

Package ‘autonomics’

July 9, 2025

Type Package

Title Unified Statistical Modeling of Omics Data

Version 1.16.0

Description This package unifies access to Statistical Modeling of Omics Data.

Across linear modeling engines (lm, lme, lmer, limma, and wilcoxon).

Across coding systems (treatment, difference, deviation, etc).

Across model formulae (with/without intercept, random effect, interaction or nesting).

Across omics platforms (microarray, rnaseq, msproteomics, affinity proteomics, metabolomics).

Across projection methods (pca, pls, sma, lda, spls, opls).

Across clustering methods (hclust, pam, cmeans).

It provides a fast enrichment analysis implementation.

And an intuitive contrastogram visualisation to summarize contrast effects in complex designs.

License GPL-3

Encoding UTF-8

LazyData true

VignetteBuilder knitr

biocViews Software, DataImport, Preprocessing, DimensionReduction,
PrincipalComponent, Regression, DifferentialExpression,
GeneSetEnrichment, Transcriptomics, Transcription,
GeneExpression, RNASeq, Microarray, Proteomics, Metabolomics,
MassSpectrometry,

BugReports

<https://gitlab.uni-marburg.de/fb20/ag-graumann/software/autonomics/issues>

RoxygenNote 7.3.1

Depends R (>= 4.0)

Imports abind, BiocFileCache, BiocGenerics, bit64, cluster,
codingMatrices, colorspace, data.table, dplyr, edgeR, ggforce,
ggplot2, ggrepel, graphics, grDevices, grid, gridExtra, limma,
magrittr, matrixStats, methods, MultiAssayExperiment, parallel,
RColorBrewer, rlang, R.utils, readxl, S4Vectors, scales, stats,
stringi, SummarizedExperiment, survival, tidyr, tidyselect,
tools, utils, vsn

Suggests affy, AnnotationDbi, AnnotationHub, apcluster, Biobase, BiocManager, BiocStyle, Biostrings, coin, diagram, DBI, e1071, ensemblDb, GenomicDataCommons, GenomicRanges, GEOquery, ggtext, hgu95av2.db, ICSNP, jsonlite, knitr, lme4, lmerTest, MASS, patchwork, mixOmics, mpm, nlme, OlinkAnalyze, org.Hs.eg.db, org.Mm.eg.db, pcaMethods, pheatmap, progeny, propagate, RCurl, RSQLite, remotes, rmarkdown, ropis, Rsubread, readODS, rtracklayer, statmod, testthat, UniProt.ws, writexl, XML

git_url <https://git.bioconductor.org/packages/autonomics>

git_branch RELEASE_3_21

git_last_commit a47f4df

git_last_commit_date 2025-04-15

Repository Bioconductor 3.21

Date/Publication 2025-07-09

Author Aditya Bhagwat [aut, cre],
Richard Cotton [ctb],
Shahina Hayat [ctb],
Laure Cougnaud [ctb],
Witold Szymanski [ctb],
Vanessa Beutgen [ctb],
Willem Ligtenberg [ctb],
Hinrich Goehlmann [ctb],
Karsten Suhre [ctb],
Johannes Graumann [aut, sad]

Maintainer Aditya Bhagwat <aditya.bhagwat@uni-marburg.de>

Contents

.coxph	7
.extract_p_features	7
.merge	10
.read_compounddiscoverer	11
.read_compounddiscoverer_masslist	11
.read_diann_precursors	12
.read_maxquant_proteingroups	14
.read_metabolon	15
.read_rectangles	17
.read_rnaseq_bams	19
.read_somascan	22
abstract_fit	24
add_adjusted_pvalues	25
add_assay_means	26
add_facetvars	27
add_opentargets_by_uniprot	28
add_psp	28

add_smiles	29
altenrich	30
analysis	31
analyze	32
annotate_compounddiscoverer	33
annotate_maxquant	34
annotate_uniprot_rest	35
assert_is_valid_sumexp	36
AUTONOMICS_DATASETS	36
bin	37
biplot	38
biplot_corrections	39
biplot_covariates	40
block2lme	42
center	43
code	44
coefs	46
collapsed_entrezg_to_symbol	47
COMPOUNDDISCOVERER_PATTERNS	48
contrast_coefs	48
contrast_subgroup_cols	49
counts	50
counts2cpm	51
counts2tpm	51
count_in	52
cpm	53
create_design	54
DATADIR	56
default_geom	57
default_sfile	58
default_subgroupvar	58
demultiplex	59
dequantify	60
dequantify_compounddiscoverer	61
DIMREDUN	62
download_gtf	62
download_mcclain21	63
dt2mat	64
enrichment	64
ens2org	66
entrezg_to_symbol	67
extract_rectangle	67
fcluster	69
fdata	70
fdr2p	72
filter_exprs_replicated_in_some_subgroup	72
filter_features	73
filter_medoid	74

filter_samples	75
fitcoefs	75
fits	76
FITSEP	77
fit_linmod	77
fix_xlgenes	81
flevels	82
fnames	83
formula2str	83
ftype	84
fvalues	85
fvars	85
genome_to_orgdb	86
group_by_level	87
guess_compounddiscoverer_quantity	88
guess_fitsep	88
guess_maxquant_quantity	89
guess_sep	90
has_multiple_levels	91
hdlproteins	93
impute	93
invert_subgroups	95
is_collapsed_subset	96
is_correlation_matrix	96
is_diann_report	97
is_fastadt	99
is_file	99
is_fraction	100
is_imputed	100
is_positive_number	101
is_scalar_subset	102
is_sig	103
is_valid_formula	104
keep_connected_blocks	105
keep_connected_features	105
keep_replicated_features	106
label2index	106
LINMODEGINES	107
list2mat	107
list_files	108
log2counts	108
log2cpm	109
log2diffs	110
log2proteins	111
log2sites	111
log2tpm	112
log2transform	113
logical2factor	114

make_alpha_palette	115
make_colors	116
make_volcano_dt	116
map_fvalues	117
matrix2sumexp	118
MAXQUANT_PATTERNS	118
mdsplot	119
merge_compounddiscoverer	120
merge_sample_excel	120
merge_sample_file	121
merge_sdata	122
message_df	124
modelvar	124
MSIGCOLLECTIONSHUMAN	130
MSIGDIR	131
nfactors	131
OPENTARGETSDIR	132
order_on_p	132
pca	134
pg_to_canonical	136
plot_coef_densities	137
plot_contrastogram	137
plot_contrast_venn	138
plot_data	139
plot_densities	140
plot_design	142
plot_detections	143
plot_exprs	145
plot_exprs_per_coef	148
plot_fit_summary	149
plot_heatmap	150
plot_joint_density	151
plot_matrix	152
plot_subgroup_points	152
plot_summary	154
plot_survival	154
plot_venn	156
plot_venn_heatmap	156
plot_violins	157
plot_volcano	159
PRECURSOR_QUANTITY	161
preprocess_rnaseq_counts	161
pull_columns	163
read_affymetrix	163
read_compounddiscoverer	164
read_fragpipe	166
read_maxquant_phosphosites	166
read_maxquant_proteingroups	168

read_msigt	170
read_olink	171
read_salmon	172
read_uniprot	172
reexports	173
reset_fit	174
rm_diann_contaminants	174
rm_missing_in_all_samples	175
rm_unmatched_samples	176
scaledlibsizes	177
scoremat	177
slevels	178
snames	179
split_samples	179
stri_any_regex	180
stri_detect_fixed_in_collapsed	181
subgroup_array	182
subtract_baseline	182
sumexplist_to_longdt	184
sumexp_to_tsv	185
sumexp_to_widedt	185
summarize_fit	186
SURVIVALENGINES	187
survival_example	188
svalues	190
svars	190
systematic_nas	191
tag_features	192
tag_hdlproteins	193
TAXON_TO_ORGNAME	193
TESTS	194
tpm	195
twofactor_sumexp	196
uncollapse	196
values	197
varlevels_dont_clash	198
venn_detects	198
weights	199
write_xl	200
X	201
zero_to_na	202

.coxph	<i>Fit onefeature survival</i>
--------	--------------------------------

Description

Fit onefeature survival

Usage

```
.coxph(timetoevent, event, expr)

.survdiff(timetoevent, event, expr)

.logrank(timetoevent, event, expr)
```

Arguments

timetoevent	numeric (time to event)
event	numeric (1=event, 0=not)
expr	numeric (.coxph) or twolevel-factor (.survdiff, .logrank_test)

Examples

```
# Prepare
  object <- survival_example()
  timetoevent <- object$timetoevent
  event <- object$event
  expr <- values(object)[1,]
  quantile <- factor(dplyr::ntile(expr, 2))
# Survival
  .coxph(timetoevent, event, expr)
  .survdiff(timetoevent, event, quantile)
  .logrank(timetoevent, event, quantile)
# Sumexp
  fit_survival(object)
```

.extract_p_features	<i>Extract coefficient features</i>
---------------------	-------------------------------------

Description

Extract coefficient features

Usage

```
.extract_p_features(  
  object,  
  coefs,  
  p = 0.05,  
  fit = fits(object),  
  combiner = "|",  
  features = NULL,  
  verbose = TRUE  
)  
  
.extract_fdr_features(  
  object,  
  coefs,  
  fdr = 0.05,  
  fit = fits(object),  
  combiner = "|",  
  features = NULL,  
  verbose = TRUE  
)  
  
.extract_effectsize_features(  
  object,  
  coefs,  
  effectsize = 1,  
  fit = fits(object),  
  combiner = "|",  
  features = NULL,  
  verbose = TRUE  
)  
  
.extract_sign_features(  
  object,  
  coefs,  
  sign,  
  fit = fits(object)[1],  
  combiner = "|",  
  features = NULL,  
  verbose = TRUE  
)  
  
.extract_n_features(  
  object,  
  coefs,  
  combiner = "|",  
  n,  
  fit = fits(object)[1],  
  features = NULL,
```

```

    verbose = TRUE
  )

  extract_coef_features(
    object,
    fit = fits(object)[1],
    coefs = autonomics::coefs(object, fit = fit),
    combiner = "|",
    decreasing = FALSE,
    p = 1,
    fdr = 1,
    effectsize = 0,
    sign = c(-1, +1),
    n = 4,
    features = NULL,
    verbose = TRUE
  )

```

Arguments

<code>object</code>	SummarizedXExperiment
<code>coefs</code>	subset of <code>coefs(object)</code>
<code>p</code>	p threshold
<code>fit</code>	subset of <code>fits(object)</code>
<code>combiner</code>	' ' or '&': how to combine multiple fits/coefs
<code>features</code>	features to include no matter what (character vector)
<code>verbose</code>	TRUE or FALSE
<code>fdr</code>	fdr threshold
<code>effectsize</code>	effectsize threshold
<code>sign</code>	effect sign
<code>n</code>	number of top features (Inf means all)
<code>decreasing</code>	TRUE or FALSE

Value

SummarizedExperiment

Examples

```

# Read and Fit
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
fdt(object) %<>% add_adjusted_pvalues('fdr')
# Single coef
object0 <- object

```

```

object %<>% .extract_p_features(      coefs = 't1-t0', p = 0.05)
object %<>% .extract_fdr_features(    coefs = 't1-t0', fdr = 0.05)
object %<>% .extract_effectsize_features(coefs = 't1-t0', effectsize = 1)
object %<>% .extract_sign_features(   coefs = 't1-t0', sign = -1)
object %<>% .extract_n_features(      coefs = 't1-t0', n = 1)
object <- object0
object %<>% extract_coef_features(
  coefs = 't1-t0', p = 0.05, fdr = 0.05, effectsize = 1, sign = -1, n = 1)
# Multiple coefs
object <- object0
object %<>% .extract_p_features(      coefs = c('t1-t0', 't2-t0'), p = 0.05)
object %<>% .extract_fdr_features(    coefs = c('t1-t0', 't2-t0'), fdr = 0.01)
object %<>% .extract_effectsize_features(coefs = c('t1-t0', 't2-t0'), effectsize = 1)
object %<>% .extract_sign_features(   coefs = c('t1-t0', 't2-t0'), sign = -1)
object %<>% .extract_n_features(      coefs = c('t1-t0', 't2-t0'), n = 1)
object <- object0
object %<>% extract_coef_features(
  coefs = c('t1-t0', 't2-t0'), p = 0.05, fdr = 0.01, effectsize = 1, sign = -1, n = 1)

```

*.merge**Clean Merge*

Description

Clean Merge

Usage

```
.merge(dt1, dt2, by)
```

Arguments

dt1	data.table
dt2	data.table
by	string

Examples

```

require(data.table)
dt1 <- data.table(feature_id = c('PG1', 'PG2'), gene = c('G1', 'G2'))
dt2 <- data.table(feature_id = c('PG1', 'PG2'), protein = c('P1', 'P2'))
dt1 %<>% .merge(dt2, by = 'feature_id')
dt1

```

.read_compounddiscoverer

Read compound discoverer files as-is

Description

Read compound discoverer files as-is

Usage

```
.read_compounddiscoverer(  
  file,  
  quantity = guess_compounddiscoverer_quantity(file),  
  colname_format = NULL,  
  mod_extract = NULL,  
  verbose = TRUE  
)
```

Arguments

file	compound discoverer file
quantity	string
colname_format	function to reformat column names
mod_extract	function to extract MS modi from sample names
verbose	TRUE / FALSE

Value

data.table

.read_compounddiscoverer_masslist

Read compound discoverer masslist files as-is

Description

Read compound discoverer masslist files as-is

Usage

```
.read_compounddiscoverer_masslist(file, verbose = TRUE)
```

Arguments

file	compound discoverer masslist file
verbose	TRUE / FALSE

Value

data.table

.read_diann_precursors

Read diann

Description

Read diann

Usage

```
.read_diann_precursors(file, Lib.PG.Q = 0.01, verbose = TRUE)
```

```
.read_diann_proteingroups(file, Lib.PG.Q = 0.01)
```

```
read_diann_proteingroups(
  file,
  Lib.PG.Q = 0.01,
  simplify_snames = TRUE,
  rm_contaminants = TRUE,
  impute = FALSE,
  plot = FALSE,
  pca = plot,
  pls = plot,
  fit = if (plot) "limma" else NULL,
  formula = as.formula("~ subgroup"),
  block = NULL,
  coefs = NULL,
  contrasts = NULL,
  palette = NULL,
  verbose = TRUE
)
```

```
read_diann(...)
```

Arguments

file	'report.tsv' file
Lib.PG.Q	Lib.PG.Q cutoff

<code>verbose</code>	TRUE or FALSE
<code>simplify_snames</code>	TRUE or FALSE: simplify (drop common parts in) samplenames ?
<code>rm_contaminants</code>	TRUE or FALSE: rm contaminants ?
<code>impute</code>	TRUE or FALSE: impute group-specific NA values ?
<code>plot</code>	TRUE or FALSE
<code>pca</code>	TRUE or FALSE: run pca ?
<code>pls</code>	TRUE or FALSE: run pls ?
<code>fit</code>	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
<code>formula</code>	model formula
<code>block</code>	model blockvar: string or NULL
<code>coefs</code>	model coefficients of interest: character vector or NULL
<code>contrasts</code>	coefficient contrasts of interest: character vector or NULL
<code>palette</code>	color palette: named string vector
<code>...</code>	used to maintain deprecated functions

Value

data.table or SummarizedExperiment

Examples

```
# Read
file <- download_data('dilution.report.tsv')
.read_diann_precursors(file)      # precursors longdt
.read_diann_proteingroups(file)   # proteingroups longdt
fdt(read_diann_proteingroups(file)) # proteingroups sumexp

# Compare
PR <- .read_diann_precursors(file)
PG <- .read_diann_proteingroups(file)
PG[intensity==top1] # matches      : 24975 (85%) proteingroups
PG[intensity!=top1] # doesnt match : 4531 (15%) proteingroups
RUN <- 'IPT_HeLa_1_DIAstd_Slot1-40_1_9997'
PR[uniprot=='Q96JP5;Q96JP5-2' & run == RUN, 1:6] # match:      8884 == 8884
PR[uniprot=='P36578' & run == RUN, 1:6] # no match: 650887 != 407978
PR[intensity != top1][feature_id == unique(feature_id)[1]][run == unique(run)[1]][1:2, 1:6]
PR[intensity != top1][feature_id == unique(feature_id)[2]][run == unique(run)[1]][1:2, 1:6]
PR[intensity != top1][feature_id == unique(feature_id)[3]][run == unique(run)[1]][1:3, 1:6]
```

```
.read_maxquant_proteingroups
```

Read proteingroups/phosphosites as-is

Description

Read proteingroups/phosphosites as-is

Usage

```
.read_maxquant_proteingroups(
  file,
  quantity = guess_maxquant_quantity(file),
  verbose = TRUE
)

.read_maxquant_phosphosites(
  file,
  profile,
  quantity = guess_maxquant_quantity(file),
  verbose = TRUE
)
```

Arguments

file	proteingroups / phosphosites file
quantity	string
verbose	TRUE / FALSE
profile	proteingroups file

Value

data.table

Examples

```
profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
prodt <- .read_maxquant_proteingroups(file = profile)
fosdt <- .read_maxquant_phosphosites( file = fosfile, profile = profile)
```

<i>.read_metabolon</i>	<i>Read metabolon xlsxfile</i>
------------------------	--------------------------------

Description

Read metabolon xlsxfile

Usage

```
.read_metabolon(
  file,
  sheet = "OrigScale",
  fidvar = "BIOCHEMICAL",
  sidvar = "(CLIENT_IDENTIFIER|Client ID)",
  sfile = NULL,
  by.x = "sample_id",
  by.y = NULL,
  groupvar = NULL,
  verbose = TRUE
)
```

```
read_metabolon(
  file,
  sheet = "OrigScale",
  fidvar = "BIOCHEMICAL",
  sidvar = "(CLIENT_IDENTIFIER|Client ID)",
  sfile = NULL,
  by.x = "sample_id",
  by.y = NULL,
  groupvar = NULL,
  fnamevar = "BIOCHEMICAL",
  kegg_pathways = FALSE,
  smiles = FALSE,
  impute = TRUE,
  plot = FALSE,
  pca = plot,
  pls = plot,
  label = "feature_id",
  fit = if (plot) "limma" else NULL,
  formula = as.formula("~ subgroup"),
  block = NULL,
  coefs = NULL,
  contrasts = NULL,
  palette = NULL,
  verbose = TRUE
)
```

Arguments

<code>file</code>	metabolon xlsx file
<code>sheet</code>	excel sheet (number or string)
<code>fidvar</code>	featureid var
<code>sidvar</code>	samplid var
<code>sfile</code>	sample file
<code>by.x</code>	'file' mergeby column
<code>by.y</code>	'sfile' mergeby column
<code>groupvar</code>	string
<code>verbose</code>	TRUE or FALSE
<code>fnamevar</code>	featurename fvar
<code>kegg_pathways</code>	TRUE or FALSE: add kegg pathways?
<code>smiles</code>	TRUE or FALSE: add smiles?
<code>impute</code>	TRUE or FALSE: impute group-specific NA values?
<code>plot</code>	TRUE or FALSE
<code>pca</code>	TRUE or FALSE
<code>pls</code>	TRUE or FALSE
<code>label</code>	fvar
<code>fit</code>	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
<code>formula</code>	model formula
<code>block</code>	model blockvar: string or NULL
<code>coefs</code>	model coefficients of interest: character vector or NULL
<code>contrasts</code>	coefficient contrasts of interest: character vector or NULL
<code>palette</code>	NULL or colorvector

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
read_metabolon(file, plot = TRUE, block = 'Subject')
```

<code>.read_rectangles</code>	<i>Read omics data from rectangular file</i>
-------------------------------	--

Description

Read omics data from rectangular file

Usage

```
.read_rectangles(  
  file,  
  sheet = 1,  
  fid_rows,  
  fid_cols,  
  sid_rows,  
  sid_cols,  
  expr_rows,  
  expr_cols,  
  fvar_rows = NULL,  
  fvar_cols = NULL,  
  svar_rows = NULL,  
  svar_cols = NULL,  
  fdata_rows = NULL,  
  fdata_cols = NULL,  
  sdata_rows = NULL,  
  sdata_cols = NULL,  
  transpose = FALSE,  
  verbose = TRUE  
)
```

```
read_rectangles(  
  file,  
  sheet = 1,  
  fid_rows,  
  fid_cols,  
  sid_rows,  
  sid_cols,  
  expr_rows,  
  expr_cols,  
  fvar_rows = NULL,  
  fvar_cols = NULL,  
  svar_rows = NULL,  
  svar_cols = NULL,  
  fdata_rows = NULL,  
  fdata_cols = NULL,  
  sdata_rows = NULL,  
  sdata_cols = NULL,
```

```

transpose = FALSE,
sfile = NULL,
sfileby = NULL,
subgroupvar = character(0),
verbose = TRUE
)

```

Arguments

file	string: name of text (txt, csv, tsv, adat) or excel (xls, xlsx) file
sheet	integer/string: only relevant for excel files
fid_rows	numeric vector: featureid rows
fid_cols	numeric vector: featureid cols
sid_rows	numeric vector: sampleid rows
sid_cols	numeric vector: sampleid cols
expr_rows	numeric vector: expr rows
expr_cols	numeric vector: expr cols
fvar_rows	numeric vector: fvar rows
fvar_cols	numeric vector: fvar cols
svar_rows	numeric vector: svar rows
svar_cols	numeric vector: svar cols
fdata_rows	numeric vector: fdata rows
fdata_cols	numeric vector: fdata cols
sdata_rows	numeric vector: sdata rows
sdata_cols	numeric vector: sdata cols
transpose	TRUE or FALSE (default)
verbose	TRUE (default) or FALSE
sfile	sample file
sfileby	sample file mergeby column
subgroupvar	subgroupvar in sfile

Value

SummarizedExperiment

Examples

```

# RNASEQ
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
read_rectangles( file, fid_rows = 2:25,    fid_cols = 2,
                  sid_rows = 1,           sid_cols = 5:28,
                  expr_rows = 2:25 ,     expr_cols = 5:28,
                  fvar_rows = 1,         fvar_cols = 1:4,
                  fdata_rows = 2:25 ,    fdata_cols = 1:4,    transpose = FALSE)

```

```

# LCMSMS PROTEINGROUPS
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
read_rectangles( file,
                  fid_rows = 2:21,    fid_cols = 383,
                  sid_rows = 1,       sid_cols = seq(124, 316, by = 6),
                  expr_rows = 2:21,   expr_cols = seq(124, 316, by = 6),
                  fvar_rows = 1,      fvar_cols = c(2, 6, 7, 383),
                  fdata_rows = 2:21,  fdata_cols = c(2, 6, 7, 383),
                  transpose = FALSE )

# SOMASCAN
file <- system.file('extdata/atkin.somascan.adat', package = 'autonomics')
read_rectangles(file, fid_rows = 30,    fid_cols = 23:42,
                  sid_rows = 42:108,    sid_cols = 4,
                  expr_rows = 42:108,   expr_cols = 23:42,
                  fvar_rows = 28:40,     fvar_cols = 22,
                  svar_rows = 41,        svar_cols = 1:21,
                  fdata_rows = 28:40,    fdata_cols = 23:42,
                  sdata_rows = 42:108,   sdata_cols = 1:21, transpose = TRUE)

# METABOLON
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
read_rectangles(file, sheet = 2,
                  fid_rows = 11:30,     fid_cols = 2,
                  sid_rows = 4,         sid_cols = 15:81,
                  expr_rows = 11:30,    expr_cols = 15:81,
                  fvar_rows = 10,       fvar_cols = 1:14,
                  svar_rows = 1:10,     svar_cols = 14,
                  fdata_rows = 11:30,   fdata_cols = 1:14,
                  sdata_rows = 1:10,    sdata_cols = 15:81,
                  transpose = FALSE )

```

<i>.read_rnaseq_bams</i>	<i>Read rnaseq counts/bams</i>
--------------------------	--------------------------------

Description

Read rnaseq counts/bams

Usage

```

.read_rnaseq_bams(
  dir,
  paired,
  genome,
  nthreads = detectCores(),
  sfile = NULL,
  by.y = NULL,
  ensdb = NULL,
  verbose = TRUE
)

```

```
.read_rnaseq_counts(  
  file,  
  fid_col = 1,  
  sfile = NULL,  
  by.y = NULL,  
  ensdb = NULL,  
  verbose = TRUE  
)  
  
read_rnaseq_bams(  
  dir,  
  paired,  
  genome,  
  nthreads = detectCores(),  
  sfile = NULL,  
  by.y = NULL,  
  block = NULL,  
  formula = as.formula("~ subgroup"),  
  min_count = 10,  
  pseudo = 0.5,  
  ensdb = NULL,  
  tpm = FALSE,  
  cpm = TRUE,  
  log2 = TRUE,  
  plot = FALSE,  
  label = "feature_id",  
  pca = plot,  
  pls = plot,  
  fit = if (plot) "limma" else NULL,  
  voom = cpm,  
  coefs = NULL,  
  contrasts = NULL,  
  palette = NULL,  
  verbose = TRUE  
)  
  
read_rnaseq_counts(  
  file,  
  fid_col = 1,  
  sfile = NULL,  
  by.y = NULL,  
  formula = as.formula("~ subgroup"),  
  block = NULL,  
  min_count = 10,  
  pseudo = 0.5,  
  tpm = FALSE,  
  ensdb = NULL,
```

```

    cpm = !tpm,
    log2 = TRUE,
    plot = FALSE,
    label = "feature_id",
    pca = plot,
    pls = plot,
    fit = if (plot) "limma" else NULL,
    voom = cpm,
    coefs = NULL,
    contrasts = NULL,
    palette = NULL,
    verbose = TRUE
)

```

Arguments

<code>dir</code>	<code>read_rnaseq_bams</code> : bam/sam dir
<code>paired</code>	<code>read_rnaseq_bams</code> : TRUE/FALSE : paired end reads ?
<code>genome</code>	<code>read_rnaseq_bams</code> : 'mm10', 'hg38', etc. or GTF file
<code>nthreads</code>	<code>read_rnaseq_bams</code> : nthreads used by Rsubread::featureCounts()
<code>sfile</code>	sample file
<code>by.y</code>	sample file mergeby column
<code>ensdb</code>	EnsDb with genesizes : e.g. AnnotationHub::AnnotationHub[['AH64923']]
<code>verbose</code>	TRUE or FALSE: message?
<code>file</code>	count file
<code>fid_col</code>	featureid column (number or string)
<code>block</code>	model blockvar: string or NULL
<code>formula</code>	model formula
<code>min_count</code>	min feature count required in some samples
<code>pseudo</code>	pseudocount added to prevent -Inf log2 values
<code>tpm</code>	TRUE or FALSE : add tpm to assays (counts / libsize / genelength) ?
<code>cpm</code>	TRUE or FALSE: add cpm to assays (counts / effectivelibsize) ?
<code>log2</code>	TRUE or FALSE: log2 transform ?
<code>plot</code>	TRUE or FALSE: plot?
<code>label</code>	fvar
<code>pca</code>	TRUE or FALSE: perform and plot pca?
<code>pls</code>	TRUE or FALSE: run pls ?
<code>fit</code>	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
<code>voom</code>	model weights to be computed? TRUE/FALSE
<code>coefs</code>	model coefficients of interest: string vector or NULL
<code>contrasts</code>	model coefficient contrasts of interest: string vector or NULL
<code>palette</code>	color palette : named string vector

Value

SummarizedExperiment

Author(s)

Aditya Bhagwat, Shahina Hayat

Examples

```
# read_rnaseq_bams
if (requireNamespace('Rsubread')){
  dir <- download_data('billing16.bam.zip')
  object <- read_rnaseq_bams(dir, paired = TRUE, genome = 'hg38')
  object <- read_rnaseq_bams(dir, paired = TRUE, genome = 'hg38', plot = TRUE)
}

# read_rnaseq_counts
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file, fit = 'limma', coefs = 'E15-E00')
object <- read_rnaseq_counts(file, fit = 'limma', coefs = 'E15-E00', voom = FALSE)
object <- read_rnaseq_counts(file, fit = 'limma', coefs = 'E15-E00', voom = FALSE, cpm = FALSE)
object <- read_rnaseq_counts(file, fit = 'limma', coefs = 'E15-E00', voom = FALSE, cpm = FALSE,
                             log2 = FALSE)
object <- read_rnaseq_counts(file, plot = TRUE)

# read_rnaseq_counts(tpm = TRUE)
## Not run:
ah <- AnnotationHub::AnnotationHub()
ensdb <- ah[['AH64923']]
object <- read_rnaseq_counts(file, fit = 'limma', coefs = 'E02-E00', tpm = TRUE, ensdb = ensdb)

## End(Not run)
```

.read_somascan	<i>Read somascan adatfile</i>
----------------	-------------------------------

Description

Read somascan adatfile

Usage

```
.read_somascan(
  file,
  fidvar = "Target",
  sidvar = "SampleId",
  sfile = NULL,
  by.x = NULL,
  by.y = NULL,
  groupvar = "SampleGroup",
```

```

    verbose = TRUE
  )

read_somascan(
  file,
  fidvar = "Target",
  sidvar = "SampleId",
  sfile = NULL,
  by.x = NULL,
  by.y = NULL,
  groupvar = "SampleGroup",
  fname_var = "EntrezGeneSymbol",
  sample_type = "Sample",
  feature_type = "Protein",
  sample_quality = c("FLAG", "PASS"),
  feature_quality = c("FLAG", "PASS"),
  rm_na_svars = FALSE,
  rm_single_value_svars = FALSE,
  plot = FALSE,
  label = "feature_id",
  pca = plot,
  pls = plot,
  fit = if (plot) "limma" else NULL,
  formula = as.formula(sprintf("~ %s", groupvar)),
  block = NULL,
  coefs = NULL,
  contrasts = NULL,
  palette = NULL,
  verbose = TRUE
)

```

Arguments

<code>file</code>	somascan (adat) file
<code>fidvar</code>	featureid var
<code>sidvar</code>	sampleid var
<code>sfile</code>	sample file
<code>by.x</code>	'file' mergeby column
<code>by.y</code>	'sfile' mergeby column
<code>groupvar</code>	string
<code>verbose</code>	TRUE or FALSE: message?
<code>fname_var</code>	featurename var: string
<code>sample_type</code>	subset of c('Sample', 'QC', 'Buffer', 'Calibrator')
<code>feature_type</code>	subset of c('Protein', 'Hybridization Control Elution', 'Rat Protein')
<code>sample_quality</code>	subset of c('PASS', 'FLAG', 'FAIL')

feature_quality	subset of c('PASS', 'FLAG', 'FAIL')
rm_na_svars	TRUE or FALSE: rm NA svars?
rm_single_value_svars	TRUE or FALSE: rm single value svars?
plot	TRUE or FALSE: plot ?
label	fvar
pca	TRUE or FALSE: run pca?
pls	TRUE or FALSE: run pls?
fit	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
formula	model formula
block	model blockvar
coefs	model coefficients of interest: character vector or NULL
contrasts	coefficient contrasts of interest: character vector or NULL
palette	character vector or NULL

Value

Summarizedexperiment

Examples

```
file <- system.file('extdata/atkin.somascan.adat', package = 'autonomics')
read_somascan(file, plot = TRUE, block = 'Subject')
```

abstract_fit

Abstract model fit

Description

Abstract model fit

Usage

```
abstract_fit(
  object,
  sep = guess_fitsep(fdt(object)),
  fit = fits(object),
  coef = coefs(object, fit = fit),
  significancevar = "p",
  significance = 0.05
)
```

Arguments

object	SummarizedExperiment
sep	string
fit	character vector
coef	character vector
significancevar	'p' or 'fdr'
significance	fraction : pvalue cutoff

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma', coef = 't3-t0')
fdt(object)
fdt(abstract_fit(object))
```

add_adjusted_pvalues *Add adjusted pvalues*

Description

Add adjusted pvalues

Usage

```
add_adjusted_pvalues(object, ...)
```

```
## S3 method for class 'data.table'
add_adjusted_pvalues(
  object,
  method = "fdr",
  fit = fits(object),
  coefs = autonomics::coefs(object, fit = fit),
  verbose = TRUE,
  ...
)
```

```
## S3 method for class 'SummarizedExperiment'
add_adjusted_pvalues(
  object,
  method = "fdr",
  fit = fits(object),
```

```
  coefs = autonomics::coefs(object, fit = fit),
  verbose = TRUE,
  ...
)
```

Arguments

object	SummarizedExperiment or (feature) data.table
...	for s3 dispatch
method	'fdr', 'bonferroni', ... (see 'p.adjust.methods')
fit	'limma', 'lm', 'lme', 'lmer'
coefs	coefficient (string)
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fdt(object) %<>% extract(, 1:2)
object %<>% fit_limma()
object %<>% extract(order(fdt(.)$`p~Adult-X30dpt~limma`), )
  fdt(object)
(fdt(object) %<>% add_adjusted_pvalues('fdr'))
(fdt(object) %<>% add_adjusted_pvalues('fdr'))      # smart enough not to add second column
(fdt(object) %>% add_adjusted_pvalues('bonferroni'))
```

add_assay_means	<i>Add assay means</i>
-----------------	------------------------

Description

Add assay means

Usage

```
add_assay_means(object, assay = assayNames(object)[1], bin = TRUE)
```

Arguments

object	SummarizedExperiment or NULL
assay	string
bin	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fdt(object) %<>% extract(, 1:2)
fdt(object)
object %<>% add_assay_means(SummarizedExperiment::assayNames(.))
fdt(object)
```

add_facetvars*Add facetvars*

Description

Add facetvars

Usage

```
add_facetvars(
  object,
  fit = fits(object)[1],
  coefs = autonomics::coefs(object, fit = fit)
)
```

Arguments

object	SummarizedExperiment
fit	string
coefs	string vector

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma')
object %<>% add_adjusted_pvalues()
fdt(object)
fdt(add_facetvars(object))
```

```
add_opentargets_by_uniprot
```

Add opentargets annotations

Description

Add opentargets annotations

Usage

```
add_opentargets_by_uniprot(
  object,
  cols = c("genesymbol", "genename", "function"),
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
cols	character vector
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
object %<>% add_opentargets_by_uniprot()
```

```
add_psp
```

Add psp

Description

Add PhosphoSitePlus literature counts

Usage

```
add_psp(
  object,
  pspfile = file.path(R_user_dir("autonomics", "cache"), "phosphositeplus",
    "Phosphorylation_site_dataset.gz")
)
```

Arguments

object	SummarizedExperiment
pspfile	phosphositeplus file

Details

Go to www.phosphosite.org
Register and Login.
Download [Phosphorylation_site_dataset.gz](#).
Save into: `file.path(R_user_dir('autonomics','cache'),'phosphositeplus')`

Value

SummarizedExperiment

Examples

```
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_phosphosites(fosfile = fosfile, profile = profile)
fdt(object)
object %<>% add_psp()
fdt(object)
```

add_smiles

Add smiles

Description

Add smiles

Usage

```
add_smiles(object)
```

Arguments

object	character/factor vector with pubchem ids
--------	--

Value

character/factor vector

References

<https://pubchemdocs.ncbi.nlm.nih.gov/pug-rest-tutorial>

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
# add_smiles(object[1:10, ]) # seems down
```

altenrich

Alternative Enrichment Analysis

Description

Alternative Enrichment Analysis

Usage

```
altenrich(
  object,
  pathwaydt,
  genevar = "gene",
  genesep = "[ ,;]",
  coef = autonomics::coefs(object)[1],
  fit = fits(object)[1],
  significancevar = "p",
  significance = 0.05,
  effectsizesize = 0,
  n = 3,
  genes = FALSE,
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
pathwaydt	data.table, e.g. read_msigt
genevar	gene fvar
genesep	string or NULL
coef	string in <code>coefs(object)</code>
fit	'limma', 'lm', 'lme', 'lmer', 'wilcoxon'
significancevar	'p' or 'fdr'
significance	significance cutoff
effectsizesize	effectsizesize cutoff
n	no of detected genes required (for geneset to be examined)
genes	whether to record genes
verbose	whether to msg

Details

This is an alternative enrichent analysis implementation. It is more modular: uses four times .enrichment(VERBOSE)? as backend. But also four times slower than enrichment, so not recommended. It is retaind for testing purposes.

This alternative enrichment implementation

See Also

[enrichment()]

analysis	<i>Get/set analysis</i>
----------	-------------------------

Description

Get/set analysis

Usage

```
analysis(object)

## S4 method for signature 'SummarizedExperiment'
analysis(object)

analysis(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,list'
analysis(object) <- value
```

Arguments

object	SummarizedExperiment
value	list

Value

analysis details (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
analysis(object)
```

analyze

*Analyze***Description**

Analyze

Usage

```
analyze(
  object,
  pca = TRUE,
  pls = TRUE,
  fit = "limma",
  formula = ~subgroup,
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  contrasts = NULL,
  coefs = contrast_coefs(object, formula = formula, drop = drop, codingfun = codingfun),
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  plot = pca & !is.null(fit),
  label = "feature_id",
  palette = NULL,
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
pca	TRUE / FALSE: perform pca ?
pls	TRUE / FALSE: perform pls ?
fit	linmod engine: 'limma', 'lm', 'lme(r)', 'lmer', 'wilcoxon'
formula	model formula
drop	TRUE / FALSE : drop varname in designmat ?
codingfun	factor coding function • contr.treatment: intercept = y0, coefi = yi - y0 • contr.treatment.explicit: intercept = y0, coefi = yi - y0 • code_control: intercept = ymean, coefi = yi - y0 • contr.diff: intercept = y0, coefi = yi - y(i-1) • code_diff: intercept = ymean, coefi = yi - y(i-1) • code_diff_forward: intercept = ymean, coefi = yi - y(i+) • code_deviation: intercept = ymean, coefi = yi - ymean (drop last) • code_deviation_first: intercept = ymean, coefi = yi - ymean (drop first)

	- code_helmert: intercept = ymean, coefi = yi - mean(y0:(yi-1)) - code_helmert_forward: intercept = ymean, coefi = yi - mean(y(i+1):yp)
contrasts	model coefficient contrasts of interest: string vector or NULL
coefs	model coefficients of interest: string vector or NULL
block	model blockvar
weightvar	NULL or name of weight matrix in assays(object)
plot	TRUE / FALSE
label	fvar
palette	NULL or colorvector
verbose	TRUE / FALSE: message?

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% analyze()
```

annotate_compounddiscoverer

Read compound discoverer output

Description

Read compound discoverer output

Usage

```
annotate_compounddiscoverer(
  x,
  dir = getwd(),
  files = list.files(path = dir, pattern = ".*masslist.*\\.xlsx$", ignore.case = TRUE,
    full.names = TRUE),
  verbose = TRUE
)
```

Arguments

x	SummarizedExperiment (read_compounddiscoverer)
dir	compound discoverer output directory
files	compound discoverer masslist files
verbose	TRUE or FALSE : message ?

Value

SummarizedExperiment

annotate_maxquant	<i>Annotate maxquant</i>
-------------------	--------------------------

Description

Annotate maxquant data.table

Usage

```
annotate_maxquant(  
  dt,  
  uniprothdrs,  
  contaminanthdrs,  
  maxquanthdrs,  
  restapi = FALSE,  
  verbose = TRUE  
)
```

Arguments

- dt data.table : output of read_maxquant_(proteingroups|phosphosites)
- uniprothdrs data.table : output of read_uniprot dt
- contaminanthdrs data.table : output of read_uniprot dt
- maxquanthdrs data.table : output of read_uniprot dt
- restapi logical(1) : use uniprot restapi to complete missing annotations ?
- verbose logical(1) : message ?

Details

Uncollapse, annotate, curate, recollapse, name

Value

data.table

Examples

```
# Fukuda 2020: contaminants + maxquanthdrs
#-----
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
dt <- .read_maxquant_proteingroups(file)
dt[, 1:2]
uniprothdrs <- NULL
contaminanthdrs <- read_contaminantdt()
maxquanthdrs <- parse_maxquant_hdrs(dt$`Fasta headers`); dt$`Fasta headers` <- NULL
dt %<>% annotate_maxquant(uniprothdrs, contaminanthdrs, maxquanthdrs)
dt[, 1:9]
dt[reverse== '+', 1:9]
dt[contaminant== '+', 1:9]

# Billing 2019: uniprothdrs + contaminants + maxquanthdrs
#-----
profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
upfile <- system.file('extdata/uniprot_hsa_20140515.fasta', package = 'autonomics')
prodt <- .read_maxquant_proteingroups(profile); prodt[, 1:2]
fosdt <- .read_maxquant_phosphosites(fosfile, profile); fosdt[, 1:3]
uniprothdrs <- read_uniprot(dt)
contaminanthdrs <- read_contaminantdt()
maxquanthdrs <- parse_maxquant_hdrs(prodt$`Fasta headers`)
annotate_maxquant(prodt, uniprothdrs, contaminanthdrs, maxquanthdrs)[, 1:8]
annotate_maxquant(fosdt, uniprothdrs, contaminanthdrs, maxquanthdrs)[, 1:8]
```

annotate_uniprot_rest *Annotate uniprot/ensp*

Description

Annotate uniprot/ensp

Usage

```
annotate_uniprot_rest(x, columns = UNIPROTCOLS, verbose = TRUE)
```

Arguments

x	character vector
columns	character vector
verbose	TRUE or FALSE

Value

data.table(dbid, uniprot, reviewed, protein, gene, canonical, isoform, fragment, existence, organism, full)

Examples

```

annotate_uniprot_rest( x = c('P00761', 'Q32MB2') )
annotate_uniprot_rest( x = c('ENSBTAP00000006074', 'ENSP00000377550') )

```

```
assert_is_valid_sumexp
```

Assert that x is a valid SummarizedExperiment

Description

Assert that x is a valid SummarizedExperiment

Usage

```
assert_is_valid_sumexp(x, .xname = get_name_in_parent(x))
```

Arguments

x	SummarizedExperiment
.xname	see get_name_in_parent

Value

TRUE or FALSE

Examples

```

# VALID
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
x <- read_metabolon(file)
assert_is_valid_sumexp(x)
# NOT VALID
rownames(SummarizedExperiment::colData(x)) <- NULL
# assert_is_valid_sumexp(x)

```

AUTONOMICS_DATASETS *Data used in examples/vignette/tests/longtests*

Description

Data used in examples/vignette/tests/longtests

Usage

```
AUTONOMICS_DATASETS
```

Format

An object of class character of length 19.

Examples

```
AUTONOMICS_DATASETS
```

bin	<i>Bin continuous variable</i>
-----	--------------------------------

Description

Bin continuous variable

Usage

```
bin(object, ...)

## S3 method for class 'logical'
bin(object, ...)

## S3 method for class 'character'
bin(object, ...)

## S3 method for class 'factor'
bin(object, ...)

## S3 method for class 'numeric'
bin(object, probs = c(0, 0.33, 0.66, 1), ...)

## S3 method for class 'SummarizedExperiment'
bin(object, fvar, probs = c(0, 0.33, 0.66, 1), ...)
```

Arguments

object	numeric or SummarizedExperiment
...	(S3 dispatch)
probs	numeric
fvar	string or NULL

Value

factor vector

Examples

```
# Numeric vector
object <- rnorm(10, 5, 1)
bin(object)

# SummarizedExperiment
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
fdt(object <- read_maxquant_proteingroups(file))
fdt(bin(object, 'pepcounts'))
```

biplot	<i>Biplot</i>
--------	---------------

Description

Biplot

Usage

```
biplot(
  object,
  method = biplot_methods(object)[1],
  by = biplot_by(object, method)[1],
  dims = biplot_dims(object, method, by)[1:2],
  color = if (method %in% DIMREDSUPER) by else "subgroup",
  shape = NULL,
  size = NULL,
  alpha = NULL,
  group = NULL,
  linetype = NULL,
  label = NULL,
  feature_label = "feature_id",
  fixed = list(shape = 15, size = 3),
  nx = 0,
  ny = 0,
  colorpalette = make_svar_palette(object, color),
  alphapalette = make_alpha_palette(object, alpha),
  title = paste0(method, guess_fitsep(fdt(object)), by),
  theme = ggplot2::theme(plot.title = element_text(hjust = 0.5), panel.grid =
    element_blank())
)
```

Arguments

object	SummarizedExperiment
method	'pca', 'pls', 'lda', 'spl', 'opls', 'sma'
by	svar

dims	numeric vector: e.g. 1:2
color	svar
shape	svar
size	svar
alpha	svar
group	svar
linetype	svar
label	svar
feature_label	fvar
fixed	fixed plot aesthetics
nx	number of x features to plot
ny	number of y features to plot
colorpalette	character vector
alphapalette	character vector
title	string
theme	ggplot2::theme output

Value

ggplot object

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% pca(ndim = 4)
object %<>% pls(ndim = 4)
biplot(object)
biplot(object, nx = 1)
biplot(object, dims = 3:4, nx = 1)
biplot(object, method = 'pls')
biplot(object, method = 'pls', dims = 3:4)
biplot(object, method = 'pls', dims = 3:4, group = 'Subject')

```

biplot_corrections	<i>Biplot batch corrections</i>
--------------------	---------------------------------

Description

Biplot batch corrections

Usage

```
biplot_corrections(  
  object,  
  method = "pca",  
  by = "sample_id",  
  color = "subgroup",  
  covariates = character(0),  
  varcols = ceiling(sqrt(1 + length(covariates))),  
  plot = TRUE  
)
```

Arguments

object	SummarizedExperiment
method	'pca', 'pls', 'lda', or 'sma'
by	svar
color	variable mapped to color (symbol)
covariates	covariates to be batch-corrected
varcols	number of covariate columns
plot	TRUE/FALSE: plot?

Value

grid object

See Also

biplot_covariates

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')  
object <- read_metabolon(file, pca = TRUE, plot = FALSE)  
biplot_corrections(object, color = 'subgroup', covariates = c('Sex', 'Diabetes', 'Subject', 'Time'))
```

biplot_covariates	<i>Biplot covariates</i>
-------------------	--------------------------

Description

Biplot covariates

Usage

```
biplot_covariates(
  object,
  method = "pca",
  by = "sample_id",
  block = NULL,
  covariates = "subgroup",
  ndim = 6,
  dimcols = 1,
  varcols = length(covariates),
  plot = TRUE
)
```

Arguments

object	SummarizedExperiment
method	'pca', 'pls', 'lda', or 'sma'
by	svar
block	svar
covariates	covariates: mapped to color or batch-corrected
ndim	number of dimensions to plot
dimcols	number of dimension columns
varcols	number of covariate columns
plot	TRUE or FALSE: whether to plot

Value

ggplot object

See Also

biplot_corrections

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, pca = TRUE)
biplot_covariates(object, covariates = 'subgroup', ndim = 12, dimcols = 3)
biplot_covariates(object, covariates = c('Sex', 'Diabetes', 'Subject', 'Time'))
biplot_covariates(object, covariates = c('Sex', 'Diabetes', 'Subject', 'Time'), ndim = 2)
biplot_covariates(object, covariates = c('subgroup'), dimcols = 3)
```

block2lme	<i>Put block in lme-compatible format</i>
-----------	---

Description

Put block in lme-compatible format

Usage

```
block2lme(block, ...)

## S3 method for class 'list'
block2lme(block, verbose = TRUE, ...)

## S3 method for class 'formula'
block2lme(block, verbose = TRUE, ...)

## S3 method for class 'character'
block2lme(block, verbose = TRUE, ...)

formula2lmer(formula, block)

formula2lm(formula, block)

block_vars(formula)
```

Arguments

block	block: character vector or formula
...	required for s3 dispatch
verbose	TRUE or FALSE
formula	formula

Examples

```
# lme: ensure lme-compatible block specification
block2lme( block = list(subject = ~1, batch = ~1))
block2lme( block = ~1|subject)
block2lme( block = c('subject', 'batch'))

# lm: integrate block into formula as random effect
formula2lm( formula = ~ subgroup, block = c('subject', 'batch') )

# lmer: integrate block into formula as fixed effect
formula2lmer( formula = ~ subgroup, block = c('subject', 'batch') )
formula2lmer( formula = ~ subgroup + (1|subject) + (1|batch) )
```

center	<i>Center samples</i>
--------	-----------------------

Description

Center samples

Usage

```
center(
  object,
  selector = rep(TRUE, nrow(object)) == TRUE,
  fun = "median",
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
selector	logical vector (length = nrow(object))
fun	aggregation function (string)
verbose	TRUE/FALSE

Value

SummarizedExperiment

Examples

```
require(matrixStats)
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fdt(object)$housekeeping <- FALSE
fdt(object)$housekeeping[order(rowVars(values(object)))[1:5]] <- TRUE
values(object)[, object$subgroup=='Adult'] %<>% magrittr::add(5)
plot_sample_densities(object)
plot_sample_densities(center(object))
plot_sample_densities(center(object, housekeeping))
```

code	<i>Contrast Code Factor</i>
------	-----------------------------

Description

Contrast Code Factor for General Linear Model

Usage

```
code(object, ...)

## S3 method for class 'factor'
code(object, codingfun, verbose = TRUE, ...)

## S3 method for class 'data.table'
code(object, codingfun, vars = names(object), verbose = TRUE, ...)

contr.treatment.explicit(n)

code_control(n)

contr.diff(n)

code_diff(n)

code_diff_forward(n)

code_deviation(n)

code_deviation_first(n)

code_helmert(n)

code_helmert_forward(n)
```

Arguments

object	factor vector
...	used for s3 dispatch
codingfun	factor coding function

- `contr.treatment`: intercept = y_0 , $\text{coef}_i = y_i - y_0$
- `contr.treatment.explicit`: intercept = y_0 , $\text{coef}_i = y_i - y_0$
- `code_control`: intercept = y_{mean} , $\text{coef}_i = y_i - y_0$
- `contr.diff`: intercept = y_0 , $\text{coef}_i = y_i - y_{(i-1)}$
- `code_diff`: intercept = y_{mean} , $\text{coef}_i = y_i - y_{(i-1)}$

	• <code>code_diff_forward</code>: intercept = ymean, coefi = $y_i - y_{i+}$ • <code>code_deviation</code>: intercept = ymean, coefi = $y_i - \text{ymean}$ (drop last) • <code>code_deviation_first</code>: intercept = ymean, coefi = $y_i - \text{ymean}$ (drop first) • <code>code_helmert</code>: intercept = ymean, coefi = $y_i - \text{mean}(y_0:(y_i-1))$ • <code>code_helmert_forward</code>: intercept = ymean, coefi = $y_i - \text{mean}(y_{i+1}:y_p)$
<code>verbose</code>	TRUE or FALSE
<code>vars</code>	svars
<code>n</code>	character vector

Details

A General Linear Model contains:

- * An Intercept Coefficient: expressing some form of sample average
- * For each numeric variable: a slope coefficient
- * For each k-leveled factor: (k-1) Contrast Coefficients.

The interpretation of (intercept and contrast) coefficients depends on the contrast coding function used. Several contrast coding functions are available in 'stats' and 'codingMatrices' But their (function and coefficient) namings are a bit confusing and unsystematic. Instead, the functions below offer an intuitive interface (to the otherwise powerful stats/codingMatrices packages). The names of these functions reflect the contrast coding used (treatment, backward, sum, or helmert contrasts). They also reflect the intercept interpretation (either first factor's first level or grand mean). They all produce intuitive coefficient names (e.g. 't1-t0' rather than just 't1'). They all have unit scaling (a coefficient of 1 means a backward of 1).

Value

(explicitly coded) factor vector

Examples

```
# Coding functions
x <- factor(paste0('t', 0:3))
xlevels <- levels(x)
contr.treatment(      xlevels)
contr.treatment.explicit(xlevels)
contr.diff(           xlevels)
code_control(         xlevels)
code_diff(            xlevels)
code_diff_forward(    xlevels)
code_deviation(       xlevels)
code_deviation_first( xlevels)
code_helmert(         xlevels)
code_helmert_forward( xlevels)

# Code
x %<>% code(contr.treatment)
x %<>% code(contr.treatment.explicit)
x %<>% code(contr.diff)
x %<>% code(code_control)
```

```

x %<>% code(code_diff)
x %<>% code(code_diff_forward)
x %<>% code(code_deviation)
x %<>% code(code_deviation_first)
x %<>% code(code_helmert)
x %<>% code(code_helmert_forward)

# Model
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma(codingfun = contr.treatment) # default
object %<>% fit_limma(codingfun = contr.treatment.explicit)
object %<>% fit_limma(codingfun = contr.diff)
object %<>% fit_limma(codingfun = code_control)
object %<>% fit_limma(codingfun = code_diff)
object %<>% fit_limma(codingfun = code_diff_forward)
object %<>% fit_limma(codingfun = code_deviation)
object %<>% fit_limma(codingfun = code_deviation_first)
object %<>% fit_limma(codingfun = code_helmert)
object %<>% fit_limma(codingfun = code_helmert_forward)

```

coefs

Get coefs

Description

Get coefs

Usage

```
coefs(object, ...)
```

```
## S3 method for class 'factor'
```

```
coefs(object, intercept = FALSE, ...)
```

```
## S3 method for class 'data.table'
```

```
coefs(object, fit = fits(object), intercept = FALSE, ...)
```

```
## S3 method for class 'SummarizedExperiment'
```

```
coefs(object, fit = fits(object), intercept = FALSE, ...)
```

Arguments

object	factor, data.table, SummarizedExperiment
...	required for s3 dispatch
intercept	TRUE or FALSE : whether to include the intercept
fit	'limma', 'lm', 'lme', 'lmer', 'wilcoxon'

Value

character vector

Examples

```
# Factor
x <- factor(c('A', 'B', 'C'))
coefs(x)
coefs(code(x, contr.treatment.explicit))
coefs(code(x, code_control))

# SummarizedExperiment
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma')
coefs(object)
coefs(object, intercept = TRUE)
```

collapsed_entrezg_to_symbol

Collapsed entrezg to genesymbol

Description

Collapsed entrezg to genesymbol

Usage

```
collapsed_entrezg_to_symbol(x, sep, orgdb)
```

Arguments

x	charactervector
sep	string
orgdb	OrgDb

Value

character vector

Examples

```
if (requireNamespace('org.Hs.eg.db', quiet = TRUE)){
  x <- c('7448/3818/727', '5034/9601/64374')
  orgdb <- org.Hs.eg.db::org.Hs.eg.db
  collapsed_entrezg_to_symbol(x, sep = '/', orgdb = orgdb)
}
```

COMPOUNDDISCOVERER_PATTERNS
<i>compound discoverer quantity patterns</i>

Description

compound discoverer quantity patterns

Usage

COMPOUNDDISCOVERER_PATTERNS

Format

An object of class character of length 2.

Examples

COMPOUNDDISCOVERER_PATTERNS

contrast_coefs	<i>Get model coefs</i>
----------------	------------------------

Description

Get model coefs

Usage

```
contrast_coefs(  
  object,  
  formula = default_formula(object),  
  drop = varlevels_dont_clash(object, all.vars(formula)),  
  codingfun = contr.treatment.explicit,  
  design = create_design(object, formula = formula, drop = drop, codingfun = codingfun,  
    verbose = FALSE)  
)  
  
model_coefs(  
  object,  
  formula = default_formula(object),  
  drop = varlevels_dont_clash(object, all.vars(formula)),  
  codingfun = contr.treatment.explicit,  
  design = create_design(object, formula = formula, drop = drop, codingfun = codingfun,  
    verbose = FALSE)  
)
```

Arguments

object	SummarizedExperiment
formula	formula
drop	TRUE or FALSE
codingfun	coding function (e.g. contr.treatment)
design	design matrix

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
model_coefs(object)
contrast_coefs(object)
```

contrast_subgroup_cols
<i>Row/Col contrasts</i>

Description

Row/Col contrasts

Usage

```
contrast_subgroup_cols(object, subgroupvar)

contrast_subgroup_rows(object, subgroupvar)
```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup svar

Value

matrix

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object$subgroup <- paste0(object$Diabetes, '.', object$Time)
subgroup_matrix(object, subgroupvar = 'subgroup')
contrast_subgroup_cols(object, subgroupvar = 'subgroup')
contrast_subgroup_rows(object, subgroupvar = 'subgroup')
```

counts	<i>Get/Set counts</i>
--------	-----------------------

Description

Get / Set counts matrix

Usage

```
counts(object)

## S4 method for signature 'SummarizedExperiment'
counts(object)

counts(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
counts(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
counts(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,NULL'
counts(object) <- value
```

Arguments

object	SummarizedExperiment
value	count matrix (features x samples)

Value

count matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
counts(object)[1:3, 1:3]
counts(object) <- values(object)
```

counts2cpm	<i>Convert between counts and cpm/tpm</i>
------------	---

Description

Convert between counts and cpm/tpm

Usage

```
counts2cpm(x, libsize = scaledlibsizes(x))  
  
cpm2counts(x, libsize)
```

Arguments

x count/cpm matrix
libsize (scaled) libsize vector

Value

cpm/tpm/count matrix

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')  
object <- read_rnaseq_counts(file)  
libsize <- scaledlibsizes(counts(object))  
tpm <- counts2tpm(counts(object), genesize = 1)  
cpm <- counts2cpm(counts(object), libsize)  
counts <- cpm2counts(cpm, libsize)  
sum(counts(object) - counts)
```

counts2tpm	<i>counts to tpm</i>
------------	----------------------

Description

counts to tpm

Usage

```
counts2tpm(x, genesize)
```

Arguments

x count matrix
genesize genesize vector (kilobase)

Value

tpm matrix

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
counts(object)[1:3, 1:3]
counts2tpm(counts(object), genesize = 1)[1:3, 1:3]
```

count_in	<i>Count/Collapse in/outside intersection</i>
----------	---

Description

Count/Collapse in/outside intersection

Usage

```
count_in(x, ...)

## S3 method for class 'character'
count_in(x, y, ...)

## S3 method for class 'factor'
count_in(x, y, ...)

## S3 method for class 'list'
count_in(x, y, ...)

collapse_in(x, ...)

## S3 method for class 'character'
collapse_in(x, y, sep, ...)

## S3 method for class 'factor'
collapse_in(x, y, sep, ...)

## S3 method for class 'list'
collapse_in(x, y, sep, ...)

count_out(x, ...)

## S3 method for class 'character'
count_out(x, y, ...)

## S3 method for class 'factor'
```

```
count_out(x, y, ...)

## S3 method for class 'list'
count_out(x, y, ...)
```

Arguments

x	character OR list
...	used for S3 dispatch
y	character
sep	string

Value

number OR numeric

Examples

```
# Sets
contrast1 <- c('a', 'b', 'c', 'd')
pathway <- c('c', 'd', 'e', 'f')
contrast2 <- c('e', 'f', 'g', 'h')

# Count outside
count_out(contrast1, pathway)
count_out(list(contrast1 = contrast1, contrast2 = contrast2), pathway)

# Count inside
count_in(contrast1, pathway)
count_in(list(contrast1 = contrast1, contrast2 = contrast2), pathway)

# Collapse inside
collapse_in(contrast1, pathway, sep = ' ')
collapse_in(list(contrast1 = contrast1, contrast2 = contrast2), pathway, sep = ' ')
```

cpm

Get/Set cpm

Description

Get / Set cpm matrix

Usage

```
cpm(object)

## S4 method for signature 'SummarizedExperiment'
cpm(object)
```

```

cpm(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
cpm(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
cpm(object) <- value

```

Arguments

object	SummarizedExperiment
value	cpm matrix (features x samples)

Value

cpm matrix (get) or updated object (set)

Examples

```

file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
cpm(object)[1:3, 1:3]
cpm(object) <- values(object)

```

create_design	<i>Create design matrix</i>
---------------	-----------------------------

Description

Create design matrix for statistical analysis

Usage

```

create_design(object, ...)

## S3 method for class 'SummarizedExperiment'
create_design(
  object,
  formula = default_formula(object),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  verbose = TRUE,
  ...
)

## S3 method for class 'data.table'
create_design(

```

```

    object,
    formula = default_formula(object),
    drop = varlevels_dont_clash(object, all.vars(formula)),
    codingfun = contr.treatment.explicit,
    verbose = TRUE,
    ...
  )

```

Arguments

object	SummarizedExperiment or data.frame
...	required to s3ify
formula	formula with svars
drop	whether to drop predictor names
codingfun	factor coding function • <code>contr.treatment</code>: $\text{intercept} = y_0$, $\text{coef}_i = y_i - y_0$ • <code>contr.treatment.explicit</code>: $\text{intercept} = y_0$, $\text{coef}_i = y_i - y_0$ • <code>code_control</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - y_0$ • <code>contr.diff</code>: $\text{intercept} = y_0$, $\text{coef}_i = y_i - y_{(i-1)}$ • <code>code_diff</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - y_{(i-1)}$ • <code>code_diff_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - y_{(i+)}$ • <code>code_deviation</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - y_{\text{mean}}$ (drop last) • <code>code_deviation_first</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - y_{\text{mean}}$ (drop first) • <code>code_helmert</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - \text{mean}(y_0:(y_{i-1}))$ • <code>code_helmert_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coef}_i = y_i - \text{mean}(y_{(i+1):y_p})$
verbose	whether to message

Value

design matrix

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
unique(create_design(object))
unique(create_design(object, ~ Time))
unique(create_design(object, ~ Time, codingfun = contr.treatment.explicit))
unique(create_design(object, ~ Time, codingfun = contr.diff))
unique(create_design(object, ~ Time + Diabetes))
unique(create_design(object, ~ Time / Diabetes))
unique(create_design(object, ~ Time * Diabetes))

```

DATADIR	<i>Download autonomics example data</i>
---------	---

Description

Download autonomics example data

Usage

```
DATADIR

download_data(
  filename = NULL,
  localdir = file.path(DATADIR, split_extract_fixed(filename, ".", 1)),
  verbose = TRUE,
  force = FALSE
)
```

Arguments

filename	file name		
	'atkin.somascan.adat'	Halama, 2018	effects of hypoglycemia
	'atkin.metabolon.xlsx'		
	'billing16.bam.zip'	Billing, 2016	stemcell comparison
	'billing16.rnacounts.txt'		
	'billing16.somascan.adat'		
	'billing16.proteingroups.txt'		
	'billing19.rnacounts.txt'	Billing, 2016	stemcell differentiation
	'billing19.proteingroups.txt'		
	'billing19.phosphosites.txt'		
	'ddglucose.proteingroups.txt'	Omics Module	glycolysis inhibitor
	'fukuda20.proteingroups.txt'	Fukuda, 2020	zebrafish development
	'halama18.metabolon.xlsx'	Halama, 2018	glutaminase inhibitor
localdir	local dir to save file to		
verbose	TRUE / FALSE		
force	TRUE / FALSE		

Format

An object of class character of length 1.

Value

local file path

Examples

```
# Show available datasets
download_data()

# atkin 2018 - hypoglycemia - pubmed 30525282
# download_data('atkin.somascan.adat')      # somascan intensities
# download_data('atkin.metabolon.xlsx')      # metabolon intensities

# billing16 - stemcell characterization - pubmed 26857143
# download_data('billing16.proteingroups.txt') # proteingroup ratios
# download_data('billing16.somascan.adat')      # somascan intensities
# download_data('billing16.rnaseq.txt')         # rnaseq counts
# download_data('billing16.bam.zip')            # rnaseq alignments

# billing19 - stemcell differentiation - pubmed 31332097
# download_data('billing19.proteingroups.txt') # proteingroup ratios
# download_data('billing19.phosphosites.txt')  # phosphosite ratios
# download_data('billing19.rnaseq.txt')         # rnaseq counts

# fukuda20 - heart regeneration - pubmed PXD016235
# download_data('fukuda20.proteingroups.txt')  # proteingroup LFQ

# halama18 - glutaminase inhibition - pubmed 30525282
# download_data('halama18.metabolon.xlsx')     # metabolon intensities
```

default_geom	<i>Default geom</i>
--------------	---------------------

Description

Default geom

Usage

```
default_geom(object, x, block = NULL)
```

Arguments

object	SummarizedExperiment
x	svar
block	svar or NULL

Value

character vector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object$Age <- runif(min = 20, max = 60, n = ncol(object))
svars(object)
default_geom(object, x = 'Age')
default_geom(object, x = c('Age', 'Diabetes'))
default_geom(object, x = c('Age', 'Diabetes'), block = 'Subject')
```

default_sfile	<i>Default sfile</i>
---------------	----------------------

Description

Default sfile

Usage

```
default_sfile(file)
```

Arguments

file data file

Value

sample file

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
default_sfile(file)
```

default_subgroupvar	<i>Create default formula</i>
---------------------	-------------------------------

Description

Create default formula

Usage

```
default_subgroupvar(object)

default_formula(object)
```

Arguments

object SummarizedExperiment

Value

formula

Examples

```
# Abundances
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
default_formula(object)
file <- download_data('billing16.proteingroups.txt')
object <- read_maxquant_proteingroups(file)
default_formula(object)
```

demultiplex	<i>Demultiplex snames</i>
-------------	---------------------------

Description

Demultiplex maxquant samplenames

Usage

```
demultiplex(x, verbose = FALSE)
```

Arguments

x character vector
verbose TRUE or FALSE

Details

WT(L).KD(H).R1{H/L} -> KD_WT.R1 WT(1).KD(2).R1{1} -> WT.R1 WT.R1 -> WT.R1

Value

character

Examples

```
# uniplexed / intensity / ratio
demultiplex(c('KD.R1','OE.R1'))
demultiplex(c('WT(L).KD(M).OE(H).R1{M}', 'WT(L).KD(M).OE(H).R1{H}'))
demultiplex(c('WT(L).KD(M).OE(H).R1{M/L}', 'WT(L).KD(M).OE(H).R1{H/L}'))
# run / replicate
demultiplex(c('WT(L).OE(H).R1{L}', 'WT(L).OE(H).R1{H}')) # run
demultiplex(c('WT.R1(L).OE.R1(H){L}', 'WT.R1(L).OE.R1(H){H}')) # repl
# label / index
demultiplex(c('WT(L).OE(H).R1{L}', 'WT(L).OE(H).R1{H}')) # label
demultiplex(c('WT(1).OE(2).R1{1}', 'WT(1).OE(2).R1{2}')) # index
# with unused channels
demultiplex('WT(1).KD(2).OE(3).R1{6}')
```

dequantify	<i>Dequantify maxquant snames</i>
------------	-----------------------------------

Description

Drop quantity ('Reporter intensity').
Encode {channel} as suffix.

Usage

```
dequantify(x, quantity = guess_maxquant_quantity(x), verbose = FALSE)
```

Arguments

x	character
quantity	'ratio', 'normalizedratio', 'LFQ intensity', 'intensity', 'labeledintensity' 'reporterintensity', 'correctedreporterintensit
verbose	TRUE or FALSE

Details

Ratio H/L WT(L).KD(H).R1 -> WT(L).KD(H).R1{H/L} LFQ intensity WT.R1 -> WT.R1
Reporter intensity 0 WT(126).KD(127).R1 -> WT(1).KD(2).R1{1}

Value

character

Examples

```
dequantify(c('Ratio H/L WT(L).KD(M).OE(H).R1',          # Ratios
             'Ratio M/L WT(L).KD(M).OE(H).R1'))
dequantify(c('Ratio H/L normalized WT(L).KD(M).OE(H).R1', # Norm. Ratios
             'Ratio M/L normalized WT(L).KD(M).OE(H).R1'))
dequantify(c('LFQ intensity WT.R1',                      # LFQ intensity
             'LFQ intensity KD.R1'))
dequantify(c('Reporter intensity 1 WT(126).KD(127).R1',  # Rep.intensities
             'Reporter intensity 2 WT(126).KD(127).R1'))
```

dequantify_compounddiscoverer
<i>dequantify_compounddiscoverer compound discoverer snames</i>

Description

Drop quantity.

Usage

```
dequantify_compounddiscoverer(
  x,
  quantity = guess_compounddiscoverer_quantity(x),
  verbose = FALSE
)
```

Arguments

x	character
quantity	'area', 'normalizedarea'
verbose	TRUE or FALSE

Details

Norm. Area: 20230908_F143_HILICNEG.raw (F11) -> 20230908_F143_HILICNEG.raw (F11)
Area: 20230908_F143_HILICNEG.raw (F11) -> 20230908_F143_HILICNEG.raw (F11)

Value

character

Examples

```
dequantify_compounddiscoverer("Norm. Area: 20230908_F143_HILICNEG.raw (F11)") # Norm. Area
dequantify_compounddiscoverer("Area: 20230908_F143_HILICNEG.raw (F11)")      # Area
```

DIMREDUN	<i>Dimension Reduction Methods</i>
Description	
Dimension Reduction Methods	
Usage	
DIMREDUN	
DIMREDSUPER	
DIMREDENGINES	
Format	
An object of class character of length 2.	
An object of class character of length 4.	
An object of class character of length 6.	
Details	
• DIMREDUN: c('pca', 'sma') • DIMREDSUPER: c('lda', 'pls', 'opls', 'spls') • DIMREDENGINES: c('pca', 'sma', 'lda', 'pls', 'opls', 'spls')	
download_gtf	<i>Download GTF file</i>

Description

Download GTF file with feature annotations

Usage

```
download_gtf(  
  organism,  
  release = 100,  
  gtffile = sprintf("%s/gtf/%s", R_user_dir("autonomics", "cache"),  
    basename(make_gtf_url(organism, release) %>% substr(1, nchar(.) - 3)))  
)
```

Arguments

organism	'Homo sapiens', 'Mus musculus' or 'Rattus norvegicus'
release	GTF release (number)
gtffile	string: path to local GTF file

Value

gtffile path

Examples

```
organism <- 'Homo sapiens'
# download_gtf(organism)
```

download_mcclain21	<i>Download mcclain21 data</i>
--------------------	--------------------------------

Description

Download mcclain21 data

Usage

```
download_mcclain21(
  counts_or_samples = "counts",
  localdir = file.path(DATADIR, "mcclain21"),
  force = FALSE
)
```

Arguments

counts_or_samples	'counts' or 'samples'
localdir	dirname
force	TRUE or FALSE

Details

Mc clain 2021: COVID19 transcriptomics:

Examples

```
download_mcclain21('counts')
download_mcclain21('samples')
```

dt2mat	<i>'data.table' to 'matrix'</i>
--------	---------------------------------

Description

Convert between 'data.table' and 'matrix'

Usage

```
dt2mat(x)

mat2dt(x, idvar)
```

Arguments

x data.table / matrix
idvar idvar string

Value

matrix / data.table

Examples

```
x <- data.table::data.table(
  gene   = c('ENSG001', 'ENSG002', 'ENSG003'),
  sampleA = c(1787, 10, 432),
  sampleB = c(1143, 3, 268))
dt2mat(x)
mat2dt(dt2mat(x), 'gene')
```

enrichment	<i>Enrichment analysis</i>
------------	----------------------------

Description

Are selected genes enriched in pathway?

Usage

```
enrichment(
  object,
  pathwaydt,
  fit = fits(object)[1],
  coef = coefs(object, fit = fit)[1],
  var = abstractvar(object, fit = fit, coef = coef),
```

```

    levels = fdt(object)[[var]] %>% base::levels() %>% extract(-1),
    genevar = "gene",
    genesep = "[ ,;]",
    n = 3,
    verbose = TRUE,
    genes = FALSE
  )

```

Arguments

object	SummarizedExperiment
pathwaydt	pathway data.table
fit	string
coef	string
var	selection fvar
levels	selection levels
genevar	gene fvar
genesep	gene separator (string)
n	number
verbose	whether to msg
genes	whether to report genes

Details

Four enrichment analyses per geneset using the Fisher Exact Test (see four pvalues). Results are returned in a data.table

in	: genes in pathway
in.det	: detected genes in pathway
in.sel	: up/downregulated genes in pathway
in.up(.genes)	: upregulated genes in pathway
in.down(.genes)	: downregulated genes in pathway
out	: genes outside pathway
det	: detected genes (in + out)
sel	: up/downregulated genes (in + out)
up	: upregulated genes (in + out)
down	: downregulated genes (in + out)
p.coef.upDET	: prob to randomly select this many (or more) upregulated genes (among detected genes)
p.coef.downDET	: prob to randomly select this many (or more) downregulated genes (among detected genes)
p.coef.selDET	: prob to randomly select this many (or more) up OR downregulated genes (among detected genes)
p.coef.selGEN	: prob to randomly select this many (or more) up OR downregulated genes (among genome genes)
p.detGEN	: prob to randomly select this many (or more) detected genes (among genome genes)

Examples

```
# Read
pathwaydt <- read_msigt(collections = 'C5:GO:BP')
file <- system.file('extdata/atkin.somascan.adat', package = 'autonomics')
object <- read_somascan(file, fit = 'limma', coefs = 't1-t0')
fvars(object) %<>% gsub('EntrezGeneSymbol', 'gene', .)
object %<>% abstract_fit()
varlevels <- c('flat', 'down', 'up')
enrichdt1 <- enrichment(object, pathwaydt, var = 't1-t0~limma') # 2:n factor
enrichdt2 <- enrichment(object, pathwaydt, var = 't1-t0~limma', levels = varlevels) # 1:n factor
enrichdt3 <- altenrich(object, pathwaydt) # alternative implementation
cols <- intersect(names(enrichdt1), names(enrichdt3))
all(enrichdt1[, cols, with = FALSE] == enrichdt3[, cols, with = FALSE]) # identical
```

ens2org	<i>taxon/ens to organism</i>
---------	------------------------------

Description

taxon/ens to organism

Usage

```
ens2org(x)

taxon2org(x)
```

Arguments

x character vector

Value

character vector

Examples

```
taxon2org( x = c('9606', '9913') )
ens2org( x = c('ENSP00000377550', 'ENSBTAP00000038329') )
```

entrezg_to_symbol	<i>Entrezg to genesymbol</i>
-------------------	------------------------------

Description

Entrezg to genesymbol

Usage

```
entrezg_to_symbol(x, orgdb)
```

Arguments

x	charactervector
orgdb	OrgDb

Value

character vector

Examples

```
if (requireNamespace('org.Hs.eg.db', quiet = TRUE)){
  orgdb <- org.Hs.eg.db::org.Hs.eg.db
  entrezg_to_symbol(x = c('7448', '3818', '727'), orgdb)
}
```

extract_rectangle	<i>Extract rectangle from omics file, data.table, or matrix</i>
-------------------	---

Description

Extract rectangle from omics file, data.table, or matrix

Usage

```
extract_rectangle(x, ...)

## S3 method for class 'character'
extract_rectangle(
  x,
  rows = seq_len(nrows(x, sheet = sheet)),
  cols = seq_len(ncols(x, sheet = sheet)),
  verbose = FALSE,
  transpose = FALSE,
  drop = FALSE,
```

```

    sheet = 1,
    ...
)

## S3 method for class 'data.table'
extract_rectangle(
  x,
  rows = seq_len(nrow(x)),
  cols = seq_len(ncol(x)),
  transpose = FALSE,
  drop = FALSE,
  ...
)

## S3 method for class 'matrix'
extract_rectangle(
  x,
  rows = seq_len(nrow(x)),
  cols = seq_len(ncol(x)),
  transpose = FALSE,
  drop = FALSE,
  ...
)

```

Arguments

x	omics datafile or datatable
...	allow for S3 method dispatch
rows	numeric vector
cols	numeric vector
verbose	logical
transpose	logical
drop	logical
sheet	numeric or string

Value

matrix

Examples

```

# FROM FILE: extract_rectangle.character
#=====
x <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
extract_rectangle(x, rows = 11:30, cols = 15:81, sheet = 2)[ 1:3, 1:3 ] # exprs
extract_rectangle(x, rows = 11:30, cols = 2, sheet = 2)[ 1:3, ] # fids
extract_rectangle(x, rows = 4, cols = 15:81, sheet = 2)[ , 1:3 ] # sids
extract_rectangle(x, rows = 10:30, cols = 1:14, sheet = 2)[ 1:3, 1:3 ] # fdt

```

```

extract_rectangle(x, rows = 1:10, cols = 14:81, sheet = 2, transpose = TRUE)[1:3, 1:3] # sdt

# FROM MATRIX: extract_rectangle.matrix
#=====
x <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
x %<>% extract_rectangle(sheet = 2)
extract_rectangle(x, rows = 11:30, cols = 15:81, sheet = 2)[ 1:3, 1:3 ] # exprs
extract_rectangle(x, rows = 11:30, cols = 2, sheet = 2)[ 1:3, ] # fids
extract_rectangle(x, rows = 4, cols = 15:81, sheet = 2)[ , 1:3 ] # sids
extract_rectangle(x, rows = 10:30, cols = 1:14, sheet = 2)[ 1:3, 1:3 ] # fdt
extract_rectangle(x, rows = 1:10, cols = 14:81, sheet = 2, transpose = TRUE)[1:3, 1:3] # sdt

```

fcluster

*Cluster features***Description**

Cluster features

Usage

```

fcluster(
  object,
  distmat = NULL,
  method = "cmeans",
  k = 2:10,
  verbose = TRUE,
  plot = TRUE,
  label = if ("gene" %in% fvars(object)) "gene" else "feature_id",
  alpha = 1,
  nrow = if (length(method) > 1) length(method) else NULL,
  ncol = NULL
)

```

Arguments

object	SummarizedExperiment
distmat	distance matrix
method	'cmeans'
k	number of clusters
verbose	TRUE or FALSE
plot	TRUE or FALSE
label	fvar
alpha	fraction
nrow	number
ncol	number

Value

SummarizedExperiment

SummarizedExperiment

Examples

```

object <- twofactor_sumexp()
dmat <- fdist(object)
fcluster(object) # membership-based colors
fcluster(object, dmat) # silhouette-based colors
fcluster(object, dmat, method = c('cmeans', 'hclust', 'pamk')) # more methods

```

fdata	<i>Get/Set sample/feature data</i>
-------	------------------------------------

Description

Get/Set sample/feature data

Usage

```

fdata(object)

sdata(object)

fdt(object)

sdt(object)

## S4 method for signature 'SummarizedExperiment'
fdata(object)

## S4 method for signature 'SummarizedExperiment'
sdata(object)

## S4 method for signature 'SummarizedExperiment'
fdt(object)

## S4 method for signature 'SummarizedExperiment'
sdt(object)

fdata(object) <- value

sdata(object) <- value

fdt(object) <- value

```

```
sdt(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,data.frame'
fdata(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,data.frame'
sdata(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,DataFrame'
sdata(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,data.table'
fdt(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,data.table'
sdt(object) <- value
```

Arguments

object	SummarizedExperiment
value	data.frame/data.table

Value

data.frame/data.table (get) or updated object (set)

Examples

```
# Read data
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
# sdt/fdt
sdt(object)[1:3, ]
fdt(object)[1:3, ]
sdt(object) %<>% cbind(b=1)
fdt(object) %<>% cbind(b=1)
sdt(object)
fdt(object)
# sdata/fdata
sdata(object)[1:3, ]
fdata(object)[1:3, ]
sdata(object) %<>% cbind(a=1)
fdata(object) %<>% cbind(a=1)
sdata(object)[1:3, ]
fdata(object)[1:3, ]
```

fdr2p	<i>fdr to p</i>
-------	-----------------

Description

fdr to p

Usage

fdr2p(fdr)

Arguments

fdr fdr values

Examples

```
# Read/Fit
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
pcol <- pvar(fdt(object), fit = 'limma', coef = 't3-t0')
object %<>% extract(order(fdt(.)[[pcol]]), )
object %<>% extract(1:10, )
fdt(object) %<>% extract(, 1)
object %<>% fit_limma()

# fdr2p
fdt(object)[[pcol]]
fdt(object)[[pcol]] %>% p.adjust(method = 'fdr')
fdt(object)[[pcol]] %>% p.adjust(method = 'fdr') %>% fdr2p()
```

filter_exprs_replicated_in_some_subgroup
<i>Filter features with replicated expression in some subgroup</i>

Description

Filter features with replicated expression in some subgroup

Usage

```
filter_exprs_replicated_in_some_subgroup(
  object,
  subgroupvar = "subgroup",
  assay = assayNames(object)[1],
  comparator = if (contains_ratios(object)) "!=" else ">",
  lod = 0,
```

```
    nsample = 2,  
    nsubgroup = 1,  
    verbose = TRUE  
  )
```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup svar
assay	string
comparator	'>' or '!='
lod	number: limit of detection
nsample	number
nsubgroup	number
verbose	TRUE or FALSE

Value

Filtered SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')  
object <- read_metabolon(file)  
object %<>% filter_exprs_replicated_in_some_subgroup()  
filter_exprs_replicated_in_some_subgroup(object, character(0))  
filter_exprs_replicated_in_some_subgroup(object, NULL)
```

filter_features	<i>Filter features on condition</i>
-----------------	-------------------------------------

Description

Filter features on condition

Usage

```
filter_features(object, condition, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
condition	filter condition
verbose	logical

Value

filtered eSet

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
filter_features(object, SUPER_PATHWAY == 'Lipid')
```

filter_medoid	<i>Filter medoid sample</i>
---------------	-----------------------------

Description

Filter medoid sample

Usage

```
filter_medoid(object, by = NULL, verbose = FALSE)
```

Arguments

object	SummarizedExperiment
by	svar
verbose	whether to message

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file, plot = FALSE)
object %<>% filter_medoid(by = 'subgroup', verbose=TRUE)
```

filter_samples	<i>Filter samples on condition</i>
----------------	------------------------------------

Description

Filter samples on condition

Usage

```
filter_samples(object, condition, verbose = TRUE, record = TRUE)
```

Arguments

object	SummarizedExperiment
condition	filter condition
verbose	TRUE/FALSE
record	TRUE/FALSE

Value

filtered SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
filter_samples(object, subgroup != 't0', verbose = TRUE)
```

fitcoefs	<i>fitcoefs</i>
----------	-----------------

Description

fitcoefs

Usage

```
fitcoefs(object)
```

Arguments

object	SummarizedExperiment
--------	----------------------

Value

string vector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
fitcoefs(object)
fitcoefs(fit_limma(object))
```

fits

Get fit models

Description

Get fit models

Usage

```
fits(object, ...)
```

S3 method for class 'data.table'

```
fits(object, ...)
```

S3 method for class 'SummarizedExperiment'

```
fits(object, ...)
```

Arguments

object	SummarizedExperiment or data.table
...	S3 dispatch

Value

character vector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma')
fits(object)
```

FITSEP	<i>Fit results separator</i>
--------	------------------------------

Description

Fit results separator

Usage

FITSEP

PPATTERN

Format

An object of class character of length 1.

An object of class character of length 1.

Examples

FITSEP

fit_linmod	<i>Fit General Linear Model</i>
------------	---------------------------------

Description

Fit General Linear Model

Usage

```
fit_linmod(
  object,
  formula = as.formula("~ subgroup"),
  engine = "limma",
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = create_design(object, formula = formula, drop = drop, codingfun = codingfun,
    verbose = FALSE),
  contrasts = NULL,
  coefs = if (is.null(contrasts)) model_coefs(design = design) else NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  sep = FITSEP,
  suffix = paste0(sep, engine),
```

```

    verbose = TRUE,
    outdir = NULL,
    writefun = "write_xl",
    volcano = FALSE,
    volcanoargs = list(),
    exprs = FALSE,
    exprargs = list(),
    ...
)

fit_limma(
  object,
  formula = as.formula("~ subgroup"),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = create_design(object, formula = formula, drop = drop, codingfun = codingfun),
  contrasts = NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  sep = FITSEP,
  suffix = paste0(sep, "limma"),
  verbose = TRUE
)

.fit_limma(
  object,
  formula = as.formula("~ subgroup"),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = create_design(object, formula = formula, drop = drop, codingfun = codingfun),
  contrasts = NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  sep = FITSEP,
  suffix = paste0(sep, "limma"),
  verbose = TRUE
)

fit_lm(
  object,
  formula = as.formula("~ subgroup"),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  sep = FITSEP,
  suffix = paste0(sep, "lm"),

```

```
    contrasts = NULL,
    verbose = TRUE
)

fit_lme(
  object,
  formula = as.formula("~ subgroup"),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  opt = "optim",
  sep = FITSEP,
  suffix = paste0(sep, "lme"),
  contrasts = NULL,
  verbose = TRUE
)

fit_lmer(
  object,
  formula = as.formula("~ subgroup"),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit,
  design = NULL,
  block = NULL,
  weightvar = if ("weights" %in% assayNames(object)) "weights" else NULL,
  sep = FITSEP,
  suffix = paste0(sep, "lmer"),
  contrasts = NULL,
  verbose = TRUE
)

fit_wilcoxon(
  object,
  formula = as.formula("~ subgroup"),
  drop = NULL,
  codingfun = contr.treatment.explicit,
  design = NULL,
  contrasts = NULL,
  block = NULL,
  weightvar = NULL,
  sep = FITSEP,
  suffix = paste0(sep, "wilcoxon"),
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
formula	model formula
engine	'limma', 'lm', 'lme', 'lmer', or 'wilcoxon'
drop	TRUE or FALSE
codingfun	factor coding function • <code>contr.treatment</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_0$ • <code>contr.treatment.explicit</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_0$ • <code>code_control</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_0$ • <code>contr.diff</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_{(i-1)}$ • <code>code_diff</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{(i-1)}$ • <code>code_diff_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{(i+)}$ • <code>code_deviation</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{\text{mean}}$ (drop last) • <code>code_deviation_first</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{\text{mean}}$ (drop first) • <code>code_helmert</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - \text{mean}(y_0:(y_i-1))$ • <code>code_helmert_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - \text{mean}(y_{(i+1):yp})$
design	design matrix
contrasts	NULL or character vector: coefficient contrasts to test
coefs	NULL or character vector: model coefs to test
block	block svar (or NULL)
weightvar	NULL or name of weight matrix in <code>assays(object)</code>
sep	string: pvar separator ("~" in "p~t2~limma")
suffix	string: pvar suffix ("limma" in "p~t2~limma")
verbose	whether to msg
outdir	NULL or dir
writefun	'write_xl' or 'write_ods'
volcano	TRUE or FALSE
volcanoargs	list: volcano args
exprs	TRUE or FALSE
exprargs	list: expr args
...	passed to <code>fit_(limmallmlmlmer)</code> functions
opt	lme options

Value

Updated SummarizedExperiment

Examples

```
# Standard usage
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_linmod() # Default
object %<>% fit_linmod( ~subgroup ) # Custom formula
object %<>% fit_linmod( ~subgroup, block = 'Subject') # Block effect
summarize_fit(object)

# Alternative engines: argument 'engine' or dedicated function
fdt(object) %<>% extract(, 'feature_id')
object %<>% fit_limma( ~subgroup, block = 'Subject') # Default engine
object %<>% fit_lm( ~subgroup, block = 'Subject') # Traditional
object %<>% fit_lme( ~subgroup, block = 'Subject') # Powerful random effects
object %<>% fit_lmer( ~subgroup, block = 'Subject') # Yet more powerful random effects
object %<>% fit_wilcoxon(~subgroup, block = 'Subject') # Non-parametric
summarize_fit(object)

# Alternative coding: backward diffs instead of baseline
fdt(object) %<>% extract(, 'feature_id')
object %<>% fit_limma( ~ subgroup, block = 'Subject', codingfun = code_diff)
object %<>% fit_lme( ~ subgroup, block = 'Subject', codingfun = code_diff)
object %<>% fit_lmer( ~ subgroup, block = 'Subject', codingfun = code_diff)
summarize_fit(object)

# Posthoc contrasts: limma-only, flexible, but sometimes approximate
fdt(object) %<>% extract(, 'feature_id')
object %<>% fit_limma( ~ subgroup, block = 'Subject', codingfun = code_control)
object %<>% fit_limma( ~ 0 + subgroup, block = 'Subject', contrasts = 't1-t0')
# flexible, but only approximate
# stat.ethz.ch/pipermail/bioconductor/2014-February/057682.html

# Custom separator
fdt(object) %<>% extract(, 'feature_id')
fdt( fit_limma(object, sep = '.'))
fdt( fit_limma(object, block = 'Subject', sep = '.') )

# Top-level function also plots and writes
fit_linmod(object, block = 'Subject', coefs = 't1-t0')
fit_linmod(object, block = 'Subject', coefs = 't1-t0', volcano = TRUE)
fit_linmod(object, block = 'Subject', coefs = 't1-t0', exprs = TRUE)
fit_linmod(object, block = 'Subject', coefs = 't1-t0', volcano = TRUE, exprs = TRUE)
fit_linmod(object, block = 'Subject', coefs = 't1-t0', volcano = TRUE, exprs = TRUE, outdir = tempdir())
fit_linmod(object, block = 'Subject', coefs = 't1-t0', volcano = TRUE, exprs = TRUE, outdir = tempdir())
```

fix_xlgenes

Fix excel genes

Description

Fix excel genes

Usage

```
fix_xlgenes(x)
```

Arguments

x character

Value

character

Examples

```
x <- c('FAM46B', '15-Sep', '2-Mar', 'MARCHF6')
x
fix_xlgenes(x)
```

flevels	<i>Get fvar levels</i>
---------	------------------------

Description

Get fvar levels

Usage

```
flevels(object, fvar)
```

Arguments

object SummarizedExperiment
fvar feature variable

Value

fvar values

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
head(flevels(object, 'feature_id'))
```

fnames	<i>Get/Set fnames</i>
--------	-----------------------

Description

Get/Set feature names

Usage

```
fnames(object)

## S4 method for signature 'SummarizedExperiment'
fnames(object)

fnames(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,character'
fnames(object) <- value
```

Arguments

object	SummarizedExperiment, eSet, or EList
value	character vector with feature names

Value

feature name vector (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fnames(object) %<>% paste0('protein_', .)
object
```

formula2str	<i>formula to string</i>
-------------	--------------------------

Description

formula to string

Usage

```
formula2str(formula)
```

Arguments

formula formula

Value

string

Examples

```
formula2str(~0+subgroup)
```

ftype	<i>Feature type</i>
-------	---------------------

Description

Feature type

Usage

```
ftype(
  object,
  formula = default_formula(object),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  fit = fits(object)[1],
  codingfun = contr.treatment.explicit
)
```

Arguments

object SummarizedExperiment
formula model formula
drop TRUE or FALSE
fit 'limma', 'lm', 'lme', 'wilcoxon'
codingfun coding function

Value

SummarizedExperiment

Examples

```
file <- download_data('atkin.metabolon.xlsx')
object <- read_metabolon(file)
object %<>% fit_limma(block = 'Subject') # model_coefs !
object %<>% ftype()                      # model_coefs not contrast_coefs !
fdt(object)                              # because intercept is required to recreate predictions
```

fvalues*Get fvalues*

Description

Get fvar values

Usage

```
fvalues(object, fvar)
```

Arguments

object	SummarizedExperiment
fvar	feature variable

Value

fvar values

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
head(fvalues(object, 'feature_id'))
fvalues(object, NULL)
```

fvars*Get/Set fvars*

Description

Get/Set feature variables

Usage

```
fvars(object)

## S4 method for signature 'SummarizedExperiment'
fvars(object)

fvars(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,character'
fvars(object) <- value
```

Arguments

object SummarizedExperiment
value character vector with feature variables

Value

feature variables vector (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fvars(object)[1] %<>% paste0('1')
fvars(object)[1]
```

genome_to_orgdb	<i>Get corresponding orgdb</i>
-----------------	--------------------------------

Description

Get corresponding orgdb

Usage

```
genome_to_orgdb(genome)
```

Arguments

genome 'hg38', 'hg19', 'mm10', or 'mm9'

Value

OrgDb

Examples

```
if (requireNamespace('org.Hs.eg.db', quiet = TRUE)){
  class(genome_to_orgdb('hg38'))
}
```

group_by_level	<i>group by level</i>
----------------	-----------------------

Description

group by level

Usage

```
group_by_level(x, ...)  
  
## S3 method for class 'character'  
group_by_level(x, ...)  
  
## S3 method for class 'factor'  
group_by_level(x, ...)  
  
## S3 method for class 'data.table'  
group_by_level(x, var, idvar, ...)
```

Arguments

x	named logical/character/factor
...	S3 dispatch
var	string
idvar	string

Value

unnamed character

Examples

```
t1 <- c( KLF5 = 'up', F11 = 'up', RIG = 'flat', ABT1 = 'down')  
dt <- data.table( gene = c( 'KL5', 'F11', 'RIG', 'ABT1' ),  
                 t1 = c( 'up', 'up', 'flat', 'down' ) )  
group_by_level(t1)           # character  
group_by_level(factor(t1))   # factor  
group_by_level(dt, 't1', 'gene') # data.table
```

guess_compounddiscoverer_quantity
<i>Guess compound discoverer quantity from snames</i>

Description

Guess compound discoverer quantity from snames

Usage

guess_compounddiscoverer_quantity(x)

Arguments

x character vector

Value

string: value from names(COMPOUNDDISCOVERER_PATTERNS)

Examples

```
## Not run:
# file
  file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
  guess_compounddiscoverer_quantity(file)

## End(Not run)

# character vector
x <- "Area: 20230908_F143_HILICNEG.raw (F11)"
guess_compounddiscoverer_quantity(x)

x <- "Norm. Area: 20230908_F143_HILICNEG.raw (F11)"
guess_compounddiscoverer_quantity(x)
```

guess_fitsep	<i>guess_fitsep</i>
--------------	---------------------

Description

guess_fitsep

Usage

```
guess_fitsep(object, ...)  
  
## S3 method for class 'data.table'  
guess_fitsep(object, ...)  
  
## S3 method for class 'SummarizedExperiment'  
guess_fitsep(object, ...)
```

Arguments

object	data.table or SummarizedExperiment
...	S3 dispatch

Value

string

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')  
object <- read_maxquant_proteingroups(file)  
object %<>% fit_limma()  
guess_fitsep(object)
```

guess_maxquant_quantity

Guess maxquant quantity from snames

Description

Guess maxquant quantity from snames

Usage

```
guess_maxquant_quantity(x)
```

Arguments

x	character vector
---	------------------

Value

string: value from names(MAXQUANT_PATTERNS)

Examples

```
# file
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
guess_maxquant_quantity(file)

# character vector
x <- "Ratio M/L normalized STD(L)_E00(M)_E01(H)_R1"
guess_maxquant_quantity(x)

x <- "Ratio M/L STD(L)_E00(M)_E01(H)_R1"
guess_maxquant_quantity(x)

x <- "LFQ intensity E00.R1"
guess_maxquant_quantity(x)

x <- "Reporter intensity corrected 0 STD(0)E00(1)E01(2)_R1"
guess_maxquant_quantity(x)

x <- "Reporter intensity 0 STD(0)E00(1)E01(2)_R1"
guess_maxquant_quantity(x)

x <- "Intensity H STD(L)_E00(M)_E01(H)_R1"
guess_maxquant_quantity(x)
```

guess_sep

Guess separator

Description

Guess separator

Usage

```
guess_sep(x, ...)
```

S3 method for class 'numeric'

```
guess_sep(x, ...)
```

S3 method for class 'character'

```
guess_sep(x, separators = c(".", "_"), verbose = FALSE, ...)
```

S3 method for class 'factor'

```
guess_sep(x, ...)
```

S3 method for class 'SummarizedExperiment'

```
guess_sep(x, var = "sample_id", separators = c(".", "_"), verbose = FALSE, ...)
```

Arguments

x character vector or SummarizedExperiment
... used for proper S3 method dispatch
separators character vector: possible separators to look for
verbose TRUE or FALSE
var svar or fvar

Value

separator (string) or NULL (if no separator could be identified)

Examples

```

# character vector
guess_sep(c('PERM_NON.R1[H/L]', 'PERM_NON.R2[H/L]'))
guess_sep(c('WT_untreated_1', 'WT_untreated_2'))
guess_sep(c('group1', 'group2.R1'))
# SummarizedExperiment
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
guess_sep(object)

```

has_multiple_levels	<i>Variable has multiple levels?</i>
---------------------	--------------------------------------

Description

Variable has multiple levels?

Usage

```

has_multiple_levels(x, ...)

## S3 method for class 'character'
has_multiple_levels(x, .xname = get_name_in_parent(x), ...)

## S3 method for class 'factor'
has_multiple_levels(x, .xname = get_name_in_parent(x), ...)

## S3 method for class 'numeric'
has_multiple_levels(x, .xname = get_name_in_parent(x), ...)

## S3 method for class 'data.table'
has_multiple_levels(
  x,
  y,

```

```

    .xname = get_name_in_parent(x),
    .yname = get_name_in_parent(y),
    ...
)

## S3 method for class 'SummarizedExperiment'
has_multiple_levels(
  x,
  y,
  .xname = get_name_in_parent(x),
  .yname = get_name_in_parent(y),
  ...
)
```

Arguments

x	vector, data.table or SummarizedExperiment
...	required for s3 dispatch
.xname	string
y	string
.yname	string

Value

TRUE or false

Examples

```

# numeric
a <- numeric(); has_multiple_levels(a)
a <- c(1, 1); has_multiple_levels(a)
a <- c(1, 2); has_multiple_levels(a)

# character
a <- character(); has_multiple_levels(a)
a <- c('A', 'A'); has_multiple_levels(a)
a <- c('A', 'B'); has_multiple_levels(a)

# factor
a <- factor(); has_multiple_levels(a)
a <- factor(c('A', 'A')); has_multiple_levels(a)
a <- factor(c('A', 'B')); has_multiple_levels(a)

# data.table
dt <- data.table(a = factor()); has_multiple_levels(dt, 'b')
dt <- data.table(a = factor()); has_multiple_levels(dt, 'a')
dt <- data.table(a = factor()); has_multiple_levels(dt, 'a')
dt <- data.table(a = factor(c('A', 'A'))); has_multiple_levels(dt, 'a')
dt <- data.table(a = factor(c('A', 'B'))); has_multiple_levels(dt, 'a')

# sumexp
object <- matrix(1:9, nrow = 3)
rownames(object) <- sprintf('%d', 1:3)
colnames(object) <- sprintf('%d', 1:3)
```

```
object <- list(exprs = object)
object %<>% SummarizedExperiment::SummarizedExperiment()
object$subgroup <- c('A', 'A', 'A');           has_multiple_levels(object, 'group')
object$subgroup <- c('A', 'A', 'A');           has_multiple_levels(object, 'subgroup')
object$subgroup <- c('A', 'B', 'A');           has_multiple_levels(object, 'subgroup')
```

hdlproteins	<i>hdl proteomewatch proteins</i>
-------------	-----------------------------------

Description

hdl proteomewatch proteins

Usage

```
hdlproteins()
```

Value

string vector: HDLProteomeWatch protein entries

Examples

```
hdlproteins()
```

impute	<i>Impute</i>
--------	---------------

Description

Impute NA values

Usage

```
impute(object, ...)
```

```
## S3 method for class 'numeric'
impute(object, shift = 2.5, width = 0.3, verbose = TRUE, plot = FALSE, ...)
```

```
## S3 method for class 'matrix'
impute(
  object,
  shift = 2.5,
  width = 0.3,
  verbose = TRUE,
  plot = FALSE,
```

```

    n = min(9, ncol(object)),
    palette = make_colors(colnames(object)),
    ...
)

## S3 method for class 'SummarizedExperiment'
impute(
  object,
  assay = assayNames(object)[1],
  by = "subgroup",
  shift = 2.5,
  width = 0.3,
  frac = 0.5,
  verbose = TRUE,
  plot = FALSE,
  palette = make_colors(colnames(object)),
  n = min(9, ncol(object)),
  ...
)

```

Arguments

<code>object</code>	numeric vector, SumExp
<code>...</code>	required for s3 dispatch
<code>shift</code>	number: sd units
<code>width</code>	number: sd units
<code>verbose</code>	TRUE or FALSE
<code>plot</code>	TRUE or FALSE
<code>n</code>	number of samples to plot
<code>palette</code>	color vector
<code>assay</code>	string
<code>by</code>	svar
<code>frac</code>	fraction: fraction of available samples should be greater than this value for a subgroup to be called available

Details

Imputes NA values from $N(\text{mean} - 2.5 \text{ sd}, 0.3 \text{ sd})$

Value

numeric vector, matrix or SumExp

Examples

```
# Simple Design
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
impute(values(object)[, 1], plot = TRUE)[1:3]           # vector
impute(values(object),      plot = TRUE)[1:3, 1:3]      # matrix
impute(object, plot = TRUE)                             # sumexp

# Complex Design
subgroups <- sprintf('%s_STD', c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'))
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, subgroups = subgroups)
impute(values(object)[1:3, 1  ] )      # vector
impute(values(object)[1:3, 1:5  ] )     # matrix
impute( object )                       # sumexp
```

invert_subgroups	<i>Invert subgroups</i>
------------------	-------------------------

Description

Invert expressions , subgroups, and sample ids

Usage

```
invert_subgroups(
  object,
  subgroups = slevels(object, "subgroup"),
  sep = guess_sep(object, "subgroup")
)
```

Arguments

object	SummarizedExperiment
subgroups	character vector: subgroup levels to be inversed
sep	string: collapsed string separator

Value

character vector or SummarizedExperiment

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
invert_subgroups(object)
```

is_collapsed_subset *Is collapsed subset*

Description

Is collapsed subset

Usage

```
is_collapsed_subset(x, y, sep = ";")
```

Arguments

x	character vector
y	character vector
sep	string

Value

character vector

Examples

```
x <- c('H3BNX8;H3BRM5', 'G5E9Y3')
y <- c('P20674;H3BNX8;H3BV69;H3BRM5', 'G5E9Y3;Q8WWN8;B4DIT1')
is_collapsed_subset(x, y)
```

is_correlation_matrix *Assert correlation matrix*

Description

Assert correlation matrix

Usage

```
is_correlation_matrix(
  x,
  .xname = get_name_in_parent(x),
  severity = getOption("assertive.severity", "stop")
)

assert_correlation_matrix(x, .xname = get_name_in_parent(x))
```

Arguments

x	correlation matrix
.xname	string
severity	'warning' or 'stop'

Value

TRUE or false

Examples

```
x <- matrix(c(1,0.7, 0.3, 1), nrow = 2)
rownames(x) <- c('gene1', 'gene2')
colnames(x) <- c('gene1', 'gene2')
is_correlation_matrix(x)
is_correlation_matrix({x[1,1] <- -2; x})
```

is_diann_report	<i>Is diann, fragpipe, proteingroups, phosphosites file?</i>
-----------------	--

Description

Is diann, fragpipe, proteingroups, phosphosites file?

Usage

```
is_diann_report(x, .xname = get_name_in_parent(x))

is_fragpipe_tsv(x, .xname = get_name_in_parent(x))

is_maxquant_proteingroups(x, .xname = get_name_in_parent(x))

is_maxquant_phosphosites(x, .xname = get_name_in_parent(x))

is_compounddiscoverer_output(x, .xname = get_name_in_parent(x))

assert_diann_report(x, .xname = get_name_in_parent(x))

assert_fragpipe_tsv(x, .xname = get_name_in_parent(x))

assert_maxquant_proteingroups(x, .xname = get_name_in_parent(x))

assert_maxquant_phosphosites(x, .xname = get_name_in_parent(x))

assert_compounddiscoverer_output(x, .xname = get_name_in_parent(x))
```

Arguments

x	file
.xname	name of x

Examples

```
file <- NULL
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- 3
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- 'blabla.tsv'
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- download_data('multiorganism.combined_protein.tsv')
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- download_data('dilution.report.tsv')
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)

file <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
is_diann_report(file)
is_fragpipe_tsv(file)
is_maxquant_proteingroups(file)
is_maxquant_phosphosites(file)
```

is_fastadt	<i>Is fastadt</i>
------------	-------------------

Description

Is fastadt

Usage

```
is_fastadt(x, .xname = get_name_in_parent(x))  
  
assert_fastadt(x, .xname = get_name_in_parent(x))
```

Arguments

x	fasta data.table
.xname	string

Examples

```
fastafile <- system.file('extdata/uniprot_hsa_20140515.fasta', package = 'autonomics')  
x <- read_uniprotDT(fastafile)  
# is_fastadt(x) # slow
```

is_file	<i>Is a file?</i>
---------	-------------------

Description

Is a file (and not a dir)

Usage

```
is_file(file)
```

Arguments

file	filepath
------	----------

Details

This function distinguishes between dir and file. Others don't: `is.file`, `fs::file_exists`, `assertive::is_existing_file`

Examples

```
dir <- tempdir(); dir.create(dir, showWarnings = FALSE)
file <- tempfile(); invisible(file.create(file))
is_file(dir)
is_file(file)
```

is_fraction	<i>Is fraction</i>
-------------	--------------------

Description

Is fraction

Usage

```
is_fraction(x, .xname = get_name_in_parent(x))

assert_is_fraction(x, .xname = get_name_in_parent(x))
```

Arguments

x	number
.xname	string

Value

TRUE or false

Examples

```
is_fraction(0.1)      # YES
is_fraction(1)        # YES
is_fraction(1.2)      # NO - more than 1
is_fraction(c(0.1, 0.2)) # NO - vector
```

is_imputed	<i>Get/set is_imputed</i>
------------	---------------------------

Description

Get/Set is_imputed

Usage

```

is_imputed(object)

## S4 method for signature 'SummarizedExperiment'
is_imputed(object)

is_imputed(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
is_imputed(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,NULL'
is_imputed(object) <- value

```

Arguments

object	SummarizedExperiment
value	matrix

Value

matrix (get) or updated object (set)

Examples

```

file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, impute = TRUE)
sum(is_imputed(object))

```

is_positive_number	<i>Is positive number</i>
--------------------	---------------------------

Description

Is positive number

Usage

```

is_positive_number(x, .xname = get_name_in_parent(x))

assert_positive_number(x, .xname = get_name_in_parent(x))

is_weakly_positive_number(x, .xname = get_name_in_parent(x))

assert_weakly_positive_number(x, .xname = get_name_in_parent(x))

```

Arguments

x	number
.xname	name of x

Value

TRUE or false

Examples

```
is_positive_number( 3)
is_positive_number(-3)
is_positive_number( 0)
is_weakly_positive_number(0)
assert_positive_number(3)
```

is_scalar_subset	<i>Is scalar subset</i>
------------------	-------------------------

Description

Is scalar subset

Usage

```
is_scalar_subset(
  x,
  y,
  .xname = get_name_in_parent(x),
  .yname = get_name_in_parent(y)
)

assert_scalar_subset(
  x,
  y,
  .xname = get_name_in_parent(x),
  .yname = get_name_in_parent(y)
)
```

Arguments

x	scalar
y	SummarizedExperiment
.xname	name of x
.yname	name of y

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
is_scalar_subset('subgroup', svars(object))
is_scalar_subset('subject', svars(object))
assert_scalar_subset('subgroup', svars(object))
```

is_sig	<i>Is significant?</i>
--------	------------------------

Description

Is significant?

Usage

```
is_sig(
  object,
  fit = fits(object)[1],
  contrast = coefs(object),
  quantity = "fdr"
)
```

Arguments

object	SummarizedExperiment
fit	subset of autonomics::TESTS
contrast	subset of colnames(metadata(object)[[fit]])
quantity	value in dimnames(metadata(object)[[fit]])[3]

Value

matrix: -1 (downregulated), +1 (upregulatd), 0 (not fdr significant)

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
object %<>% fit_lm()
object %<>% fit_limma()
issig <- is_sig(object, fit = c('lm','limma'), contrast = 'Adult-X30dpt')
plot_contrast_venn(issig)
```

is_valid_formula	<i>Is valid formula</i>
------------------	-------------------------

Description

Is valid formula

Usage

```
is_valid_formula(  
  x,  
  y,  
  .xname = get_name_in_parent(x),  
  .yname = get_name_in_parent(y)  
)  
  
assert_valid_formula(  
  x,  
  y,  
  .xname = get_name_in_parent(x),  
  .yname = get_name_in_parent(y)  
)
```

Arguments

x	formula
y	SummarizedExperiment
.xname	string
.yname	string

Value

TRUE or false

Examples

```
object <- matrix(1:9, nrow = 3)  
rownames(object) <- sprintf('f%d', 1:3)  
colnames(object) <- sprintf('s%d', 1:3)  
object <- list(exprs = object)  
object %<>% SummarizedExperiment::SummarizedExperiment()  
object$group <- 'group0'  
object$subgroup <- c('A', 'B', 'C')  
svars(object)  
  is_valid_formula( 'condition', object) # not formula  
  is_valid_formula( ~condition, object) # not svar  
  is_valid_formula( ~group, object) # not multilevel
```

```

is_valid_formula( ~subgroup,    object) # TRUE
is_valid_formula( ~0+subgroup,  object) # TRUE
is_valid_formula( ~1,          object) # TRUE
assert_valid_formula( ~subgroup, object)

```

keep_connected_blocks *Keep fully connected blocks*

Description

Keep fully connected blocks

Usage

```
keep_connected_blocks(object, block, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
block	svar
verbose	TRUE or FALSE

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% keep_connected_blocks( block = 'Subject')

```

keep_connected_features
Keep features with n+ connected blocks

Description

Keep features with n+ connected blocks

Usage

```
keep_connected_features(object, block, n = 2, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
block	svar
n	number
verbose	TRUE or FALSE

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% keep_connected_blocks( block = 'Subject')
object %<>% keep_connected_features(block = 'Subject')
```

keep_replicated_features
<i>Keep replicated features</i>

Description

Keep features replicated for each slevel

Usage

```
keep_replicated_features(object, formula = ~1, n = 3, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
formula	formula
n	min replications required
verbose	TRUE or FALSE

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% keep_replicated_features()
object %<>% keep_replicated_features(~ subgroup)
```

label2index	<i>Convert labels into indices</i>
-------------	------------------------------------

Description

Convert labels into indices

Usage

```
label2index(x)
```

Arguments

x	‘character’
---	-------------

Examples

```
label2index(x = 'Reporter intensity 0 WT(0).KD(1).OE(2).R1')
label2index(x = 'Reporter intensity 1 WT(1).KD(2).OE(3).R1')
label2index(x = 'Reporter intensity 0 WT(126).KD(127).OE(128).R1')
label2index(x = 'Reporter intensity 1 WT(126).KD(127).OE(128).R1')
label2index(x = 'Reporter intensity 1 Mix1')
```

LINMODENGINES	<i>Linear Modeling Engines</i>
---------------	--------------------------------

Description

Linear Modeling Engines

Usage

```
LINMODENGINES
```

Format

An object of class character of length 5.

Examples

```
LINMODENGINES
```

list2mat	<i>list to matrix</i>
----------	-----------------------

Description

list to matrix

Usage

```
list2mat(x)
```

Arguments

x list

Value

matrix

Examples

```
x <- list(roundfruit = c('apple', 'orange'), redfruit = c('apple', 'strawberry'))
list2mat(x)
```

list_files	<i>list files</i>
------------	-------------------

Description

list.files for programming

Usage

```
list_files(dir, full.names)
```

Arguments

dir	directory
full.names	TRUE or FALSE

Details

Adds a small layer on list.files. Returning NULL rather than character(0) when no files. Making it better suited for programming.

log2counts	<i>Get/Set log2counts</i>
------------	---------------------------

Description

Get / Set log2counts matrix

Usage

```
log2counts(object)

## S4 method for signature 'SummarizedExperiment'
log2counts(object)

log2counts(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
log2counts(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
log2counts(object) <- value
```

Arguments

object SummarizedExperiment
value log2count matrix (features x samples)

Value

log2count matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')  
object <- read_rnaseq_counts(file)  
log2counts(object)[1:3, 1:3]  
log2counts(object) <- values(object)
```

log2cpm	<i>Get/Set log2cpm</i>
---------	------------------------

Description

Get / Set log2cpm matrix

Usage

```
log2cpm(object)  
  
## S4 method for signature 'SummarizedExperiment'  
log2cpm(object)  
  
log2cpm(object) <- value  
  
## S4 replacement method for signature 'SummarizedExperiment,matrix'  
log2cpm(object) <- value  
  
## S4 replacement method for signature 'SummarizedExperiment,numeric'  
log2cpm(object) <- value
```

Arguments

object SummarizedExperiment
value log2cpm matrix (features x samples)

Value

log2cpm matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
log2cpm(object)[1:3, 1:3]
log2cpm(object) <- values(object)
```

log2diffs	<i>Get/Set log2diffs</i>
-----------	--------------------------

Description

Get/Set log2diffs

Usage

```
log2diffs(object)

## S4 method for signature 'SummarizedExperiment'
log2diffs(object)

log2diffs(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
log2diffs(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
log2diffs(object) <- value
```

Arguments

object	SummarizedExperiment
value	occupancy matrix (features x samples)

Value

occupancy matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
log2diffs(object)[1:3, 1:3]
```

log2proteins	<i>Get/Set log2proteins</i>
--------------	-----------------------------

Description

Get/Set log2proteins

Usage

```
log2proteins(object)

## S4 method for signature 'SummarizedExperiment'
log2proteins(object)

log2proteins(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
log2proteins(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
log2proteins(object) <- value
```

Arguments

object	SummarizedExperiment
value	occupancy matrix (features x samples)

Value

occupancy matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
log2proteins(object)[1:3, 1:3]
```

log2sites	<i>Get/Set log2sites</i>
-----------	--------------------------

Description

Get/Set log2sites

Usage

```
log2sites(object)

## S4 method for signature 'SummarizedExperiment'
log2sites(object)

log2sites(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
log2sites(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
log2sites(object) <- value
```

Arguments

object	SummarizedExperiment
value	occupancy matrix (features x samples)

Value

occupancy matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
log2sites(object)[1:3, 1:3]
```

log2tpm

Get/Set log2tpm

Description

Get / Set log2tpm matrix

Usage

```
log2tpm(object)

## S4 method for signature 'SummarizedExperiment'
log2tpm(object)

log2tpm(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
log2tpm(object) <- value
```

```
## S4 replacement method for signature 'SummarizedExperiment,numeric'
log2tpm(object) <- value
```

Arguments

object SummarizedExperiment
value log2tpm matrix (features x samples)

Value

log2tpm matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
log2tpm(object) <- values(object)
log2tpm(object)[1:3, 1:3]
```

log2transform	<i>Transform values</i>
---------------	-------------------------

Description

Transform values

Usage

```
log2transform(  
  object,  
  assay = assayNames(object)[1],  
  pseudo = 0,  
  verbose = FALSE  
)  
  
exp2(object, verbose = FALSE)  
  
zscore(object, verbose = FALSE)  
  
sscale(mat, verbose = FALSE)  
  
fscale(mat, verbose = FALSE)  
  
quantnorm(object, verbose = FALSE)  
  
invnorm(object, verbose = FALSE)  
  
vsn(object, verbose = FALSE, delog = TRUE)
```

Arguments

object	SummarizedExperiment
assay	character vector : assays for which to perform transformation
pseudo	number : pseudo value to be added prior to transformation
verbose	TRUE or FALSE : whether to msg
mat	matrix
delog	TRUE or FALSE (vsn)

Value

Transformed sumexp

Examples

```

file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)

object %>% plot_sample_densities()
invnorm(object) %>% plot_sample_densities()

object %>% plot_sample_densities()
quantnorm(object) %>% plot_sample_densities()

object %>% plot_sample_densities()
#vsn(object) %>% plot_sample_densities() # dataset too small

object %>% plot_sample_densities()
zscore(object) %>% plot_sample_densities()

object %>% plot_sample_densities()
exp2(object) %>% plot_sample_densities()
log2transform(exp2(object)) %>% plot_sample_densities()

```

logical2factor

logical to factor

Description

logical to factor

Usage

```

logical2factor(x, true = get_name_in_parent(x), false = paste0("not", true))

factor2logical(x)

```

Arguments

x	logical vector
true	string : truelevel
false	string : falselevel

Value

factor

Examples

```
t1up <- c( TRUE,  FALSE,  TRUE)
t1   <- c('flat', 'down', 'up' ) %>% factor(., .)
t1up
logical2factor(t1up)
factor2logical(t1)
```

make_alpha_palette	<i>Make alpha palette</i>
--------------------	---------------------------

Description

Make alpha palette

Usage

```
make_alpha_palette(object, alpha)
```

Arguments

object	SummarizedExperiment
alpha	string

Value

character vector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
make_alpha_palette(object, 'Time')
```

make_colors	<i>Make colors</i>
-------------	--------------------

Description

Make colors

Usage

```
make_colors(  
  varlevels,  
  sep = guess_sep(varlevels),  
  show = FALSE,  
  verbose = FALSE  
)
```

Arguments

- varlevels character vector
- sep string
- show TRUE or FALSE: whether to plot
- verbose TRUE or FALSE: whether to msg

Examples

```
make_colors(c('A', 'B', 'C', 'D' ), show = TRUE)  
make_colors(c('A.1', 'B.1', 'A.2', 'B.2'), show = TRUE)
```

make_volcano_dt	<i>Create volcano datatable</i>
-----------------	---------------------------------

Description

Create volcano datatable

Usage

```
make_volcano_dt(  
  object,  
  fit = fits(object)[1],  
  coefs = coefs(object, fit = fit)[1],  
  shape = "imputed",  
  size = NULL,  
  alpha = NULL,  
  label = "feature_id"  
)
```

Arguments

object	SummarizedExperiment
fit	'limma', 'lme', 'lm', 'wilcoxon'
coefs	character vector: coefs for which to plot volcanoes
shape	fvar or NULL
size	fvar or NULL
alpha	fvar or NULL
label	fvar or NULL

Value

data.table

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, impute = TRUE, fit = 'limma')
make_volcano_dt(object, fit = 'limma', coefs = 'Adult-X30dpt')
```

map_fvalues	<i>Map fvalues</i>
-------------	--------------------

Description

Map fvalues

Usage

```
map_fvalues(object, fvalues, from = "uniprot", to = "feature_id", sep = ";")
```

Arguments

object	SummarizedExperiment
fvalues	uncollapsed string vector
from	string (fvar)
to	string (svar)
sep	collapse separator

Value

string vector

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
fdt(object)
map_fvalues(object, c('Q6DHL5', 'Q6PFS7'), from = 'uniprot', to = 'feature_id', sep = ';')
```

matrix2sumexp	<i>Convert matrix into SummarizedExperiment</i>
---------------	---

Description

Convert matrix into SummarizedExperiment

Usage

```
matrix2sumexp(x, verbose = TRUE)
```

Arguments

x	matrix
verbose	TRUE/FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
x <- values(read_metabolon(file))
object <- matrix2sumexp(x)
object %<>% pca()
biplot(object, color = 'subgroup')
```

MAXQUANT_PATTERNS	<i>maxquant quantity patterns</i>
-------------------	-----------------------------------

Description

maxquant quantity patterns

Usage

```
MAXQUANT_PATTERNS
```

Format

An object of class character of length 7.

Examples

MAXQUANT_PATTERNS

mdsplot	<i>Feature correlations/distances</i>
---------	---------------------------------------

Description

Feature correlations/distances

Usage

```
mdsplot(distmat, title = NULL)

fcor(object, verbose = TRUE)

scor(object, verbose = TRUE)

fdist(object, method = "cor")

sdist(object, method = "cor")
```

Arguments

distmat	distance matrix
title	NULL or string
object	SummarizedExperiment
verbose	TRUE or FALSE
method	'cor', 'euclidian', etc

Value

matrix

Examples

```
# Correlations
object <- twofactor_sumexp()
scor(object) %>% pheatmap::pheatmap()
fcor(object) %>% pheatmap::pheatmap()
# Distances
sdist(object, 'cor') %>% mdsplot('samples: cor')
sdist(object, 'euclidian') %>% mdsplot('samples: euclidian')
```

```
fdist(object, 'cor')           %>% mdsplot('features: cor')
fdist(object, 'euclidian') %>% mdsplot('features: euclidian')
```

merge_compounddiscoverer	<i>merge compound discoverer files</i>
--------------------------	--

Description

merge compound discoverer files

Usage

```
merge_compounddiscoverer(x, quantity = NULL, verbose = TRUE)
```

Arguments

x	‘list‘
quantity	‘‘area’, ‘normalizedarea’‘
verbose	‘TRUE‘ or ‘FALSE‘

Value

‘data.table‘

merge_sample_excel	<i>Merge sample excel</i>
--------------------	---------------------------

Description

Merge sample excel

Usage

```
merge_sample_excel(
  object,
  sfile,
  range = NULL,
  by.x = "sample_id",
  by.y = "sample_id"
)
```

Arguments

object	SummarizedExperiment
sfile	sample file
range	string
by.x	string
by.y	string

Value

SummarizedExperiment

merge_sample_file	<i>Merge sample / feature file</i>
-------------------	------------------------------------

Description

Merge sample / feature file

Usage

```
merge_sample_file(  
  object,  
  sfile = NULL,  
  by.x = "sample_id",  
  by.y = "sample_id",  
  all.x = TRUE,  
  select = NULL,  
  stringsAsFactors = FALSE,  
  verbose = TRUE  
)  
  
merge_ffile(  
  object,  
  ffile = NULL,  
  by.x = "feature_id",  
  by.y = "feature_id",  
  all.x = TRUE,  
  select = NULL,  
  stringsAsFactors = FALSE,  
  verbose = TRUE  
)
```

Arguments

object	SummarizedExperiment
sfile	string : sample file path
by.x	string : object mergevar
by.y	string : file mergevvar
all.x	TRUE / FALSE : whether to keep samples / feature without annotation
select	character : [sf]file columns to select
stringsAsFactors	TRUE / FALSE
verbose	TRUE / FALSE
ffile	string : ffile path

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
subgroups <- c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00')
subgroups %<>% paste0('_STD')
object <- read_maxquant_proteingroups(file, subgroups = subgroups)
sfile <- paste0(tempdir(), '/', basename(tools::file_path_sans_ext(file)))
sfile %<>% paste0('.samples.txt')
dt <- data.table(sample_id = object$sample_id,
                 day = split_extract_fixed(object$subgroup, '_', 1))
data.table::fwrite(dt, sfile)
sdt(object)
sdt(merge_sample_file(object, sfile))
```

merge_sdata	<i>Merge sample/feature dt</i>
-------------	--------------------------------

Description

Merge sample/feature dt

Usage

```
merge_sdata(
  object,
  dt,
  by.x = "sample_id",
  by.y = names(dt)[1],
  all.x = TRUE,
```

```
    verbose = TRUE
  )

merge_sdt(
  object,
  dt,
  by.x = "sample_id",
  by.y = "sample_id",
  all.x = TRUE,
  verbose = TRUE
)

merge_fdata(
  object,
  dt,
  by.x = "feature_id",
  by.y = names(dt)[1],
  all.x = TRUE,
  verbose = TRUE
)

merge_fdt(
  object,
  dt,
  by.x = "feature_id",
  by.y = "feature_id",
  all.x = TRUE,
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
dt	data.frame, data.table, DataFrame
by.x	string : object mergevar
by.y	string : df mergevar
all.x	TRUE / FALSE : whether to keep samples / features without annotation
verbose	TRUE / FALSE : whether to msg

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
sdt(object)
```

```
sdt(merge_sdt(object, data.table(sample_id = object$sample_id,
                                number = seq_along(object$sample_id))))
```

message_df	<i>message dataframe</i>
------------	--------------------------

Description

message dataframe using sprintf syntax. Use place holder ' '

Usage

```
message_df(format_string, x)
```

Arguments

- format_string sprintf style format string
- x data.frame

Value

nothing returned

Examples

```
x <- data.frame(feature_id = c('F001', 'F002'), symbol = c('FEAT1', 'FEAT2'))
message_df('\t%s', x)

x <- c(rep('PASS', 25), rep('FAIL', 25))
message_df(format_string = '%s', table(x))
```

modelvar	<i>Get model variable</i>
----------	---------------------------

Description

Get model variable

Usage

```

modelvar(object, ...)

## S3 method for class 'data.table'
modelvar(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),
  ...
)

## S3 method for class 'SummarizedExperiment'
modelvar(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),
  ...
)

effectvar(
  object,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit)
)

tvar(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

pvar(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

fdrvar(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

abstractvar(object, ...)

## S3 method for class 'data.table'
abstractvar(
  object,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),
  ...
)

## S3 method for class 'SummarizedExperiment'
abstractvar(
  object,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),

```

```

    ...
)

modelvec(object, ...)

## S3 method for class 'data.table'
modelvec(
  object,
  quantity,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  ...
)

## S3 method for class 'SummarizedExperiment'
modelvec(
  object,
  quantity,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  ...
)

effectvec(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object)[1],
  fvar = "feature_id"
)

tvec(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id"
)

pvec(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id"
)

fdrvec(
  object,

```

```

    fit = fits(object)[1],
    coef = autonomics::coefs(object, fit = fit)[1],
    fvar = "feature_id"
  )

  abstractvec(object, ...)

## S3 method for class 'data.table'
abstractvec(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  ...
)

## S3 method for class 'SummarizedExperiment'
abstractvec(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  ...
)

modeldt(object, ...)

## S3 method for class 'data.table'
modeldt(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),
  ...
)

## S3 method for class 'SummarizedExperiment'
modeldt(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit),
  ...
)

effectdt(
  object,
  fit = fits(object),

```

```

    coef = autonomics::coefs(object, fit = fit)
  )

tdt(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

pdt(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

modelmat(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit)
)

modelmat(
  object,
  quantity,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit)
)

effectmat(
  object,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit)
)

effectsize(
  object,
  fit = fits(object),
  coef = autonomics::coefs(object, fit = fit)
)

tmat(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

pmat(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

fdrmat(object, fit = fits(object), coef = autonomics::coefs(object, fit = fit))

modelfeatures(object, ...)

## S3 method for class 'data.table'
modelfeatures(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  significancevar = "p",

```

```

    significance = 0.05,
    effectdirection = "<>",
    effectsizesize = 0,
    ...
)

## S3 method for class 'SummarizedExperiment'
modelfeatures(object, ...)

upfeatures(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  significancevar = "p",
  significance = 0.05,
  effectsizesize = 0
)

downfeatures(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  fvar = "feature_id",
  significancevar = "p",
  significance = 0.05,
  effectsizesize = 0
)

```

Arguments

object	data.table or SummarizedExperiment
...	S3 dispatch
quantity	'p', 'effect', 'fdr', 't', or 'se'
fit	string (vector)
coef	string (vector)
fvar	'feature_id' or other fvar for values (pvec) or names (upfeatures)
significancevar	'p' or 'fdr'
significance	p or fdr cutoff (fractional number)
effectdirection	'<>', '<' or '>'
effectsizesize	effectsizesize cutoff (positive number)

Value

string (tvar), matrix (tmat), numeric vector (tvec), character vector (tfeatures)

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
object %<>% fit_lm()

effectvar(object)
effectvec(object)[1:3]
effecttdt(object)[1:3, ]
effectmat(object)[1:3, ]

tvar(object)
tvec(object)[1:3]
tdt(object)[1:3, ]
tmat(object)[1:3, ]

pvar(object)
pvec(object)[1:3]
pdt(object)[1:3, ]
pmat(object)[1:3, ]

modelfeatures(object)
downfeatures(object)
upfeatures(object)

```

MSIGCOLLECTIONSHUMAN *Human/Mouse Msigdb Collections*

Description

Human/Mouse Msigdb Collections

Usage

MSIGCOLLECTIONSHUMAN

MSIGCOLLECTIONSMOUSE

Format

An object of class character of length 25.

An object of class character of length 13.

MSIGDIR	<i>local msigdb dir</i>
---------	-------------------------

Description

local msigdb dir

Usage

MSIGDIR

Format

An object of class character of length 1.

nfactors	<i>stri_split and extract</i>
----------	-------------------------------

Description

stri_split and extract

Usage

nfactors(x, sep = guess_sep(x))

split_extract_fixed(x, sep, i)

split_extract_regex(x, sep, i)

split_extract(x, i, sep = guess_sep(x))

Arguments

x	character vector
sep	string
i	integer

Value

character vector

Examples

```
# Read
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
x <- object$sample_id[1:5]
nfactors(x)

# Split
split_extract_fixed(x, '.', 1:2)
split_extract_fixed(x, '.', seq_len(nfactors(x)-1))
split_extract_fixed(x, '.', nfactors(x))
split_extract_fixed(fdt(object)$PUBCHEM, ';', 1) # with NA values
```

OPENTARGETSDIR	<i>opentargets dir</i>
----------------	------------------------

Description

opentargets dir

Usage

OPENTARGETSDIR

Format

An object of class character of length 1.

order_on_p	<i>Order on p</i>
------------	-------------------

Description

Order on p

Usage

```
order_on_p(
  object,
  fit = autonomics::fits(object),
  coefs = autonomics::coefs(object, fit = fit),
  combiner = "|",
  decreasing = FALSE,
  verbose = TRUE
)

order_on_t(
```

```

    object,
    fit = autonomics::fits(object),
    coefs = autonomics::coefs(object, fit = fit),
    combiner = "|",
    decreasing = FALSE,
    verbose = TRUE
)

order_on_effect(
  object,
  fit = autonomics::fits(object),
  coefs = autonomics::coefs(object, fit = fit),
  combiner = "|",
  verbose = TRUE
)

```

Arguments

object	SummarizedExperiment
fit	string vector: subset of 'fits(object)'
coefs	string vector: subset of 'coefs(object)'
combiner	' ' or '&'
decreasing	TRUE or FALSE
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```

# Linmod
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
order_on_p(object)
object %<>% fit_limma()
order_on_p(object)

# Survival
object <- survival_example()
object %<>% fit_survival()
order_on_p(object)

```

pca

PCA, SMA, LDA, PLS, SPLS, OPLS

Description

Perform a dimension reduction. Store sample scores, feature loadings, and dimension variances.

Usage

```
pca(  
  object,  
  by = "sample_id",  
  assay = assayNames(object)[1],  
  ndim = 2,  
  sep = FITSEP,  
  minvar = 0,  
  center_samples = TRUE,  
  verbose = TRUE,  
  plot = FALSE,  
  ...  
)
```

```
pls(  
  object,  
  by = "subgroup",  
  assay = assayNames(object)[1],  
  ndim = 2,  
  sep = FITSEP,  
  minvar = 0,  
  verbose = FALSE,  
  plot = FALSE,  
  ...  
)
```

```
sma(  
  object,  
  by = "sample_id",  
  assay = assayNames(object)[1],  
  ndim = 2,  
  sep = FITSEP,  
  minvar = 0,  
  verbose = TRUE,  
  plot = FALSE,  
  ...  
)
```

```
lda(  
  ...  
)
```

```

    object,
    assay = assayNames(object)[1],
    by = "subgroup",
    ndim = 2,
    sep = FITSEP,
    minvar = 0,
    verbose = TRUE,
    plot = FALSE,
    ...
)

spls(
  object,
  assay = assayNames(object)[1],
  by = "subgroup",
  ndim = 2,
  sep = FITSEP,
  minvar = 0,
  plot = FALSE,
  ...
)

opls(
  object,
  by = "subgroup",
  assay = assayNames(object)[1],
  ndim = 2,
  sep = FITSEP,
  minvar = 0,
  verbose = FALSE,
  plot = FALSE,
  ...
)

```

Arguments

object	SummarizedExperiment
by	svar or NULL
assay	string
ndim	number
sep	string
minvar	number
center_samples	TRUE/FALSE: center samples prior to pca ?
verbose	TRUE/FALSE: message ?
plot	TRUE/FALSE: plot ?
...	passed to biplot

Value

SummarizedExperiment

Author(s)

Aditya Bhagwat, Laure Cougnaud (LDA)

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
pca(object, plot = TRUE) # Principal Component Analysis
pls(object, plot = TRUE) # Partial Least Squares
lda(object, plot = TRUE) # Linear Discriminant Analysis
sma(object, plot = TRUE) # Spectral Map Analysis
spls(object, plot = TRUE) # Sparse PLS
# opl(s(object, plot = TRUE) # OPLS # outcommented because it produces a file named FALSE
```

pg_to_canonical	<i>proteingroup to isoforms</i>
-----------------	---------------------------------

Description

proteingroup to isoforms

Usage

```
pg_to_canonical(x, unique = TRUE)

pg_to_isoforms(x, unique = TRUE)
```

Arguments

x	proteingroups string vector
unique	whether to remove duplicates

Value

string vector

Examples

```
(x <- c('Q96JP5;Q96JP5-2', 'Q96JP5', 'Q96JP5-2;P86791'))
pg_to_isoforms(x)
pg_to_canonical(x)
pg_to_isoforms(x, unique = FALSE)
pg_to_canonical(x, unique = FALSE)
# .pg_to_isoforms(x[1]) # unexported dot functions
# .pg_to_canonical(x[1]) # operate on scalars
```

plot_coef_densities	<i>Plot contrast densities</i>
---------------------	--------------------------------

Description

Plot contrast densities

Usage

```
plot_coef_densities(  
  object,  
  fit = fits(object)[1],  
  coefs = autonomics::coefs(object, fit = fit),  
  sep = FITSEP,  
  label = "feature_id"  
)
```

Arguments

object	SummarizedExperiment
fit	'limma', 'lm', 'lme', 'lmer', or 'wilcoxon'
coefs	character vector
sep	string
label	svar

Value

ggplot

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')  
object <- read_metabolon(file)  
object %<>% fit_limma(~subgroup, block = 'Subject')  
plot_coef_densities(object)
```

plot_contrastogram	<i>Plot contrastogram</i>
--------------------	---------------------------

Description

Plot contrastogram

Usage

```
plot_contrastogram(  
  object,  
  subgroupvar,  
  formula = as.formula(paste0("~ 0 +", subgroupvar)),  
  colors = make_colors(slevels(object, subgroupvar), guess_sep(object)),  
  curve = 0.1  
)
```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup svar
formula	formula
colors	named color vector (names = subgroups)
curve	arrow curvature

Value

list returned by [plotmat](#)

Examples

```
if (requireNamespace('diagram', quietly = TRUE)){  
  file <- download_data('halama18.metabolon.xlsx')  
  object <- read_metabolon(file)  
  plot_contrastogram(object, subgroupvar = 'subgroup')  
}
```

plot_contrast_venn	<i>Plot contrast venn</i>
--------------------	---------------------------

Description

Plot contrast venn

Usage

```
plot_contrast_venn(issig, colors = NULL)
```

Arguments

issig	matrix(nrow, ncontrast): -1 (down), +1 (up)
colors	NULL or colorvector

Value

nothing returned

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_wilcoxon(~ subgroup, block = 'Subject')
object %<>% fit_limma( ~ subgroup, block = 'Subject', codingfun = contr.treatment.explicit)
isfdr <- is_sig(object, contrast = 't3-t0', quantity = 'p', fit = fits(object))
plot_contrast_venn(isfdr)
```

plot_data	<i>Plot data</i>
-----------	------------------

Description

Plot data

Usage

```
plot_data(
  data,
  geom = geom_point,
  color = NULL,
  fill = NULL,
  linetype = NULL,
  ...,
  palette = NULL,
  fixed = list(),
  theme = list()
)
```

Arguments

data	data.frame'
geom	geom_point, etc.
color	variable mapped to color (symbol)
fill	variable mapped to fill (symbol)
linetype	variable mapped to linetype (symbol)
...	mapped aesthetics
palette	color palette (named character vector)
fixed	fixed aesthetics (list)
theme	list with ggplot theme specifications

Value

ggplot object

Author(s)

Aditya Bhagwat, Johannes Graumann

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% pca()
data <- sdt(object)
plot_data(data, x = `t~sample_id~pca1`, y = `t~sample_id~pca2`)
plot_data(data, x = `t~sample_id~pca1`, y = `t~sample_id~pca2`, color = subgroup)
plot_data(data, x = `t~sample_id~pca1`, y = `t~sample_id~pca2`, color = NULL)
fixed <- list(shape = 15, size = 3)
plot_data(data, x = `t~sample_id~pca1`, y = `t~sample_id~pca2`, fixed = fixed)
```

plot_densities	<i>Plot sample/feature distributions</i>
----------------	--

Description

Plot sample/feature distributions

Usage

```
plot_densities(
  object,
  assay = assayNames(object)[1],
  group,
  fill,
  color = NULL,
  linetype = NULL,
  facet = NULL,
  nrow = NULL,
  ncol = NULL,
  dir = "h",
  scales = "free_y",
  labeller = label_value,
  palette = NULL,
  fixed = list(alpha = 0.8, na.rm = TRUE)
)

plot_sample_densities(
  object,
  assay = assayNames(object)[1],
```

```

    group = "sample_id",
    fill = if ("subgroup" %in% svars(object)) "subgroup" else "sample_id",
    color = NULL,
    linetype = NULL,
    n = 100,
    facet = NULL,
    nrow = NULL,
    ncol = NULL,
    dir = "h",
    scales = "free_y",
    labeller = label_value,
    palette = NULL,
    fixed = list(alpha = 0.8, na.rm = TRUE)
  )

plot_feature_densities(
  object,
  assay = assayNames(object)[1],
  fill = "feature_id",
  group = fill,
  color = NULL,
  linetype = NULL,
  n = 9,
  facet = NULL,
  nrow = NULL,
  ncol = NULL,
  dir = "h",
  scales = "free",
  labeller = label_value,
  palette = NULL,
  fixed = list(alpha = 0.8, na.rm = TRUE)
)

```

Arguments

object	SummarizedExperiment
assay	string
group	svar (string)
fill	svar (string)
color	svar (string)
linetype	svar (string)
facet	svar (character vector)
nrow	number of facet rows
ncol	number of facet cols
dir	'h' (horizontal) or 'v' (vertical)
scales	'free', 'fixed', 'free_y'

labeller	e.g. label_value
palette	named character vector
fixed	fixed aesthetics
n	number

Value

ggplot object

See Also

[plot_sample_violins](#), [plot_sample_boxplots](#)

Examples

```
# Data
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% extract(, order(.$subgroup))

# Sample distributions
plot_sample_densities(object)
plot_sample_violins( object, facet = 'Time')
plot_sample_boxplots(object)
plot_exprs(object)
plot_exprs(object, dim = 'samples', x = 'subgroup', facet = 'Time')

# Feature distributions
plot_feature_densities(object)
plot_feature_violins( object)
plot_feature_boxplots( object)
```

plot_design	<i>Plot model</i>
-------------	-------------------

Description

Plot model

Usage

```
plot_design(object, codingfun = contr.treatment.explicit)
```

Arguments

object	<code>SummarizedExperiment</code>
codingfun	factor coding function <code>contr.treatment</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_0$ <code>contr.treatment.explicit</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_0$ <code>code_control</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_0$ <code>contr.diff</code>: $\text{intercept} = y_0$, $\text{coefi} = y_i - y_{(i-1)}$ <code>code_diff</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{(i-1)}$ <code>code_diff_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{(i+)}$ <code>code_deviation</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{\text{mean}}$ (drop last) <code>code_deviation_first</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - y_{\text{mean}}$ (drop first) <code>code_helmert</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - \text{mean}(y_0:(y_{i-1}))$ <code>code_helmert_forward</code>: $\text{intercept} = y_{\text{mean}}$, $\text{coefi} = y_i - \text{mean}(y_{(i+1):y_p})$

Value

ggplot

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
subgroups <- paste0(c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'), '_STD')
object <- read_maxquant_proteingroups(file, subgroups = subgroups)
object$subgroup %<>% substr(1,3)
plot_design(object)
```

plot_detections	<i>Plot missingness per sample / subgroup</i>
-----------------	---

Description

`plot_sample_nas` shows systematic and random missingness (white), and full detection (bright color) at sample resolution. Imputations are also shown (light color).

Usage

```
plot_detections(...)

plot_summarized_detections(...)

plot_sample_nas(
  object,
  by = "subgroup",
  fill = by,
  palette = make_svar_palette(object, fill),
```

```

    axis.text.y = element_blank()
  )

  plot_subgroup_nas(
    object,
    by = "subgroup",
    fill = by,
    palette = NULL,
    na_imputes = TRUE
  )

```

Arguments

...	used to maintain deprecated functions
object	SummarizedExperiment
by	svar (string)
fill	svar (string)
palette	color vector (names = levels, values = colors)
axis.text.y	passed to ggplot2::theme
na_imputes	TRUE or FALSE

Details

plot_subgroup_nas shows systematic missingness at subgroup resolution. Random missingness and full detection are shown together (bright color). Imputations are also shown (light color).

Value

ggplot object

Examples

```

file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
plot_sample_nas(object)
plot_sample_nas(impute(object))
plot_subgroup_nas(object)
plot_subgroup_nas(impute(object))

subgroups <- sprintf('%s_STD', c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'))
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, subgroups = subgroups)
plot_subgroup_nas(object)
plot_subgroup_nas(object, 'subgroup')
plot_sample_nas(object)
plot_sample_nas(object, 'subgroup')

```

plot_exprs	<i>Plot exprs for coef</i>
------------	----------------------------

Description

Plot exprs for coef

Usage

```
plot_exprs(
  object,
  dim = "both",
  assay = assayNames(object)[1],
  features = NULL,
  fit = fits(object)[1],
  coefs = autonomics::coefs(object, fit = fit),
  block = NULL,
  x = default_x(object, dim),
  geom = default_geom(object, x = x, block = block),
  color = x,
  fill = x,
  shape = NULL,
  size = NULL,
  alpha = NULL,
  linetype = NULL,
  highlight = NULL,
  combiner = "|",
  p = 1,
  fdr = 1,
  facet = if (dim == "both") "feature_id" else NULL,
  file = NULL,
  width = 7,
  height = 7,
  n = if (is.null(file)) 4 else 12,
  ncol = if (is.null(file)) NULL else 3,
  nrow = if (is.null(file)) NULL else 4,
  scales = "free_y",
  labeller = "label_value",
  pointsize = if (is.null(block)) 0 else 0.5,
  jitter = if (is.null(block)) 0.1 else 0,
  fillpalette = make_var_palette(object, fill),
  colorpalette = make_var_palette(object, color),
  hlevels = NULL,
  title = switch(dim, both = x, features = "Feature Boxplots", samples =
    "Sample Boxplots"),
  subtitle = if (!is.null(fit)) coefs else "",
  xlab = x,
```

```

    ylab = "value",
    theme = ggplot2::theme(plot.title = element_text(hjust = 0.5)),
    verbose = TRUE
)

plot_sample_boxplots(
  object,
  fill = if ("subgroup" %in% svars(object)) "subgroup" else "sample_id",
  n = min(ncol(object), 16),
  ...
)

plot_feature_boxplots(object, ...)

```

Arguments

object	SummarizedExperiment
dim	'samples' (per-sample distribution across features), 'features' (per-feature distribution across samples) or 'both' (subgroup distribution faceted per feature)
assay	string: value in assayNames(object)
features	features to plot no matter what (character vector)
fit	'limma', 'lm', 'lme', 'lmer', 'wilcoxon'
coefs	subset of coefs(object) to consider in selecting top
block	group svar
x	x svar
geom	'boxplot' or 'point'
color	color svar: points, lines
fill	fill svar: boxplots
shape	shape svar
size	size svar
alpha	alpha svar
linetype	linetype svar
highlight	highlight svar
combiner	'&' or ' '
p	fraction: p cutoff
fdr	fraction: fdr cutoff
facet	string: fvar mapped to facet
file	NULL or filepath
width	inches
height	inches

n	number of samples (dim = 'samples') or features (dim = 'features' or 'both') to plot
ncol	number of cols in faceted plot (if dim = 'both')
nrow	number of rows in faceted plot (if dim = 'both')
scales	'free_y', 'free_x', 'fixed'
labeller	string or function
pointsize	number
jitter	jitter width (number)
fillpalette	named character vector: fill palette
colorpalette	named character vector: color palette
hlevels	xlevels for which to plot hlines
title	string
subtitle	string
xlab	string
ylab	string
theme	ggplot2::theme(...) or NULL
verbose	TRUE or FALSE
...	used to maintain deprecated functions

Value

ggplot object

See Also

[plot_sample_densities](#), [plot_sample_violins](#)

Examples

```
# Without limma
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
plot_exprs(object, block = 'Subject', title = 'Subgroup Boxplots')
plot_exprs(object, dim = 'samples')
plot_exprs(object, dim = 'features', block = 'sample_id')

# With limma
object %<>% fit_limma(block = 'Subject')
plot_exprs(object, block = 'Subject')
plot_exprs(object, block = 'Subject', coefs = c('t1-t0', 't2-t0', 't3-t0'))
plot_exprs_per_coef(object, x = 'Time', block = 'Subject')

# Points
plot_exprs(object, geom = 'point', block = 'Subject')

# Add highlights
controlfeatures <- c('biotin', 'phosphate')
fdt(object) %<>% cbind(control = .$feature_name %in% controlfeatures)
plot_exprs(object, dim = 'samples', highlight = 'control')

# Multiple pages
plot_exprs(object, block = 'Subject', n = 4, nrow = 1, ncol = 2)
```

plot_exprs_per_coef	<i>Plot exprs per coef</i>
---------------------	----------------------------

Description

Plot exprs per coef

Usage

```
plot_exprs_per_coef(  
  object,  
  fit = fits(object)[1],  
  coefs = autonomics::coefs(object, fit = fit),  
  x = default_x(object),  
  block = NULL,  
  geom = default_geom(object, x, block = block),  
  orderbyp = FALSE,  
  title = x,  
  subtitle = default_subtitle(fit, x, coefs),  
  n = 1,  
  nrow = 1,  
  ncol = NULL,  
  theme = ggplot2::theme(legend.position = "bottom", legend.title = element_blank(),  
    plot.title = element_text(hjust = 0.5), plot.subtitle = element_text(hjust = 0.5))  
)
```

Arguments

object	SummarizedExperiment
fit	'limma', 'lm', 'lme', 'lmer', 'wilcoxon'
coefs	subset of coefs(object) to consider in selecting top
x	x svar
block	group svar
geom	'boxplot' or 'point'
orderbyp	TRUE or FALSE
title	string
subtitle	string
n	number
nrow	number of rows in faceted plot
ncol	number of cols in faceted plot
theme	ggplot2::theme(...) or NULL

Value

ggplot object

See Also

[plot_sample_densities](#), [plot_sample_violins](#)

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
object %<>% pls(by = 'subgroup')
object %<>% pls(by = 'Diabetes')
object %<>% pls(by = 'Subject')
plot_exprs_per_coef(object)
plot_exprs_per_coef(object, orderbyp = TRUE)
plot_exprs_per_coef(object, fit = 'pls1', block = 'Subject')
```

plot_fit_summary	<i>Plot fit summary</i>
------------------	-------------------------

Description

Plot fit summary

Usage

```
plot_fit_summary(sumdt, nrow = NULL, ncol = NULL, order = FALSE)
```

Arguments

sumdt	data.table
nrow	number
ncol	number
order	TRUE or FALSE

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_lm()
object %<>% fit_limma(block = 'Subject')
sumdt <- summarize_fit(object, coefs = c('t1-t0', 't2-t0', 't3-t0'))
plot_fit_summary(sumdt)
```

plot_heatmap	<i>Plot heatmap</i>
--------------	---------------------

Description

Plot heatmap

Usage

```
plot_heatmap(
  object,
  fit = fits(object)[1],
  coef = autonomics::coefs(object, fit = fit)[1],
  effectsize = 0,
  p = 1,
  fdr = 0.05,
  n = 100,
  assay = assayNames(object)[1],
  cluster_features = FALSE,
  cluster_samples = FALSE,
  flabel = intersect(c("gene", "feature_id"), fvars(object))[1],
  group = "subgroup",
  verbose = TRUE,
  title = NULL
)
```

Arguments

object	SummarizedExperiment
fit	'limma', 'lm', 'lme(r)', 'wilcoxon'
coef	string: one of coefs(object)
effectsize	number: effectsize filter
p	number: p filter
fdr	number: fdr filter
n	number: n filter
assay	string: one of assayNames(object)
cluster_features	TRUE or FALSE
cluster_samples	TRUE or FALSE
flabel	string: feature label
group	sample groupvar
verbose	TRUE or FALSE
title	string

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, fit = 'limma')
plot_heatmap(object)
```

plot_joint_density	<i>Plot joint density</i>
--------------------	---------------------------

Description

Plot joint density

Usage

```
plot_joint_density(
  object,
  xvar,
  yvar,
  color = TRUE,
  contour = TRUE,
  smooth = TRUE
)
```

Arguments

object	SummarizedExperiment
xvar	svar
yvar	svar
color	TRUE or FALSE
contour	TRUE or FALSE
smooth	TRUE or FALSE

Value

ggplot

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
set.seed(20)
object$Height <- rnorm(ncol(object), mean = 176)
object$Weight <- rnorm(ncol(object), mean = 85.4)
plot_joint_density(object, 'Height', 'Weight')
plot_joint_density(object, 'Height', 'Weight', smooth = TRUE)
plot_joint_density(object, 'Height', 'Weight', color = TRUE)
plot_joint_density(object, 'Height', 'Weight', contour = TRUE)
```

plot_matrix	<i>Plot binary matrix</i>
-------------	---------------------------

Description

Plot binary matrix

Usage

plot_matrix(mat)

Arguments

mat matrix

Value

no return (base R plot)

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
mat <- sdt(object)[, .(Subject, subgroup)]
mat$present <- 1
mat %<>% data.table::dcast(Subject ~ subgroup, value.var = 'present', fill = 0)
mat %<>% dt2mat()
plot_matrix(mat)
```

plot_subgroup_points	<i>Plot features</i>
----------------------	----------------------

Description

Plot features

Usage

```
plot_subgroup_points(
  object,
  subgroup = "subgroup",
  block = NULL,
  x = subgroup,
  color = subgroup,
  group = block,
  facet = "feature_id",
```

```

nrow = NULL,
scales = "free_y",
...,
palette = NULL,
fixed = list(na.rm = TRUE),
theme = list(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1))
)

```

Arguments

object	SummarizedExperiment
subgroup	subgroup svar
block	block svar
x	svar mapped to x
color	svar mapped to color
group	svar mapped to group
facet	svar mapped to facets
nrow	number of rows
scales	'free_y' etc.
...	mapped aesthetics
palette	color palette (named character vector)
fixed	fixed aesthetics
theme	ggplot theme specifications

Value

ggplot object

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma')
idx <- order(fdata(object)$`p~t1-t0~limma`)[1:9]
object %<>% extract(idx, )
plot_sample_boxplots( object)
plot_feature_boxplots( object)
plot_sample_boxplots(object, x = 'Time')
plot_subgroup_points( object, subgroup = 'Time')
plot_subgroup_points( object, subgroup = 'Time', block = 'Subject')

```

plot_summary	<i>Plot summary</i>
--------------	---------------------

Description

Plot summary

Usage

```
plot_summary(  
  object,  
  fit = "limma",  
  formula = default_formula(object),  
  block = NULL,  
  label = "feature_id",  
  palette = make_svar_palette(object, svar = svar)  
)
```

Arguments

object	SummarizedExperiment
fit	linmod engine : 'limma', 'lm', 'lme', 'lmer' or 'wilcoxon'
formula	model formula
block	NULL or svar
label	fvar
palette	NULL or colorvector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')  
object <- read_metabolon(file)  
object %<>% pca()  
object %<>% pls(by = 'subgroup')  
object %<>% fit_limma()  
plot_summary(object, block = 'Subject')
```

plot_survival	<i>Plot survival</i>
---------------	----------------------

Description

Plot survival

Usage

```
plot_survival(
  object,
  assay = assayNames(object)[1],
  engine = intersect(fits(object), c("coxph", "survdif", "logrank")),
  ntile = 2,
  title = sprintf("surv ~ expr"),
  subtitle = sprintf("%s", paste0(engine, collapse = " ")),
  file = NULL,
  width = 7,
  height = 7,
  n = min(nrow(object), 9),
  ncol = 3,
  nrow = 3
)
```

Arguments

object	SummarizedExperiment
assay	value in assayNames(object)
engine	'coxph', 'survdif' or 'logrank'
ntile	number of quantiles
title	string
subtitle	string
file	filepath
width	number
height	number
n	number of features to plot
ncol	number of columns
nrow	number of rows

Value

ggplot

Examples

```
# Defaults
object <- survival_example()
object %<>% fit_survival()
plot_survival(object)

# Engines
object <- survival_example()
object %<>% fit_survival(engine = c('coxph', 'survdif', 'logrank'))
plot_survival(object)

# Pdf
# plot_survival(object, file = file.path('testdir', 'survival', 'survival.pdf'))
```

plot_venn	<i>Plot venn</i>
-----------	------------------

Description

Plot venn

Usage

plot_venn(x)

Arguments

x list

Examples

```
x <- list(roundfruit = c('apple', 'orange'), redfruit = c('apple', 'strawberry'))
plot_venn(x)
```

plot_venn_heatmap	<i>Plot venn heatmap</i>
-------------------	--------------------------

Description

Plot venn heatmap

Usage

plot_venn_heatmap(x)

Arguments

x list

Examples

```
x <- list(roundfruit = c('apple', 'orange'), redfruit = c('apple', 'strawberry'))
plot_venn_heatmap(x)
```

plot_violins	<i>Plot sample/feature violins</i>
--------------	------------------------------------

Description

Plot sample/feature violins

Usage

```
plot_violins(  
  object,  
  assay = assayNames(object)[1],  
  x,  
  fill,  
  color = NULL,  
  group = NULL,  
  facet = NULL,  
  nrow = NULL,  
  ncol = NULL,  
  dir = "h",  
  scales = "free",  
  labeller = label_value,  
  highlight = NULL,  
  palette = NULL,  
  fixed = list(na.rm = TRUE)  
)
```

```
plot_feature_violins(  
  object,  
  assay = assayNames(object)[1],  
  x = "feature_id",  
  fill = "feature_id",  
  color = NULL,  
  n = 9,  
  facet = NULL,  
  nrow = NULL,  
  ncol = NULL,  
  dir = "h",  
  scales = "free",  
  labeller = label_value,  
  highlight = NULL,  
  fixed = list(na.rm = TRUE)  
)
```

```
plot_sample_violins(  
  object,  
  assay = assayNames(object)[1],
```

```

    x = "sample_id",
    fill = if ("subgroup" %in% svars(object)) "subgroup" else "sample_id",
    color = NULL,
    n = 100,
    facet = NULL,
    nrow = NULL,
    ncol = NULL,
    dir = "h",
    scales = "free",
    labeller = label_value,
    highlight = NULL,
    fixed = list(na.rm = TRUE)
  )

plot_subgroup_violins(
  object,
  assay = assayNames(object)[1],
  subgroup,
  x = "subgroup",
  fill = "subgroup",
  color = NULL,
  highlight = NULL,
  facet = "feature_id",
  fixed = list(na.rm = TRUE)
)

```

Arguments

object	SummarizedExperiment
assay	string
x	svar (string)
fill	svar (string)
color	svar (string)
group	svar (string)
facet	svar (character vector)
nrow	NULL or number
ncol	NULL or number
dir	'h' or 'v' : are facets filled horizontally or vertically ?
scales	'free', 'free_x', 'free_y', or 'fixed'
labeller	label_both or label_value
highlight	fvar expressing which feature should be highlighted (string)
palette	named color vector (character vector)
fixed	fixed aesthetics
n	number
subgroup	subgroup svar

Value

ggplot object

See Also

[plot_exprs](#), [plot_densities](#)

Examples

```
# data
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% extract(, order(.$subgroup))
control_features <- c('biotin','phosphate')
fdata(object) %<>% cbind(control = .$feature_name %in% control_features)

# plot
plot_violins(object[1:12, ], x = 'feature_id', fill = 'feature_id')
plot_feature_violins(object[1:12, ])
plot_sample_violins(object[, 1:12], highlight = 'control')
plot_subgroup_violins(object[1:4, ], subgroup = 'subgroup')
```

plot_volcano	<i>Plot volcano</i>
--------------	---------------------

Description

Plot volcano

Usage

```
plot_volcano(
  object,
  fit = fits(object)[1],
  coefs = autonomics::coefs(object, fit = fit)[1],
  facet = if (is_scalar(fit)) "coef" else c("fit", "coef"),
  scales = "fixed",
  shape = if ("imputed" %in% fvars(object)) "imputed" else NULL,
  size = NULL,
  alpha = NULL,
  label = "feature_id",
  max.overlaps = 10,
  features = NULL,
  nrow = length(fit),
  p = 0.05,
  fdr = 0.05,
  n = Inf,
  xndown = NULL,
  xnup = NULL,
```

```

    title = NULL,
    file = NULL,
    width = 7,
    height = 7,
    verbose = TRUE
  )

```

Arguments

object	SummarizedExperiment
fit	'limma', 'lme', 'lm', 'wilcoxon'
coefs	character vector
facet	character vector
scales	'free', 'fixed', etc.
shape	fvar (string)
size	fvar (string)
alpha	fvar (string)
label	fvar (string)
max.overlaps	number: passed to ggrepel
features	feature ids (character vector): features to encircle
nrow	number: no of rows in plot
p	number: p cutoff for labeling
fdr	number: fdr cutoff for labeling
n	number: n cutoff for labeling
xndown	x position of ndown labels
xnup	x position of nup labels
title	string or NULL
file	filename
width	number
height	number
verbose	TRUE or FALSE

Value

ggplot object

Examples

```

# Regular Usage
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
object %<>% fit_lm()

```

```

plot_volcano(object, coefs = 't3-t0', fit = 'limma')           # single contrast
plot_volcano(object, coefs = c('t2-t0', 't3-t0'), fit = 'limma') # multip contrasts
plot_volcano(object, coefs = c('t2-t0', 't3-t0'), fit = c('limma', 'lm')) # multip contrs & methods

# When nothing passes FDR
fdr(object) %<>% add_adjusted_pvalues('fdr', fit = 'limma', coefs = 't3-t0')
object %<>% extract( fdrvec(object, fit = 'limma', coef = 't3-t0') > 0.05, )
plot_volcano(object, coefs = 't3-t0', fit = 'limma')

# Additional mappings
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, impute = TRUE)
object %<>% fit_limma()
plot_volcano(object)
plot_volcano(object, label = 'gene')
plot_volcano(object, label = 'gene', size = 'log2maxlfq')
plot_volcano(object, label = 'gene', size = 'log2maxlfq', alpha = 'pepcounts')
plot_volcano(object, label = 'gene', features = c('Q503D2_DANRE'))
plot_volcano(object, label = 'gene', features = list(c('Q503D2_DANRE', 'Q6DGK4_DANRE'),
c('Q6DGK4_DANRE', 'F1Q7L0_DANRE'))))

```

```
PRECURSOR_QUANTITY      diann precursor quantity
```

Description

diann precursor quantity

Usage

```
PRECURSOR_QUANTITY
```

Format

An object of class character of length 1.

```
preprocess_rnaseq_counts
      Preprocess RNAseq counts
```

Description

Preprocess RNAseq counts

Usage

```
preprocess_rnaseq_counts(  
  object,  
  formula = ~subgroup,  
  block = NULL,  
  min_count = 10,  
  pseudo = 0.5,  
  tpm = FALSE,  
  cpm = TRUE,  
  voom = TRUE,  
  log2 = TRUE,  
  verbose = TRUE,  
  plot = TRUE  
)
```

Arguments

object	SummarizedExperiment
formula	designmat formula
block	block svar
min_count	min count required in some samples
pseudo	added pseudocount to avoid $\log(x)=-\infty$
tpm	TRUE or FALSE : tpm normalize?
cpm	TRUE or FALSE : cpm normalize? (counts per million (scaled) reads)
voom	TRUE or FALSE : voom weight?
log2	TRUE or FALSE : log2 transform?
verbose	TRUE or FALSE : msg?
plot	TRUE or FALSE : plot?

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')  
object <- .read_rnaseq_counts(file)  
object$subgroup  
object %<>% preprocess_rnaseq_counts()
```

pull_columns	<i>Pull columns in a dataframe to the front</i>
--------------	---

Description

Pull columns in a dataframe to the front

Usage

```
pull_columns(df, first_cols, verbose = TRUE)
```

Arguments

df	data.frame
first_cols	character vector: columns to be pulled to the front
verbose	TRUE (default) or FALSE

Value

dataframe with re-ordered columns

Examples

```
df <- data.frame(
  symbol = c('A1BG', 'A2M'),
  id     = c('1', '2'),
  name   = c('alpha-1-B glycoprotein', 'alpha-2-macroglobulin'),
  type   = c('proteinencoding', 'proteinencoding'))
first_cols <- c('id', 'symbol', 'location', 'uniprot')
pull_columns(df, first_cols)
```

read_affymetrix	<i>Read affymetrix microarray</i>
-----------------	-----------------------------------

Description

Read affymetrix microarray

Usage

```
read_affymetrix(celfiles)
```

Arguments

celfiles	string vector: CEL file paths
----------	-------------------------------

Value

RangedSummarizedExperiment

Examples

```
# Downloading example dataset fails 600s limit - example outcommented.
# url <- paste0('http://www.bioconductor.org/help/publications/2003/Chiaretti/chiaretti2/T33.tgz')
# localdir <- file.path(tools::R_user_dir('autonomics', 'cache'), 'T33')
# dir.create(localdir, showWarnings = FALSE)
# localfile <- file.path(localdir, basename(url))
# if (!file.exists(localfile)){ download.file(url, destfile = localfile)
#                               untar(localfile, exdir = path.expand(localdir)) }
# localfile %<>% substr(1, nchar(.)-4)
# if (!requireNamespace("BiocManager", quietly = TRUE)) install.packages('BiocManager')
# if (!requireNamespace("hgu95av2.db", quietly = TRUE)) BiocManager::install('hgu95av2.db')
# read_affymetrix(celfiles = list.files(localfile, full.names = TRUE))
```

read_compounddiscoverer

Read compound discoverer output

Description

Read compound discoverer output

Usage

```
read_compounddiscoverer(
  dir = getwd(),
  files = list.files(path = dir, pattern = "(RP|HILIC).*\\.csv$", full.names = TRUE),
  colname_regex = "^(.*)\\d{8,8}_+(.*)+((HILIC|RP)(NEG|POS))\\.raw.*$",
  colname_format = function(x) stringi::stri_replace_first_regex(x, colname_regex,
    "$1$2", opts_regex = stringi::stri_opts_regex(case_insensitive = TRUE)),
  mod_extract = function(x) stringi::stri_subset_regex(x, colname_regex, opts_regex =
    stringi::stri_opts_regex(case_insensitive = TRUE)) %>%
    stringi::stri_replace_first_regex(colname_regex, "$3", opts_regex =
    stringi::stri_opts_regex(case_insensitive = TRUE)),
  quantity = NULL,
  nonames = FALSE,
  exclude_sname_pattern = "(blank|QC|RS)",
  subgroups = NULL,
  logbase = 2,
  impute = FALSE,
  plot = FALSE,
  label = "feature_id",
  pca = plot,
  pls = plot,
```

```

    fit = if (plot) "limma" else NULL,
    formula = ~subgroup,
    block = NULL,
    coefs = NULL,
    contrasts = NULL,
    palette = NULL,
    verbose = TRUE
  )

```

Arguments

dir	compound discoverer output directory
files	compound discoverer output files
colname_regex	regular expression to parse sample names from column names
colname_format	function to reformat column names
mod_extract	function to extract MS modi from sample names
quantity	'area', 'normalizedarea' or NULL
nonames	TRUE or FALSE: retain compounds without Names?
exclude_sname_pattern	regular expression of sample names to exclude
subgroups	NULL or string vector : subgroups to retain
logbase	base for logarithmization of the data
impute	TRUE or FALSE: impute group-specific NA values?
plot	TRUE or FALSE: plot ?
label	fvar
pca	TRUE or FALSE: run pca ?
pls	TRUE or FALSE: run pls ?
fit	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
formula	model formula
block	model blockvar: string or NULL
coefs	model coefficients of interest: character vector or NULL
contrasts	coefficient contrasts of interest: character vector or NULL
palette	color palette : named character vector
verbose	TRUE or FALSE : message ?

Value

SummarizedExperiment

read_fragpipe	<i>Read fragpipe</i>
---------------	----------------------

Description

Read fragpipe

Usage

```
read_fragpipe(  
  dir = getwd(),  
  file = if (is_file(dir)) dir else file.path(dir, "combined_protein.tsv"),  
  contaminants = FALSE,  
  verbose = TRUE  
)
```

Arguments

- dir directory with 'combined_protein.tsv'
- file 'combined_protein.tsv' (full path)
- contaminants whether to include contaminants
- verbose whether to msg

Value

SummarizedExperiment

Examples

```
file <- download_data('multiorganism.combined_protein.tsv')  
object <- read_fragpipe(file = file)  
object  
fdt(object)  
sdt(object)
```

read_maxquant_phosphosites	<i>Read maxquant phosphosites</i>
----------------------------	-----------------------------------

Description

Read maxquant phosphosites

Usage

```

read_maxquant_phosphosites(
  dir = getwd(),
  fosfile = if (is_file(dir)) dir else file.path(dir, "phospho (STY)Sites.txt"),
  profile = file.path(dirname(fosfile), "proteinGroups.txt"),
  fastafile = NULL,
  restapi = FALSE,
  quantity = NULL,
  subgroups = NULL,
  invert = character(0),
  rm_contaminants = TRUE,
  rm_reverse = TRUE,
  rm_missing_in_all_samples = TRUE,
  localization = 0.75,
  impute = FALSE,
  plot = FALSE,
  label = "feature_id",
  pca = plot,
  pls = plot,
  fit = if (plot) "limma" else NULL,
  formula = as.formula("~ subgroup"),
  block = NULL,
  coefs = NULL,
  contrasts = NULL,
  palette = NULL,
  verbose = TRUE
)

read_phosphosites(...)

```

Arguments

dir	proteingroups directory
fosfile	phosphosites file
profile	proteingroups file
fastafile	uniprot fastafile
restapi	TRUE or FALSE : annotate non-fastadt uniprot using uniprot restapi
quantity	'normalizedratio', 'ratio', 'correctedreporterintensity', 'reporterintensity', 'maxlfq', 'labeledintensity', 'intensity' or NULL
subgroups	NULL or string vector : subgroups to retain
invert	string vector: subgroups which require inversion
rm_contaminants	TRUE or FALSE: rm contaminants ?
rm_reverse	TRUE or FALSE: rm reverse proteins ?
rm_missing_in_all_samples	TRUE or FALSE

localization	number: min localization probability (for phosphosites)
impute	TRUE or FALSE: impute group-specific NA values?
plot	TRUE or FALSE
label	fvar
pca	TRUE or FALSE: run pca ?
pls	TRUE or FALSE: run pls ?
fit	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
formula	model formula
block	model blockvar: string or NULL
coefs	model coefficients of interest: string vector or NULL
contrasts	model coefficient contrasts of interest: string vector or NULL
palette	color palette: named string vector
verbose	TRUE or FALSE: message ?
...	maintain deprecated functions

Value

SummarizedExperiment

Examples

```
profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
fastafile <- system.file('extdata/uniprot_hsa_20140515.fasta', package = 'autonomics')
subgroups <- sprintf('%s_STD', c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'))
pro <- read_maxquant_proteingroups(file = profile, subgroups = subgroups)
fos <- read_maxquant_phosphosites(fosfile = fosfile, profile = profile, subgroups = subgroups)
fos <- read_maxquant_phosphosites(fosfile = fosfile, profile = profile, fastafile = fastafile, subgroups = subgroups)
```

read_maxquant_proteingroups
<i>Read maxquant proteingroups</i>

Description

Read maxquant proteingroups

Usage

```

read_maxquant_proteingroups(
  dir = getwd(),
  file = if (is_file(dir)) dir else file.path(dir, "proteinGroups.txt"),
  fastafile = NULL,
  restapi = FALSE,
  quantity = NULL,
  subgroups = NULL,
  invert = character(0),
  rm_contaminants = TRUE,
  rm_reverse = TRUE,
  rm_missing_in_all_samples = TRUE,
  impute = FALSE,
  plot = FALSE,
  label = "feature_id",
  pca = plot,
  pls = plot,
  fit = if (plot) "limma" else NULL,
  formula = as.formula("~ subgroup"),
  block = NULL,
  coefs = NULL,
  contrasts = NULL,
  palette = NULL,
  verbose = TRUE
)

read_proteingroups(...)

```

Arguments

dir	proteingroups directory
file	proteingroups file
fastafile	uniprot fastafile
restapi	TRUE or FALSE : use uniprot restapi to annotate uniprot not in fastadt ?
quantity	'normalizedratio', 'ratio', 'correctedreporterintensity', 'reporterintensity', 'maxlfq', 'labeledintensity', 'intensity' or NULL
subgroups	NULL or string vector : subgroups to retain
invert	string vector : subgroups which require inversion
rm_contaminants	TRUE or FALSE : rm contaminants ?
rm_reverse	TRUE or FALSE : rm reverse proteins ?
rm_missing_in_all_samples	TRUE or FALSE
impute	TRUE or FALSE: impute group-specific NA values?
plot	TRUE or FALSE: plot ?

label	fvar
pca	TRUE or FALSE: run pca ?
pls	TRUE or FALSE: run pls ?
fit	model engine: 'limma', 'lm', 'lme(r)', 'wilcoxon' or NULL
formula	model formula
block	model blockvar: string or NULL
coefs	model coefficients of interest: character vector or NULL
contrasts	coefficient contrasts of interest: character vector or NULL
palette	color palette : named character vector
verbose	TRUE or FALSE : message ?
...	maintain deprecated functions

Value

SummarizedExperiment

Examples

```
# fukuda20 - LFQ
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
pro <- read_maxquant_proteingroups(file = file)

# billing19 - Normalized Ratios
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fastafile <- system.file('extdata/uniprot_hsa_20140515.fasta', package = 'autonomics')
subgroups <- sprintf('%s_STD', c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'))
pro <- read_maxquant_proteingroups(file = file, subgroups = subgroups)
pro <- read_maxquant_proteingroups(file = file, fastafile = fastafile, subgroups = subgroups)
```

read_msigdt	<i>Read msigdb datatable</i>
-------------	------------------------------

Description

Read msigdb datatable

Usage

```
read_msigdt(
  file = list_files(MSIGDIR, full.names = TRUE)[1],
  collections = if (is.null(file)) NULL else switch(basename(file) %>% substr(nchar(.)
    - 4, nchar(.) - 3), Hs = c("C2:CP:REACTOME", "C5:GO:BP", "C5:GO:MF", "C5:GO:CC"), Mm
    = c("M2:CP:REACTOME", "M5:GO:BP", "M5:GO:MF", "M5:GO:CC"))
)
```

Arguments

file msigdb file: one of the files in dir(MSIGDB).
collections subset of names(MSIGCOLLECTIONS)

Examples

```
read_msigdt()
```

read_olink	<i>Read olink file</i>
------------	------------------------

Description

Read olink file

Usage

```
read_olink(file, sample_excel = NULL, sample_tsv = NULL, by.y = "SampleID")
```

Arguments

file olinkfile
sample_excel sample excel
sample_tsv sample tsv
by.y sample tsv mergeby column

Value

SummarizedExperiment

Examples

```
# Example data
npxdt <- data.table::data.table(OlinkAnalyze::npx_data1)[, c(1:11, 17)]
sampledt <- data.table::data.table(OlinkAnalyze::npx_data1)[, c(1, 12:15)]
sampledt %<>% extract(!grepl('CONTROL', SampleID))
sampledt %<>% unique()
# Write to file
file <- paste0(tempfile(), '.olink.csv')
samplefile <- paste0(tempfile(), '.samples.xlsx')
data.table::fwrite(npxdt, file)
writexl::write_xlsx(sampledt, samplefile)
# Read
object <- read_olink(file, sample_excel = samplefile)
biplot(pca(object), color = 'Time', group = 'Subject', shape = 'Treatment')
```

read_salmon	<i>Read salmon</i>
-------------	--------------------

Description

Read salmon

Usage

```
read_salmon(dir, sfile = NULL, by = NULL, ensdb = NULL)
```

Arguments

dir	salmon results rootdir
sfile	samplefile
by	samplefile column to merge by
ensdb	EnsDb object

Value

SummarizedExperiment

Examples

```
# dir <- '../bh/salmon_quants'
# sfile <- '../bh/samplesheet.csv'
# by <- 'salmonDir'
# ah <- AnnotationHub::AnnotationHub()
# ensdb <- ah[['AH98078']]
# read_salmon(dir, sfile = sfile, by = 'salmonDir', ensdb = ensdb)
```

read_uniprot	<i>Read fasta hdrs</i>
--------------	------------------------

Description

Read fasta hdrs

Usage

```
read_uniprot(fastafile, fastafields = FASTAFIELDS, verbose = TRUE)

parse_maxquant_hdrs(fastahdrs)

read_contaminantdt(force = FALSE, verbose = TRUE)
```

Arguments

fastafile	string (or character vector)
fastafields	character vector : which fastahdr fields to extract ?
verbose	bool
fastahdrs	character vector
force	whether to overwrite existing file

Value

data.table(uniprot, protein, gene, uniprot, reviewed, existence)

Note

existence values are always those of the canonical isoform (no isoform-level resolution for this field)

Examples

```
# uniprot hdrs
fastafile <- system.file('extdata/uniprot_hsa_20140515.fasta', package = 'autonomics')
read_uniprot_dt(fastafile)

# maxquant hdrs
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
dt <- .read_maxquant_proteingroups(file)
parse_maxquant_hdrs(dt$`Fasta headers`)

profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
prodt <- .read_maxquant_proteingroups(profile)
fosdt <- .read_maxquant_phosphosites(fosfile, profile)
parse_maxquant_hdrs(prodt$`Fasta headers`)
parse_maxquant_hdrs(fosdt$`Fasta headers`)

# contaminant hdrs
read_contaminant_dt()
```

reexports

Objects exported from other packages

Description

These objects are imported from other packages. Follow the links below to see their documentation.

data.table [data.table](#)

magrittr [%<>%](#), [%>%](#), [extract](#)

reset_fit	<i>Reset fit</i>
-----------	------------------

Description

Reset fit

Usage

```
reset_fit(  
  object,  
  fit = fits(object),  
  coefs = autonomics::coefs(object, fit = fit),  
  verbose = TRUE  
)
```

Arguments

object	SummarizedExperiment
fit	character vector
coefs	character vector
verbose	TRUE or FALSE

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')  
(object <- read_metabolon(file))  
object %<>% reset_fit()  
object %<>% fit_limma() %>% reset_fit()  
object %<>% fit_limma() %>% fit_lm() %>% reset_fit()  
object %<>% fit_limma() %>% fit_lm() %>% reset_fit('limma')
```

rm_diann_contaminants	<i>Rm contaminants</i>
-----------------------	------------------------

Description

Rm contaminants from DIA-NN SumExp

Usage

```
rm_diann_contaminants(object, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- download_data('dilution.report.tsv')
object <- read_diann_proteingroups(file)
object %<>% rm_diann_contaminants()
```

rm_missing_in_all_samples

Rm features missing in some samples

Description

Rm features missing in some samples

Usage

```
rm_missing_in_all_samples(object, verbose = TRUE)

rm_missing_in_some_samples(object, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
verbose	TRUE (default) or FALSE

Value

updated object

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
rm_missing_in_all_samples( object)
rm_missing_in_some_samples(object)
```

rm_unmatched_samples *rm unmatched/singleton samples*

Description

rm unmatched/singleton samples

Usage

```
rm_unmatched_samples(  
  object,  
  subgroupvar = "subgroup",  
  subgroupctr = slevels(object, subgroupvar)[1],  
  block,  
  verbose = TRUE  
)  
  
rm_singleton_samples(object, subgroupvar = "subgroup", verbose = TRUE)
```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup variable (string)
subgroupctr	control subgroup (string)
block	block variable (string)
verbose	TRUE/FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.somascan.adat', package = 'autonomics')  
object <- read_somascan(file)  
object %<>% filter_samples(subgroup %in% c('t1', 't2'), verbose = TRUE)  
rm_singleton_samples(object, subgroupvar = 'Subject')  
rm_unmatched_samples(object, subgroupvar = 'subgroup', block = 'Subject')
```

scaledlibsizes	<i>Get tmm-scaled libsizes</i>
----------------	--------------------------------

Description

Get tmm-scaled libsizes

Usage

scaledlibsizes(counts)

Arguments

counts counts matri

Value

scaled libsize vector

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
scaledlibsizes(counts(object))
```

scoremat	<i>Extract scores/loadings</i>
----------	--------------------------------

Description

Extract scores/loadings

Usage

```
scoremat(object, method = "pca", by = biplot_by(object, method), dim = 1:2)

scores(object, method = "pca", by = biplot_by(object, method), dim = 1)

loadingmat(object, method = "pca", by = biplot_by(object, method), dim = 1:2)

loadings(object, method = "pca", by = biplot_by(object, method), dim = 1)
```

Arguments

object	SummarizedExperiment
method	'pca', 'pls', etc.
by	svar (string)
dim	numeric vector

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% pca()
  scores(object)[1:2]
  loadings(object)[1:2]
  scoremat(object)[1:2, ]
  loadingmat(object)[1:2, ]
```

slevels	<i>Get slevels</i>
---------	--------------------

Description

Get svar levels

Usage

```
slevels(object, svar)

subgroup_levels(object)
```

Arguments

object	SummarizedExperiment, eSet, or eList
svar	sample var (character)

Value

svar values (character)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
slevels(object, 'subgroup')
subgroup_levels(object)
```

snames	<i>Get/Set snames</i>
--------	-----------------------

Description

Get/Set sample names

Usage

```
snames(object)

## S4 method for signature 'SummarizedExperiment'
snames(object)

snames(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,character'
snames(object) <- value
```

Arguments

object	SummarizedExperiment
value	string vector with sample names

Value

sample names vector (get) or updated eSet (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
head(snames(object))
head(snames(object) %<>% paste0('SAMPLE_', .))
```

split_samples	<i>Split samples</i>
---------------	----------------------

Description

Split samples by svar

Usage

```
split_samples(object, by = "subgroup")

cbind_imputed(objlist)

split_features(object, by)
```

Arguments

object	SummarizedExperiment
by	svar to split by (string)
objlist	SummarizedExperiment list

Value

SummarizedExperiment list

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
objlist <- split_features(object, by = 'PLATFORM')
objlist <- split_samples(object, 'Diabetes')
objlist %<>% Map(impute, .)
object <- cbind_imputed(objlist)
```

stri_any_regex	<i>Does any string have a regex</i>
----------------	-------------------------------------

Description

Does any string have a regex

Usage

```
stri_any_regex(str, pattern)
```

Arguments

str	string vector
pattern	string

Value

TRUE or FALSE

Examples

```
str <- c('s1 Spectral Count', 's1 Unique Spectral Count')
patterns <- c('Spectral Count', '(?!Unique) Spectral Count', 'Intensity')
stringi::stri_detect_regex(str, pattern = patterns[1])
stringi::stri_detect_regex(str, pattern = patterns[2])
stringi::stri_detect_regex(str, pattern = patterns[3])
stri_any_regex(str, pattern = patterns)
```

stri_detect_fixed_in_collapsed

Detect fixed patterns in collapsed strings

Description

Detect fixed patterns in collapsed strings

Usage

```
stri_detect_fixed_in_collapsed(x, patterns, sep)
```

Arguments

x	vector with collapsed strings
patterns	vector with fixed patterns (strings)
sep	collapse separator (string) or NULL (if uncollapsed)

Value

boolean vector

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
x <- fdt(object)$uniprot
patterns <- c('A0A0R4IKT8', 'Q7T3G6')
table(stri_detect_fixed_in_collapsed(x = x, patterns = patterns, sep = ';'))
```

subgroup_array	<i>Get subgroup matrix</i>
----------------	----------------------------

Description

Arrange (subgroup)levels in matrix

Usage

```
subgroup_array(object, subgroupvar)
subgroup_matrix(object, subgroupvar)
```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup svar

Value

matrix

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object$subgroup <- paste0(object$Diabetes, '.', object$subgroup)
subgroup_matrix(object, 'subgroup')
```

subtract_baseline	<i>Subtract baseline</i>
-------------------	--------------------------

Description

Subtract baseline level within block

Usage

```
subtract_baseline(
  object,
  subgroupvar,
  subgroupctr = slevels(object, subgroupvar)[1],
  block = NULL,
  assaynames = setdiff(assayNames(object), c("weights", "pepcounts")),
  verbose = TRUE
)
```

```

subtract_pairs(
  object,
  subgroupvar = "subgroup",
  subgroupctr = slevels(object, subgroupvar)[1],
  block,
  assaynames = assayNames(object)[1],
  verbose = TRUE
)

subtract_differences(object, block, subgroupvar, verbose = TRUE)

```

Arguments

object	SummarizedExperiment
subgroupvar	subgroup svar
subgroupctr	control subgroup
block	block svar (within which subtraction is performed)
assaynames	which assays to subtract for
verbose	TRUE/FALSE

Details

subtract_baseline subtracts baseline levels within block, using the medoid baseline sample if multiple exist.

subtract_pairs also subtracts baseline level within block. It cannot handle multiple baseline samples, but has instead been optimized for many blocks

subtract_differences subtracts differences between subsequent levels, again within block

Value

SummarizedExperiment

Examples

```

# read
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object0 <- read_metabolon(file)
pca(object0, plot = TRUE, color = 'Time')

# subtract_baseline: takes medoid of baseline samples if multiple
object <- subtract_baseline(object0, block = 'Subject', subgroupvar = 'Time')
pca(object, plot = TRUE, color = 'Time')

# subtract_pairs: optimized for many blocks
object <- subtract_pairs(object0, block = 'Subject', subgroupvar = 'Time')

```

```

pca(object, plot = TRUE, color = 'Time')

# subtract_differences
object <- subtract_differences(object0, block = 'Subject', subgroupvar = 'Time')
values(object) %<>% na_to_zero()
pca(object, plot = TRUE, color = 'Time')

```

sumexplist_to_longdt *SummarizedExperiment list to long data.table*

Description

SummarizedExperiment list to long data.table

Usage

```

sumexplist_to_longdt(
  sumexplist,
  svars = intersect("subgroup", autonomics::svars(sumexplist[[1]])),
  fvars = intersect("gene", autonomics::fvars(sumexplist[[1]])),
  setvarname = "set"
)

```

Arguments

sumexplist	list of SummarizedExperiments
svars	character vector
fvars	character vector
setvarname	string

Value

data.table

Examples

```

subgroups <- paste0(c('E00', 'E01', 'E02', 'E05', 'E15', 'E30', 'M00'), '_STD')
rnafile <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
profile <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
fosfile <- system.file('extdata/billing19.phosphosites.txt', package = 'autonomics')
rna <- read_rnaseq_counts(rnafile)
pro <- read_maxquant_proteingroups(file = profile, subgroups = subgroups)
fos <- read_maxquant_phosphosites(fosfile = fosfile, profile = profile, subgroups = subgroups)
pro$subgroup %<>% stringi::stri_replace_first_fixed('_STD', '')
fos$subgroup %<>% stringi::stri_replace_first_fixed('_STD', '')

sumexplist <- list(rna = rna, pro = pro, fos = fos)
dt <- sumexplist_to_longdt(sumexplist, setvarname = 'platform')
dt %<>% extract(gene %in% c('TNMD', 'TSPAN6'))

```

sumexp_to_tsv	<i>Write sumexp to tsv</i>
---------------	----------------------------

Description

Write sumexp to tsv

Usage

```
sumexp_to_tsv(object, assay = assayNames(object)[1], file)
```

Arguments

object	SummarizedExperiment
assay	string
file	filename

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file, fit = 'limma')
tsv <- file.path(tempdir(), 'fukuda20.proteingroups.tsv')
sumexp_to_tsv(object, file = tsv)
```

sumexp_to_widedt	<i>SummarizedExperiment to data.table</i>
------------------	---

Description

SummarizedExperiment to data.table

Usage

```
sumexp_to_widedt(
  object,
  fvars = autonomics::fvars(object),
  assay = assayNames(object)[1]
)

sumexp_to_longdt(
  object,
  fvars = intersect("feature_name", autonomics::fvars(object)),
  svars = intersect("subgroup", autonomics::svars(object)),
  assay = assayNames(object) %>% intersect(c(.[1], "is_imputed"))
)

sumexp_to_subrep_dt(object, subgroup = subgroup)
```

Arguments

object	sumexp
fvars	additional fvars to include in table
assay	matrix in assays(object) to be used
svars	additional svars to include in table
subgroup	subgroup (sym)

Details

- sumexp_to_widedt: feature x sample
- sumexp_to_subrep_dt: feature.subgroup x replicate
- sumexp_to_longdt: feature.sample

Value

data.table

Examples

```
# Atkin Hypoglycemia
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
sumexp_to_widedt(object)
sumexp_to_longdt(object)
sumexp_to_subrep_dt(object)

# Fukuda
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
values(object)
fdt(object)
object %<>% impute()
table(fdt(object)$imputed)
sumexp_to_longdt(object)
sumexp_to_widedt(object)
sumexp_to_longdt(object)
```

summarize_fit	<i>Summarize fit</i>
---------------	----------------------

Description

Summarize fit

Usage

```

summarize_fit(object, ...)

## S3 method for class 'data.table'
summarize_fit(
  object,
  fit = fits(object),
  coefs = autonomics::coefs(object, fit = fit),
  ...
)

## S3 method for class 'SummarizedExperiment'
summarize_fit(
  object,
  fit = fits(object),
  coefs = autonomics::coefs(object, fit = fit),
  ...
)

```

Arguments

object	SummarizedExperiment or data.table
...	S3 dispatch
fit	'limma', 'lme', 'lm', 'lme', 'wilcoxon' or NULL
coefs	string vector

Value

data.table(contrast, nup, ndown)

Examples

```

file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma()
object %<>% fit_lm()
summarize_fit(object, coefs = c('t1-t0', 't2-t0', 't3-t0'))

```

SURVIVALENGINES

Survival engines

Description

Survival engines

Usage

SURVIVALENGINES

Format

An object of class character of length 3.

Examples

SURVIVALENGINES

survival_example	<i>Fit survival</i>
------------------	---------------------

Description

Investigates association between expression and survival

Usage

```
survival_example()

fit_survival(
  object,
  ntile = 2,
  engine = c("survdif", "coxph", "logrank")[1:2],
  assay = assayNames(object)[1],
  sep = FITSEP,
  verbose = TRUE,
  outdir = NULL,
  plot = if (is.null(outdir)) FALSE else TRUE,
  width = 7,
  height = 7,
  n = min(nrow(object), 9),
  ncol = 3,
  nrow = 3,
  writefunname = "write_xl"
)
```

Arguments

object	SummarizedExperiment
ntile	number
engine	'coxph' (survival), 'survdif' (survival), 'logrank' (coin)
assay	string
sep	fvar string separator : e.g. '~' gives p~surv~LR50

verbose	TRUE or FALSE
outdir	dir
plot	TRUE or FALSE
width	number
height	number
n	number of features to plot
ncol	number of cols
nrow	number of rows
writefunname	'write_xl' or 'write_ods'

Details

Investigates association between expression and survival.
 Continuous for coxph.
 Categorical for survdiff or logrank
 Samples are split into ntile expression groups.
 Survival is compared between highest and lowest expressors.

Three statistics recorded per engine

p
 effect: coef(coxph)
 mean survival difference (survdiff, logrank)
 t: z(coxph)
 χ^2 (survdiff, logrank)
 sign reflects whether expression
 increases (positive) or decreases (negative) survival

Value

SummarizedExperiment

Examples

```
# Defaults
object <- survival_example()
fit_survival(object)

# Engines
fit_survival(object, engine = c('coxph', 'survdiff'))
fit_survival(object, engine = c('coxph', 'survdiff', 'logrank'))

# Quantiles
fit_survival(object, engine = 'logrank')
fit_survival(object, engine = 'logrank', ntile = 4)

# Plot
fit_survival(object)
fit_survival(object, plot = TRUE)
fit_survival(object, engine = c('coxph', 'survdiff', 'logrank'), plot = TRUE)
```

svalues	<i>Get/Set svalues</i>
---------	------------------------

Description

Get/Set svar values

Usage

```
svalues(object, svar)

subgroup_values(object)

sampleid_values(object)

svalues(object, svar) <- value

## S4 replacement method for signature 'SummarizedExperiment,character'
svalues(object, svar) <- value
```

Arguments

object	SummarizedExperiment
svar	sample var (character)
value	value vector

Value

character vector (get) or SummarizedExperiment (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
svalues(object, 'subgroup')
subgroup_values(object)
```

svars	<i>Get/Set svars</i>
-------	----------------------

Description

Get/Set sample variables

Usage

```

svars(object)

## S4 method for signature 'SummarizedExperiment'
svars(object)

## S4 method for signature 'MultiAssayExperiment'
svars(object)

svars(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,character'
svars(object) <- value

## S4 replacement method for signature 'MultiAssayExperiment,character'
svars(object) <- value

```

Arguments

object	SummarizedExperiment
value	string vector with variable names

Value

sample variable names (get) or updated SummarizedExperiment

Examples

```

file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
svars(object)[1]
(svars(object)[1] %<>% paste0('1'))

```

systematic_nas	<i>Is systematic/random/full NA</i>
----------------	-------------------------------------

Description

Is systematic/random/full NA

Usage

```

systematic_nas(object, by = "subgroup", frac = 0.5)

random_nas(object, by = "subgroup")

no_nas(object)

```

Arguments

object	SummarizedExperiment
by	svar (string)
frac	fraction

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
table(systematic_nas(object)) # missing in some subgroups, present in others
table(random_nas(object))    # missing in some samples, independent of subgroup
table(no_nas(object))        # missing in no samples
```

tag_features	<i>Tag features</i>
--------------	---------------------

Description

Tag features

Usage

```
tag_features(
  object,
  keyvar,
  sep,
  features,
  tagvar = get_name_in_parent(features),
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
keyvar	string : intersection fvar
sep	string : keyvar collapse separator
features	character vector : intersection set
tagvar	string :
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/atkin.somascan.adat', package = 'autonomics')
object <- read_somascan(file)
features <- AnnotationDbi::keys(org.Hs.eg.db::org.Hs.eg.db, keytype = 'SYMBOL')
object %<>% tag_features(keyvar = 'EntrezGeneSymbol', sep = ' ', features)
table(fdt(object)$features)
```

tag_hdlproteins	<i>Tag hdlproteins</i>
-----------------	------------------------

Description

Tag hdlproteins

Usage

```
tag_hdlproteins(object, verbose = TRUE)
```

Arguments

object	SummarizedExperiment
verbose	TRUE or FALSE

Value

SummarizedExperiment

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
object %<>% tag_hdlproteins()
fdt(object)
```

TAXON_TO_ORGNAME	<i>Annotation Maps</i>
------------------	------------------------

Description

Annotation Maps

Usage

TAXON_TO_ORGNAME

ABBREV_TO_ORGNAME

REVIEWED_TO_NUMBER

EXISTENCE_TO_NUMBER

Format

- An object of class character of length 7.
- An object of class character of length 4.
- An object of class character of length 2.
- An object of class numeric of length 4.

Examples

```
TAXON_TO_ORGNAME['9606']  
ABBREV_TO_ORGNAME['HSA']  
REVIEWED_TO_NUMBER['reviewed']  
EXISTENCE_TO_NUMBER['Evidence at protein level']
```

TESTS	<i>Statistical models supported in autonomics</i>
-------	---

Description

Statistical models supported in autonomics

Usage

TESTS

Format

- An object of class character of length 5.

Examples

TESTS

tpm*Get/Set tpm*

Description

Get / Set tpm matrix

Usage

```
tpm(object)

## S4 method for signature 'SummarizedExperiment'
tpm(object)

tpm(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
tpm(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
tpm(object) <- value
```

Arguments

object	SummarizedExperiment
value	tpm matrix (features x samples)

Value

tpm matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file, plot=FALSE)
tpm(object) <- values(object)
tpm(object)[1:3, 1:3]
```

twofactor_sumexp	<i>twofactor sumexp</i>
------------------	-------------------------

Description

twofactor sumexp

Usage

```
twofactor_sumexp()
```

Value

SummarizedExperiment

uncollapse	<i>Uncollapse/Recollapse</i>
------------	------------------------------

Description

Uncollapse data.table cols

Usage

```
uncollapse(dt, ..., sep = ";")  
  
recollapse(dt, by, sep = ";")
```

Arguments

dt	data.table
...	cols
sep	string
by	string

Examples

```
# Example data  
(dt <- data.table::data.table(  
  uniprot = 'Q9BQL6;Q96AC1;Q96AC1-3',  
  protein = 'FERM1_HUMAN;FERM2_HUMAN',  
  gene    = 'FERMT1;FERMT2',  
  family  = 'FERM'))  
  
# Uncollapse  
uncollapse(dt, protein, gene, sep = ';')  
recollapse(uncollapse(dt, protein, gene, sep = ';'), by = 'uniprot')
```

```
# Unchanged when no sep
  uncollapse(dt, family, sep = ';')
  uncollapse(dt, family, sep = 'NOSEP')
```

values

Get/Set expr values

Description

Get/Set value matrix

Usage

```
values(object)

## S4 method for signature 'SummarizedExperiment'
values(object)

values(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,matrix'
values(object) <- value

## S4 replacement method for signature 'SummarizedExperiment,numeric'
values(object) <- value
```

Arguments

object	SummarizedExperiment
value	ratio matrix (features x samples)

Value

value matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.proteingroups.txt', package = 'autonomics')
object <- read_maxquant_proteingroups(file)
values(object)[1:3, 1:3]
values(object) <- 0
values(object)[1:3, 1:3]
```

varlevels_dont_clash	<i>Are varlevels unique</i>
----------------------	-----------------------------

Description

Are varlevels unique

Usage

```
varlevels_dont_clash(object, ...)

## S3 method for class 'data.table'
varlevels_dont_clash(object, vars = names(object), ...)

## S3 method for class 'SummarizedExperiment'
varlevels_dont_clash(object, vars = svars(object), ...)
```

Arguments

object	SummarizedExperiment or data.table
...	required for s3 dispatch
vars	character vector

Value

TRUE or FALSE

Examples

```
require(data.table)
object1 <- data.table(expand.grid(genome = c('WT', 'MUT'), treat = c('control', 'drug')))
object2 <- data.table(expand.grid(mutant = c('YES', 'NO'), treated = c('YES', 'NO')))
varlevels_dont_clash(object1)
varlevels_dont_clash(object2)
```

venn_detects	<i>Venn detects</i>
--------------	---------------------

Description

Venn diagram full/consistent/random detects

Usage

```
venn_detects(object, by = "subgroup")
```

Arguments

object SummarizedExperiment
by svar (string)

Value

NULL

Examples

```
file <- system.file('extdata/fukuda20.proteingroups.txt', package = 'autonomics')  
object <- read_maxquant_proteingroups(file)  
venn_detects(object, 'subgroup')
```

weights	<i>Get/Set weights</i>
---------	------------------------

Description

Get/Set weight matrix

Usage

```
weights(object, ...)  
  
## S4 method for signature 'SummarizedExperiment'  
weights(object)  
  
weights(object) <- value  
  
## S4 replacement method for signature 'SummarizedExperiment,matrix'  
weights(object) <- value  
  
## S4 replacement method for signature 'SummarizedExperiment,numeric'  
weights(object) <- value  
  
## S4 replacement method for signature 'SummarizedExperiment,NULL '  
weights(object) <- value
```

Arguments

object SummarizedExperiment
... additional params
value ratio matrix (features x samples)

Value

weight matrix (get) or updated object (set)

Examples

```
file <- system.file('extdata/billing19.rnacounts.txt', package = 'autonomics')
object <- read_rnaseq_counts(file)
weights(object)[1:3, 1:2]
weights(object) <- 1
weights(object)[1:3, 1:2]
```

write_xl	<i>Write xl/ods</i>
----------	---------------------

Description

Write xl/ods

Usage

```
write_xl(
  object,
  xlfile,
  fitcoefs = autonomics::fitcoefs(object),
  verbose = TRUE
)

write_ods(
  object,
  odsfile,
  fitcoefs = autonomics::fitcoefs(object),
  verbose = TRUE
)
```

Arguments

object	SummarizedExperiment
xlfile	file
fitcoefs	character vector
verbose	TRUE or FALSE
odsfile	file

Value

filepath

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file, fit = 'limma')
xlfile <- file.path(tempdir(), 'fukuda20.proteingroups.fdt.xlsx')
odsfile <- file.path(tempdir(), 'fukuda20.proteingroups.fdt.ods')
# write_xl(object, xlfile)
# write_ods(object, odsfile)
```

X	<i>Model based prediction</i>
---	-------------------------------

Description

Model based prediction

Usage

```
X(
  object,
  formula = default_formula(object),
  drop = varlevels_dont_clash(object, all.vars(formula)),
  codingfun = contr.treatment.explicit
)

beta(object, fit = fits(object)[1])
```

Arguments

object	SummarizedExperiment or data.frame
formula	formula
drop	TRUE or FALSE
codingfun	function
fit	'limma', 'lm', 'lme', 'wilcoxon'

Value

beta matrix (nlevel x nfeature)

Examples

```
file <- system.file('extdata/atkin.metabolon.xlsx', package = 'autonomics')
object <- read_metabolon(file)
object %<>% fit_limma(block = 'Subject') # intercept required!
beta(object) # betas : nlevel x nfeature
X(object) # design : nlevel x nlevel
X(object) %*% beta(object) # response : nlevel x nfeature
```

zero_to_na	<i>Change nondetect representation</i>
------------	--

Description

Change nondetect representation

Usage

```
zero_to_na(x, verbose = FALSE)
nan_to_na(x, verbose = FALSE)
na_to_zero(x, verbose = FALSE)
inf_to_na(x, verbose = FALSE)
minusrinf_to_na(x, verbose = FALSE)
na_to_string(x)
```

Arguments

x	matrix
verbose	logical(1)

Value

Updated matrix

Examples

```
matrix(c(0, 7), nrow=1)
matrix(c(0, 7), nrow=1) %>% zero_to_na(verbose=TRUE)

matrix(c(NA, 7), nrow=1)
matrix(c(NA, 7), nrow=1) %>% na_to_zero(verbose=TRUE)

matrix(c(NaN, 7), nrow=1)
matrix(c(NaN, 7), nrow=1) %>% nan_to_na(verbose=TRUE)

matrix(c(Inf, 7), nrow=1)
matrix(c(Inf, 7), nrow=1) %>% inf_to_na(verbose=TRUE)

matrix(c(-Inf, 7), nrow=1)
matrix(c(-Inf, 7), nrow=1) %>% minusrinf_to_na(verbose=TRUE)
```

Index

* datasets

AUTONOMICS_DATASETS, 36
COMPOUNDDISCOVERER_PATTERNS, 48
DATADIR, 56
DIMREDUN, 62
FITSEP, 77
LINMODENGINES, 107
MAXQUANT_PATTERNS, 118
MSIGCOLLECTIONSHUMAN, 130
MSIGDIR, 131
OPENTARGETSDIR, 132
PRECURSOR_QUANTITY, 161
SURVIVALENGINES, 187
TAXON_TO_ORGNAME, 193
TESTS, 194

* internal

reexports, 173
.coxph, 7
.extract_effectsize_features
 (.extract_p_features), 7
.extract_fdr_features
 (.extract_p_features), 7
.extract_n_features
 (.extract_p_features), 7
.extract_p_features, 7
.extract_sign_features
 (.extract_p_features), 7
.fit_limma(fit_linmod), 77
.logrank(.coxph), 7
.merge, 10
.read_compounddiscoverer, 11
.read_compounddiscoverer_masslist, 11
.read_diann_precursors, 12
.read_diann_proteingroups
 (.read_diann_precursors), 12
.read_maxquant_phosphosites
 (.read_maxquant_proteingroups),
 14
.read_maxquant_proteingroups, 14

.read_metabolon, 15
.read_rectangles, 17
.read_rnaseq_bams, 19
.read_rnaseq_counts
 (.read_rnaseq_bams), 19
.read_somascan, 22
.survdiff(.coxph), 7
%<>%(reexports), 173
%>%(reexports), 173
%<>%, 173
%>%, 173
ABBREV_TO_ORGNAME (TAXON_TO_ORGNAME),
 193
abstract_fit, 24
abstractvar(modelvar), 124
abstractvec(modelvar), 124
add_adjusted_pvalues, 25
add_assay_means, 26
add_facetvars, 27
add_opentargets_by_uniprot, 28
add_psp, 28
add_smiles, 29
altenrich, 30
analysis, 31
analysis, SummarizedExperiment-method
 (analysis), 31
analysis<- (analysis), 31
analysis<-, SummarizedExperiment, list-method
 (analysis), 31
analyze, 32
annotate_compounddiscoverer, 33
annotate_maxquant, 34
annotate_uniprot_rest, 35
assert_compounddiscoverer_output
 (is_diann_report), 97
assert_correlation_matrix
 (is_correlation_matrix), 96
assert_diann_report(is_diann_report),
 97

- assert_fastadt (is_fastadt), 99
- assert_fragpipe_tsv (is_diann_report), 97
- assert_is_fraction (is_fraction), 100
- assert_is_valid_sumexp, 36
- assert_maxquant_phosphosites (is_diann_report), 97
- assert_maxquant_proteingroups (is_diann_report), 97
- assert_positive_number (is_positive_number), 101
- assert_scalar_subset (is_scalar_subset), 102
- assert_valid_formula (is_valid_formula), 104
- assert_weakly_positive_number (is_positive_number), 101
- AUTONOMICS_DATASETS, 36
- beta (X), 201
- bin, 37
- biplot, 38
- biplot_corrections, 39
- biplot_covariates, 40
- block2lme, 42
- block_vars (block2lme), 42
- cbind_imputed (split_samples), 179
- center, 43
- code, 44
- code_control (code), 44
- code_deviation (code), 44
- code_deviation_first (code), 44
- code_diff (code), 44
- code_diff_forward (code), 44
- code_helmert (code), 44
- code_helmert_forward (code), 44
- coefs, 46
- collapse_in (count_in), 52
- collapsed_entrezg_to_symbol, 47
- COMPOUNDDISCOVERER_PATTERNS, 48
- contr.diff (code), 44
- contr.treatment.explicit (code), 44
- contrast_coefs, 48
- contrast_subgroup_cols, 49
- contrast_subgroup_rows (contrast_subgroup_cols), 49
- count_in, 52
- count_out (count_in), 52
- counts, 50
- counts, SummarizedExperiment-method (counts), 50
- counts2cpm, 51
- counts2tpm, 51
- counts<- (counts), 50
- counts<-, SummarizedExperiment, matrix-method (counts), 50
- counts<-, SummarizedExperiment, NULL-method (counts), 50
- counts<-, SummarizedExperiment, numeric-method (counts), 50
- cpm, 53
- cpm, SummarizedExperiment-method (cpm), 53
- cpm2counts (counts2cpm), 51
- cpm<- (cpm), 53
- cpm<-, SummarizedExperiment, matrix-method (cpm), 53
- cpm<-, SummarizedExperiment, numeric-method (cpm), 53
- create_design, 54
- data.table, 173
- data.table (reexports), 173
- DATADIR, 56
- default_formula (default_subgroupvar), 58
- default_geom, 57
- default_sfile, 58
- default_subgroupvar, 58
- demultiplex, 59
- dequantify, 60
- dequantify_compounddiscoverer, 61
- DIMREDENGINES (DIMREDUN), 62
- DIMREDSUPER (DIMREDUN), 62
- DIMREDUN, 62
- downfeatures (modelvar), 124
- download_data (DATADIR), 56
- download_gtf, 62
- download_mcclain21, 63
- dt2mat, 64
- effectdt (modelvar), 124
- effectmat (modelvar), 124
- effectsizemat (modelvar), 124
- effectvar (modelvar), 124
- effectvec (modelvar), 124
- enrichment, 64

- ens2org, 66
- entrezg_to_symbol, 67
- EXISTENCE_TO_NUMBER (TAXON_TO_ORGNAME), 193
- exp2 (log2transform), 113
- extract, 173
- extract (reexports), 173
- extract_coef_features
 (.extract_p_features), 7
- extract_rectangle, 67
- factor2logical (logical2factor), 114
- fcluster, 69
- fcor (mdsplot), 119
- fdata, 70
- fdata, SummarizedExperiment-method
 (fdata), 70
- fdata<- (fdata), 70
- fdata<-, SummarizedExperiment, data.frame-method
 (fdata), 70
- fdist (mdsplot), 119
- fdr2p, 72
- fdrmat (modelvar), 124
- fdrvar (modelvar), 124
- fdrvec (modelvar), 124
- fdt (fdata), 70
- fdt, SummarizedExperiment-method
 (fdata), 70
- fdt<- (fdata), 70
- fdt<-, SummarizedExperiment, data.table-method
 (fdata), 70
- filter_exprs_replicated_in_some_subgroup, 72
- filter_features, 73
- filter_medoid, 74
- filter_samples, 75
- fit_limma (fit_linmod), 77
- fit_linmod, 77
- fit_lm (fit_linmod), 77
- fit_lme (fit_linmod), 77
- fit_lmer (fit_linmod), 77
- fit_survival (survival_example), 188
- fit_wilcoxon (fit_linmod), 77
- fitcoefs, 75
- fits, 76
- FITSEP, 77
- fix_xlgenes, 81
- flevels, 82
- fnames, 83
- fnames, SummarizedExperiment-method
 (fnames), 83
- fnames<- (fnames), 83
- fnames<-, SummarizedExperiment, character-method
 (fnames), 83
- formula2lm (block2lme), 42
- formula2lmer (block2lme), 42
- formula2str, 83
- fscale (log2transform), 113
- ftype, 84
- fvalues, 85
- fvars, 85
- fvars, SummarizedExperiment-method
 (fvars), 85
- fvars<- (fvars), 85
- fvars<-, SummarizedExperiment, character-method
 (fvars), 85
- genome_to_orfdb, 86
- group_by_level, 87
- guess_compounddiscoverer_quantity, 88
- guess_fitsep, 88
- guess_maxquant_quantity, 89
- guess_sep, 90
- has_multiple_levels, 91
- hdlproteins, 93
- impute, 93
- inf_to_na (zero_to_na), 202
- invert_subgroups, 95
- invnorm (log2transform), 113
- is_collapsed_subset, 96
- is_compounddiscoverer_output
 (is_diann_report), 97
- is_correlation_matrix, 96
- is_diann_report, 97
- is_fastadt, 99
- is_file, 99
- is_fraction, 100
- is_fragpipe_tsv (is_diann_report), 97
- is_imputed, 100
- is_imputed, SummarizedExperiment-method
 (is_imputed), 100
- is_imputed<- (is_imputed), 100
- is_imputed<-, SummarizedExperiment, matrix-method
 (is_imputed), 100
- is_imputed<-, SummarizedExperiment, NULL-method
 (is_imputed), 100

- is_maxquant_phosphosites
 (is_diann_report), 97
- is_maxquant_proteingroups
 (is_diann_report), 97
- is_positive_number, 101
- is_scalar_subset, 102
- is_sig, 103
- is_valid_formula, 104
- is_weakly_positive_number
 (is_positive_number), 101

- keep_connected_blocks, 105
- keep_connected_features, 105
- keep_replicated_features, 106

- label2index, 106
- lda(pca), 134
- LINMODEGINES, 107
- list2mat, 107
- list_files, 108
- loadingmat(scoremat), 177
- loadings(scoremat), 177
- log2counts, 108
- log2counts, SummarizedExperiment-method
 (log2counts), 108
- log2counts<- (log2counts), 108
- log2counts<- , SummarizedExperiment, matrix-method
 (log2counts), 108
- log2counts<- , SummarizedExperiment, numeric-method
 (log2counts), 108
- log2cpm, 109
- log2cpm, SummarizedExperiment-method
 (log2cpm), 109
- log2cpm<- (log2cpm), 109
- log2cpm<- , SummarizedExperiment, matrix-method
 (log2cpm), 109
- log2cpm<- , SummarizedExperiment, numeric-method
 (log2cpm), 109
- log2diffs, 110
- log2diffs, SummarizedExperiment-method
 (log2diffs), 110
- log2diffs<- (log2diffs), 110
- log2diffs<- , SummarizedExperiment, matrix-method
 (log2diffs), 110
- log2diffs<- , SummarizedExperiment, numeric-method
 (log2diffs), 110
- log2proteins, 111
- log2proteins, SummarizedExperiment-method
 (log2proteins), 111
- log2proteins<- (log2proteins), 111
- log2proteins<- , SummarizedExperiment, matrix-method
 (log2proteins), 111
- log2proteins<- , SummarizedExperiment, numeric-method
 (log2proteins), 111
- log2sites, 111
- log2sites, SummarizedExperiment-method
 (log2sites), 111
- log2sites<- (log2sites), 111
- log2sites<- , SummarizedExperiment, matrix-method
 (log2sites), 111
- log2sites<- , SummarizedExperiment, numeric-method
 (log2sites), 111
- log2tpm, 112
- log2tpm, SummarizedExperiment-method
 (log2tpm), 112
- log2tpm<- (log2tpm), 112
- log2tpm<- , SummarizedExperiment, matrix-method
 (log2tpm), 112
- log2tpm<- , SummarizedExperiment, numeric-method
 (log2tpm), 112
- log2transform, 113
- logical2factor, 114

- make_alpha_palette, 115
- make_colors, 116
- make_volcano_dt, 116
- map_fvalues, 117
- mat2dt(dt2mat), 64
- matrix2sumexp, 118
- MAXQUANT_PATTERNS, 118
- mdsplot, 119
- merge_compounddiscoverer, 120
- merge_fdata(merge_sdata), 122
- merge_fdt(merge_sdata), 122
- merge_ffile(merge_sample_file), 121
- merge_sample_excel, 120
- merge_sample_file, 121
- merge_sdata, 122
- merge_sdt(merge_sdata), 122
- message_df, 124
- minusinf_to_na(zero_to_na), 202
- model_coefs(contrast_coefs), 48
- modeldt(modelvar), 124
- modelfeatures(modelvar), 124
- modelmat(modelvar), 124
- modelvar, 124
- modelvec(modelvar), 124
- MSIGCOLLECTIONSHUMAN, 130

- MSIGCOLLECTIONSMOUSE
 - (MSIGCOLLECTIONSHUMAN), 130
- MSIGDIR, 131
- na_to_string(zero_to_na), 202
- na_to_zero(zero_to_na), 202
- nan_to_na(zero_to_na), 202
- nfactors, 131
- no_nas(systematic_nas), 191
- OPENTARGETSDIR, 132
- opls(pca), 134
- order_on_effect(order_on_p), 132
- order_on_p, 132
- order_on_t(order_on_p), 132
- parse_maxquant_hdrs(read_uniprot), 172
- pca, 134
- pdt(modelvar), 124
- pg_to_canonical, 136
- pg_to_isoforms(pg_to_canonical), 136
- plot_coef_densities, 137
- plot_contrast_venn, 138
- plot_contrastogram, 137
- plot_data, 139
- plot_densities, 140, 159
- plot_design, 142
- plot_detections, 143
- plot_exprs, 145, 159
- plot_exprs_per_coef, 148
- plot_feature_boxplots(plot_exprs), 145
- plot_feature_densities
 - (plot_densities), 140
- plot_feature_violins(plot_violins), 157
- plot_fit_summary, 149
- plot_heatmap, 150
- plot_joint_density, 151
- plot_matrix, 152
- plot_sample_boxplots, 142
- plot_sample_boxplots(plot_exprs), 145
- plot_sample_densities, 147, 149
- plot_sample_densities(plot_densities), 140
- plot_sample_nas(plot_detections), 143
- plot_sample_violins, 142, 147, 149
- plot_sample_violins(plot_violins), 157
- plot_subgroup_nas(plot_detections), 143
- plot_subgroup_points, 152
- plot_subgroup_violins(plot_violins), 157
- plot_summarized_detections
 - (plot_detections), 143
- plot_summary, 154
- plot_survival, 154
- plot_venn, 156
- plot_venn_heatmap, 156
- plot_violins, 157
- plot_volcano, 159
- plotmat, 138
- pls(pca), 134
- pmat(modelvar), 124
- PPATTERN(FITSEP), 77
- PRECURSOR_QUANTITY, 161
- preprocess_rnaseq_counts, 161
- pull_columns, 163
- pvar(modelvar), 124
- pvec(modelvar), 124
- quantnorm(log2transform), 113
- random_nas(systematic_nas), 191
- read_affymetrix, 163
- read_compounddiscoverer, 164
- read_contaminantdt(read_uniprot), 172
- read_diann(.read_diann_precursors), 12
- read_diann_proteingroups
 - (.read_diann_precursors), 12
- read_fragpipe, 166
- read_maxquant_phosphosites, 166
- read_maxquant_proteingroups, 168
- read_metabolon(.read_metabolon), 15
- read_msigdt, 30, 170
- read_olink, 171
- read_phosphosites
 - (read_maxquant_phosphosites), 166
- read_proteingroups
 - (read_maxquant_proteingroups), 168
- read_rectangles(.read_rectangles), 17
- read_rnaseq_bams(.read_rnaseq_bams), 19
- read_rnaseq_counts(.read_rnaseq_bams), 19
- read_salmon, 172
- read_somscan(.read_somscan), 22
- read_uniprot, 172
- recollapse(uncollapse), 196

- reexports, 173
- reset_fit, 174
- REVIEWED_TO_NUMBER (TAXON_TO_ORGNAME), 193
- rm_diann_contaminants, 174
- rm_missing_in_all_samples, 175
- rm_missing_in_some_samples (rm_missing_in_all_samples), 175
- rm_singleton_samples (rm_unmatched_samples), 176
- rm_unmatched_samples, 176
- sampleid_values (svalues), 190
- scaledlibsizes, 177
- scor (mdsplot), 119
- scoremat, 177
- scores (scoremat), 177
- sdata (fdata), 70
- sdata, SummarizedExperiment-method (fdata), 70
- sdata<- (fdata), 70
- sdata<-, SummarizedExperiment, data.frame-method (fdata), 70
- sdata<-, SummarizedExperiment, DataFrame-method (fdata), 70
- sdist (mdsplot), 119
- sdt (fdata), 70
- sdt, SummarizedExperiment-method (fdata), 70
- sdt<- (fdata), 70
- sdt<-, SummarizedExperiment, data.table-method (fdata), 70
- slevels, 178
- sma (pca), 134
- snames, 179
- snames, SummarizedExperiment-method (snames), 179
- snames<- (snames), 179
- snames<-, SummarizedExperiment, character-method (snames), 179
- split_extract (nfactors), 131
- split_extract_fixed (nfactors), 131
- split_extract_regex (nfactors), 131
- split_features (split_samples), 179
- split_samples, 179
- spls (pca), 134
- sscale (log2transform), 113
- stri_any_regex, 180
- stri_detect_fixed_in_collapsed, 181
- subgroup_array, 182
- subgroup_levels (slevels), 178
- subgroup_matrix (subgroup_array), 182
- subgroup_values (svalues), 190
- subtract_baseline, 182
- subtract_differences (subtract_baseline), 182
- subtract_pairs (subtract_baseline), 182
- sumexp_to_longdt (sumexp_to_widedt), 185
- sumexp_to_subrep_dt (sumexp_to_widedt), 185
- sumexp_to_tsv, 185
- sumexp_to_widedt, 185
- sumexplist_to_longdt, 184
- summarize_fit, 186
- survival_example, 188
- SURVIVALENGINES, 187
- svalues, 190
- svalues<- (svalues), 190
- svalues<-, SummarizedExperiment, character-method (svalues), 190
- svars, 190
- svars, MultiAssayExperiment-method (svars), 190
- svars, SummarizedExperiment-method (svars), 190
- svars<- (svars), 190
- svars<-, MultiAssayExperiment, character-method (svars), 190
- svars<-, SummarizedExperiment, character-method (svars), 190
- systematic_nas, 191
- tag_features, 192
- tag_hdlproteins, 193
- taxon2org (ens2org), 66
- TAXON_TO_ORGNAME, 193
- tdt (modelvar), 124
- TESTS, 194
- tmat (modelvar), 124
- tpm, 195
- tpm, SummarizedExperiment-method (tpm), 195
- tpm<- (tpm), 195
- tpm<-, SummarizedExperiment, matrix-method (tpm), 195
- tpm<-, SummarizedExperiment, numeric-method (tpm), 195

tvar (modelvar), [124](#)
tvec (modelvar), [124](#)
twofactor_sumexp, [196](#)

uncollapse, [196](#)
upfeatures (modelvar), [124](#)

values, [197](#)
values, SummarizedExperiment-method
 (values), [197](#)
values<- (values), [197](#)
values<-, SummarizedExperiment, matrix-method
 (values), [197](#)
values<-, SummarizedExperiment, numeric-method
 (values), [197](#)
varlevels_dont_clash, [198](#)
venn_detects, [198](#)
vsn (log2transform), [113](#)

weights, [199](#)
weights, SummarizedExperiment-method
 (weights), [199](#)
weights<- (weights), [199](#)
weights<-, SummarizedExperiment, matrix-method
 (weights), [199](#)
weights<-, SummarizedExperiment, NULL-method
 (weights), [199](#)
weights<-, SummarizedExperiment, numeric-method
 (weights), [199](#)
write_ods (write_xl), [200](#)
write_xl, [200](#)

X, [201](#)

zero_to_na, [202](#)
zscore (log2transform), [113](#)