report overview

### Usage

```
CreateEnrichOverview()
```

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichProcessDoc

Create report of analyses (Met Enrichment)

# Description

Report generation using Sweave Metabolite enrichment analysis report enrichment process

# Usage

```
CreateEnrichProcessDoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichQEAdoc

Create report of analyses (Met Enrichment)

### **Description**

Report generation using Sweave Metabolite enrichment analysis report Quantitative enrichment analysis

### Usage

```
CreateEnrichQEAdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichRnwReport Create report of analyses (Met Enrichment)

# Description

Report generation using Sweave Metabolite enrichment analysis report

# Usage

```
CreateEnrichRnwReport(mSetObj, usrName)
```

#### **Arguments**

mSetObj Inp

Input the name of the created mSetObj (see InitDataObjects)

usrName Inpu

Input the name of the user

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateEnrichSSPdoc 39

CreateEnrichSSPdoc

Create report of analyses (Met Enrichment)

# Description

Report generation using Sweave Metabolite enrichment analysis report Single sampling profiling

### Usage

```
CreateEnrichSSPdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateFooter

Create report of analyses (Met Enrichment)

### **Description**

Report generation using Sweave Metabolite enrichment analysis report footer

### Usage

```
CreateFooter()
```

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

40 CreateGSEAAnalTable

CreateGraph

Create igraph from the edgelist saved from graph DB and decompose into subnets

# Description

Function for the network explorer module, prepares user's data for network exploration.

# Usage

```
CreateGraph(mSetObj = NA)
```

### **Arguments**

mSetObj

Input name of the created mSet Object

CreateGSEAAnalTable

Create Mummichog report of analyses

# Description

Report generation using Sweave Function to create a summary table of mummichog analysis

### Usage

```
CreateGSEAAnalTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

CreateHCdoc 41

CreateHCdoc

Create report of analyses

### **Description**

Report generation using Sweave Create hierarchical clustering document

### Usage

```
CreateHCdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateHeatmap2doc

Create report of analyses

# Description

Report generation using Sweave 2-way heatmap

### Usage

```
CreateHeatmap2doc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt CreateIntegPathwayAnalysisRnwReport}$ 

Create report of analyses (IntegPathwayAnalysis)

# **Description**

Report generation using Sweave Puts together the analysis report

### Usage

CreateIntegPathwayAnalysisRnwReport(mSetObj, usrName)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateIntegratedPathwayAnalInputDoc

Create integrated pathway report: Data Input

### **Description**

Report generation using Sweave integrated pathway report, data input documentation.

### Usage

CreateIntegratedPathwayAnalInputDoc(mSetObj = NA)

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

CreateIntegratedPathwayAnalIntr

Create integrated pathway analysis report: Introduction

# Description

Report generation using Sweave Integrated pathwayr analysis report introduction

## Usage

CreateIntegratedPathwayAnalIntr()

# Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateIntegratedPathwayDoc

Create integrated pathway analysis report

# Description

Report generation using Sweave Biomarker analysis report, ROC Curve Based Model Creation and Evaluation

#### Usage

CreateIntegratedPathwayDoc(mSetObj = NA)

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

 ${\tt CreateIntegratedPathwayGeneMapTable}$ 

Create a x-table for gene name mapping

# Description

Report generation using Sweave Function to create a table for gene name mapping

# Usage

```
CreateIntegratedPathwayGeneMapTable(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateIntegratedPathwayNameMapTable

Create a x-table for compound name mapping

# Description

Report generation using Sweave Function to create a table for compound name mapping

### Usage

```
CreateIntegratedPathwayNameMapTable(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

 ${\tt CreateIntegratedPathwayResultsTable}$ 

Create a x-table for pathway results

### **Description**

Report generation using Sweave Function to create a table for pathway results

#### **Usage**

```
CreateIntegratedPathwayResultsTable(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateiPCAdoc}$ 

Create report of analyses

### **Description**

Report generation using Sweave For Interactive PCA

# Usage

```
CreateiPCAdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

46 CreateLadder

CreateKMdoc

Create report of analyses

### **Description**

Report generation using Sweave Create Kmeans partitional clustering document

### Usage

```
CreateKMdoc(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateLadder

R-code for R-SVM

### **Description**

use leave-one-out / Nfold or bootstrape to permute data for external CV build SVM model and use mean-balanced weight to sort genes on training set and recursive elimination of least important genes

# Usage

```
CreateLadder(Ntotal, Nmin = 5)
```

#### **Arguments**

Ntotal

Total number

Nmin

Minimum number, default set to 5

### Author(s)

Dr. Xin Lu, Research Scientist Biostatistics Department, Harvard School of Public Health create a decreasing ladder for recursive feature elimination

CreateLibFromKEGG 47

CreateLibFromKEGG

Creates cpd.tree

#### **Description**

Creates cpd.tree

#### Usage

```
CreateLibFromKEGG(cpd.lib, pathways, org)
```

 ${\tt Create Mapping Result Table}$ 

Creates the mapping result table

# Description

Creates the mapping result table

# Usage

```
CreateMappingResultTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

CreateMBdoc

Create report of analyses

### **Description**

Report generation using Sweave Multivariate Bayes

### Usage

```
CreateMBdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateMetaAnalTable

Create Mummichog report of analyses

# Description

Report generation using Sweave Function to create a summary table of mummichog analysis

#### Usage

```
CreateMetaAnalTable(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateMetaAnalysisDEdoc

Create MetaAnalysis analysis report: Data Normalization

### **Description**

Report generation using Sweave Meta-Analysis, data normalization documentation.

# Usage

```
CreateMetaAnalysisDEdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

CreateMetaAnalysisInputDoc

Create MetaAnalysis analysis report: Data Input

# Description

Report generation using Sweave Power analysis report, data input documentation.

# Usage

```
CreateMetaAnalysisInputDoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateMetaAnalysisIntr

Create MetaAnalysis analysis report: Introduction

# Description

Report generation using Sweave MetaAnalysis analysis report introduction

### Usage

```
CreateMetaAnalysisIntr()
```

# Author(s)

CreateMetaAnalysisNORMdoc

Create MetaAnalysis analysis report: Data Normalization

# Description

Report generation using Sweave Meta-Analysis, data normalization documentation.

# Usage

```
CreateMetaAnalysisNORMdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateMetaAnalysisOutput}$ 

Create MetaAnalysis analysis report: Data Normalization

# Description

Report generation using Sweave MetaAnalysis analysis, data normalization documentation.

### Usage

```
CreateMetaAnalysisOutput(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

CreateMetaAnalysisOverview

Create MetaAnalysis analysis report: Overview

# Description

Report generation using Sweave Power analysis report overview

# Usage

CreateMetaAnalysisOverview()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt Create Meta Analysis Rnw Report}$ 

Create report of analyses (Meta-Analysis)

# Description

Report generation using Sweave Puts together the analysis report

### Usage

CreateMetaAnalysisRnwReport(mSetObj, usrName)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

### Author(s)

CreateMetaTable

Create MetaAnalysis table of results

### **Description**

Report generation using Sweave Function to create a table containing meta-analysis results.

### Usage

```
CreateMetaTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateModelBiomarkersDoc

Create biomarker analysis report: ROC Curve Based Model Creation and Evaluation

### **Description**

Report generation using Sweave Biomarker analysis report, ROC Curve Based Model Creation and Evaluation

### Usage

```
CreateModelBiomarkersDoc(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

CreateMultiBiomarkersDoc

53

CreateMultiBiomarkersDoc

Create biomarker analysis report: Multivariate Biomarker Analysis

# Description

Report generation using Sweave Biomarker analysis report, Multivariate Biomarker Analysis

# Usage

```
CreateMultiBiomarkersDoc(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateMummichogAnalTable}$ 

Create Mummichog report of analyses

### **Description**

Report generation using Sweave Function to create a summary table of mummichog analysis

### Usage

```
CreateMummichogAnalTable(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

 ${\tt CreateMummichogAnalysisDoc}$ 

Create mummichog analysis report

# Description

Report generation using Sweave Mummichog analysis report

# Usage

```
CreateMummichogAnalysisDoc(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateMummichogInputDoc}$ 

Create Mummichog analysis report: Data Input

# Description

Report generation using Sweave Mummichog analysis report, data input documentation.

# Usage

```
CreateMummichogInputDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

CreateMummichogIntro Create mummichog analysis report: Introduction

### Description

Report generation using Sweave Mummichog analysis report introduction

### Usage

CreateMummichogIntro()

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateMummichogLibs

Create Mummichog Libraries from KEGG

### Description

Function to create mummichog libraries from MetaboAnalyst pathway libraries (metpa). Outputs the RDS files in the current working directory. RDS files are saved using the KEGG organism code.

### Usage

CreateMummichogLibs("~/Desktop/MetaboAnalyst/mummichog/2020\_mummichog\_libs/test", kegg\_compounds\_20

# **Arguments**

folder Input the path of the folder containing the metpa rda files.

kegg\_compounds Input the name of the KEGG dictionary containing the KEGG compound IDs,

KEGG compopund names, and molecular weight.

CreateMummichogOverview

Create Mummichog analysis report: Overview

# Description

Report generation using Sweave Mummichog analysis report overview

# Usage

CreateMummichogOverview()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateMummichogRnwReport}$ 

Create report of analyses (Biomarker)

# Description

Report generation using Sweave Puts together the analysis report

# Usage

CreateMummichogRnwReport(mSetObj, usrName)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

# Author(s)

 ${\tt CreateNetworkExplorerDoc}$ 

Create integrated pathway analysis report

# Description

Report generation using Sweave Biomarker analysis report, ROC Curve Based Model Creation and Evaluation

### Usage

```
CreateNetworkExplorerDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateNetworkExplorerInputDoc

Create network explorer: Data Input

### **Description**

Report generation using Sweave network explorer report, data input documentation.

#### Usage

```
CreateNetworkExplorerInputDoc(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

 ${\tt CreateNetworkExplorerIntr}$ 

Create integrated pathway analysis report: Introduction

# Description

Report generation using Sweave Network explorer report introduction

### Usage

CreateNetworkExplorerIntr()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateNetworkExplorerOverview

Create network explorer report: Overview

# Description

Report generation using Sweave for the network explorer report overview

# Usage

CreateNetworkExplorerOverview()

### Author(s)

 ${\tt CreateNetworkExplorerRnwReport}$ 

Create report of analyses (Network Explorer)

# **Description**

Report generation using Sweave Puts together the analysis report

#### Usage

CreateNetworkExplorerRnwReport(mSetObj, usrName)

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateNetworkGeneMapTable

Create a x-table for gene name mapping

# Description

Report generation using Sweave Function to create a table for gene name mapping

# Usage

CreateNetworkGeneMapTable(mSetObj = NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

60 CreateNORMdoc

 ${\tt CreateNetworkNameMapTable}$ 

Create a x-table for compound name mapping

### **Description**

Report generation using Sweave Function to create a table for compound name mapping

#### Usage

```
CreateNetworkNameMapTable(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateNORMdoc

Create report of analyses

### **Description**

Report generation using Sweave Create normalization document

# Usage

```
CreateNORMdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateOPLSDAdoc 61

CreateOPLSDAdoc

Create report of analyses

### **Description**

Report generation using Sweave Create OPLSDA document

### Usage

```
CreateOPLSDAdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreatePathAnalDoc

Create report of analyses (Met Pathway)

# Description

Report generation using Sweave Metabolomic pathway analysis Create pathway analysis doc

### Usage

```
CreatePathAnalDoc(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

62 CreatePathProcessDoc

CreatePathInputDoc

Create report of analyses (Met Pathway)

### **Description**

Report generation using Sweave Metabolomic pathway analysis Create data input doc

### Usage

```
CreatePathInputDoc()
```

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreatePathIntr

Create report of analyses (Met Pathway)

### **Description**

Report generation using Sweave Metabolomic pathway analysis Introduction

#### Usage

```
CreatePathIntr()
```

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreatePathProcessDoc Create report of analyses (Met Pathway)

### **Description**

Report generation using Sweave Metabolomic pathway analysis Create MetPA process

### Usage

```
CreatePathProcessDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

CreatePathResultDoc 63

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreatePathResultDoc

Create report of analyses (Met Pathway)

### **Description**

Report generation using Sweave Metabolomic pathway analysis Create MetPA results doc

### Usage

```
CreatePathResultDoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreatePathRnwReport}$ 

Create report of analyses (Met Pathway)

# Description

Report generation using Sweave Metabolomic pathway analysis write .Rnw file template

### Usage

```
CreatePathRnwReport(mSetObj, usrName)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

usrName

Input the name of the user

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

64 CreatePLSdoc

CreatePCAdoc

Create report of analyses

### **Description**

Report generation using Sweave Create PCA document

### Usage

```
CreatePCAdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreatePLSdoc

Create report of analyses

# Description

Report generation using Sweave Create PLS document

### Usage

```
CreatePLSdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreatePowerAnalDoc 65

CreatePowerAnalDoc

Create power analysis report: Power Analysis

### **Description**

Report generation using Sweave Power analysis report, analysis

### Usage

```
CreatePowerAnalDoc(mSetObj)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreatePowerInputDoc

Create power analysis report: Data Input

# Description

Report generation using Sweave Power analysis report, data input documentation.

### Usage

```
CreatePowerInputDoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

CreatePowerIntr

Create power analysis report: Introduction

# Description

Report generation using Sweave Power analysis report introduction

### Usage

CreatePowerIntr()

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreatePowerOverview

Create power analysis report: Overview

# Description

Report generation using Sweave Power analysis report overview

### Usage

CreatePowerOverview()

#### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreatePowerParametersDoc

Create power analysis report: Power Parameter Selection

### **Description**

Report generation using Sweave Power analysis report, parameter selection

### Usage

CreatePowerParametersDoc(mSetObj = NA)

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreatePowerRnwReport Create report of analyses (Power)

### **Description**

Report generation using Sweave Put together the analysis report

### Usage

CreatePowerRnwReport(mSetObj, usrName)

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CreateRatioTable Create report of analyses

### **Description**

Report generation using Sweave Function to create a summary table for biomarker analysis: included metabolite ratios

#### Usage

CreateRatioTable(mSetObj = NA)

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

 ${\tt CreateRFdoc}$ 

Create report of analyses

# Description

Report generation using Sweave Create Random Forest document

### Usage

```
CreateRFdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateRHistAppendix

Create report of analyses

### **Description**

Report generation using Sweave Create footer

### Usage

```
CreateRHistAppendix()
```

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateROCLabelsTable 69

CreateROCLabelsTable Create a x-table for newly classified samples

### **Description**

Report generation using Sweave Function to create a table for newly classified samples

### Usage

```
CreateROCLabelsTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

 ${\tt CreateSAMdoc}$ 

Create report of analyses

# Description

Report generation using Sweave Create SAM document

# Usage

```
CreateSAMdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

70 CreateSOMdoc

CreateSemiTransColors Create semitransparant colors

### **Description**

Create semitransparant colors for a given class label

# Usage

```
CreateSemiTransColors(cls)
```

### **Arguments**

cls

Input class labels

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateSOMdoc

Create report of analyses

# Description

Report generation using Sweave Create SOM partitional clustering document

### Usage

```
CreateSOMdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateSPLSDAdoc 71

 ${\tt CreateSPLSDAdoc}$ 

Create report of analyses

# Description

Report generation using Sweave Create sPLS-DA document

### Usage

```
CreateSPLSDAdoc(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateStatIntr

Create report of analyses

### **Description**

Report generation using Sweave Create header

### Usage

```
CreateStatIntr()
```

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateStatIOdoc

Create report of analyses

# Description

Report generation using Sweave Read and process raw data

### Usage

```
CreateStatIOdoc(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt CreateStatRnwReport}$ 

Create report for statistical analysis module

### **Description**

Report generation using Sweave Write .Rnw file template

# Usage

```
CreateStatRnwReport(mSetObj, usrName)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

usrName

Input the name of the user

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateSummaryTable 73

CreateSummaryTable

Create report of analyses

# Description

Report generation using Sweave Create a summary table for each type of uploaded data csv table has 5 col: sampleID, feature #, zero, missing #

### Usage

```
CreateSummaryTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateSVMdoc

Create report of analyses

## Description

Report generation using Sweave Create R-SVM document

## Usage

```
CreateSVMdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

74 CreateTimeSeriesIOdoc

 ${\tt CreateTimeSeriesAnalNullMsg}$ 

Create null analysis message for time-series sweave report

## Description

Creates empty time-series analysis message

### Usage

CreateTimeSeriesAnalNullMsg()

CreateTimeSeriesIOdoc Create report of analyses (Met Pathway)

## Description

Report generation using Sweave Metabolomic pathway analysis, time-series Read and process the raw data

## Usage

```
CreateTimeSeriesIOdoc(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateTimeSeriesRnwReport

Create report of analyses (Met Pathway)

### Description

Report generation using Sweave Metabolomic pathway analysis Create timeseries .Rnw file template

#### Usage

CreateTimeSeriesRnwReport(mSetObj, usrName)

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

usrName Input the name of the user

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CreateUnivarBiomarkersDoc

Create power analysis report: Biomarker Univariate Analysis

## Description

Report generation using Sweave Biomarker analysis report, Univariate Analysis

### Usage

CreateUnivarBiomarkersDoc(mSetObj = NA)

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

76 CreateUnivROCTable

CreateUNIVdoc

Create report of analyses

## Description

Report generation using Sweave Create univariate analyses document

## Usage

```
CreateUNIVdoc(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

CreateUnivROCTable

Create summary table for univariate ROC analysis

### **Description**

Report generation using Sweave Function to create a summary table for univariate biomarker analysis

## Usage

```
CreateUnivROCTable()
```

## Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

Create VennMetaTable 77

CreateVennMetaTable

Create MetaAnalysis table of results for Venn Diagram

### **Description**

Report generation using Sweave Function to create a table containing meta-analysis results.

## Usage

```
CreateVennMetaTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

CrossReferencing

Various functions for mapping b/w names & database identifiers Given a list of compound names or ids, find matched name or ids from selected databases

### **Description**

Given a list of compound names or ids find matched name or IDs from selected databases

# Usage

```
CrossReferencing(
  mSetObj = NA,
  q.type,
  hmdb = T,
  pubchem = T,
  chebi = F,
  kegg = T,
  metlin = F
)
```

78 CVTest.LRmodel

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects).

q.type Input the query type, "name" for compound names, "hmdb" for HMDB IDs, "kegg" for KEGG IDs, "pubchem" for PubChem CIDs, "chebi" for ChEBI IDs,

"metlin" for METLIN IDs, and "hmdb\_kegg" for a both KEGG and HMDB IDs.

hmdb Logical, T to cross reference to HMDB, F to not.

pubchem Logical, T to cross reference to PubChem, F to not.

chebi Logical, T to cross reference to CheBI, F to not.

kegg Logical, T to cross reference to KEGG, F to not.

metlin Logical, T to cross reference to MetLin, F to not.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

CVTest.LRmodel Calculate ROC performance with CV

### Description

Calculate ROC performance with CV

#### Usage

```
CVTest.LRmodel(data.in, fmla.in, kfold = 10, run.stepwise = FALSE)
```

### **Arguments**

data.in Input matrix of data

fmla.in Input for generalized linear model

kfold Numeric run.stepwise Logical

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

descendMin 79

descendMin

Perform utilities for peak grouping

## Description

Perform various utilities for peak grouping

## Usage

```
descendMin(y, istart = which.max(y))
```

# Arguments

y Input peaks

istart Performs which.max on y

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt doCompoundMapping}$ 

Perform compound mapping

## Description

Perform compound mapping

## Usage

```
doCompoundMapping(cmpd.vec, q.type)
```

# Arguments

 ${\tt cmpd.vec} \qquad \qquad {\tt Input\ compound\ vector}$ 

q. type Query type

doGeneIDMapping

Convert different gene IDs into entrez IDs for downstream analysis

## Description

Gene ID mapping, gene annotation, compound mapping, KEGG mapping

## Usage

```
doGeneIDMapping(q.vec, org, type)
```

## Arguments

q.vec Input the query

org Input the organism type

type Input the type of data to annotate

## Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

doKEGG2NameMapping

Perform KEGG to compound name mapping

# Description

Perform KEGG to compound name mapping

## Usage

```
doKEGG2NameMapping(kegg.vec)
```

### **Arguments**

kegg.vec

Input vector of KEGG compounds

doKOFiltering 81

doKOFiltering

Utility function

### **Description**

Returns matched KO in the same order (NA if no match)

# Usage

```
doKOFiltering(ko.vec, type)
```

## Arguments

ko.vec

Input the vector containing KOs

type

Input the type

EBAM.Init

For EBAM analysis

### **Description**

deteriming a0, only applicable for z.ebam (default)

## Usage

```
EBAM.Init(
   mSetObj = NA,
   isPaired,
   isVarEq,
   nonPar,
   A0 = -99,
   delta,
   imgA0,
   imgSig
)
```

## Arguments

mSetObj Input name of the created mSet Object

isPaired Logical isVarEq Logical

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

FC.Anal.paired

ExtractMS2data	Extract MS2 Data
----------------	------------------

### **Description**

This function returns a list of spectra that matches with a user specified precursor m/z.

### Usage

```
ExtractMS2data(filename, peakParams, mzmin, mzmax)
```

### **Arguments**

filename Name of the file (e.g. mzML, mzXML)

peakParams Object containing parameters for peak picking.

mzmin Minimum m/z when selecting a precursor from peak list mzmax Maximum m/z when selecting a precursor from peak list

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

FC.Anal.paired Fold change analysis, paired

### Description

Perform paired fold change analysis

### Usage

```
FC.Anal.paired(
   mSetObj = NA,
   fc.thresh = 2,
   percent.thresh = 0.75,
   cmp.type = 0
)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

fc. thresh Fold-change threshold, numeric input

percent.thresh Numeric input, from 0 to 1 to indicate the significant count threshold cmp.type Comparison type, 0 for group 1 minus group 2, and 1 for group

FC.Anal.unpaired 83

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

FC.Anal.unpaired

Fold change analysis, unpaired

#### **Description**

Perform fold change analysis, method can be mean or median

#### Usage

```
FC.Anal.unpaired(mSetObj, fc.thresh=2, cmp.type = 0)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

fc. thresh Fold-change threshold, numeric input

cmp. type Comparison type, 0 for group 1 minus group 2, and 1 for group 1 minus group 2

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

FeatureCorrelation

Pattern hunter

#### **Description**

Calculate correlation of all other feature to a given feature name

#### Usage

```
FeatureCorrelation(mSetObj = NA, dist.name, varName)
```

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

dist.name Input the name of the distance measure

varName Input the variable name

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

84 FilterVariable

fgsea2

Pre-ranked gsea adapted for untargeted metabolomics

### **Description**

Pre-ranked gsea adapted for untargeted metabolomics

### Usage

```
fgsea2(
  mSetObj,
  pathways,
  stats,
  ranks,
  nperm,
  minSize = 1,
  maxSize = Inf,
  nproc = 0,
  gseaParam = 1,
  BPPARAM = NULL
)
```

fillpathways

Fill in the pathways

### **Description**

Fill in the pathways

### Usage

fillpathways(f)

FilterVariable

Methods for non-specific filtering of variables

### **Description**

This is a function that filters the dataset, dependent on the user-specified method for filtering. The function applies a filtering method, ranks the variables within the dataset, and removes variables based on its rank. The final dataset should contain no more than than 5000 variables for effective computing.

findEqualGreaterM 85

### Usage

```
FilterVariable(mSetObj=NA, filter, qcFilter, rsd)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

filter Select the filter option, "rsd" which is the relative standard deviation, "nrsd"

which is the non-parametric relative standard deviation, "mean" which is the mean, "sd" which is the standard deviation, "mad" which is the median absolute

deviation, or "iqr" which is the interquantile range.

qcFilter Filter the variables based on QC samples - True (T), or use non-QC based filter-

ing - False (F).

rsd Define the relative standard deviation cut-off. Variables with a RSD greater than

this number will be removed from the dataset. It is only necessary to specify this argument if qcFilter is True (T). Otherwise, it will not be used in the function.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

findEqualGreaterM Perform utilities for peak grouping

### Description

Perform various utilities for peak grouping

## Usage

findEqualGreaterM(x, values)

## **Arguments**

x Input the datavalues Input the values

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

86 FormatPeakList

FisherLSD

Fisher for ANOVA

### **Description**

Perform Fisher LSD for ANOVA, used in higher function

## Usage

```
FisherLSD(aov.obj, thresh)
```

#### **Arguments**

aov.obj Input the anova object

thresh Numeric, input the alpha threshold

#### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

FormatPeakList

Format Peak List

## Description

This function formats the CAMERA output to a usable format for MetaboAanlyst.

## Usage

```
FormatPeakList(
   annotPeaks,
   annParams,
   filtIso = TRUE,
   filtAdducts = FALSE,
   missPercent = 0.5
)
```

## Arguments

annotPeaks The object created using the PerformPeakAnnotation.

annParams The object created using the SetAnnotationParam function, containing user's

specified or default parameters for downstream raw MS data pre-processing.

filtIso Logical, filter out all isotopes except for [M]+ for positive ion mode and [M]-

for negative ion mode. By default it is set to true.

genLogisticRegMdl 87

filtAdducts Logical, filter out all adducts except [M+H]+ for positive ion more and [M-H]-

for negative ion mode. By default it is set to false.

missPercent Numeric, specify the threshold to remove features missing in X% of samples.

For instance, 0.5 specifies to remove features that are missing from 50% of all

samples per group. Method is only valid when there are two groups.

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 $\begin{array}{ll} {\it genLogisticRegMdl} & {\it Develop\ a\ Logistic\ Regression\ Model\ with\ all\ of\ the\ combined\ k-fold} \\ {\it CV\ subsets} \end{array}$ 

### Description

Develop a Logistic Regression Model with all of the combined k-fold CV subsets

#### Usage

```
genLogisticRegMdl(x.train, y.train, x.test, y.test)
```

# Arguments

x.train Input the X training sety.train Input the Y training setx.test Input the X test sety.test Input the Y test set

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

88 Get.bwss

Get.asca.tss

Function for ASCA permutation

### Description

Dummy is used only for the purpose to maintain lapply API this is used for permutation on ANOVA paritions, not on the SCA/PCA part, so the number of selected components is not applicable in this step

### Usage

```
Get.asca.tss(dummy, perm = T)
```

#### Arguments

dummy Dummy variable

perm Logical, TRUE by default

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Get.bwss

Compute within group and between group sum of squares (BSS/WSS) for each row of a matrix which may have NA

### **Description**

Columns have labels, x is a numeric vector, cl is consecutive integers

### Usage

```
Get.bwss(x, cl)
```

#### **Arguments**

x Numeric vector

cl Columns

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Get.ConcRef 89

Get.ConcRef

Get the concentration reference

## Description

Get the concentration reference

## Usage

```
Get.ConcRef(mSetObj = NA, cmpd.nm)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

cmpd.nm

Input the compound name

Get.Leverage

Fast leverage calculation for permutation purpose

# Description

note, the leverage combines all components the importance feature is for the factor not per components

## Usage

```
Get.Leverage(XKw, Fac)
```

### **Arguments**

XKw Features
Fac Factor

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

90 Get.pred

. pAUC

Calculate partial area under ROC curve

## Description

Calculate partial area under ROC curve

## Usage

```
Get.pAUC(x, y, focus, cutoff)
```

### **Arguments**

X	Input X
у	Input Y
focus	Method
cutoff	Numerio

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Get.	nred
oc c.	pı cu

Get predicted class probability

## Description

Get predicted class probability, used in higher function

### Usage

```
Get.pred(x.train, y.train, x.test, y.test, clsMethod = "pls")
```

### **Arguments**

x.train	Training X
y.train	Training Y
x.test	Test X
y.test	Test Y

clsMethod Method to predict class, by default it is PLS

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Get.rpart.summary 91

Get.rpart.summary

Get the text description of a recursive partitioning (rpart) result

## Description

x must be an rpart object

#### Usage

```
Get.rpart.summary(x)
```

## Arguments

Х

An Rpart object

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Get.VIP

Calculate variable importance of projection (VIP) score for PLS object

### **Description**

Users give a pls object ('oscorespls'=T), function calculates VIP score usually one VIP for each component, return is the average of all VIP

#### Usage

```
Get.VIP(pls.obj, comp = 2)
```

### **Arguments**

pls.obj Input the PLS object

comp Numeric, input the number of components, by default it is 2

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

92 GetAllDataNames

GetAbundanceLabel

Determine value label for plotting

## Description

Concentration or intensity data type

### Usage

```
GetAbundanceLabel(data.type)
```

### **Arguments**

data.type

Input concentration or intensity data

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt GetAccuracyInfo}$ 

Export biomarker accuracy information

### **Description**

Export biomarker accuracy information

### Usage

```
GetAccuracyInfo(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetAllDataNames

Get all meta-analysis name data

### **Description**

Get all meta-analysis name data

#### Usage

```
GetAllDataNames()
```

GetAllKMClusterMembers 93

GetAllKMClusterMembers

K-means analysis - cluster

## Description

K-means analysis - cluster

## Usage

```
GetAllKMClusterMembers(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetAllSOMClusterMembers

SOM analysis

## Description

Get members for given cluster index, return a character string

# Usage

```
GetAllSOMClusterMembers(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

94 GetCircleInfo

 ${\tt GetCandidateList}$ 

Get all candidate compound names for a given index

## Description

Returns 3 coloumns - inx, name, score

### Usage

```
GetCandidateList(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetCircleInfo

Export information about selected circle

### **Description**

Export information about selected circle

### Usage

```
GetCircleInfo(mSetObj = NA)
```

## Arguments

mSetObj

Input name of the created mSet Object

GetCIs 95

GetCIs

Get confidence intervals

## Description

For non-parametric tests, use quantiles, use normal (1.96\*std.err) if parametric

### Usage

```
GetCIs(data, param = F)
```

# **Arguments**

data Input data matrix

param Logical, False by default

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetCMD

Retrieve last command from the Rhistory.R file

## Description

Fetches the last command from the Rhistory.R file

### Usage

```
GetCMD(regexp)
```

## Arguments

regexp

Retrieve last command from Rhistory file

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

96 GetConvertFullPath

GetCompoundDetails

Function to get adduct details from a specified compound

## Description

Function to get adduct details from a specified compound. The results will be both printed in the console as well as saved as a csv file. Note that performing this function multiple times will overwrite previous queries.

### Usage

```
GetCompoundDetails(mSetObj = NA, cmpd.id)
```

## Arguments

mSetObj Input the name of the created mSetObj object.

 ${\sf cmpd.id} \qquad \qquad {\sf Input\ the\ name\ of\ the\ selected\ compound.}$ 

 ${\tt GetConvertFullPath}$ 

Perform utilities for cropping images

# Description

Obtain the full path to convert (from imagemagik) for cropping images

### Usage

```
GetConvertFullPath()
```

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

getDataFromTextArea 97

getDataFromTextArea

Transform two column text to data matrix

### **Description**

Transform two column input text to data matrix (single column data frame)

## Usage

```
getDataFromTextArea(txtInput, sep.type = "space")
```

### Arguments

txtInput

Input text

sep.type

Indicate the seperator type for input text. Default set to "space"

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetExtendRange

Extend axis

### **Description**

Extends the axis range to both ends vec is the values for that axis unit is the width to extend, 10 will increase by 1/10 of the range

### Usage

```
GetExtendRange(vec, unit = 10)
```

#### **Arguments**

vec

Input the vector

unit

Numeric

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

98 GetFeatureNumbers

GetFC

Used by higher functions to calculate fold change

## Description

Utility method to calculate FC, used in higher function

### Usage

```
GetFC(mSetObj = NA, paired = FALSE, cmpType)
```

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

paired Logical, true of false cmpType Numeric, 0 or 1

#### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetFeatureNumbers

Numbers for subset selection

## Description

Return a series of number for subsets selection

## Usage

```
GetFeatureNumbers(feat.len)
```

### **Arguments**

feat.len Input the feature length

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetFinalNameMap 99

# Description

Returns three columns: original name, HMDB name and KEGG ID, for enrichment and pathway analysis, respectively

### Usage

```
GetFinalNameMap(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

## Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetFisherPvalue

Get fisher p-values

### **Description**

Get fisher p-values

## Usage

```
GetFisherPvalue(numSigMembers, numSigAll, numMembers, numAllMembers)
```

# Arguments

numSigMembers Number of significant members numSigAll Number of all significant features

numMembers Number of members
numAllMembers Number of all members

100 GetHTMLPathSet

GetHTMLMetSet Given a metset inx, return hmtl highlighted metset cmpds and refer-

ences

### **Description**

Given a metset inx, return hmtl highlighted metset cmpds and references

## Usage

```
GetHTMLMetSet(mSetObj = NA, msetNm)
```

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

msetNm Input the name of the metabolite set

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetHTMLPathSet

Given a metset inx, return hmtl highlighted pathway cmpds

### Description

Given a metset inx, return hmtl highlighted pathway cmpds

#### Usage

```
GetHTMLPathSet(mSetObj = NA, msetNm)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

msetNm Input the name of the metabolite set

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetImpFeatureMat 101

natrix	Get important feature matrix	GetImpFeatureMat
--------	------------------------------	------------------

### **Description**

feat.outp is a list that contains the ranked features in each cross validation (CV) and returns a two column matrix, col 1 = median ranking and col 2 = mean importance measure

### Usage

```
GetImpFeatureMat(mSetObj = NA, feat.outp, bestFeatNum)
```

## Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

feat.outp Input the list that contains the ranked features in each cross validation (CV)

and returns a two column matrix, col 1 = median ranking and col 2 = mean

importance measure

bestFeatNum Numeric

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetKEGGNodeInfo Retrieves KEGG node information	
---	--

### **Description**

Retrieves KEGG node information

#### Usage

```
GetKEGGNodeInfo(pathName, g, width, height, usr = par("usr"))
```

# Arguments

g	Input data
width	Input the width
height	Input the height
usr	Input the user
path.id	Input the path ID

102 GetLassoFreqs

GetKMClusterMembers

K-means analysis - cluster

## Description

Get the cluster members for given index add HTML color to the names based on its group membership

### Usage

```
GetKMClusterMembers(mSetObj = NA, i)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

i Input the cluster index

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetLassoFreqs

Compute lasso frequency

## **Description**

Not part of default, need to perform function to compute lasso frequency msg: There are more than 500 variables and n<m You may wish to restart and set use.Gram=FALSE

#### Usage

```
GetLassoFreqs(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetLimmaResTable 103

GetLimmaResTable

Get result table from eBayes fit object

## Description

Get result table from eBayes fit object

## Usage

```
GetLimmaResTable(fit.obj)
```

### Arguments

fit.obj

eBayes fit object to parse to a table

GetMapTable

Get mapping table

## Description

Return results from compound name mapping in a table

### Usage

```
GetMapTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetMaxPCAComp

For plotting PCA, selects max top 9 components

# Description

Rotate PCA analysis

### Usage

```
GetMaxPCAComp(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

104 GetMetaResultMatrix

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetMeanROC

Compute data points on the ROC curve

### **Description**

perf is the performance object from ROCR

# Usage

```
GetMeanROC(perf)
```

#### **Arguments**

perf

Performance object from ROCR

## Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetMetaResultMatrix

Single.type return logFC or p value for individual data analysis

# Description

Single.type return logFC or p value for individual data analysis

### Usage

```
GetMetaResultMatrix(mSetObj = NA, single.type = "fc")
```

### **Arguments**

mSetObj Input name of the created mSet Object

single.type Default is "fc"

GetMetaSigHitsTable 105

 ${\tt GetMetaSigHitsTable}$ 

Export the significant hits from meta-analysis

# Description

Export the significant hits from meta-analysis

## Usage

```
GetMetaSigHitsTable(mSetObj = NA)
```

# Arguments

mSetObj

Input name of the created mSet Object

GetMetSetName

Given a metset inx, give its name

## Description

Given a metset inx, give its name

# Usage

```
GetMetSetName(mSetObj = NA, msetInx)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

msetInx

Input the index of the metabolite set

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

106 GetMsetLibSearchResult

GetMsetLibCheckMsg

Get the library check messages

## Description

Get the library check messages

### Usage

```
GetMsetLibCheckMsg(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetMsetLibSearchResult}$ 

Return metset search results

## Description

since String[][] is not supported, have to return as 1D vector, matrix can be directly convert to vector, note default will be column first

### Usage

```
GetMsetLibSearchResult(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetMsetNames 107

GetMsetNames

Return the selected metset library to java for display

### **Description**

Return the selected metset library to java for display

### Usage

```
GetMsetNames(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetMummichogPathSetDetails

Function to get compound details from a specified pathway

### Description

Function to get compound details from a specified pathway. The results will be both printed in the console as well as saved as a csv file. Note that performing this function multiple times will overwrite previous queries.

### Usage

```
GetMummichogPathSetDetails(mSetObj = NA, msetNm)
```

# Arguments

mSetObj Input the name of the created mSetObj object.

msetNm Input the name of the pathway

**GetNewSampleNames** 

 ${\tt GetNetworkGeneMappingResultTable}$ 

Exports Gene-Mapping result into a table

## Description

Exports Gene-Mapping result into a table

### Usage

```
GetNetworkGeneMappingResultTable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input name of the created mSet Object

GetNewSampleNames

Obtain sample names and their class labels

## Description

Obtain sample names and their class labels

## Usage

```
GetNewSampleNames(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetORA.pathNames 109

GetORA.pathNames

Export pathway names from ORA analysis

#### **Description**

Export pathway names from ORA analysis

### Usage

```
GetORA.pathNames(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetORA.smpdbIDs

Only for human pathways (SMPDB)

### **Description**

Only for human pathways + ath, eco, mmu & sce

### Usage

```
GetORA.smpdbIDs(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt GetORATable}$ 

Get ORA table

#### **Description**

Get ORA table

# Usage

```
GetORATable(mSetObj = NA)
```

#### **Arguments**

mSetObj

110 GetQEA.pathNames

GetQEA.keggIDs

Only for human pathways (KEGG)

# Description

Only for human pathways + ath, eco, mmu & sce

### Usage

```
GetQEA.keggIDs(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetQEA.pathNames

Export pathway names from QEA analysis

### **Description**

Export pathway names from QEA analysis

### Usage

```
GetQEA.pathNames(mSetObj = NA)
```

### **Arguments**

mSetObj

GetQEATable 111

GetQEATable

QEA table

# Description

QEA table

### Usage

```
GetQEATable(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetRCommandHistory}$ 

Export R Command History

### **Description**

Export R Command History

### Usage

```
GetRCommandHistory(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetRFConf.Table

Classification performance table for random forest analysis

# Description

Classification performance table for random forest analysis

### Usage

```
GetRFConf.Table(mSetObj = NA)
```

### **Arguments**

mSetObj

GetRFOOB

GetRFConfMat

Random Forest Confusion Matrix

# Description

Return double confusion matrix

### Usage

```
GetRFConfMat(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetRF00B

Random Forest OOB

# Description

Get the OOB error for the last signif

### Usage

```
GetRFOOB(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetRFSigMat 113

_			
Ca:	tRF:	Siσ	Mat.

Random Forest Significance matrix

### Description

Significance measure, double brackets

### Usage

```
GetRFSigMat(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetROC.coords

Return ROC corodinates with confidence intervals

### **Description**

Return ROC corodinates with confidence intervals

### Usage

```
GetROC.coords(mSetObj = NA, fld.nm, val, plot = TRUE, imgNm)
```

### **Arguments**

mSetObi	Input the name of the created mSetObj (see InitDataObjects)	

fld.nm The kind of input coordinate
val The coordinates to look for
plot Logical, by default set to TRUE

imgNm Input the image name

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

114 GetROCTtestP

GetROCLassoFreq

Get p-values from lasso

# Description

Get p-values from lasso

# Usage

```
GetROCLassoFreq(data, cls)
```

# Arguments

data Input data

cls Input class labels

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetROCTtestP

Get p-values for ROC

# Description

ROC p-vaues, used in higher function

### Usage

```
GetROCTtestP(data, cls)
```

# Arguments

data Input data

cls Input class labels

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetSampleSizeLadder115

 ${\tt GetSampleSizeLadder}$ 

Retrieve sample size ladder

### **Description**

Return sample size ladder, used in higher functions

# Usage

GetSampleSizeLadder(maxNum)

### **Arguments**

maxNum

Numeric

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetSelectedDataNames Retrieve data names

# Description

Retrieve data names

### Usage

GetSelectedDataNames(mSetObj = NA)

# **Arguments**

mSetObj

Input name of the created mSet Object

GetSelectedDataNumber Retrieve selected data numbers

# **Description**

Retrieve selected data numbers

#### Usage

```
GetSelectedDataNumber(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input name of the created mSet Object

116 GetSigTable.Anova

 ${\tt GetSigTable}$ 

Create Latex table

# Description

generate Latex table

# Usage

```
GetSigTable(mat, method, data.type)
```

# Arguments

mat Input matrix

method Input method to create table

data.type Input the data type

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetSigTable.Anova

Sig Table for Anova

# Description

Sig Table for Anova

# Usage

```
GetSigTable.Anova(mSetObj = NA)
```

### **Arguments**

mSetObj

GetSigTable.Aov2

GetSigTable.Aov2

Sig table for AOV2

# Description

Sig table for AOV2

# Usage

```
GetSigTable.Aov2(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetSigTable.ASCA}$ 

Table of features well modelled by ASCA

# Description

Table of features well modelled by ASCA

# Usage

```
GetSigTable.ASCA(mSetObj = NA, nm)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

nm

Input the name of the well modelled features

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

118 GetSigTable.FC

GetSigTable.Corr

Sig table for Correlation Analysis

# Description

Sig table for Correlation Analysis

### Usage

```
GetSigTable.Corr(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetSigTable.EBAM}$ 

Sig table for EBAM

### **Description**

Sig table for EBAM

#### Usage

```
GetSigTable.EBAM(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetSigTable.FC

Sig Table for Fold-Change Analysis

# Description

Sig Table for Fold-Change Analysis

### Usage

```
GetSigTable.FC(mSetObj = NA)
```

### **Arguments**

mSetObj

GetSigTable.MB

 ${\tt GetSigTable.MB}$ 

Sig table for MB analysis

# Description

Sig table for MB analysis

### Usage

```
GetSigTable.MB(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetSigTable.RF}$ 

Sig table for random forest analysis

### **Description**

Sig table for random forest analysis

#### Usage

```
GetSigTable.RF(mSetObj = NA)
```

### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

GetSigTable.SAM

Sig table for SAM

# Description

Sig table for SAM

### Usage

```
GetSigTable.SAM(mSetObj = NA)
```

### **Arguments**

mSetObj

120 GetSigTable.Volcano

 ${\tt GetSigTable.SVM}$ 

Sig table for SVM

# Description

Sig table for SVM

### Usage

```
GetSigTable.SVM(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetSigTable.TT}$ 

Sig Table for T-test Analysis

### **Description**

Sig Table for T-test Analysis

#### Usage

```
GetSigTable.TT(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt GetSigTable.Volcano}$ 

Sig table for Volcano Analysis

# Description

Sig table for Volcano Analysis

### Usage

```
GetSigTable.Volcano(mSetObj = NA)
```

# Arguments

mSetObj

GetSOMClusterMembers 121

GetSOMClusterMembers SOM analysis

### **Description**

Get members for given cluster index, return a character string

### Usage

```
GetSOMClusterMembers(mSetObj = NA, i, j)
```

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)

i Index of Xj Index of Y

# Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetSSPTable Replace the last column of the ssp.mat with the final selection from users

### **Description**

Replace the last column of the ssp.mat with the final selection from users

# Usage

```
GetSSPTable(mSetObj = NA)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

122 GetSVMSigMat

```
GetSuggestedSAMDelta For SAM analysis
```

### **Description**

obtain a default delta with reasonable number of sig features and decent FDR

### Usage

```
GetSuggestedSAMDelta(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt GetSVMSigMat}$ 

Recursive Support Vector Machine (R-SVM) Significance Measure

# Description

Return significance measure, double[][]

### Usage

```
GetSVMSigMat(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetTopInx 123

GetTopInx Volcano indices

# Description

Get indices of top n largest/smallest number

### Usage

```
GetTopInx(vec, n, dec = T)
```

### **Arguments**

vec Vector containing volcano indices

n Numeric

dec Logical, default set to TRUE

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetTrainTestSplitMat Make random partitions

### **Description**

Make random partitions, returns matrices indicating whether the observation is in train/test for each run note: try to get a balanced sampling for each group (classification) or each quantile (regression). This is very useful for unbalanced data

### Usage

```
GetTrainTestSplitMat(y, propTraining = 2/3, nRuns = 30)
```

#### **Arguments**

y Input the data

propTraining By default set to 2/3 nRuns By default set to 30

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

124 GetTTSigMat

GetTtestRes

Retrieve T-test p-values

### **Description**

Utility method to get p values

# Usage

```
GetTtestRes(mSetObj = NA, paired = FALSE, equal.var = TRUE, nonpar = F)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

paired Default set to FALSE equal.var Default set to TRUE

nonpar Use non-parametric tests, default is set to FALSE

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

GetTTSigMat

T-test matrix

# Description

Return a double matrix with 2 columns - p values and lod

# Usage

```
GetTTSigMat(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetUnivReport 125

GetUnivReport

Utility method to perform the univariate analysis automatically

# Description

The approach is computationally expensive, and fails more often get around: make it lazy unless users request, otherwise the default t-test will also be affected

### Usage

```
GetUnivReport(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

# Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GetVariableLabel

Determine variable label for plotting

# Description

Determine data type, binned spectra, nmr peak, or ms peak

# Usage

```
GetVariableLabel(data.type)
```

### **Arguments**

data.type

Input the data type

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

126 GetXYCluster

GetVennGeneNames

Get Venn names

# Description

Get Venn names

# Usage

```
GetVennGeneNames(mSetObj = NA, areas)
```

# Arguments

mSetObj

Input name of the created mSet Object

areas

Input areas to retrieve names

 ${\tt GetXYCluster}$ 

Determine row/column number for plotting

# Description

Determine the number of rows and columns for a given total number of plots (used by Kmeans and SOM plots)

### Usage

```
GetXYCluster(total)
```

# Arguments

total

Input the total

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

GroupPeakList 127

### Description

Group peaks from the peak list based on position using the XCMS grouping algorithm (align peaks wrt, rt, and mz). For NMR peaks, need to change ppm -> mz and add dummy rt. If the data is 2-column MS, first need to add dummy rt. If the data is 3-column MS, the data can be used directly. The default mzwid for MS is 0.25 m/z, and for NMR is 0.03 ppm. The default bw is 30 for LCMS, and 5 for GCMS.

### Usage

GroupPeakList(mSetObj=NA, mzwid, bw, minfrac, minsamp, max)

### Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
mzwid,	define the width of overlapping m/z slices to use for creating peak density chromatograms and grouping peaks across samples
bw,	define the bandwidth (standard deviation or half width at half maximum) of gaussian smoothing kernel to apply to the peak density chromatogram
minfrac,	define the minimum fraction of samples necessary in at least one of the sample groups for it to be a valid group
minsamp,	define the minimum number of samples necessary in at least one of the sample groups for it to be a valid group
max,	define the maximum number of groups to identify in a single m/z slice

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

t <i>Heck</i>	t algorithm	
	O	

# Description

function to calculate tick mark based on Heckbert algorithm available in the "labeling" package implemented by Justin Talbot adapted from the imagemap package Heckbert's labeling algorithm Heckbert, P. S. (1990) Nice numbers for graph labels, Graphics Gems I, Academic Press Professional, Inc.

#### Usage

```
heckbert(dmin, dmax, m)
```

128 HMDBID2Name

#### **Arguments**

dmin Heckbert
dmax Heckbert
m Heckbert

#### Author(s)

Justin Talbot <jtalbot@stanford.edu>

HMDBID2KEGGID

Given a vector of HMDBIDs, return a vector of KEGG IDs

#### **Description**

This function, when given a vector of HMDBIDs, returns a vector of KEGG ID. HMDB standing for the Human Metabolome Database.

### Usage

HMDBID2KEGGID(ids)

#### **Arguments**

ids

Input the vector of HMDB Ids

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

HMDBID2Name

Given a vector of HMDBIDs, return a vector of HMDB compound names

# Description

This function, when given a vector of HMDBIDs, return a vector of HMDB compound names. HMDB standing for the Human Metabolome Database.

### Usage

HMDBID2Name(ids)

### **Arguments**

ids

Input the vector of HMDB Ids

ImportRawMSData 129

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

ImportRawMSData

Import raw MS data

#### **Description**

This function handles the reading in of raw MS data (.mzML, .CDF and .mzXML). Users must set their working directory to the folder containing their raw data, divided into two subfolders named their desired group labels. The function will output two chromatograms into the user's working directory, a base peak intensity chromatogram (BPIC) and a total ion chromatogram (TIC). Further, this function sets the number of cores to be used for parallel processing. It first determines the number of cores within a user's computer and then sets it that number/2.

#### Usage

```
ImportRawMSData(
  foldername,
  format = "png",
  dpi = 72,
  width = 9,
  par.cores = TRUE,
  plot = TRUE,
  plot.opts = "default"
)
```

issues.

#### **Arguments**

foldername	Character, input the file path to the folder containing the raw MS spectra to be processed.
format	Character, input the format of the image to create.
dpi	Numeric, input the dpi of the image to create.
width	Numeric, input the width of the image to create.
par.cores	Logical, if true, the function will automatically set the number of parallel cores. If false, it will not.
plot	Logical, if true the function will create BPIS and TICS plots.
plot.opts	By default, it will create BPIS and TICS plots using up to 10 samples per group.

# Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Set to "all" to create plots using all samples, though this may cause memory

# Description

This function handles the reading in of raw MS data (.mzML, .CDF and .mzXML). Users must provide a matrix with meta information about file such that each file has the name, file path, group class and extension type. The function will output two chromatograms into the user's working directory, a base peak intensity chromatogram (BPIC) and a total ion chromatogram (TIC). Further, this function sets the number of cores to be used for parallel processing. It first determines the number of cores within a user's computer and then sets it that number/2.

### Usage

```
ImportRawMSDataList(
  dataset.meta,
  format = "png",
  dpi = 72,
  width = 9,
  par.cores = TRUE,
  plot = TRUE,
  bpis_name = "BPIS_",
  tics_name = "TICS_"
)
```

### Arguments

dataset.meta	Matrix, input the meta	lata for files containing the raw	MS spectra to be pro-
	4		

cessed.

format Character, input the format of the image to create.

dpi Numeric, input the dpi of the image to create.

width Numeric, input the width of the image to create.

par. cores Logical, if true, the function will automatically set the number of parallel cores.

If false, it will not.

plot Logical, if true the function will create BPIS and TICS plots.

bpis\_name Character, input the name of the BPIS image to create.

tics\_name Character, input the name of the TICS image to create.

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Impute Var

T	
ImputeVar	Data processing: Replace missing variables
•	

#### **Description**

Replace missing variables by min/mean/median/KNN/BPCA/PPCA/svdImpute.

#### Usage

ImputeVar(mSetObj, method)

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

method Select the option to replace missing variables, either replacement based on the

minimum ("min), the mean ("mean"), or the median ("median") value of each feature columns, or several options to impute the missing values, using k-nearest neighbour ("KNN"), probabilistic PCA ("PPCA"), Bayesian PCA ("BPCA")

method, or Singular Value Decomposition ("svdImpute")

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

InitDataObjects Constructs a dataSet object for storing data	
--	--

### **Description**

This functions handles the construction of a mSetObj object for storing data for further processing and analysis. It is necessary to utilize this function to specify to MetaboAnalystR the type of data and the type of analysis you will perform.

#### Usage

InitDataObjects(data.type, anal.type, paired=FALSE)

#### **Arguments**

data.type	The type of data, either list (Compound lists), conc (Compound concentration data), specbin (Binned spectra data), pktable (Peak intensity table), nmrpeak (NMR peak lists), mspeak (MS peak lists), or msspec (MS spectra data)
anal.type	Indicate the analysis module to be performed: stat, pathora, pathqea, msetora, msetssp, msetqea, ts, cmpdmap, smpmap, or pathinteg
paired	Indicate if the data is paired or not. Logical, default set to FALSE

132 InitStatAnalMode

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

InitPowerAnal

Function for power analysis

### **Description**

Perform power analysis, requires the SSPA R package.

### Usage

InitPowerAnal(mSetObj, clsOpts)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

clsOpts For data with >2 groups, specify the two classes on which to perform power

analysis, otherwise for data with 2 groups, "NA" will automatically select the 2

groups.

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

InitStatAnalMode

Introduction for statistical analysis module report Initialize Statistical Analysis Report

# Description

Introduction for statistical analysis module report Initialize Statistical Analysis Report

#### Usage

InitStatAnalMode()

InitTimeSeriesAnal 133

InitTimeSeriesAnal

Create report of analyses (Met Pathway)

#### **Description**

Report generation using Sweave Metabolomic pathway analysis, time-series analysis

#### Usage

```
InitTimeSeriesAnal()
```

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

iPCA.Anal

Perform PCA analysis, prepare file for interactive liveGraphics3D

### **Description**

Perform PCA analysis, prepares a JSON file for interactive liveGraphics3D, as well as interactive 3D PCA score and loading plots using the plotly R package. These plots are saved in the created mSetObj; to view these, type "mSetObj\$imgSet\$time\$score3d" to view the interactive score plot, and "mSetObj\$imgSet\$time\$load3d" to view the interactive loading plot.

#### Usage

```
iPCA.Anal(mSetObj, fileNm)
```

### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

fileNm select a file name

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

134 IsSmallSmplSize

 $is {\tt EmptyMatrix}$ 

Sig table matrix is empty

# Description

Test if a sig table matrix is empty

# Usage

```
isEmptyMatrix(mat)
```

### **Arguments**

mat

Matrix to test if empty

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

IsSmallSmplSize

Check if the sample size is small

# Description

Returns whether or not the sanity check found that there were too many groups in the dataset containing too few samples. It will return a 0 if the data passes the check, or will return a 1 if the data does not.

### Usage

```
IsSmallSmplSize(mSetObj=NA)
```

# Arguments

mSetObj

Input name of the created mSet Object

IsSpectraProcessingOK 135

IsSpectraProcessingOK Check if the spectra processing is ok

# Description

Check if the spectra processing is ok

# Usage

```
IsSpectraProcessingOK(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects).

KEGGID2HMDBID

Given a vector of KEGGIDs, return a vector of HMDB ID

### **Description**

This functionn, when given a vector of KEGGIDs, returns a vector of HMDB IDs. HMDB standing for the Human Metabolome Database.

# Usage

KEGGID2HMDBID(ids)

# Arguments

ids

Vector of KEGG ids

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

KEGGID2Name

Given a vector containing KEGGIDs, returns a vector of KEGG compound names

#### **Description**

This function, given a vector containing KEGGIDs, returns a vector of KEGG compound names.

### Usage

KEGGID2Name(ids)

# Arguments

ids

Vector of KEGG ids

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

KEGGPATHID2SMPDBIDs

Given a vector containing KEGG pathway IDs, return a vector containing SMPDB IDs (only for hsa)

# Description

This function, when given a vector of KEGG pathway IDs, return a vector of SMPDB IDs (only for hsa). SMPDB standing for the Small Molecule Pathway Database, and hsa standing for human serum albumin.

### Usage

KEGGPATHID2SMPDBIDs(ids)

### **Arguments**

ids

Vector of KEGG pathway IDs

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Kmeans.Anal

Kmeans.Anal

K-means analysis

### **Description**

Perform K-means analysis

### Usage

```
Kmeans.Anal(mSetObj = NA, clust.num)
```

# Arguments

mSetObj Input name of the created mSet Object

clust.num Numeric, input the number of clusters for K-means analysis

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

kwtest

Kruskal-Wallis

# Description

Perform Kruskal-Wallis Test

### Usage

```
kwtest(x, cls)
```

# Arguments

x Input data to perform Kruskal-Wallis

cls Input class labels

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

138 LoadSmpLib

LoadKEGGKO\_lib

Utility function for PerformKOEnrichAnalysis\_KO01100

# Description

Utility function for PerformKOEnrichAnalysis\_KO01100

# Usage

```
LoadKEGGKO_lib(category)
```

# Arguments

category

Module or pathway

LoadKEGGLib

Load KEGG library

### **Description**

Load different libraries

### Usage

```
LoadKEGGLib(libType, libNm)
```

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

LoadSmpLib

Load pathway library

# Description

Load pathway library

# Usage

```
LoadSmpLib(mSetObj = NA)
```

# Arguments

mSetObj

Input name of the created mSet Object

LogNorm 139

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

LogNorm

Column-wise Normalization

### **Description**

Column-wise norm methods, when x is a column Options for log, zero mean and unit variance, and several zero mean and variance/SE

#### Usage

```
LogNorm(x, min.val)
```

### **Arguments**

x Input data

min.val Input minimum value

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada

LSD.test

Calculate Fisher's Least Significant Difference (LSD)

### **Description**

Adapted from the 'agricolae' package

### Usage

```
LSD.test(y, trt, alpha = 0.05)
```

#### **Arguments**

 $\begin{array}{ccc} y & & Input \ Y \\ trt & & Input \ trt \end{array}$ 

alpha Numeric, default is 0.05

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

make\_cpdlist

make\_cpdlib

Gets names and exact mass of all cpds (cpd.lib)

# Description

Gets names and exact mass of all cpds (cpd.lib)

### Usage

```
make_cpdlib(org)
```

make\_cpdlist

Utility function to create compound lists for permutation analysis

# Description

From a vector of m/z features, this function outputs a vector of compounds.

### Usage

```
make_cpdlist(mSetObj=NA, input_mzs)
```

### **Arguments**

mSetObj Input the name of the created mSetObj

```
Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

map 141

map

sPLS-DA Map

# Description

map variable for (s)plsda

# Usage

map(Y)

# Arguments

Υ

Input data

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

MapCmpd2KEGGNodes

Utility function for PrepareQueryJson

# Description

Utility function for PrepareQueryJson

# Usage

```
MapCmpd2KEGGNodes(cmpds, net = "ko01100")
```

# Arguments

cmpds Input the compounds
net Input the network name

Match.Pattern

MapK02KEGGEdg	29
Hapitoziteoolag	CO

Utility function for PrepareQueryJson

# Description

Utility function for PrepareQueryJson

### Usage

```
MapKO2KEGGEdges(kos, net = "ko01100")
```

# Arguments

kos Input the KOs

net Input the name of the network

Match.Pattern

Match pattern for correlation analysis

# Description

Match pattern for correlation analysis

# Usage

```
Match.Pattern(mSetObj = NA, dist.name = "pearson", pattern = NULL)
```

# Arguments

mSetObj Input the name of the created mSetObj

dist.name Input the distance method, default is set to pearson

pattern Set the pattern, default is set to NULL

MergeDatasets 143

Utility function for PrepareQueryJson

# Description

Utility function for PrepareQueryJson

# Usage

```
MergeDatasets(dataSet1, dataSet2)
```

# Arguments

dataSet1	Input the first dataset
dataSet2	Input the second dataset

 ${\tt MergeDuplicates}$ 

Merge duplicated columns or rows by their mean

# Description

```
\dim 1 \Rightarrow \text{row}, \dim 2 \Rightarrow \text{column}
```

# Usage

```
MergeDuplicates(data, dim = 2)
```

# Arguments

data Input the data

dim Numeric, input the dimensions, default is set to 2

144 MSspec.fillPeaks

MetaboliteMappingExact

Mapping from different metabolite IDs

#### **Description**

For compound names to other ids, can do exact or approximate matches For other IDs, except HMDB ID, all others may return multiple/non-unique hits Multiple hits or non-unique hits will allow users to manually select

### Usage

```
MetaboliteMappingExact(mSetObj = NA, q.type)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj.

q. type Inpute the query-type, "name" for compound names, "hmdb" for HMDB IDs,

"kegg" for KEGG IDs, "pubchem" for PubChem CIDs, "chebi" for ChEBI IDs, "metlin" for METLIN IDs, and "hmdb\_kegg" for a both KEGG and HMDB IDs.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

MSspec.fillPeaks

Function to fill in missing peaks

#### **Description**

For each sample in the processed MS spectra data, this function will fill in missing peaks using the fillPeaks function from the XCMS package. First, the function will identify any peak groups that are missing any peaks from the samples and will then fill in those peaks by rereading the raw data and integrating signals at those regions to create a new peak.

#### Usage

```
MSspec.fillPeaks(mSetObj=NA)
```

# Arguments

mSetObj

MSspec.rtCorrection 145

MSspec.rtCorrection

Retention time correction for LC/GC-MS spectra

# Description

Performs retention time correction for LC/GC-MS spectra using the XCMS package. Following retention time correction, the object dataSet will be regrouped.

### Usage

```
MSspec.rtCorrection(mSetObj=NA, bw=30)
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

bw Numeric, define the bandwidth (standard deviation or half width at half maxi-

mum) of gaussian smoothing kernel to apply to the peak density chromatogram

multi.stat

Get multiple category statistics

# Description

Get multiple category statistics

# Usage

```
multi.stat(pred, resp)
```

# Arguments

pred Input predictions
resp Input responses

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Normalization Normalization

Normalization	Normalization		
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# Description

This function performs row-wise normalization, transformation, and scaling of your metabolomic data.

# Usage

Normalization(mSetObj, rowNorm, transNorm, scaleNorm, ref=NULL, ratio=FALSE, ratioNum=20)

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
rowNorm	Select the option for row-wise normalization, "QuantileNorm" for Quantile Normalization, "ProbNormT" for Probabilistic Quotient Normalization without using a reference sample, "ProbNormF" for Probabilistic Quotient Normalization based on a reference sample, "CompNorm" for Normalization by a reference feature, "SumNorm" for Normalization to constant sum, "MedianNorm" for Normalization to sample median, and "SpecNorm" for Normalization by a sample-specific factor.
transNorm	Select option to transform the data, "LogNorm" for Log Normalization, and "CrNorm" for Cubic Root Transformation.
scaleNorm	Select option for scaling the data, "MeanCenter" for Mean Centering, "AutoNorm" for Autoscaling, "ParetoNorm" for Pareto Scaling, amd "RangeNorm" for Range Scaling.
ref	Input the name of the reference sample or the reference feature, use " " around the name.
ratio	This option is only for biomarker analysis.
ratioNum	Relevant only for biomarker analysis.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca>, Jasmine Chong McGill University, Canada

OPLSDA.Permut 147

OPLSDA.Permut

Perform OPLS-DA permutation

# Description

Orthogonal PLS-DA (from ropls) perform permutation, using training classification accuracy as indicator, for two or multi-groups

## Usage

```
OPLSDA.Permut(mSetObj = NA, num = 100)
```

### **Arguments**

mSet0bj Input name of the created mSet Object

num Input the number of permutations, default is set to 100.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

OPLSR.Anal

Perform OPLS-DA

# **Description**

Orthogonal PLS-DA (from ropls) Add reg (regression i.e. if class order matters)

## Usage

```
OPLSR.Anal(mSetObj = NA, reg = FALSE)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

reg Logical

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

148 parseTukey

parse	Fi	she	r
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Return only the signicant comparison names

# Description

Return only the signicant comparison names, used in higher function

# Usage

```
parseFisher(fisher, cut.off)
```

## **Arguments**

fisher Input fisher object cut.off Numeric, set cut-off

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

parseTukey

Return only the signicant comparison names

## **Description**

Return only the signicant comparison names, used in higher function

## Usage

```
parseTukey(tukey, cut.off)
```

# Arguments

tukey Input tukey output cut.off Input numeric cut-off

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PCA.Anal

PCA.Anal

Perform PCA analysis

# Description

Perform PCA analysis, obtain variance explained, store item to PCA object

#### Usage

```
PCA.Anal(mSetObj = NA)
```

## **Arguments**

mSetObj

Input name of the created mSet Object McGill University, Canada License: GNU GPL (>= 2)

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca>
```

PCA.Flip

Rotate PCA analysis

# Description

Rotate PCA analysis

# Usage

```
PCA.Flip(mSetObj = NA, axisOpt)
```

## **Arguments**

mSetObj

Input name of the created mSet Object

axisOpt

Input the axis option

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

150 Perform.ASCA

PCA.GENES	Obtain principal components into a matrix that has more variables than individuals

# Description

X is a matrix that has as columns the compounds that were considered as variables in the PCA analysis. First we center the matrix by columns (Xoff) and then we obtain the eigenvalues and the eigenvectors of the matrix Xoff use the equivalences between the loadings and scores to obtain the solution

#### Usage

PCA.GENES(X)

#### **Arguments**

Χ

Input matrix that has as columns the compounds that were considered as variables in the PCA analysis

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Perform.ASCA

Perform ASCA

# Description

The ASCA algorithm was adapted from the ASCA-genes method (analysis of variance (ANOVA) simultaneous component analysis) by Maria Jose Nueda (mj.nueda@ua.es) and Ana Conesa (aconesa@ivia.es)

## Usage

```
Perform.ASCA(mSetObj = NA, a = 1, b = 2, x = 2, res = 2)
```

#### **Arguments**

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
а	specify the number of components for facA
b	specify the number of components for facB
X	specify the number of components for interaction AB
res	specify the number of model residuals type is string, indicating the type of analysis "abc" separately "aab" facA joins with AB "bab" facB joins with AB

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Perform. ASCA. permute Perform ASCA model validation by permutation

#### **Description**

Perform ASCA model validation by permutation we use Manly's unrestricted permutation of observations which esentially permutes the data over all cells in the designed experiment, then calculates the score for each main factor or interaction components. This will get the null distribution for all effects in one go

#### **Usage**

Perform.ASCA.permute(mSetObj=NA, perm.num)

#### **Arguments**

mSet0bj Input name of the created mSet Object

perm. num Select the number of permutations, default is 20

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Perform.Permut Perform permutation tests only for ROC Tester

### **Description**

Perform permutation tests for the ROC Curve Based Model Creation and Evaluation module

# Usage

Perform.Permut(mSetObj=NA, perf.measure, perm.num, propTraining = 2/3)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

perf.measure Input the performance measure to rate the performance of the model, either the

area under the ROC curve ("auroc") or the predictive accuracy ("accu")

perm. num Input the number of permutations to perform

propTraining Numeric, input the fraction of samples to set aside for training. Default is set to

2/3.

152 Perform.UnivROC

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Perform.permutation Permutation

# Description

Perform permutation, options to change number of cores used

#### Usage

Perform.permutation(perm.num, fun)

#### Arguments

perm. num Numeric, input the number of permutations to perform

fun Dummy function

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Perform. UnivROC Perform Classical Univariate ROC

# **Description**

Perform Classical Univariate ROC

#### Usage

Perform.UnivROC(mSetObj=NA, feat.nm, imgName, format="png", dpi=72, isAUC, isOpt, optMethod, isPartia

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

feat.nm Input the name of the feature to perform univariate ROC analysis

imgName Input a name for the plot

format Select the image format, png, of pdf.

dpi Input the dpi. If the image format is pdf, users need not define the dpi. For png

images, the default dpi is 72. It is suggested that for high-resolution images,

select a dpi of 300.

i a ALIC	Lagical calcat T	to commute the O5 ma	maamt aamfidamaa	interval hand and "I" to
isAUC	Logical, select 1	to combute the 93 be	rcent confidence	interval band and "F" to

not

isOpt Logical, show the optimal cutoff, T to show it and F to not

optMethod Select the optimal cutoff by using either closest.topleft for closest to top-left

corner or youden for farthest to the diagonal line (Youden)

isPartial Logical, input T to calculate a partial ROC curve, and F to not

measure Select the parameter to limit the calculation of the partial ROC curve, se for

the X-axis (maximum false-positive rate) and sp for the Y-axis, representing the

minimum true positive-rate

cutoff Input the threshold to limit the calculation of the partial ROC curve, the number

must be between 0 and 1.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformAdductMapping Read Adduct List

#### **Description**

This function reads in the user's adduct list and saves it as a matrix.

#### Usage

Read.AdductData(mSetObj=NA, adductList)

## **Arguments**

mSetObj Input the name of the created mSetObj object

adductList Input the name of the adduct list

# Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

154 PerformBatchCorrection

PerformApproxMatch

Perform approximate compound matches

# Description

Given a query, perform approximate compound matching

# Usage

```
PerformApproxMatch(mSetObj = NA, q)
```

## Arguments

mSetObj Input the name of the created mSetObj.

q Input the q vector.

PerformBatchCorrection

Set up two matrixes

# Description

One is a batch containing summed concentrations of each sample the other contains the features aligned across all samples

# Usage

```
PerformBatchCorrection(mSetObj = NA, imgName)
```

# Arguments

mSetObj Input name of the created mSet Object imgName Input the name of the plot to create

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PerformCurrencyMapping

Map currency metabolites to KEGG & BioCyc

# Description

This function maps the user selected list of compounds to its corresponding KEGG IDs and BioCyc IDs

#### Usage

PerformCurrencyMapping(mSetObj = NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj object

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformCV.explore

Perform Monte-Carlo Cross Validation (MCCV)

# Description

Classification MCCV, aims to find the best feature subsets using default model parameters

#### Usage

PerformCV.explore(mSetObj, cls.method, rank.method="auroc", lvNum=2, propTraining=2/3)

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
cls.method	Select the classification method, "rf" for random forest classification, "pls" for PLS-DA, and "svm" for support vector machine
rank.method	Select the ranking method, "rf" for random forest mean decrease accuracy, "fisher" for Fisher's univariate ranking based on area under the curve "auroc" for univariate ranking based on area under the curve, "tt" for T-test univariate ranking based on area under the curve, "pls" for partial least squares, and "svm" for support vector machine
l∨Num	Input the number of latent variables to include in the analyis, only for PLS-DA classification
propTraining	Input the proportion of samples to use for training

156 PerformDetailMatch

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformCV.test

Perform MCCV for manually selected features

### **Description**

MCCV for manually selected features (no additional feature selection)

#### Usage

PerformCV.test(mSetObj, method, lvNum, propTraining=2/3, nRuns=100)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

method Select the classification method, "rf" for random forest classification, "pls" for

PLS-DA, and "svm" for support vector machine

1vNum Input the number of latent variables to include in the analyis, only for PLS-DA

classification

propTraining Input the proportion of samples to use for training, by default it is 2/3

nRuns Input the number of MCCV runs, by default it is 100

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformDetailMatch

Perform detailed name match

# **Description**

Given a query, perform compound matching.

## Usage

PerformDetailMatch(mSetObj = NA, q)

#### **Arguments**

mSet0bj Input name of the created mSet Object.

q Input the query.

PerformEachDEAnal 157

PerformEachDEAnal	Doufoung	differential	avnuaggion	analysis	an individual d	ata
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#### **Description**

This function performs DE analysis on individual data using the common matrix, which will be used/compared in later steps of the analysis (according to the p-value). The DE for each feature may be adjusted using the p-value.

#### Usage

```
PerformEachDEAnal(mSetObj = NA)
```

## **Arguments**

mSetObj Input name of the created mSet Object

#### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PerformIndNormalization

Perform normalization for individually-uploaded datasets for metaanalysis

## **Description**

This function performs normalization of individuall-uploaded datasets prior to meta-analysis.

# Usage

```
PerformIndNormalization(mSetObj = NA, dataName, norm.opt, auto.opt)
```

### **Arguments**

mSetObj	Input name of the created mSet Object
dataName	Input the name of the individual dataset for normalization.
norm.opt	Performs log2 normalization "log", or no normalization "none".
auto.opt	Performs auto-scaling of data (1), or no (0).

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PerformIntegCmpdMapping

Perform compound mapping for integrative analysis methods

# **Description**

Perform compound mapping

## Usage

PerformIntegCmpdMapping(mSetObj = NA, cmpdIDs, org, idType)

## Arguments

mSetObj Input name of the created mSet Object

cmpdIDs Input the list of compound IDs

org Input the organism code

idType Input the ID type

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformIntegGeneMapping

Perform integrated gene mapping

## **Description**

Used for the pathinteg module

# Usage

PerformIntegGeneMapping(mSetObj = NA, geneIDs, org, idType)

## **Arguments**

mSetObj Input name of the created mSet Object

geneIDs Input the list of gene IDs org Input the organism code

idType Input the ID type

PerformIntegPathwayAnalysis

Perform integrative pathway analysis

## **Description**

used for integrative analysis as well as general pathways analysis for meta-analysis results

## Usage

PerformIntegPathwayAnalysis(mSetObj, topo="dc", enrich="hyper", libOpt="integ")

# **Arguments**

mSetObj	Input name of the created mSet Object
topo	Select the mode for topology analysis: Degree Centrality ("dc") measures the number of links that connect to a node (representing either a gene or metabolite) within a pathway; Closeness Centrality ("cc") measures the overall distance from a given node to all other nodes in a pathway; Betweenness Centrality ("bc")measures the number of shortest paths from all nodes to all the others that pass through a given node within a pathway.
enrich	Method to perform over-representation analysis (ORA) based on either hypergenometrics analysis ("hyper") or Fisher's exact method ("fisher").
libOpt	Select the different modes of pathways, either the gene-metabolite mode ("integ") which allows for joint-analysis and visualization of both significant genes and metabolites or the gene-centric ("genetic") and metabolite-centric mode ("metab") which allows users to identify enriched pathways driven by significant genes or metabolites, respectively.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformKOEnrichAnalysis\_KO01100

Performs KO enrichment analysis based on the KO01100 map

## **Description**

This function performs KO enrichment analysis based on the KO01100 map and saves the .JSON file

# Usage

PerformKOEnrichAnalysis\_KO01100(mSetObj = NA, category, file.nm)

160 PerformLimmaDE

# Arguments

mSet0bj Input name of the created mSet Object

category Input the option to perform enrichment analysis, "pathway"

file.nm Input name of file to save

#### Author(s)

Othman Soufan, Jeff Xia <jeff.xia@mcgill.ca>, othman.soufan@mcgill.ca McGill University, Canada License: GNU GPL (>= 2)

PerformKOEnrichAnalysis\_List

Utility function for PerformKOEnrichAnalysis\_KO01100

## **Description**

Please note: only return hits in map KO01100

# Usage

PerformKOEnrichAnalysis\_List(file.nm)

#### **Arguments**

file.nm Input the file name

PerformLimmaDE Perform differential expression analysis using Limma for individually-

uploaded data.

# Description

This function performs DE analysis of individually-uploaded data prior to meta-analysis.

## Usage

```
PerformLimmaDE(mSetObj = NA, dataName, p.lvl = 0.1, fc.lvl = 0)
```

# Arguments

mSetObj	Input name of the created mSet	Object
---------	--------------------------------	--------

dataName Input the name of the individual dataset for normalization.

p.1vl Numeric, input the p-value (FDR) cutoff.fc.1vl Numeric, input the fold-change (FC) cutoff.

PerformMapping 161

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformMapping Utility function for PrepareQueryJson geneIDs is text one string, need

to make to vector

## **Description**

Utility function for PrepareQueryJson geneIDs is text one string, need to make to vector

# Usage

PerformMapping(inputIDs, type)

# Arguments

inputIDs Input list of IDs type Input the type of IDs

performMB Timecourse analysis

## **Description**

Adapted from the timecourse package by Yu Chuan Tai This method is only applicable for timeseries, not for general case two/multiple factor analysis

#### Usage

```
performMB(mSetObj, topPerc)
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

topPerc select the cut-off, default is 10

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

162 PerformMultiMatch

PerformMetaMerge

Meta-Analysis Method: Direct merging of datasets

# **Description**

This function is one of three methods to perform meta-analysis. Direct merging of individual data into a mega-dataset results in an analysis of that mega-dataset as if the individual data were derived from the same experiment. This method thereby ignores any inherent bias and heterogeneity between the different data. Because of this, there exists several confounders such as different experimental protocols, technical platforms, and raw data processing procedures that can mask true underlying differences. It is therefore highly suggested that this approach be used only when individual data are very similar (i.e. from the same lab, same platform, without batch effects)."

## Usage

```
PerformMetaMerge(mSetObj = NA, BHth = 0.05)
```

### Arguments

mSetObj Input name of the created mSet Object.

BHth Numeric input to set the significance level. By default it is 0.05.

# Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PerformMultiMatch

Perform multiple name matches

# Description

Given a query, performs compound name matching.

## Usage

```
PerformMultiMatch(mSetObj = NA, q)
```

### **Arguments**

mSetObj Input name of the created mSet Object.

q Input the query.

PerformPeakAnnotation 163

PerformPeakAnnotation Perform peak annotation

#### **Description**

This function performs peak annotation on the xset object created using the PerformPeakPicking function.

# Usage

PerformPeakAnnotation(xset, annParams)

## **Arguments**

xset The object created using the PerformPeakPicking function, containing the peak

picked MS data.

annParams The object created using the SetAnnotationParam function, containing user's

specified or default parameters for downstream raw MS data pre-processing.

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformPeakProfiling

Perform peak annotation This function performs feature extraction of user's raw MS data using the rawData object created using the InspectRawMSData function.

## **Description**

Perform peak annotation This function performs feature extraction of user's raw MS data using the rawData object created using the InspectRawMSData function.

## Usage

```
PerformPeakProfiling(
  rawData,
  peakParams,
  rtPlot = TRUE,
  pcaPlot = TRUE,
  labels = TRUE,
  format = "png",
  dpi = 72,
  width = 9,
```

```
rtplot_name = "RT_adjustment",
pcaplot_name = "PCA_plot"
)
```

#### Arguments

rawData The object created using the InspectRawMSData function, containing the raw

MS data.

peakParams The object created using the SetPeakParam function, containing user's specified

or default parameters for downstream raw MS data pre-processing.

rtPlot Logical, if true creates a plot of retention time correction. Defaut is set to true.

pcaPlot Logical, if true creates a PCA plot to evaluate the sample grouping. Default is

set to true.

labels Logical, if true, the PCA plot will be annotated with sample names.

format Character, input the format of the image to create.

dpi Numeric, input the dpi of the image to create.

width Numeric, input the width of the image to create.

rtplot\_name Character, input the name of the RT adjustment image to create.

pcaplot\_name Character, input the name of the PCA image to create.

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformPowerProfiling Perform power profiling

# Description

Perform power profiling of data

#### Usage

```
PerformPowerProfiling(mSetObj=NA, fdr.lvl, smplSize)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

fdr.lvl Specify the false-discovery rate level.

smplSize Specify the maximum sample size, the number must be between 60-1000.

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PerformPSEA 165

PerformPSEA	Function to perform peak set enrichment analysis	
-------------	--	--

#### **Description**

This is the main function that performs either the mummichog algorithm, GSEA, or both for peak set enrichment analysis.

#### Usage

```
PerformPSEA(mSetObj=NA, lib, libVersion, permNum = 100)
```

#### Arguments

mSetObj Input the name of the created mSetObj object.

lib Input the name of the organism library, default is hsa\_mfn.

libVersion Input the version of the KEGG pathway libraries ("current" or "old").

PermNum Numeric, input the number of permutations to perform. Default is 100.

#### Author(s)

```
Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PerformPvalCombination

Meta-Analysis Method: Combining p-values

# Description

This function is one of three methods to perform meta-analysis. Here, p-values are combined using either the Fisher's method or the Stouffer's method.

#### Usage

```
PerformPvalCombination(mSetObj = NA, method = "stouffer", BHth = 0.05)
```

## Arguments

mSetObj Input name of the created mSet Object.

method Method of p-value combination. By default it is "stouffer", else it is "fisher".

BHth Numeric input to set the significance level. By default it is 0.05.

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

166 Plot.Permutation

PerformVoteCounting Meta-Analysis Method: Vote Counting

## **Description**

This function is one of three methods to perform meta-analysis. Here, significant features are selected based on a selected criteria (i.e. an adjusted p-value <0.05 and the same direction of FC) for each dataset. The votes are then calculated for each feature by counting the total of number of times a feature is significant across all included datasets. However, this method is statistically inefficient and should be considered the last resort in situations where other methods to perform meta-analysis cannot be applied.

# Usage

PerformVoteCounting(mSetObj = NA, BHth = 0.05, minVote)

# **Arguments**

mSet0bj Input name of the created mSet Object.

BHth Numeric input to set the significance level. By default it is 0.05.

minVote Numeric input to set the minimum vote-count.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Plot.Permutation Plot results of permutation tests

#### **Description**

Plot results of permutation tests

# Usage

Plot.Permutation(mSetObj=NA, imgName, format="png", dpi=72)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format elect the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

PlotAccuracy 167

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotAccuracy Plot classification performance using different features for Multi-Biomarker

# **Description**

Plot of the accuracy of classification with an increasing number of features.

## Usage

PlotAccuracy(mSetObj=NA, imgName, format="png", dpi=72)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotANOVA Plot ANOVA

# Description

Plot ANOVA

## Usage

PlotANOVA(mSetObj=NA, imgName, format="png", dpi=72, width=NA)

168 PlotANOVA2

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotANOVA2 Plot Venn diagram of ANOVA results

### Description

Plot Venn diagram of ANOVA results

## Usage

PlotANOVA2(mSetObj, imgName, format="png", dpi=72, width=NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia < jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotASCA.Permutation 169

PlotASCA.Permutation Plot ASCA permutation

#### **Description**

Plot plsda classification performance using different components

#### Usage

PlotASCA.Permutation(mSetObj=NA, imgName, format="png", dpi=72, width=NA)

## **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotAscaImpVar Plot the important variables for each factor

## **Description**

Plot the important variables for each factor

#### Usage

PlotAscaImpVar(mSetObj=NA, imgName, format, dpi, width=NA, type)

170 PlotASCAModel

### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

type select model a, b, or ab

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotASCAModel Plot score plots of each ASCA model for component 1 against time

### **Description**

Plot score plots of each ASCA model for component 1 against time

# Usage

PlotASCAModel(mSetObj=NA, imgName, format="png", dpi=72, width=NA, type, colorBW=FALSE)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the ASCA score plot format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

type select model a or b

colorBW Logical, use black/white coloring (T) or not (F)

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotBoxPlot 171

PlotBoxPlot

Plot a boxplot view of a selected compound

## **Description**

Plots a boxplot of the selected compound's concentration between the groups.

# Usage

```
PlotBoxPlot(
   mSetObj,
   feat.nm,
   imgName,
   format = "png",
   dpi = 72,
   isOpt,
   isQuery
)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

feat.nm Input the name of the selected compound.

imgName Input a name for the plot

format Select the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotCmpdSummary

Plot compound summary change to use dataSet\$proc instead of dataSet\$orig in case of too many NAs

### Description

Plot compound summary change to use dataSet\$proc instead of dataSet\$orig in case of too many NAs

#### Usage

```
PlotCmpdSummary(mSetObj = NA, cmpdNm, format = "png", dpi = 72, width = NA)
```

172 PlotCmpdView

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

cmpdNm Input the name of the compound to plot format Input the format of the image to create dpi Input the dpi of the image to create width Input the width of the image to create

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotCmpdView Plot Compound View

# Description

Plots a bar-graph of selected compound over groups

#### Usage

PlotCmpdView(mSetObj=NA, cmpdNm, format="png", dpi=72, width=NA)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

cmpdNm Input a name for the compound

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotConcRange 173

PlotConcRange	Plot the compound concentration data compared to the reference concentration range
	O .

## **Description**

Plot the compound concentration data compared to the reference concentration range

## Usage

```
PlotConcRange(mSetObj, nm, format="png", dpi=72, width=NA)
```

# **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

nm of the input compound

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotCorr	Pattern hunter, correlation plot

## **Description**

Plot correlation

## Usage

```
PlotCorr(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

174 PlotCorrHeatMap

## **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotCorrHeatMap

Pattern hunter, corr heatmap

## **Description**

Plot correlation heatmap

### Usage

```
PlotCorrHeatMap(
    mSetObj = NA,
    imgName,
    format = "png",
    dpi = 72,
    width = NA,
    target,
    cor.method,
    colors,
    viewOpt,
    fix.col,
    no.clst,
    top,
    topNum
)
```

# Arguments

mSetObj Input name of the created mSet Object.

imgName Input the name of the image to create
format Select the image format, "png", or "pdf".

PlotDetailROC 175

dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
target	Input "row" to select features, or "col" to select samples.
cor.method	Indicate the correlation method, 'pearson', 'spearman', or 'kendall'.
colors	Indicate the colors for the heatmap, "bwm" for default, "gbr" for red/green, "heat" for heat colors, "topo" for topo colors, and "gray" for gray scale.
viewOpt	Indicate "overview" to get an overview of the heatmap, and "detail" to get a detailed view of the heatmap.
fix.col	Logical, fix colors (TRUE) or not (FALSE).
no.clst	Logical, indicate if the correlations should be clustered (TRUE) or not (FALSE).
top	View top
topNum	Numeric, view top McGill University, Canada License: GNU GPL (>= 2)

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca>

# Description

Plot detailed ROC

# Usage

PlotDetailROC(mSetObj = NA, imgName, thresh, sp, se, dpi = 72, format = "png")

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
imgName	Input a name for the plot
thresh	Input the threshold
sp	Specificity
se	Sensitivity
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
format	Select the image format, "png", or "pdf".

176 PlotEIC

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotEBAM.Cmpd

Plot EBAM

# Description

Plot EBAM

### Usage

PlotEBAM.Cmpd(mSetObj=NA, imgName, format, dpi, width)

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotEIC Plot EIC

#### **Description**

This function creates an extracted ion chromatogram (EIC) for a specific m/z and retention time. This is used for quality-control of raw m/s data.

PlotEnrichNet.Overview 177

# Usage

```
PlotEIC(
   raw_data,
   rt_mn,
   rt_mx,
   mz_mn,
   mz_mx,
   aggreg = "sum",
   format = "png",
   dpi = 72,
   width = 9
)
```

# Arguments

MS data.	aw
rt_mn Numeric, specify the minimum bound of the retention time range.	
rt_mx Numeric, specify the maximum bound of the retention time range.	
mz_mn Numeric, specify the minimum bound of the m/z range.	
mz_mx Numeric, specify the maximum bound of the m/z range.	
aggreg Character, if "sum", creates a total ion chromatogram. If "max", creates a bapeak chromatogram. By default it is set to "sum".	ise
format Character, input the format of the image to create.	
dpi Numeric, input the dpi of the image to create.	
width Numeric, input the width of the image to create.	

PlotEnrichNet.Overview

Barplot height is enrichment fold change

# Description

Used in higher functions, the color is based on p values

## Usage

```
PlotEnrichNet.Overview(folds, pvals, layoutOpt = layout.fruchterman.reingold)
```

# Arguments

folds	Input fold-change for bar plot
pvals	Input p-values for bar plot

layoutOpt Input the layout option, default is set to layout.fruchterman.reingold

178 PlotHCTree

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotFC

Plot fold change

## **Description**

Plot fold change analysis

#### Usage

PlotFC(mSetObj=NA, imgName, format="png", dpi=72, width=NA)

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotHCTree

Plot Dendrogram

## **Description**

Dendogram

PlotHeatMap 179

#### Usage

```
PlotHCTree(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   smplDist,
   clstDist
)
```

#### **Arguments**

mSet0bj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

smplDist Method to calculate sample distance

clstDist Method to calculate clustering distance

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotHeatMap

Create Heat Map Plot

#### **Description**

Plot a heatmap based on results from t-tests/ANOVA, VIP or randomforest

# Usage

```
PlotHeatMap(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   dataOpt,
```

180 PlotHeatMap

```
scaleOpt,
smplDist,
clstDist,
palette,
viewOpt = "detail",
rowV = T,
colV = T,
var.inx = NA,
border = T,
grp.ave = F
)
```

## Arguments

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

dataOpt Set data options

scaleOpt Set the image scale

smplDist Input the sample distance method

clstDist Input the clustering distance method

palette Input color palette choice

viewOpt Set heatmap options, default is set to "detail"

rowV Default is set to T

colV Default is set to T

var.inx Default is set to NA

border Indicate whether or not to show cell-borders, default is set to T

grp.ave Logical, default is set to F

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotHeatMap2 181

PlotHeatMap2	Plot heatmap visualization for time-series data

# Description

Plot heatmap visualization for time-series data

# Usage

PlotHeatMap2(mSetObj=NA, imgName, format="png", dpi=72, width=NA, smplDist="pearson", clstDist="average"

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
smplDist	Select distance measure: euclidean, pearson, or minkowski
clstDist	Select clustering algorithm: ward, average, complete, single
colors	Select heatmap colors: bwm, gray
viewOpt	Select overview or detailed view: overview or detail
hiRes	Select high-resolution or not: logical, default set to F
sortInx	Sort by index
useSigFeature	Use significant features only: F or T (default false)
drawBorder	Show cell borders: F or T (default F)

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

182 PlotImpVars

PlotImpVar	Plot PLS important variables,

## **Description**

Plot PLS important variables, BHan: added bgcolor parameter for B/W color

# Usage

```
PlotImpVar(mSetObj = NA, imp.vec, xlbl, feat.num = 15, color.BW = FALSE)
```

## **Arguments**

mSetObj Input name of the created mSet Object imp.vec Input the vector of important variables

xlbl Input the x-label

feat.num Numeric, set the feature numbers, default is set to 15

color.BW Use black-white for plot (T) or colors (F)

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotImpVars	Plot selected compounds by their percentage frequency

#### **Description**

Plot the important variables of single biomarker model ranked by order of importance

## Usage

```
PlotImpVars(mSetObj=NA, imgName, format="png", dpi=72, mdl.inx, measure = "freq", feat.num = 15)
```

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format elect the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

mdl.inx Model index, -1 selects the model with the best AUC, input 1-6 to view the

important features of one of the top six models

PlotInmexGraph 183

measure Choose to rank features by the frequency of being selected "freq", or the mean

importance measure "mean"

feat . num Input the number of features to include in the plot, by default it is 15.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotInmexGraph Plot an igraph object and return the node information (position and labels)

## **Description**

Plot an igraph object and return the node information (position and labels) Used in a higher function

## Usage

```
PlotInmexGraph(
   mSetObj,
   pathName,
   g,
   width = NA,
   height = NA,
   bg.color = NULL,
   line.color = NULL,
   format = "png",
   dpi = NULL
)
```

## Arguments

mSetObj Input name of the created mSet Object

g Input the graph

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

height Input the height of the graph to create

bg.color Set the background color, default is set to NULL

line.color Set the line color, default is set to NULL

path.id Input the pathway id

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

184 PlotIntegPaths

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РΙ	OT. I	nmex	(Path	1

Plot integrated methods pathway analysis

## **Description**

Only update the background info for matched node

## Usage

```
PlotInmexPath(mSetObj=NA, path.id, width, height)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

height Input the height of the image to create.

path.id Input the ID of the pathway to plot.

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotIntegPaths

**PlotIntegPaths** 

## **Description**

Plots both the original mummichog and the GSEA results by combining p-values using the Fisher's method (sumlog).

## Usage

```
PlotIntegPaths(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = 9,
   labels = "default",
   labels.x = 5,
   labels.y = 5
)
```

PlotInteraction 185

#### **Arguments**

mSetObj Input the name of the created mSetObj object

imgName Input a name for the plot

format Character, input the format of the image to create.

dpi Numeric, input the dpi of the image to create.

width Numeric, input the width of the image to create.

labels.x Numeric, indicate the number of top-ranked pathways using the fGSEA algo-

rithm to annotate on the plot.

labels.y Numeric, indicate the number of top-ranked pathways using the original mum-

michog algorithm to annotate on the plot.

Labels Character, indicate if the plot should be labeled. By default it is set to "default",

and the 5 top-ranked pathways per each algorithm will be plotted. Users can adjust the number of pathways to be annotated per pathway using the "labels.x" and "labels.y" parameters. Users can set this to "none" for no annotations, or

"all" to annotate all pathways.

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotInteraction	Plot ASCA interaction plots	

## Description

Plot ASCA interaction plots

## Usage

PlotInteraction(mSetObj=NA, imgName, format="png", dpi=72, colorBW=FALSE, width=NA)

# Arguments

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

colorBW Logical, use black and white (TRUE) or colors (FALSE)

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

186 PlotKmeans

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotKEGGPath

Plot metabolome pathway

## **Description**

Orthogonal PLS-DA (from ropls)

# Usage

```
PlotKEGGPath(
  mSetObj = NA,
  pathName,
  width = NA,
  height = NA,
  format = "png",
  dpi = NULL
)
```

## **Arguments**

mSetObj Input name of the created mSet Object pathName Input the name of the selected pathway

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

height Input the height of the created plot.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotKmeans

Plot K-means analysis

## **Description**

Plot K-means analysis

## Usage

```
PlotKmeans(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

PlotLoadingCmpd 187

#### **Arguments**

mSet0bj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotLoadingCmpd Plot loading compounds

## **Description**

Plot loading compounds

## Usage

PlotLoadingCmpd(mSetObj = NA, cmpdNm, format = "png", dpi = 72, width = NA)

# **Arguments**

mSetObj Input name of the created mSet Object cmpdNm Input the name of the selected compound format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

188 PlotMetaVenn

PlotMBTimeProfile

Plot MB Time Profile

#### Description

Plot MB Time Profile

#### **Usage**

```
PlotMBTimeProfile(mSetObj = NA, cmpdNm, format = "png", dpi = 72, width = NA)
```

## Arguments

mSetObj Input name of the created mSet Object cmpdNm Input the name of the compound format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia < jeff . xia@mcgill . ca> McGill University, Canada License: GNU GPL (>= 2)

PlotMetaVenn

Meta-Analysis: Plot Venn Diagram

#### **Description**

This function plots a venn diagram of the individual studies.

#### Usage

```
PlotMetaVenn(mSetObj = NA, imgNM = NA)
```

#### **Arguments**

mSetObj Input name of the created mSet Object.
imgNM Input the name of the created Venn Diagram

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotMetpaPath 189

PlotMetpaPath

Plot KEGG pathway

## **Description**

Plot KEGG pathway

#### Usage

```
PlotMetpaPath(
   mSetObj = NA,
   pathName,
   width = NA,
   height = NA,
   format = "png",
   dpi = NULL
)
```

# Arguments

mSetObj Input name of the created mSet Object

pathName Input the name of the selected KEGG pathway

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt PlotModelScree}$ 

Plot scree plots for each model in ASCA

#### **Description**

Plot scree plots for each model in ASCA

#### Usage

```
PlotModelScree(mSetObj, imgName, format="png", dpi=72, width=NA)
```

190 PlotMS.RT

## **Arguments**

mSetObj Input name of the created mSet Object.

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

# Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotMS.RT Plot rentention time corrected spectra

## **Description**

Plot the retention time corrected spectra

## Usage

```
PlotMS.RT(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

# **Arguments**

imgName Input name of the created mSet Object
Input the name for the created plot
format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

PlotMS2Spectra 191

PlotMS2Spectra

Plot selected M2 spectra for an m/z feature

## **Description**

This function creates a plot of the user selected precursor m/z.

## Usage

```
PlotMS2Spectra(ms2, spectra = 1)
```

## **Arguments**

ms2

Spectrum2 class object containing all of the spectra for the selected m/z feature.

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotMSEA.Overview

Plot MSEA overview

#### **Description**

Barplot height is enrichment fold change color is based on p values, used in higher functions

## Usage

```
PlotMSEA.Overview(folds, pvals)
```

#### Arguments

folds Input the fold-change values

pvals Input the p-values

## Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

192 PlotNormSummary

PlotMSPeaksPerm

Plot MS Peaks to Paths mummichog permutation p-values

# Description

Plots the mummichog permutation p-values

#### Usage

```
PlotMSPeaksPerm(
  mSetObj = NA,
  pathway,
  imgName,
  format = "png",
  dpi = 72,
  width = NA
)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

pathway Input the name of the pathway

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotNormSummary

Two plot summary plot: Feature View of before and after normaliza-

tion

## Description

For each plot, the top is a box plot, bottom is a density plot

PlotOPLS.MDL 193

## Usage

PlotNormSummary(mSetObj, imgName, format, dpi, width)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia < jeff.xia@mcgill.ca>, Jasmine Chong McGill University, Canada

PlotOPLS.MDL Plot OF	PLS
----------------------	-----

## **Description**

Plot OPLS

# Usage

```
PlotOPLS.MDL(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

## Arguments

mSetObi	Input name	of the	created	mSet	Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

194 PlotOPLS.Splot

```
PlotOPLS.Permutation Plot OPLS-DA permutation
```

## **Description**

Orthogonal PLS-DA (from ropls) perform permutation, using training classification accuracy as indicator, for two or multi-groups

# Usage

```
PlotOPLS.Permutation(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA
)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia < jeff . xia@mcgill . ca> McGill University, Canada License: GNU GPL (>= 2)

PlotOPLS.Splot S-plot for OPLS-DA

#### Description

Orthogonal PLS-DA (from ropls) S-plot for important features from OPLS-DA

PlotOPLS2DScore 195

## Usage

```
PlotOPLS.Splot(
   mSetObj = NA,
   imgName,
   plotType = "all",
   format = "png",
   dpi = 72,
   width = NA
)
```

## Arguments

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotOPLS2DScore

Create OPLS-DA score plot

#### **Description**

Orthogonal PLS-DA (from ropls) score plot

## Usage

```
PlotOPLS2DScore(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   inx1,
   inx2,
   reg = 0.95,
   show = 1,
   grey.scale = 0
)
```

196 PlotORA

#### **Arguments**

imgName Input name of the created mSet Object
imgName Input a name for the plot

Select the image format, "png", or "pdf".

Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

inx1 Numeric, indicate the number of the principal component for the x-axis of the

loading plot.

inx2 Numeric, indicate the number of the principal component for the y-axis of the

loading plot.

reg Numeric

show Show variable labels, 1 or O

grey.scale Numeric, indicate grey-scale, 0 for no, and 1 for yes

#### Author(s)

Jeff Xia < jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotORA Plot over-representation analysis (ORA)

#### **Description**

Plot over-representation analysis (ORA)

#### Usage

PlotORA(mSetObj=NA, imgName, imgOpt, format="png", dpi=72, width=NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

imgOpt "net"

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

PlotPathSummary 197

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPathSummary

Plot a scatterplot (circle) overview of the matched pathways

# Description

x axis is the pathway impact factor y axis is the p value (from ORA or globaltest) return the circle information

# Usage

```
PlotPathSummary(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   x,
   y
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5.
x	Input the X
У	Input the Y

## Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

198 PlotPCA.overview

PlotPathwayMZHits	Plot m/z hits in a pathway
-------------------	----------------------------

## **Description**

Function to create a boxplot of m/z features within a specific pathway. m/z features used by the original mummichog algorithm are highlighted with an asterisk.

# Usage

```
PlotPathwayMZHits(mSetObj = NA, msetNM, format = "png", dpi = 300, width = 10)
```

# **Arguments**

mSetObj	Input the name of the created mSetObj object.
msetNM	Character, input the name of the pathway.
format	Character, input the format of the image to create.
dpi	Numeric, input the dpi of the image to create. Default is set to 300.
width	Numeric, input the width of the image to create. Default is set to 10.

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCA.overview Scatter plot colored by different batches

# Description

Scatter plot colored by different batches

## Usage

```
PlotPCA.overview(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.

PlotPCA2DScore 199

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCA2DScore	Create 2D PCA score plot	
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# Description

Rotate PCA analysis

# Usage

PlotPCA2DScore(mSetObj=NA, imgName, format="png", dpi=72, width=NA, pcx, pcy, reg = 0.95, show=1, green to the state of th

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
рсх	Specify the principal component on the x-axis
рсу	Specify the principal component on the y-axis
reg	Numeric, input a number between 0 and 1, 0.95 will display the 95 percent confidence regions, and 0 will not.
show	Display sample names, $1 =$ show names, $0 =$ do not show names.
grey.scale	Use grey-scale colors, 1 = grey-scale, 0 = not grey-scale.

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCA3DScoreImg

_		
Ρ	otPCA3DScor	Θ.

Create 3D PCA score plot

# Description

Rotate PCA analysis

## Usage

```
PlotPCA3DScore(mSetObj=NA, imgName, format="json", inx1, inx2, inx3)
```

## **Arguments**

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCA3DScoreImg	Create 3D PCA score plot
-------------------	--------------------------

# Description

This function creates both a static 3D PCA score plot as well as an interactive 3D PCA score plot using the plotly R package. The 3D PCA score plot is stored in the mSetObj (mSetObj\$imgSet\$pca.3d). To view the plot, if your mSetObj is named mSet, type "mSet\$imgSet\$pca.3d" inro your R console, and the 3D plot will appear.

## Usage

```
PlotPCA3DScoreImg(mSetObj=NA, imgName, format="png", dpi=72, width=NA, inx1, inx2, inx3, ang1)

mSet <- PlotPCA3DScore(mSetObj=NA, imgName, format="json", dpi=72, width=NA, inx1, inx2, inx3, ang1)
```

PlotPCABiplot 201

# Arguments

mSetObj	Input name of the created mSet Object.
imgName	Input a name for the plot.
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.
angl	Input the angle

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCABiplot	Create PCA Biplot, set $xpd = T$ to plot outside margin	
·	1 / 1 1	

# Description

Rotate PCA analysis

# Usage

PlotPCABiplot(mSetObj=NA, imgName, format="png", dpi=72, width=NA, inx1, inx2)

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width

202 PlotPCALoading

inx1 Numeric, indicate the number of the principal component for the x-axis of the

loading plot.

inx2 Numeric, indicate the number of the principal component for the y-axis of the

loading plot.

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCALoading Plot PCA loadings and also set up the matrix for display

# Description

Rotate PCA analysis

## Usage

PlotPCALoading(mSetObj=NA, imgName, format="png", dpi=72, width=NA, inx1, inx2, plotType, lbl.feat=1)

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

loading plot.

PlotPCAPairSummary 203

D1 (D0(D ) 0	DI - DG1 10 10 10 10 10
PlotPCAPairSummary	Plot PCA pair summary, format image in png, tiff, pdf, ps, svg

# Description

Rotate PCA analysis

# Usage

PlotPCAPairSummary(mSetObj=NA, imgName, format="png", dpi=72, width=NA, pc.num)

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.
pc.num	Numeric, input a number to indicate the number of principal components to display in the pairwise score plot.

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPCAScree	Plot PCA scree plot

# Description

Rotate PCA analysis

# Usage

PlotPCAScree(mSetObj=NA, imgName, format="png", dpi=72, width=NA, scree.num)

204 PlotPeaks2Paths

## **Arguments**

mSet0bj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

scree.num Numeric, input a number to indicate the number of principal components to

display in the scree plot.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPeaks2Paths

PlotPeaks2Paths

## **Description**

Plots either the original mummichog or GSEA results.

## Usage

```
PlotPeaks2Paths(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = 9,
   labels = "default",
   num_annot = 5
)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj object

imgName Input a name for the plot

format Character, input the format of the image to create.

dpi Numeric, input the dpi of the image to create.

width Numeric, input the width of the image to create.

Labels Character, indicate if the plot should be labeled. By default it is set to "default",

and the 5 top-ranked pathways per each algorithm will be plotted. Users can adjust the number of pathways to be annotated per pathway using the "num\_annot"

parameter.

PlotPLS.Classification 205

#### Author(s)

```
Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotPLS.Classification

Plot PLS-DA classification performance using different components

## **Description**

Plot plsda classification performance using different components

# Usage

```
PlotPLS.Classification(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA
)
```

## **Arguments**

mSetObj	Input name of the	created mSet Object
---------	-------------------	---------------------

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

206 PlotPLS.Imp

PlotPLS.Imp

Plot PLS important features

## **Description**

Plot PLS important features, BHan: added bgcolor parameter for B/W color

Input name of the created mSet Object

# Usage

```
PlotPLS.Imp(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   type,
   feat.nm,
   feat.num,
   color.BW = FALSE
)
```

## Arguments

mSetObj

imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.
type	Indicate the type variables of importance to use, "vip" to use VIp scores, or "type" for coefficients
feat.nm	Feature name

# Author(s)

feat.num

color.BW

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Logical, true to use black and white, or false to not

Feature numbers

PlotPLS.Permutation 207

PlotPLS.Permutation

Plot PLS-DA classification performance using different components, permutation

## **Description**

Plot plsda classification performance using different components

# Usage

```
PlotPLS.Permutation(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA
)
```

## Arguments

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPLS2DScore

Plot PLS score plot

#### **Description**

Plot PLS score plot

208 PlotPLS2DScore

# Usage

```
PlotPLS2DScore(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   inx1,
   inx2,
   reg = 0.95,
   show = 1,
   grey.scale = 0,
   use.sparse = FALSE
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
reg	Numeric, default is 0.95
show	Show labels, 1 or 0
grey.scale	Numeric, use a grey scale (0) or not (1)
use.sparse	Logical, use a sparse algorithm (T) or not (F)

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPLS3DScore 209

PlotPLS3DScore	Plot PLS 3D score plot	
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# Description

Plot PLS 3D score plot

## Usage

```
PlotPLS3DScore(mSetObj = NA, imgName, format = "json", inx1, inx2, inx3)
```

## **Arguments**

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotPLS3DScoreImg	Plot PLS 3D score plot	

# Description

This function creates two 3D PLS-DA score plots, the first is static for Analysis Report purposes, as well as an interactive 3D plot using the plotly R package. The 3D score plot is saved in the created mSetObj (mSetObj\$imgSet\$plsda.3d). To view the score plot, if the name of your mSetObj is mSet, enter "mSet\$imgSet\$plsda.3d" to view the interactive score plot.

210 PlotPLSLoading

# Usage

```
PlotPLS3DScoreImg(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   inx1,
   inx2,
   inx3,
   ang1
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.
angl	Input the angle

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt PlotPLS loading \ plot, \ also \ set \ the \ loading \ matrix \ for \ display}$ 

# Description

Plot PLS loading plot, also set the loading matrix for display

PlotPLSPairSummary 211

#### Usage

```
PlotPLSLoading(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
  width = NA,
  inx1,
  inx2
)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

inx1 Numeric, indicate the number of the principal component for the x-axis of the

loading plot.

inx2 Numeric, indicate the number of the principal component for the y-axis of the

loading plot.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPLSPairSummary

Plot PLS pairwise summary

#### Description

Plot PLS pairwise summary

# Usage

```
PlotPLSPairSummary(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
```

212 PlotPowerProfile

```
width = NA,
pc.num
)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

pc.num Numeric, indicate the number of principal components

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPowerProfile Plot power profile

## **Description**

Plot power profile, specifying FDR level and sample size. It will return the image as well as the predicted power at various sample sizes.

## Usage

PlotPowerProfile(mSetObj=NA, fdr.lvl, smplSize, imgName, format, dpi, width)

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
fdr.lvl	Specify the false-discovery rate level.
smplSize	Specify the maximum sample size, the number must be between 60-1000.
imgName	Specify the name to save the image as.
format	Specify the format of the image to save it as, either "png" or "pdf".
dpi	Specify the dots-per-inch (dpi). By default it is 72, for publications the recommended dpi is 300.
width	Specify the width of the image. NA specifies a width of 9, 0 specifies a width of 7, otherwise input a chosen width.

PlotPowerStat 213

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotPowerStat Plot power statistics

## **Description**

Create plot for power statistics

#### Usage

PlotPowerStat(mSetObj, imgName, format="png", dpi=72, width=NA)

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
imgName	Specify the name to save the image as.
format	Specify the format of the image to save it as, either "png" or "pdf"
dpi	Specify the dots-per-inch (dpi). By default it is 72, for publications the recommended dpi is 300.
width	Specify the width of the image. NA or 0 specifies a width of 10, otherwise input a chosen width.

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotProbView	Plot a summary view of the classification result	

#### **Description**

Plot of predicted class probabilities. On the x-axis is the proability, and the y-axis is the index of each predicted sample based on the probility. The samples are turned into separations at the x-axis. This plot can be created for multivariate ROC curve analysis using SVM, PLS, and RandomForest. Please note that sometimes, not all samples will be tested, instead they will be plotted at the 0.5 neutral line.

## Usage

PlotProbView(mSetObj=NA, imgName, format="png", dpi=72, mdl.inx, show, showPred)

214 plotProfile

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

mdl.inx Model index, 0 means to compare all models, -1 means to use the best model,

input 1-6 to plot a ROC curve for one of the top six models

show 1 or 0, if 1, label samples classified to the wrong groups

showPred Show predicted samples

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

# Description

Colored by experimental conditions, used in higher function

#### Usage

```
plotProfile(mSetObj = NA, varName)
```

# Arguments

mSetObj Input name of the created mSet Object

varName Input the name of the variable

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotQEA.MetSet 215

PlotQEA.MetSet	View individual compounds related to a given metabolite set	

# Description

View individual compounds related to a given metabolite set Functions for various plots for enrichment analysis

# Usage

```
PlotQEA.MetSet(mSetObj=NA, setNM, format="png", dpi=72, width=NA)
```

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)	
setNM	Input the name of the metabolite set	
format	Select the image format, "png", or "pdf".	
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.	
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.	

# Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotQEA.Overview	Plot QEA overview	

# Description

Plot QEA overview

## Usage

```
PlotQEA.Overview(mSetObj=NA, imgName, imgOpt, format="png", dpi=72, width=NA)
```

216 PlotRF.Classify

## Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

imgOpt "net"

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotRF.Classify

Plot Random Forest

## **Description**

Random Forest plot

#### Usage

PlotRF.Classify(mSetObj, imgName, format, dpi, width)

#### Arguments

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotRF.Outlier 217

_				
P1	$\wedge$ t	RF	.Outi	lior

Plot Random Forest outliers

### **Description**

Random Forest plot of outliers

### Usage

```
PlotRF.Outlier(mSetObj=NA, imgName, format="png", dpi=72, width=NA)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotRF.VIP

Plot Random Forest variable importance

### **Description**

Random Forest plot of variable importance ranked by MeanDecreaseAccuracy

#### Usage

```
PlotRF.VIP(mSetObj=NA, imgName, format, dpi, width)
```

218 PlotROC

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotROC Plot ROC

### **Description**

Pred and auroc are lists containing predictions and labels from different cross-validations

#### Usage

PlotROC(mSetObj=NA, imgName, format="png", dpi=72, mdl.inx, avg.method, show.conf, show.holdout, focu

## Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
imgName	Input a name for the plot
format	Select the image format, "png", of "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
mdl.inx	Model index, 0 means to compare all models, input 1-6 to plot a ROC curve for one of the top six models
avg.method	Input the method to compute the average ROC curve, either "threshold", "vertical" or "horizontal"
show.conf	Logical, if 1, show confidence interval, if 0 do not show
show.holdout	Logical, if 1, show the ROC curve for hold-out validation, if 0 do not show
focus	"fpr"
cutoff	Input the threshold to limit the calculation of the ROC curve, the number must be between 0 and 1.

PlotROC.LRmodel 219

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotROC.LRmodel

Plot ROC for the logistic regression model

# Description

Plot ROC for the logistic regression model

## Usage

```
PlotROC.LRmodel(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   show.conf = FALSE,
   sp.bin = 0.01
)
```

### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

show.conf Logical, show confidence intervals

sp.bin Numeric, default is set to 0.01.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

220 PlotROCTest

PlotROCTest	Plot ROC for the ROC Curve Based Model Creation and Evaluation module

# Description

Plot the ROC curve of the biomarker model created using a user-selected subset of features. Pred and auroc are lists containing predictions and labels from different cross-validations.

## Usage

PlotROCTest(mSetObj=NA, imgName, format="png", dpi=72, mdl.inx, avg.method, show.conf, show.holdout,

## Arguments

Input the name of the created mSetObj (see InitDataObjects)
Input a name for the plot
Select the image format, "png", of "pdf".
Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
Model index, $0$ means to compare all models, input 1-6 to plot a ROC curve for one of the top six models
Input the method to compute the average ROC curve, either "threshold", "vertical" or "horizontal"
Logical, if 1, show confidence interval, if 0 do not show
Logical, if 1, show the ROC curve for hold-out validation, if 0 do not show
"fpr"
Input the threshold to limit the calculation of the ROC curve, the number must be between 0 and 1.

# Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotRSVM.Classification 221

PlotRSVM.Classification

Recursive Support Vector Machine (R-SVM) plot

## Description

Plot recursive SVM classification

#### Usage

PlotRSVM.Classification(mSetObj, imgName, format, dpi, width)

### Arguments

mSet0bj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotRSVM.Cmpd Recursive Support Vector Machine (R-SVM) plot of important variables

**Description** 

Plot recursive SVM variables of importance if too many, plot top 15

#### Usage

PlotRSVM.Cmpd(mSetObj=NA, imgName, format="png", dpi=72, width=NA)

222 PlotSAM.Cmpd

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSAM.Cmpd Plot SAM

#### **Description**

Plot SAM with positive and negative metabolite sets

## Usage

```
PlotSAM.Cmpd(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSAM.FDR 223

PlotSAM.	FDR

Plot SAM Delta Plot

## Description

Plot SAM Delta Plot (FDR)

## Usage

```
PlotSAM.FDR(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

## Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is $10.5$ . The second default is width = 0, where the width is $7.2$ . Otherwise users can input their own width.
delta	Input the delta

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotSampleNormSummary Two plot summary plot: Sample View of before and after normalization

## Description

For each plot, the top is a density plot and the bottom is a box plot.

### Usage

```
PlotSampleNormSummary(mSetObj=NA, imgName, format="png", dpi=72, width=NA)
```

224 PlotSelectedFeature

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca>, Jasmine Chong McGill University, Canada

datasets

### **Description**

This function plots a box-plot of the expression pattern of a user-selected feature across the different datasets included in meta-analysis.

## Usage

PlotSelectedFeature(mSetObj = NA, gene.id)

#### **Arguments**

mSetObj Input name of the created mSet Object.
gene.id Input the name of the selected feature.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSigVar 225

PlotSigVar	Supporting function for plotting important variables for each factor

### **Description**

Supporting function for plotting important variables for each factor note, by control xpd to plot legend outside the plotting area without using layout

## Usage

```
PlotSigVar(x, y, xline, yline, title)
```

### **Arguments**

X	Input the X variable
У	Input the Y variable
xline	Input the xline
yline	Input the yline
title	Input the title

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

lot	SOM Plot	PlotSOM
-----	----------	---------

# Description

Plot SOM map for less than 20 clusters

## Usage

```
PlotSOM(mSetObj = NA, imgName, format = "png", dpi = 72, width = NA)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.

226 PlotSPLS2DScore

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSPLS2DScore

Score Plot SPLS-DA

# Description

Sparse PLS-DA (from mixOmics) score plot

# Usage

```
PlotSPLS2DScore(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA,
   inx1,
   inx2,
   reg = 0.95,
   show = 1,
   grey.scale = 0
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is $10.5$ . The second default is width = $0$ , where the width is $7.2$ . Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
reg	Numeric, between 1 and 0
show	Numeric, 1 or 0
grey.scale	Numeric, use grey-scale, 0 for no, and 1 for yes.

PlotSPLS3DScore 227

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotSPLS3DScore

3D SPLS-DA score plot

# Description

Sparse PLS-DA (from mixOmics) 3D score plot

## Usage

```
PlotSPLS3DScore(
   mSetObj = NA,
   imgName,
   format = "json",
   inx1 = 1,
   inx2 = 2,
   inx3 = 3
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotSPLS3DScoreImg

Plot sPLS-DA 3D score plot

### **Description**

This function creates two 3D sPLS-DA score plots, the first is static for Analysis Report purposes, as well as an interactive 3D plot using the plotly R package. The 3D score plot is saved in the created mSetObj (mSetObj\$imgSet\$splsda.3d). To view the score plot, if the name of your mSetObj is mSet, enter "mSet\$imgSet\$splsda.3d" to view the interactive score plot.

This function creates two 3D sPLS-DA score plots, the first is static for Analysis Report purposes, as well as an interactive 3D plot using the plotly R package. The 3D score plot is saved in the created mSetObj (mSetObj\$imgSet\$splsda.3d). To view the score plot, if the name of your mSetObj is mSet, enter "mSet\$imgSet\$splsda.3d" to view the interactive score plot.

### Usage

```
PlotSPLS3DScoreImg(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
  width = NA,
  inx1,
  inx2,
  inx3,
  angl
PlotSPLS3DScoreImg(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
  width = NA,
  inx1,
  inx2,
  inx3,
  angl
)
```

### **Arguments**

mSetObj Input name of the created mSet Object imgName Input a name for the plot format Select the image format, "png", or "pdf".

dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The second default is width = $0$ , where the width is 7.2. Otherwise users can input their own width.
inx1	Numeric, indicate the number of the principal component for the x-axis of the loading plot.
inx2	Numeric, indicate the number of the principal component for the y-axis of the loading plot.
inx3	Numeric, indicate the number of the principal component for the z-axis of the loading plot.
angl	Input the angle

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PlotSPLSDA.Classification

Create SPLS-DA classification plot

## **Description**

Sparse PLS-DA (from mixOmics) plot of classification performance using different components

### Usage

```
PlotSPLSDA.Classification(
   mSetObj = NA,
   imgName,
   format = "png",
   dpi = 72,
   width = NA
)
```

### **Arguments**

mSetObj Input name of the created mSet Object
imgName Input a name for the plot

Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.

230 PlotSPLSLoading

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

validOpt "Mfold"

#### Author(s)

Jeff Xia < jeff . xia@mcgill . ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSPLSLoading

Create SPLS-DA loading plot

## Description

Sparse PLS-DA (from mixOmics) loading plot

### Usage

```
PlotSPLSLoading(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
  width = NA,
  inx,
  viewOpt = "detail"
)
```

### **Arguments**

mSet0bj Input name of the created mSet Object

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

inx Input the model index viewOpt Detailed view "detail"

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotSPLSPairSummary Plot SPLS-DA

## Description

Sparse PLS-DA (from mixOmics) pairwise summary plot

# Usage

```
PlotSPLSPairSummary(
  mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
  width = NA,
  pc.num
)
```

# Arguments

mSetObj	Input name of the created mSet Object
imgName	Input a name for the plot
format	Select the image format, "png", or "pdf".
dpi	Input the dpi. If the image format is "pdf", users need not define the dpi. For "png" images, the default dpi is 72. It is suggested that for high-resolution images, select a dpi of 300.
width	Input the width, there are 2 default widths, the first, width = $NULL$ , is 10.5. The second default is width = 0, where the width is 7.2. Otherwise users can input their own width.
pc.num	Numeric, indicate the number of principle components

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

## Description

Plot a sub heatmap based on results from t-tests/ANOVA, VIP or randomforest

232 PlotSubHeatMap

#### Usage

```
PlotSubHeatMap(
 mSetObj = NA,
  imgName,
  format = "png",
  dpi = 72,
 width = NA,
  dataOpt,
  scaleOpt,
  smplDist,
  clstDist,
  palette,
 method.nm,
  top.num,
  viewOpt,
  rowV = T,
  colV = T,
  border = T,
  grp.ave = F
)
```

#### **Arguments**

mSetObj Input name of the created mSet Object

imgName Input a name for the plot

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

dataOpt Set data options scaleOpt Set the image scale

smplDist Input the sample distance method clstDist Input the clustering distance method

palette Input color palette choice

method.nm Input the method for sub-heat map

top.num Input the top number

viewOpt Set heatmap options, default is set to "detail"

rowV Default is set to T colV Default is set to T

border Indicate whether or not to show cell-borders, default is set to T

grp. ave Logical, default is set to F

PlotTestAccuracy 233

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotTestAccuracy Plot classification performance using different features for Biomarker
Tester

### **Description**

Plot of the accuracy of classification with an increasing number of features.

### Usage

PlotTestAccuracy(mSetObj=NA, imgName, format="png", dpi=72)

## Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", of "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotTT Plot t-test

# Description

Plot t-test

### Usage

```
PlotTT(mSetObj=NA, imgName, format="png", dpi=72, width=NA)
```

234 PlotVolcano

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

imgName Input a name for the plot

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PlotVolcano Create volcano plot

### **Description**

For labelling interesting points, it is defined by the following rules: need to be signficant (sig.inx) and or 2. top 5 p, or 2. top 5 left, or 3. top 5 right.

#### Usage

PlotVolcano(mSetObj=NA, imgName, plotLbl, format="png", dpi=72, width=NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects) imgName Input a name for the plot

plotLbl Logical, plot labels, 1 for yes and 0 for no.

format Select the image format, "png", or "pdf".

dpi Input the dpi. If the image format is "pdf", users need not define the dpi. For

"png" images, the default dpi is 72. It is suggested that for high-resolution im-

ages, select a dpi of 300.

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PLSDA.CV 235

PLSDA.CV

PLS-DA classification and feature selection

## **Description**

PLS-DA classification and feature selection

## Usage

```
PLSDA.CV(
   mSetObj = NA,
   methodName = "T",
   compNum = GetDefaultPLSCVComp(mSetObj),
   choice = "Q2"
)
```

### **Arguments**

mSetObj Input name of the created mSet Object

methodName Logical, by default set to TRUE

compNum GetDefaultPLSCVComp()

choice Input the choice, by default it is Q2

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PLSDA.Permut

Perform PLS-DA permutation

### **Description**

Perform PLS-DA permutation using training classification accuracy as indicator, for two or multigroups

## Usage

```
PLSDA.Permut(mSetObj = NA, num = 100, type = "accu")
```

## Arguments

mSetObj Input name of the created mSet Object num Numeric, input the number of permutations

type Type of accuracy, if "accu" indicate prediction accuracy, else "sep" is separation

distance

236 Predict.class

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PLSR.Anal PLS analysis using oscorespls (Orthogonal scores algorithm) so that VIP can be calculated note: the VIP is calculated only after PLSDA-

CV is performed to determine the best # of comp. used for VIP

### **Description**

PLS analysis using oscorespls

## Usage

```
PLSR.Anal(mSetObj = NA, reg = FALSE)
```

## **Arguments**

mSetObj Input name of the created mSet Object

reg Logical

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Predict.class

Get predicted class probability

# Description

Get predicted class probability

## Usage

```
Predict.class(x.train, y.train, x.test, clsMethod = "pls", lvNum, imp.out = F)
```

## Arguments

x.train	Input the x training samples
y.train	Input the y training samples
x.test	Input the x testing samples

clsMethod Se the classification method, default is set to pls

1vNum Input the number of levelsimp.out Logical, set to F by default

PrepareIntegData 237

### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

PrepareIntegData

Prepare integrated data

## Description

Used for the pathinteg module.

## Usage

```
PrepareIntegData(mSetObj = NA)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

 ${\tt Prepare Network Data}$ 

Prepare data for network exploration

## Description

Function for the network explorer module, prepares user's data for network exploration.

## Usage

```
PrepareNetworkData(mSetObj = NA)
```

## Arguments

mSetObj

Input name of the created mSet Object

238 PreparePermResult

PreparePDFReport

Create report of analyses

## Description

Report generation using Sweave Note: most analyses were already performed, only need to embed the results to the right place without rerunning the whole analysis through Sweave. Only some auxilliary info (i.e. time, version etc need to run in R through Sweave

#### Usage

```
PreparePDFReport(mSetObj = NA, usrName)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

usrName

Input the name of the user

### Author(s)

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PreparePermResult

Prepare report for permutation tests

### Description

Function to prepare a report for permutation tests, used in higher functions

#### Usage

```
PreparePermResult(perm.vec)
```

#### **Arguments**

perm.vec

Input permutation vector

```
Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PreparePrenormData 239

PreparePrenormData

Prepare data for normalization

### **Description**

Function should always be initialized (new or overwrite previous prenorm object).

## Usage

```
PreparePrenormData(mSetObj = NA)
```

### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

PrepareQueryJson

Prepare user's query for mapping KEGG Global Metabolic Network

## Description

This function prepares the user's data for the KEGG Global Metabolic Network

### Usage

```
PrepareQueryJson(mSetObj = NA)
```

### **Arguments**

mSetObj

Input name of the created mSet Object

## Author(s)

Othman Soufan, Jeff Xia <jeff.xia@mcgill.ca>, othman.soufan@mcgill.ca McGill University, Canada License: GNU GPL (>= 2)

240 PrepareROCDetails

PrepareROCData

Prepare data for ROC analysis

## Description

Prepare data for ROC analysis

## Usage

```
PrepareROCData(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

PrepareROCDetails

ROC with CI for AUC

## Description

ROC with CI for AUC

# Usage

```
PrepareROCDetails(mSetObj = NA, feat.nm)
```

## **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

feat.nm

Input the feature name

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Prepare VennData 241

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Prepare data for Venn diagram

## Description

Prepare data for Venn diagram

# Usage

```
PrepareVennData(mSetObj = NA)
```

### **Arguments**

mSetObj

Input name of the created mSet Object

RankFeatures

Rank features based on different importance measures

# Description

Ranks features based on various importance measures, return imp.vec which contains the importance measures of unordered features

## Usage

```
RankFeatures(x.in, y.in, method, lvNum)
```

## Arguments

x.in Input the X features y.in Input the Y features method Input the method

1vNum Input the number of levels

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

242 Read.BatchCSVdata

rda2list

Utility function Make list of KEGG rda files

## Description

Utility function Make list of KEGG rda files

### Usage

```
rda2list(file)
```

Read.BatchCSVdata

Data I/O for batch effect checking

## Description

Read multiple user uploaded CSV data one by one format: row, col

## Usage

```
Read.BatchCSVdata(mSetObj = NA, filePath, format, label)
```

## Arguments

mSetObj Input name of the created mSet Object

filePath Input the path to the batch files

format Input the format of the batch files

label Input the label-type of the files

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Read.MSspec 243

Read.MSspec Read LC/GC-MS spectra (.netCDF, .mzXML, mzData)
---

### Description

This function handles reading in LC/GC-MS spectra files and fills in the dataSet object. It uses functions from the XCMS package to perform peak detection and alignment (grouping).

## Usage

Read.MSspec(mSetObj, folderName, profmethod, fwhm, bw)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

folderName the name of the folder containing the MS spectra

profmethod specify the method to use for profile generation, supports "bin", "binlin", "bin-

linbase" and "intlin"

fwhm specify the full width at half maximum of the matched filtration gaussian model

peak

bw define the bandwidth (standard deviation of the smoothing kernel) to be used

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Read.PeakList	Read peak list files	

### **Description**

This function reads peak list files and fills the data into a dataSet object. For NMR peak lists, the input should be formatted as two-columns containing numeric values (ppm, int). Further, this function will change ppm to mz, and add a dummy 'rt'. For MS peak data, the lists can be formatted as two-columns (mz, int), in which case the function will add a dummy 'rt', or the lists can be formatted as three-columns (mz, rt, int).

#### Usage

```
Read.PeakList(mSetObj=NA, foldername)
```

### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects).

foldername Name of the folder containing the NMR or MS peak list files to read.

244 Read.TextData

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Read.PeakListData

Constructor to read uploaded user files into the mummichog object

#### **Description**

This function handles reading in CSV or TXT files and filling in the mSet object for mummichog analysis. It makes sure that all necessary columns are present.

### Usage

```
Read.PeakListData(mSetObj=NA, filename = NA)
```

### **Arguments**

Input the name of the created mSetObj. mSetObj

filename Input the path name for the CSV/TXT files to read.

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Read.TextData

Constructor to read uploaded CSV or TXT files into the dataSet object

## **Description**

This function handles reading in CSV or TXT files and filling in the dataSet object created using "InitDataObjects".

#### Usage

```
Read.TextData(mSetObj=NA, filePath, format, lbl.type)
```

## **Arguments**

mSetObj	Input the name of the created mSetObj (see InitDataObjects).	
filePath	Input the path name for the CSV/TXT files to read.	
format	Specify if samples are paired and in rows (rowp), unpaired and in rows (rowu), in solumns and paired (colp.)	
	in columns and paired (colp), or in columns and unpaired (colu).	

Specify the data label type, either discrete (disc) or continuous (cont). lbl.type

ReadIndData 245

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca>, Jasmine Chong McGill University, Canada License: GNU GPL (>= 2)

ReadIndData

Read in individual data

### **Description**

This function determines reads in user's individual data for meta-analysis.

### Usage

```
ReadIndData(mSetObj = NA, dataName, format = "colu")
```

#### **Arguments**

mSetObj Input name of the created mSet Object

dataName Name of inputted dataset.

format Specify if samples are paired and in rows (rowp), unpaired and in rows (rowu),

in columns and paired (colp), or in columns and unpaired (colu).

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

ReadPairFile

Read paired peak or spectra files

## Description

This function reads paired peak lists or spectra files. The pair information is stored in a file where each line is a pair and names are separated by ":".

#### Usage

```
ReadPairFile(filePath = "pairs.txt")
```

### **Arguments**

filePath

Set file path

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

246 rectUnique

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Record R Commands

## Description

Record R Commands

## Usage

```
RecordRCommand(mSetObj = NA, cmd)
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

cmd Commands

rectUnique

Perform utilities for peak grouping

## Description

Perform various utilities for peak grouping

### Usage

```
rectUnique(m, order = seq(length = nrow(m)), xdiff = 0, ydiff = 0)
```

# Arguments

```
m Peaks
```

order Performs seq(length = nrow(m))

xdiff Default set to 0 ydiff Default set to 0

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

RegisterData 247

|--|--|--|--|

# Description

When there are multiple datasets, record their name and save the inputted data as a .RDS file to save memory. Note, the memory will only contain one mSetObj\$dataSet object. By default the last one will be the most recent/current dataSet object. Users can switch which data to load into memory.

## Usage

```
RegisterData(mSetObj = NA, dataSet)
```

## **Arguments**

mSetObj	Input name of the created mSet Object
dataSet	Input dataset to be registered in R.

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

RemoveCmpd	Remove selected compounds	

## Description

Remove compounds

## Usage

```
RemoveCmpd(mSetObj = NA, inx)
```

## Arguments

mSetObj	Input name of the created mSet Object
inx	Input the index of compound to remove

248 RemoveDuplicates

RemoveData

Remove data object, the current dataSet will be the last one by default

## Description

Remove data object, the current dataSet will be the last one by default

## Usage

RemoveData(dataName)

# Arguments

dataName

Input name of data to remove

 ${\tt RemoveDuplicates}$ 

Given a data with duplicates, remove duplicates

# Description

Dups is the one with duplicates

### Usage

```
RemoveDuplicates(data, lvlOpt = "mean", quiet = T)
```

#### **Arguments**

data Input data to remove duplicates

1v10pt Set options, default is mean

quiet Set to quiet, logical, default is T

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

RemoveFile 249

RemoveFile

Remove file

# Description

Remove file

## Usage

RemoveFile(fileName)

### **Arguments**

fileName

Input name of file to remove

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

RemoveFolder

Remove folder

## Description

Remove folder

### Usage

RemoveFolder(folderName)

## **Arguments**

folderName

Input name of folder to remove

### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Dam		C ~ .	
Rem	ove	GEI	ıe

Remove selected genes

## Description

Remove selected genes based on an index

### Usage

```
RemoveGene(mSetObj = NA, inx)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

inx Input compound index

RemoveMissingPercent Data processing: remove variables with missing values

## Description

Remove variables based upon a user-defined percentage cut-off of missing values. If a user specifies a threshold of 20 in at least 20

## Usage

```
RemoveMissingPercent(mSetObj, percent)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

percent Input the percentage cut-off you wish to use. For instance, 50 percent is repre-

sented by percent=0.5.

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

ReplaceMin 251

ReplaceMin	Replace missing or zero values	
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### Description

This function will replace zero/missing values by half of the smallest positive value in the original dataset. This method will be called after all missing value imputations are conducted. Also, it directly modifies the mSet\$dataSet\$proc if executed after normalization, or the mSet\$dataSet\$norm if before normalization.

#### Usage

```
ReplaceMin(mSetObj=NA)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jeff Xia < jeff . xia@mcgill . ca> McGill University, Canada License: GNU GPL (>= 2)

RerenderMetPAGraph Redraw current graph for zooming or clipping then return a value

#### **Description**

Redraw current graph for zooming or clipping then return a value

#### Usage

```
RerenderMetPAGraph(mSetObj = NA, imgName, width, height, zoom.factor = NA)
```

#### **Arguments**

mSet0bj Input name of the created mSet Object

imgName Input the name of the plot

width Input the width, there are 2 default widths, the first, width = NULL, is 10.5. The

second default is width = 0, where the width is 7.2. Otherwise users can input

their own width.

height Input the height of the created plot.

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

RF.Anal

Perform Random Forest Analysis

## Description

Perform Random Forest

### Usage

```
RF.Anal(mSetObj = NA, treeNum = 500, tryNum = 7, randomOn = 1)
```

### **Arguments**

mSet0bj Input name of the created mSet Object

treeNum Input the number of trees to create, default is set to 500

tryNum Set number of tries, default is 7

randomOn Set random, default is 1

### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

ROCPredSamplesTable

Create a table of newly classified samples

### **Description**

Function to create the table of newly classified samples

#### Usage

```
ROCPredSamplesTable(mSetObj = NA)
```

## Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects) Function to create the table of newly classified samples

RSVM 253

RSVM R-SVM core code

#### Description

Core code to perform R-SVM

#### Usage

```
RSVM(x, y, ladder, CVtype, CVnum = 0)
```

#### **Arguments**

x Row matrix of data

y Class label: 1 / -1 for 2 classes

ladder Input the ladder

CVtype Integer (N fold CV), "LOO" leave-one-out CV, "bootstrape" bootstrape CV

CVnum Number of CVs, LOO: defined as sample size, Nfold and bootstrape: user de-

fined, default as sample size outputs a named list Error: a vector of CV error on each level SelFreq: a matrix for the frequency of each gene being selected in each level with each column corresponds to a level of selection and each row for

a gene The top important gene in each level are those high-freqent ones

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

RSVM.Anal

Recursive Support Vector Machine (R-SVM)

#### **Description**

recursive SVM for feature selection and classification

#### Usage

```
RSVM.Anal(mSetObj = NA, cvType)
```

## Arguments

mSetObj Input name of the created mSet Object

cvType Cross-validation type

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

254 SanityCheckData

SAM.Anal

Perform Signifiance Analysis of Microarrays (SAM) analysis

#### **Description**

Perform SAM

#### Usage

```
SAM.Anal(
  mSetObj = NA,
  method = "d.stat",
  paired = FALSE,
  varequal = TRUE,
  delta = 0,
  imgName
)
```

## **Arguments**

mSet0bj Input name of the created mSet Object

method Method for SAM analysis, default is set to "d.stat", alternative is "wilc.stat" paired Logical, indicates if samples are paired or not. Default is set to FALSE

varequal Logical, indicates if variance is equal. Default is set to TRUE

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SanityCheckData

Sanity Check Data

## **Description**

SanityCheckData is used for data processing, and performs a basic sanity check of the uploaded content, ensuring that the data is suitable for further analysis. The function will return a message if the data has successfully passed the check and is deemed suitable for further analysis. If it fails, the function will return a 0. The function will perform the check directly onto the mSet\$dataSet object, and must be performed immediately after reading in data. The sanity check function evaluates the accuracy of sample and class labels, data structure, deals with non-numeric values, removes columns that are constant across all samples (variance = 0), and by default replaces missing values with half of the original minimal positive value in your dataset.

SanityCheckIndData 255

#### Usage

SanityCheckData(mSetObj=NA)

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SanityCheckIndData

Sanity check of individual datasets for meta-analysis

#### **Description**

Performs a sanity check on each-uploaded dataset for meta-analysis. Briefly, this function will exclude empty rows, check class labels, ensure only 2 groups are being compared within the dataset, ensure sample names are unique, remove low quality samples/features, and replace missing values.

#### Usage

```
SanityCheckIndData(mSetObj = NA, dataName)
```

#### Arguments

mSet0bj Input name of the created mSet Object

dataName Input name of the dataset to perform the sanity check.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SanityCheckMummichogData

Sanity Check Data

## Description

SanityCheckData is used for data processing, and performs a basic sanity check of the uploaded data, ensuring that the data is suitable for further analysis. The function ensure that all parameters are properly set based on updated parameters.

#### Usage

SanityCheckMummichogData(mSetObj=NA)

256 SearchByCompound

#### Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects).

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt SaveTransformedData}$ 

Save the processed data with class names

## Description

This function saves the processed data with class names as CSV files. Several files may be generated, the original data, processed data, peak normalized, and/or normalized data.

#### Usage

```
SaveTransformedData(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input name of the created mSet Object

#### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

SearchByCompound

Search for compound from all member compounds of metabolite set

## **Description**

Search for compound from all member compounds of metabolite set

## Usage

```
SearchByCompound(mSetObj = NA, query)
```

# Arguments

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

query

Input the query to search

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

SearchByName 257

SearchByName

Given a metabolite set name, search its index

#### **Description**

Given a metabolite set name, search its index

## Usage

```
SearchByName(mSetObj = NA, query)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

query Input the query to search

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

 ${\tt SearchMsetLibraries}$ 

Search metabolite set libraries

# Description

Search metabolite set libraries

#### Usage

```
SearchMsetLibraries(mSetObj = NA, query, type)
```

#### **Arguments**

mSet0bj Input name of the created mSet Object

query Input the query to search

type Input the data type (name or compound)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

258 SelectMultiData

SearchNetDB	Perform mapping of user's data to interaction network
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#### **Description**

This function performs mapping of user's data to the internal network to create a network from the seed nodes

## Usage

```
SearchNetDB(
  mSetObj = NA,
  db.type,
  table.nm,
  require.exp = TRUE,
  min.score = 900
)
```

#### **Arguments**

mSe	tObj	Input name of the created mSet Object
db.	type	Input the database type
tab	le.nm	Input the organism code for the sqlite table (ppi). For chemical type, the table.nm is drugbank of ctd
req	uire.exp	Logical, only used for the STRING database
min	.score	Input the minimal score, only used for the STRING database

#### Author(s)

Othman Soufan, Jeff Xia <jeff.xia@mcgill.ca>, othman.soufan@mcgill.ca McGill University, Canada License: GNU GPL (>= 2)

```
SelectMultiData Select one or more datasets for meta-analysis
```

## Description

This function selects one or more datasets to be used for meta-analysis. 1 is used to indicate that a dataset is selected and by default, all datasets will be selected for meta-analysis.

# Usage

```
SelectMultiData(mSetObj = NA)
```

SetAnalysisMode 259

#### **Arguments**

mSetObj

Input name of the created mSet Object

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetAnalysisMode

Set biomarker analysis mode

## Description

**ROC** utilities

#### Usage

```
SetAnalysisMode(mSetObj, mode)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)s

mode

Input the selected mode for biomarker analysis, "univ" for univariate ROC curve analysis, "explore" for multivariate ROC curve analysis, and "test" for ROC curve based model creation and evaluation. McGill University, Canada License:

GNU GPL (>= 2)

## Author(s)

```
Jeff Xia < jeff.xia@mcgill.ca>
```

SetAnnotationParam

Set annotation parameters

# Description

This function sets the parameters for peak annotation.

#### Usage

```
SetAnnotationParam(
  polarity = "positive",
  perc_fwhm = 0.6,
  mz_abs_iso = 0.005,
  max_charge = 2,
  max_iso = 2,
  corr_eic_th = 0.85,
  mz_abs_add = 0.001
)
```

260 SetCachexiaSetUsed

# Arguments

polarity	Character, specify the polarity of the MS instrument. Either "negative" or "positive".
perc_fwhm	Numeric, set the percentage of the width of the FWHM for peak grouping. Default is set to 0.6.
mz_abs_iso	Numeric, set the allowed variance for the search (for isotope annotation). The default is set to $0.005$ .
max_charge	Numeric, set the maximum number of the isotope charge. For example, the default is 2, therefore the max isotope charge is 2+/
max_iso	Numeric, set the maximum number of isotope peaks. For example, the default is 2, therefore the max number of isotopes per peaks is 2.
corr_eic_th	Numeric, set the threshold for intensity correlations across samples. Default is set to $0.85$ .
mz_abs_add	Numeric, set the allowed variance for the search (for adduct annotation). The default is set to 0.001.

## Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetCachexiaSetUsed	Set the cachexia set used

# Description

Set cachexia set used

# Usage

SetCachexiaSetUsed(mSetObj = NA, used)

## **Arguments**

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
used	Set data to be used

SetCandidate 261

C - 1	O		-1	- 4	_
Set	∖.ar	naı	a	аτ	e

Set matched name based on user selection from all potential hits

## Description

Note: to change object in the enclosing environment, use "«-"

## Usage

```
SetCandidate(mSetObj = NA, query_nm, can_nm)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects).

query\_nm Input the query name.

can\_nm Input the candidate name.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetClass

Set class information for MS data

## Description

This function sets the class information for preprocessing MS data.

#### Usage

```
SetClass(class)
```

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

262 SetCurrentMsetLib

SetCurrentGroups To choose from two groups

#### Description

Choose two groups (when more than two groups uploaded)

# Usage

```
SetCurrentGroups(mSetObj = NA, grps)
```

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

grps Input the groups

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetCurrentMsetLib Set current user selected metset library for search

#### **Description**

if enrichment analysis, also prepare lib by creating a list of metabolite sets

#### Usage

```
SetCurrentMsetLib(mSetObj=NA, lib.type, excludeNum)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

(metabolitesets < excludeNum)

lib.type Input user selected name of library, "self", "kegg\_pathway", "smpdb\_pathway",

"blood", "urine", "csf", "snp", "predicted", "location", and "drug".

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetCustomData 263

SetCustomData

Set custom data

## Description

The "selected.cmpds" should be for extraction

## Usage

```
SetCustomData(mSetObj = NA, selected.cmpds, selected.smpls)
```

# Arguments

```
mSetObj Input the name of the created mSetObj (see InitDataObjects)
selected.cmpds Input the vector containing the compounds
selected.smpls Input the vector containing the samples
```

 ${\tt SetDesignType}$ 

For two factor time series only

# Description

For two factor time series only

# Usage

```
SetDesignType(mSetObj = NA, design)
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

design Input the design type

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

264 SetMetabolomeFilter

SetKEGG.PathLib

Set KEGG pathway library

#### **Description**

note, this process can be long, need to return a value to force Java to wait

## Usage

```
SetKEGG.PathLib(mSetObj = NA, libNm, lib.version)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

lib.version Input the KEGG pathway version. "current" for the latest KEGG pathway li-

brary or "v2018" for the KEGG pathway library version prior to November

2019.

kegg.rda Input the name of the KEGG library

#### Author(s)

Jeff Xia < jeff. xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetMetabolomeFilter

Set metabolome filter

# Description

Set metabolome filter

#### Usage

```
SetMetabolomeFilter(mSetObj = NA, TorF)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

TorF Input metabolome filter

SetMummichogPval 265

SetMummichogPval

Set the cutoff for mummichog analysis

## **Description**

Set the p-value cutoff for mummichog analysis.

#### Usage

```
SetMummichogPval(mSetObj = NA, cutoff)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj.

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

 ${\tt SetMummichogPvalFromPercent}$ 

Set the cutoff for mummichog analysis

## **Description**

Set the p-value cutoff for mummichog analysis.

#### Usage

```
SetMummichogPvalFromPercent(mSetObj = NA, fraction)
```

## **Arguments**

mSetObj

Input the name of the created mSetObj.

```
Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

266 SetPeakEnrichMethod

SetOrganism

Set organism for further analysis

## Description

Set organism for further analysis

## Usage

```
SetOrganism(mSetObj = NA, org)
```

# Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

org Set organism ID

SetPeakEnrichMethod

Set the peak enrichment method for the MS Peaks to Paths module

## Description

This function sets the peak enrichment method.

#### Usage

```
SetPeakEnrichMethod(mSetObj = NA, algOpt)
```

## **Arguments**

mSetObj

Input the name of the created mSetObj.

```
Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

SetPeakFormat 267

SetPeakFormat

Set the peak format for the mummichog analysis

## Description

Set the peak format for mummichog analysis.

## Usage

```
SetPeakFormat(type)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj.

## Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetPeakList.GroupValues

Set peak list group values

# Description

Set peak list group values

#### Usage

```
SetPeakList.GroupValues(mSetObj = NA)
```

# Arguments

mSetObj

Input name of mSetObj, the data used is the nmr.xcmsSet object

268 SetPeakParam

SetPeakParam

Set parameters for peak picking using XCMS and CAMERA

## Description

This function sets all the parameters used for downstream pre-processing of user's raw MS data.

## Usage

```
SetPeakParam(
  alg = "centwave",
  ppm = 10,
  min_pkw = 10,
  max_pkw = 60,
  sn_{thresh} = 6,
  mzdiff = 0.01,
  bw = 5,
  min_frac = 0.5,
  min_sample_num = 1,
  max_feats = 100,
  peakgroup = FALSE,
  bin_size = 1,
  min_frac_retcor = 0.9,
  rt_filt = FALSE,
  rt_min = 200,
  rt_max = 1000
)
```

## Arguments

alg	Character, specify the algorithm to perform peak detection. "centwave" to use the CentWave algorithm, and "match_filt" to use the MatchedFilter algorithm.
ppm	Numeric, specify the mass error in ppm.
min_pkw	Numeric, specify the minimum peak width in seconds.
max_pkw	Numeric, specify the maximum peak width in seconds.
sn_thresh	Numeric, specify the signal to noise threshold.
mzdiff	Numeric, specify the minimum m/z difference for signals to be considered as different features when retention times are overlapping.
bw	Numeric, specify the band width (sd or half width at half maximum) of gaussian smoothing kernel to be applied during peak grouping.
min_frac	Numeric, specify fraction of samples in each group that contain the feature for it to be grouped.
min_sample_num	Numeric, specify minimum number of sample(s) in each group that contain the feature for it to be included.

SetSMPDB.PathLib 269

Numeric, specify the maximum number of features to be identified. max\_feats Boolean, if true, PeakGroup algorithm is used for peak alignment; if false, Obipeakgroup warp method is used. bin\_size Numeric, specify the bin size (in m/z) to be used for the profile matrix generation used for peak alignment (Obiwarp method). min\_frac\_retcor Numeric, specify fraction of samples in all groups that contain the peaks for them to be aligned (PeakGroup method). rt\_filt Boolean, if true, users must specify the minimum and maximum retention time to be included in the analysis. By default this is set to 200 - 1000. Numeric, specify the minimum retention time. rt\_min Numeric, specify the maximum retention time. rt\_max

#### Author(s)

Jasmine Chong <jasmine.chong@mail.mcgill.ca>, Mai Yamamoto <yamamoto.mai@mail.mcgill.ca>, and Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

	SetSMPDB.PathLib	Set SMPDB pathway library	
--	------------------	---------------------------	--

## Description

note, this process can be long, need to return a value to force Java to wait

#### Usage

```
SetSMPDB.PathLib(mSetObj = NA, libNm)
```

## **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)
smpdb.rda Input the name of the SMPDB library (e.g. hsa or mmu)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

270 Setup.BiofluidType

Setup.AdductData

Save adduct names for mapping

## Description

Save adduct names for mapping

## Usage

```
Setup.AdductData(mSetObj = NA, qvec)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

qvec Input the vector to query

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Setup.BiofluidType

Save biofluid type for SSP

## Description

Save biofluid type for SSP

#### Usage

```
Setup.BiofluidType(mSetObj = NA, type)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

type Input the biofluid type

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Setup.ConcData 271

Setup.ConcData

Save concentration data

#### **Description**

Save concentration data

## Usage

```
Setup.ConcData(mSetObj = NA, conc)
```

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

conc Input the concentration data

#### Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Setup.HMDBReferenceMetabolome

Read user uploaded metabolome as a list of HMDB compound names

## **Description**

Read user uploaded metabolome as a list of HMDB compound names

## Usage

```
Setup.HMDBReferenceMetabolome(mSetObj = NA, filePath)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)
filePath Input the path to the user's list of HMDB compound names

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

272 Setup.MapData

Setup.KEGGReferenceMetabolome

Read user uploaded metabolome as a list of KEGG pathway ids

## **Description**

Read user uploaded metabolome as a list of KEGG pathway ids

#### Usage

```
Setup.KEGGReferenceMetabolome(mSetObj = NA, filePath)
```

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects)

filePath Input the path to the user's list of KEGG pathway ids

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Setup.MapData

Save compound name for mapping

## Description

Save compound name for mapping

## Usage

```
Setup.MapData(mSetObj = NA, qvec)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)

qvec Input the vector to query

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

Setup.UserMsetLibData Read user upload metabolite set library file

#### **Description**

Return two col csv file, first name, second cmpd list

#### Usage

```
Setup.UserMsetLibData(mSetObj = NA, filePath)
```

#### **Arguments**

mSetObj Input the name of the created mSetObj (see InitDataObjects)
filePath Input the path to the user's uploaded metabolite set library

#### Author(s)

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

SetupKEGGLinks

Only works for human (hsa.rda) data

# Description

Only works for human (hsa.rda) data 2018 - works for ath, eco, mmu, sce

#### Usage

```
SetupKEGGLinks(smpdb.ids)
```

# Arguments

kegg.ids

Input the list of KEGG ids to add SMPDB links

```
Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

274 SetupSMPDBLinks

SetupMSdataMatrix

Create a MS spectra data matrix of peak values for each group

#### **Description**

This function sets up a MS spectra data matrix using the 'groupval' function from XCMS. This generates a usable matrix of peak values for analysis where columns represent peak groups and rows represent samples. Collisions where more than one peak from a single sample are in the same group get resolved utilizing "medret", which uses the peak closest to the median retention time.

#### Usage

SetupMSdataMatrix(mSetObj, intvalue)

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

intvalue

name of peak column to enter into the returned matrix, if intvalue = 'into', use integrated area of original (raw) peak intensities, if intvalue = 'intf', use integrated area of filtered peak intensities, if intvalue = 'intb', use baseline corrected integrated peak intensities, if intvalue = 'maxo', use the maximum intensity of original (raw) peaks, or if intvalue = 'maxf' use the maximum intensity of fil-

tered peaks

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SetupSMPDBLinks

Only works for human (hsa.rda) data

#### **Description**

Only works for human (hsa.rda) data 2018 - works for ath, eco, mmu, sce

#### Usage

```
SetupSMPDBLinks(kegg.ids)
```

#### **Arguments**

kegg.ids

Input the list of KEGG ids to add SMPDB links

## Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SOM.Anal 275

SOM. Anal

SOM analysis

## Description

SOM analysis

#### Usage

```
SOM.Anal(mSetObj = NA, x.dim, y.dim, initMethod, neigb = "gaussian")
```

#### **Arguments**

mSetObj Input name of the created mSet Object x.dim Input X dimension for SOM analysis y.dim Input Y dimension for SOM analysis initMethod Input the method

neigb Default is set to 'gaussian'

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

```
sparse.mint.block_iteration
```

Perform Sparse Generalized Canonical Correlation (sgccak)

#### **Description**

Runs sgccak() modified from RGCCA

## Usage

```
sparse.mint.block_iteration(
   A,
   design,
   study = NULL,
   keepA.constraint = NULL,
   keepA = NULL,
   scheme = "horst",
   init = "svd",
   max.iter = 100,
   tol = 1e-06,
   verbose = TRUE,
   bias = FALSE,
   penalty = NULL
)
```

276 splsda

#### **Arguments**

Data design Set design study Default set to NULL keepA.constraint Default set to NULL keepA Default set to NULL scheme Scheme, default set to "horst" init Init mode, default set to "svd" max.iter Max number of iterations, numeric, default set to 100 tol Tolerance, numeric, default set to 1e-06 Default set to TRUE verbose Default set to FALSE bias

Default set to NULL

#### Author(s)

penalty

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

splsda Perform sPLS-DA

#### **Description**

Sparse PLS functions (adapted from mixOmics package for web-based usage) this function is a particular setting of internal\_mint.block the formatting of the input is checked in internal\_wrapper.mint

#### Usage

```
splsda(
   X,
   Y,
   ncomp = 2,
   mode = c("regression", "canonical", "invariant", "classic"),
   keepX,
   keepX.constraint = NULL,
   scale = TRUE,
   tol = 1e-06,
   max.iter = 100,
   near.zero.var = FALSE,
   logratio = "none",
   multilevel = NULL
)
```

SPLSR.Anal 277

#### **Arguments**

X numeric matrix of predictors

Y a factor or a class vector for the discrete outcome

ncomp the number of components to include in the model. Default to 2.

mode Default set to c("regression", "canonical", "invariant", "classic")

keepX Number of X variables kept in the model on the last components (once all

keepX.constraint[[i]] are used).

keepX.constraint

A list containing which variables of X are to be kept on each of the first PLS-

components.

scale Boleean. If scale = TRUE, each block is standardized to zero means and unit

variances (default: TRUE).

tol Convergence stopping value.

max.iter integer, the maximum number of iterations.

near.zero.var boolean, see the internal nearZeroVar function (should be set to TRUE in par-

ticular for data with many zero values). Setting this argument to FALSE (when

appropriate) will speed up the computations

logratio "None" by default, or "CLR"

multilevel Designate multilevel design, "NULL" by default

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

SPLSR. Anal Perform SPLS-DA

#### **Description**

Sparse PLS-DA (from mixOmics)

#### Usage

```
SPLSR.Anal(mSetObj = NA, comp.num, var.num, compVarOpt, validOpt = "Mfold")
```

#### **Arguments**

mSetObj Input name of the created mSet Object comp.num Input the number of computations to run

var.num Input the number of variables

compVarOpt Input the option to perform SPLS-DA

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

278 template.match

SumNorm

Row-wise Normalization

#### **Description**

Row-wise norm methods, when x is a row. Normalize by a sum of each sample, assume constant sum (1000). Options for normalize by sum median, reference sample, reference reference (compound), or quantile normalization

## Usage

SumNorm(x)

#### **Arguments**

Х

Input data to normalize

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada

template.match

Pattern hunter

#### **Description**

Run template on all the high region effect genes

#### Usage

```
template.match(x, template, dist.name)
```

## Arguments

x Input data template Input template

dist.name Input distance method

# Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

Ttests.Anal 279

Ttests.Anal

Perform t-test analysis

#### **Description**

This function is used to perform t-test analysis.

# Usage

```
Ttests.Anal(
  mSetObj = NA,
  nonpar = F,
  threshp = 0.05,
  paired = FALSE,
  equal.var = TRUE,
  all_results = FALSE)
```

## Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
nonpar	Logical, use a non-parametric test, T or F. False is default.
threshp	Numeric, enter the adjusted p-value (FDR) cutoff
paired	Logical, is data paired (T) or not (F).
equal.var	Logical, evaluates if the group variance is equal (T) or not (F).
all_results	Logical, if TRUE, returns T-Test analysis results for all compounds.

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

UnzipUploadedFile

Unzip .zip files

#### **Description**

Unzips uploaded .zip files, removes the uploaded file, checks for success

# Usage

```
UnzipUploadedFile(inPath, outPath, rmFile = T)
```

#### **Arguments**

inPath Input the path of the zipped files

outPath Input the path to directory where the unzipped files will be deposited

rmFile Logical, input whether or not to remove files. Default set to T

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

UpdateData

Update data for filtering

#### **Description**

Function to update the mSetObj after removing features or samples.

#### Usage

```
UpdateData(mSetObj = NA)
```

#### **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

UpdateGraphSettings

Update graph settings

# Description

Function to update the graph settings.

## Usage

```
UpdateGraphSettings(mSetObj = NA, colVec, shapeVec)
```

## **Arguments**

mSetObj

Input the name of the created mSetObj (see InitDataObjects)

UpdateInstrumentParameters

Update the mSetObj with user-selected parameters for MS Peaks to Pathways.

## Description

This functions handles updating the mSet object for mummichog analysis. It is necessary to utilize this function to specify to the organism's pathways to use (libOpt), the mass-spec mode (msModeOpt) and mass-spec instrument (instrumentOpt).

#### Usage

UpdateInstrumentParameters(mSetObj=NA, instrumentOpt, msModeOpt, custom=FALSE)

#### Arguments

mSetObj Input the name of the created mSetObj (see InitDataObjects).

instrumentOpt Define the mass-spec instrument used to perform untargeted metabolomics.

msModeOpt Define the mass-spec mode of the instrument used to perform untargeted metabolomics.

custom Logical, select adducts for mummichog to consider.

#### Author(s)

Jasmine Chong, Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

UpdateIntegPathwayAnalysis

Update integrative pathway analysis for new input list

#### **Description**

used for integrative analysis as well as general pathways analysis for meta-analysis results

#### Usage

UpdateIntegPathwayAnalysis(mSetObj=NA, qids, file.nm, topo="dc", enrich="hyper", libOpt="integ")

282 UpdateOPLS.Splot

#### **Arguments**

mSetObj Input name of the created mSet Object

qids Input the query IDs

file.nm Input the name of the file

topo Select the mode for topology analysis: Degree Centrality ("dc") measures the

number of links that connect to a node (representing either a gene or metabolite) within a pathway; Closeness Centrality ("cc") measures the overall distance from a given node to all other nodes in a pathway; Betweenness Centrality ("bc")measures the number of shortest paths from all nodes to all the others that

pass through a given node within a pathway.

enrich Method to perform over-representation analysis (ORA) based on either hyper-

genometrics analysis ("hyper") or Fisher's exact method ("fisher").

libOpt Select the different modes of pathways, either the gene-metabolite mode ("in-

teg") which allows for joint-analysis and visualization of both significant genes and metabolites or the gene-centric ("genetic") and metabolite-centric mode ("metab") which allows users to identify enriched pathways driven by signifi-

cant genes or metabolites, respectively.

#### Author(s)

Jeff Xia <jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

UpdateOPLS.Splot

Update OPLS loadings

#### Description

Update the OPLS loadings

#### Usage

```
UpdateOPLS.Splot(mSetObj = NA, plotType)
```

#### Arguments

mSetObj Input name of the created mSet Object

plotType Set annotation type, "all" to label all variables and "none" to label no variables.

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

UpdatePCA.Loading 283

UpdatePCA.Loading

Update PCA loadings

## Description

Update the PCA loadings

## Usage

```
UpdatePCA.Loading(mSetObj = NA, plotType)
```

#### **Arguments**

mSet0bj Input name of the created mSet Object

plotType Set annotation type, "all" to label all variables and "none" to label no variables.

## Author(s)

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

UpdatePLS.Loading

Update PLS loadings

#### **Description**

Update the PLS loadings

#### Usage

```
UpdatePLS.Loading(mSetObj = NA, plotType)
```

## Arguments

mSet0bj Input name of the created mSet Object

plotType Set annotation type, "all" to label all variables and "none" to label no variables.

```
Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)
```

284 Volcano.Anal

## Description

Convert user coords (as used in current plot) to pixels in a png adapted from the imagemap package

## Usage

```
usr2png(xy, im)
```

# Arguments

xy	Input coordinates
im	Input coordinates

## Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

|--|

# Description

Perform volcano analysis

#### Usage

Volcano.Anal(mSetObj=NA, paired=FALSE, fcthresh, cmpType, percent.thresh, nonpar=F, threshp, equal.va

# Arguments

mSetObj	Input the name of the created mSetObj (see InitDataObjects)
paired	Logical, T if data is paired, F if data is not.
fcthresh	Numeric, input the fold change threshold
cmpType	Comparison type, 1 indicates group 1 vs group 2, and 2 indicates group 2 vs group 1 $$
${\tt percent.thresh}$	Only for paired data, numeric, indicate the significant count threshold
nonpar	Logical, indicate if a non-parametric test should be used (T or F)
threshp	Numeric, indicate the p-value threshold
equal.var	Logical, indicates if the group variance is equal (T) or unequal (F)
pval.type	To indicate raw p-values, use "raw". To indicate FDR-adjusted p-values, use "fdr".

XSet2MSet 285

#### Author(s)

Jeff Xia<jeff.xia@mcgill.ca> McGill University, Canada License: GNU GPL (>= 2)

XSet2MSet Converts xset object from XCMS to mSet object for MetaboAnalyst

# Description

This function converts processed raw LC/MS data from XCMS to a usable data object (mSet) for MetaboAnalyst. The immediate next step following using this function is to perform a SanityCheck, and then further data processing and analysis can continue.

## Usage

XSet2MSet(xset, dataType, analType, paired = F, format, lbl.type)

#### **Arguments**

xset	The name of the xcmsSet object created.
dataType	The type of data, either list (Compound lists), conc (Compound concentration data), specbin (Binned spectra data), pktable (Peak intensity table), nmrpeak (NMR peak lists), mspeak (MS peak lists), or msspec (MS spectra data).
analType	Indicate the analysis module to be performed: stat, pathora, pathqea, msetora, msetssp, msetqea, ts, cmpdmap, smpmap, or pathinteg.
paired	Logical, is data paired (T) or not (F).
format	Specify if samples are paired and in rows (rowp), unpaired and in rows (rowu), in columns and paired (colp), or in columns and unpaired (colu).
lbl.type	Specify the data label type, either discrete (disc) or continuous (cont).

# **Index**

.read.metaboanalyst.lib, 13	CreateBiomarkerInputDoc, 31
.readDataTable, 13	CreateBiomarkerIntr, 32
	CreateBiomarkerOverview, 32
AddErrMsg, 14	CreateBiomarkerRatioOverview, 32
analyze.lipids, 14	CreateBiomarkerRnwReport, 33
ANOVA.Anal, 15	CreateCorrDoc, 33
ANOVA2.Anal, 15	createCVset, 34
aof, 16	CreateEBAMdoc, 34
aov.between, 16	CreateEnrichAnalDoc, 35
aov.repeated, 17	CreateEnrichInputDoc, 35
aov.within, 17	CreateEnrichIntr, 36
ASCAfun.res, 18	CreateEnrichORAdoc, 36
ASCAfun1, 18	CreateEnrichOverview, 37
ASCAfun2, 19	CreateEnrichProcessDoc, 37
	CreateEnrichQEAdoc, 38
calculateConcISO, 19	CreateEnrichRnwReport, 38
CalculateFeatureRanking, 20	CreateEnrichSSPdoc, 39
CalculateGlobalTestScore, 20	CreateFooter, 39
CalculateHyperScore, 21	CreateGraph, 40
CalculateImpVarCutoff, 21	CreateGSEAAnalTable, 40
CalculateOraScore, 22	CreateHCdoc, 41
CalculatePairwiseDiff, 22	CreateHeatmap2doc, 41
CalculateQeaScore, 23	CreateIntegPathwayAnalysisRnwReport,
CalculateSSP, 23	42
CheckMetaDataConsistency, 24	CreateIntegratedPathwayAnalInputDoc,
CleanData, 24	42
CleanDataMatrix, 25	CreateIntegratedPathwayAnalIntr, 43
CleanNumber, 25	CreateIntegratedPathwayDoc, 43
ClearNegatives, 26	CreateIntegratedPathwayGeneMapTable,
ClearStrings, 26 ClearUserDir, 27	44
Compound_function_mzlist, 27	CreateIntegratedPathwayNameMapTable,
ComputeAverageCurve, 27	44
computeConc, 28	CreateIntegratedPathwayResultsTable,
ComputeHighLow, 28	45
Convert2Mummichog, 29	CreateiPCAdoc, 45
CreateAnalNullMsg, 29	CreateKMdoc, 46
CreateANOVAdoc, 30	CreateLadder, 46
CreateA0V2doc, 30	CreateLibFromKEGG, 47
CreateASCAdoc, 31	CreateMappingResultTable, 47
or carendonauc, J1	or cateriappingnesuitrable, 47

CreateMBdoc, 47	CreateSOMdoc, 70
CreateMetaAnalTable, 48	CreateSPLSDAdoc, 71
CreateMetaAnalysisDEdoc,48	CreateStatIntr, 71
CreateMetaAnalysisInputDoc, 49	CreateStatIOdoc, 72
CreateMetaAnalysisIntr, 49	CreateStatRnwReport, 72
CreateMetaAnalysisNORMdoc, 50	CreateSummaryTable, 73
CreateMetaAnalysisOutput, 50	CreateSVMdoc, 73
CreateMetaAnalysisOverview, 51	CreateTimeSeriesAnalNullMsg,74
CreateMetaAnalysisRnwReport, 51	CreateTimeSeriesIOdoc, 74
CreateMetaTable, 52	CreateTimeSeriesRnwReport, 75
CreateModelBiomarkersDoc, 52	CreateUnivarBiomarkersDoc, 75
CreateMultiBiomarkersDoc, 53	CreateUNIVdoc, 76
CreateMummichogAnalTable, 53	CreateUnivROCTable, 76
CreateMummichogAnalysisDoc, 54	CreateVennMetaTable, 77
CreateMummichogInputDoc, 54	CrossReferencing, 77
CreateMummichogIntro, 55	CVTest.LRmodel, 78
CreateMummichogLibs, 55	
CreateMummichogOverview, 56	descendMin, 79
CreateMummichogRnwReport, 56	doCompoundMapping, 79
CreateNetworkExplorerDoc, 57	doGeneIDMapping, 80
CreateNetworkExplorerInputDoc, 57	doKEGG2NameMapping, 80
CreateNetworkExplorerIntr, 58	doKOFiltering,81
CreateNetworkExplorerOverview, 58	FDAM To:+ 01
CreateNetworkExplorerRnwReport, 59	EBAM. Init, 81
CreateNetworkGeneMapTable, 59	ExtractMS2data, 82
CreateNetworkNameMapTable, 60	FC.Anal.paired, 82
CreateNORMdoc, 60	FC.Anal.unpaired, 83
CreateOPLSDAdoc, 61	FeatureCorrelation, 83
CreatePathAnalDoc, 61	fgsea2, 84
CreatePathInputDoc, 62	fillpathways, 84
CreatePathIntr, 62	FilterVariable, 84
CreatePathProcessDoc, 62	findEqualGreaterM, 85
CreatePathResultDoc, 63	FisherLSD, 86
CreatePathRnwReport, 63	FormatPeakList, 86
CreatePCAdoc, 64	Tormatr canzist, oo
CreatePLSdoc, 64	genLogisticRegMdl,87
CreatePowerAnalDoc, 65	Get.asca.tss, 88
CreatePowerInputDoc, 65	Get.bwss, 88
CreatePowerIntr, 66	Get.ConcRef, 89
CreatePowerOverview, 66	Get.Leverage, 89
CreatePowerParametersDoc, 66	Get.pAUC, 90
CreatePowerRnwReport, 67	Get.pred, 90
CreateRatioTable, 67	Get.rpart.summary, 91
CreateRFdoc, 68	Get.VIP, 91
CreateRHistAppendix, 68	GetAbundanceLabel, 92
CreateROCLabelsTable, 69	GetAccuracyInfo, 92
CreateSAMdoc, 69	GetAllDataNames, 92
CreateSemiTransColors, 70	GetAllKMClusterMembers, 93

GetAllSOMClusterMembers, 93	GetSelectedDataNumber, 115
GetCandidateList, 94	GetSigTable, 116
GetCircleInfo, 94	GetSigTable.Anova, 116
GetCIs, 95	GetSigTable.Aov2, 117
GetCMD, 95	GetSigTable.ASCA, 117
GetCompoundDetails, 96	GetSigTable.Corr, 118
GetConvertFullPath, 96	GetSigTable.EBAM, 118
getDataFromTextArea, 97	GetSigTable.FC, 118
GetExtendRange, 97	GetSigTable.MB, 119
GetFC, 98	GetSigTable.RF, 119
GetFeatureNumbers, 98	GetSigTable.SAM, 119
GetFinalNameMap, 99	GetSigTable.SVM, 120
GetFisherPvalue, 99	GetSigTable.TT, 120
GetHTMLMetSet, 100	GetSigTable.Volcano, 120
GetHTMLPathSet, 100	GetSOMClusterMembers, 121
GetImpFeatureMat, 101	GetSSPTable, 121
GetKEGGNodeInfo, 101	GetSuggestedSAMDelta, 122
GetKMClusterMembers, 102	GetSVMSigMat, 122
GetLassoFreqs, 102	GetTopInx, 123
GetLimmaResTable, 103	<pre>GetTrainTestSplitMat, 123</pre>
GetMapTable, 103	GetTtestRes, 124
GetMaxPCAComp, 103	GetTTSigMat, 124
GetMeanROC, 104	GetUnivReport, 125
	GetVariableLabel, 125
GetMetaResultMatrix, 104	GetVennGeneNames, 126
GetMetaSigHitsTable, 105	GetXYCluster, 126
GetMetSetName, 105	GroupPeakList, 127
GetMsetLibCheckMsg, 106	
GetMsetLibSearchResult, 106	heckbert, 127
GetMsetNames, 107	HMDBID2KEGGID, 128
GetMummichogPathSetDetails, 107	HMDBID2Name, 128
GetNetworkGeneMappingResultTable, 108	
GetNewSampleNames, 108	ImportRawMSData, 129
GetORA.pathNames, 109	<pre>ImportRawMSDataList, 130</pre>
GetORA.smpdbIDs, 109	ImputeVar, 131
GetORATable, 109	InitDataObjects, 131
GetQEA.keggIDs, 110	InitPowerAnal, 132
GetQEA.pathNames, 110	<pre>InitStatAnalMode, 132</pre>
GetQEATable, 111	<pre>InitTimeSeriesAnal, 133</pre>
GetRCommandHistory, 111	iPCA.Anal, 133
GetRFConf.Table, 111	isEmptyMatrix, 134
GetRFConfMat, 112	IsSmallSmplSize, 134
GetRF00B, 112	IsSpectraProcessingOK, 135
GetRFSigMat, 113	
GetROC.coords, 113	KEGGID2HMDBID, 135
GetROCLassoFreq, 114	KEGGID2Name, 136
GetROCTtestP, 114	KEGGPATHID2SMPDBIDs, 136
GetSampleSizeLadder, 115	Kmeans.Anal, 137
GetSelectedDataNames, 115	kwtest, 137

LoadKEGGKO_lib, 138	PerformLimmaDE, 160
LoadKEGGLib, 138	PerformMapping, 161
LoadSmpLib, 138	performMB, 161
LogNorm, 139	PerformMetaMerge, 162
LSD.test, 139	PerformMultiMatch, 162
	PerformPeakAnnotation, 163
make_cpdlib, 140	PerformPeakProfiling, 163
make_cpdlist, 140	PerformPowerProfiling, 164
map, 141	PerformPSEA, 165
MapCmpd2KEGGNodes, 141	PerformPvalCombination, 165
MapK02KEGGEdges, 142	PerformVoteCounting, 166
Match.Pattern, 142	Plot.Permutation, 166
MergeDatasets, 143	PlotAccuracy, 167
MergeDuplicates, 143	PlotANOVA, 167
MetaboliteMappingExact, 144	PlotANOVA2, 168
MSspec.fillPeaks, 144	PlotASCA.Permutation, 169
MSspec.rtCorrection, 145	PlotAscaImpVar, 169
multi.stat, 145	PlotASCAModel, 170
	PlotBoxPlot, 171
nearZeroVar, 277	PlotCmpdSummary, 171
Normalization, 146	PlotCmpdView, 172
OPLSDA.Permut, 147	PlotConcRange, 173
OPLSR.Anal, 147	PlotCorr, 173
	PlotCorrHeatMap, 174
parseFisher, 148	PlotDetailROC, 175
parseTukey, 148	PlotEBAM. Cmpd, 176
PCA.Anal, 149	PlotEIC, 176
PCA.Flip, 149	PlotEnrichNet.Overview, 177
PCA.GENES, 150	PlotFC, 178
Perform. ASCA, 150	PlotHCTree, 178
Perform. ASCA. permute, 151	PlotHeatMap, 179
Perform.Permut, 151	PlotHeatMap2, 181
Perform.permutation, 152	PlotImpVar, 182
Perform.UnivROC, 152	PlotImpVars, 182
PerformAdductMapping, 153	PlotInmexGraph, 183
PerformApproxMatch, 154	PlotInmexPath, 184
PerformBatchCorrection, 154	PlotIntegPaths, 184
PerformCurrencyMapping, 155	PlotInteraction, 185
PerformCV.explore, 155	PlotKEGGPath, 186
PerformCV.test, 156	PlotKmeans, 186
PerformDetailMatch, 156	PlotLoadingCmpd, 187
PerformEachDEAnal, 157	PlotMBTimeProfile, 188
PerformIndNormalization, 157	PlotMetaVenn, 188
PerformIntegCmpdMapping, 158	PlotMetpaPath, 189
PerformIntegGeneMapping, 158	PlotModelScree, 189
PerformIntegPathwayAnalysis, 159	PlotMS.RT, 190
PerformKOEnrichAnalysis_KO01100, 159	PlotMS2Spectra, 191
PerformKOEnrichAnalysis_List, 160	PlotMSEA.Overview, 191
·	•

DI ANCDA LA DANNA 102	D1 - + CD1 C2DC 1 220
PlotMSPeaksPerm, 192	PlotSPLS3DScoreImg, 228
PlotNormSummary, 192	PlotSPLSDA.Classification, 229
PlotOPLS. MDL, 193	PlotSPLSLoading, 230
PlotOPLS. Permutation, 194	PlotSPLSPairSummary, 231
PlotOPLS.Splot, 194	PlotSubHeatMap, 231
Plot0PLS2DScore, 195	PlotTestAccuracy, 233
Plot0RA, 196	PlotTT, 233
PlotPathSummary, 197	PlotVolcano, 234
PlotPathwayMZHits, 198	PLSDA CV, 235
PlotPCA.overview, 198	PLSDA.Permut, 235
PlotPCA2DScore, 199	PLSR. Anal, 236
PlotPCA3DScore, 200	Predict.class, 236
PlotPCA3DScoreImg, 200	Prepare Network Data 237
PlotPCABiplot, 201	PrepareNetworkData, 237
PlotPCALoading, 202	PreparePormPosult 238
PlotPCAPairSummary, 203	PreparePrenermPata 230
PlotPCAScree, 203	PreparePrenormData, 239
PlotPeaks2Paths, 204	PrepareQueryJson, 239
PlotPLS.Classification, 205	PrepareROCDataile 240
PlotPLS.Imp, 206	PrepareNormPate 241
PlotPLS.Permutation, 207	PrepareVennData, 241
PlotPLS2DScore, 207	RankFeatures, 241
PlotPLS3DScore, 209	rda2list, 242
PlotPLS3DScoreImg, 209	Read.BatchCSVdata, 242
PlotPLSLoading, 210	Read. MSspec, 243
PlotPLSPairSummary, 211	Read.PeakList, 243
PlotPowerProfile, 212	Read.PeakListData, 244
PlotPowerStat, 213	Read.TextData, 244
PlotProbView, 213	ReadIndData, 245
plotProfile, 214	ReadPairFile, 245
PlotQEA.MetSet, 215	RecordRCommand, 246
PlotQEA.Overview, 215	rectUnique, 246
PlotRF.Classify, 216	RegisterData, 247
PlotRF.Outlier, 217	RemoveCmpd, 247
PlotRF.VIP, 217	RemoveData, 248
PlotROC, 218	RemoveDuplicates, 248
PlotROC.LRmodel, 219	RemoveFile, 249
PlotROCTest, 220	RemoveFolder, 249
PlotRSVM.Classification, 221	RemoveGene, 250
PlotRSVM.Cmpd, 221	RemovedissingPercent, 250
PlotSAM.Cmpd, 222	ReplaceMin, 251
PlotSAM.FDR, 223	RerenderMetPAGraph, 251
PlotSampleNormSummary, 223	RF. Anal, 252
PlotSelectedFeature, 224	ROCPredSamplesTable, 252
PlotSigVar, 225	RSVM, 253
PlotSOM, 225	RSVM. Anal, 253
PlotSPLS2DScore, 226	nominat, 200
PlotSPLS3DScore, 227	SAM. Anal, 254
	- , -

SanityCheckData, 254	UpdateGraphSettings, 280
SanityCheckIndData, 255	UpdateInstrumentParameters, 28
SanityCheckMummichogData, 255	UpdateIntegPathwayAnalysis, 28
SaveTransformedData, 256	UpdateOPLS.Splot, 282
SearchByCompound, 256	UpdatePCA.Loading, 283
SearchByName, 257	UpdatePLS.Loading, 283
SearchMsetLibraries, 257	usr2png, 284
SearchNetDB, 258	, 6,
SelectMultiData, 258	Volcano.Anal, 284
SetAnalysisMode, 259	
SetAnnotationParam, 259	XSet2MSet, 285
SetCachexiaSetUsed, 260	
SetCandidate, 261	
SetClass, 261	
SetCurrentGroups, 262	
SetCurrentMsetLib, 262	
SetCustomData, 263	
SetDesignType, 263	
SetKEGG.PathLib, 264	
SetMetabolomeFilter, 264	
SetMummichogPval, 265	
SetMummichogPvalFromPercent, 265	
SetOrganism, 266	
SetPeakEnrichMethod, 266	
SetPeakFormat, 267	
SetPeakList.GroupValues, 267	
SetPeakParam, 268	
SetSMPDB.PathLib, 269	
Setup.AdductData, 270	
Setup.BiofluidType, 270	
Setup.ConcData, 271	
Setup.HMDBReferenceMetabolome, 271	
Setup.KEGGReferenceMetabolome, 272	
Setup.MapData, 272	
Setup.UserMsetLibData, 273	
SetupKEGGLinks, 273	
SetupMSdataMatrix, 274	
SetupSMPDBLinks, 274	
SOM. Anal, 275	
<pre>sparse.mint.block_iteration, 275</pre>	
splsda, 276	
SPLSR.Anal, 277	
SumNorm, 278	
template.match, 278	
Ttests.Anal, 279	
UnzipUploadedFile, 279	
UpdateData, 280	