

Package ‘ProteinBatcher’

September 15, 2026

Title An end-to-end proteomics workflow with condition-aware imputation, flexible statistical modelling and interactive visualization

Version 0.99.7

Description ProteinBatcher provides utilities to read, validate, and harmonize quantitative proteomics result tables generated by DIA-NN from data-independent acquisition mass spectrometry experiments. The package enables batch processing of multiple DIA-NN output files, consistent handling of protein identifiers and associated metadata, and transformation of tabular data into analysis-ready formats. It facilitates downstream differential abundance analysis using linear modeling frameworks based on limma, with explicit support for multivariable experimental designs including batch effects and other confounding covariates, as well as correlation-aware designs via duplicateCorrelation.

License MIT + file LICENSE

URL <https://github.com/DataScienceRD-Almirall/ProteinBatcher>

BugReports <https://github.com/DataScienceRD-Almirall/ProteinBatcher/issues>

Encoding UTF-8

Roxygen list(markdown = TRUE)

Imports data.table, ggplot2, ggrepel, grDevices, grid, limma, methods, S4Vectors, stats, SummarizedExperiment, utils

Suggests knitr, rmarkdown, testthat (>= 3.0.0)

VignetteBuilder knitr

biocViews Proteomics, MassSpectrometry, DataImport, DataRepresentation, Software, BatchEffect

Config/testthat/edition 3

Config/roxygen2/version 8.1.0

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

git_branch devel

git_last_commit 1c6df35

git_last_commit_date 2026-08-21

Repository Bioconductor 3.24

Date/Publication 2026-09-14

Author Aitor Moruno-Cuenca [aut, cre] (ORCID: <https://orcid.org/0009-0009-8133-2552>),
 Dr. Sergi Sayols-Puig [ctb] (ORCID: <https://orcid.org/0000-0002-3877-4170>),
 Dr. Bruna Oriol-Tordera [ctb] (ORCID: <https://orcid.org/0000-0002-2714-9097>),
 Dr. Hiba Salim [dtc] (ORCID: <https://orcid.org/0000-0002-2168-0258>),
 Dr. Francesc Fernández-Albert [ths] (ORCID: <https://orcid.org/0000-0001-5561-0701>),
 Prof. Dr. Alexandre Perera-Lluna [ths] (ORCID: <https://orcid.org/0000-0001-6427-851X>),
 Dr. Guadalupe Espadas-García [ctb, dtc] (ORCID: <https://orcid.org/0000-0002-6415-8013>),
 Dr. Elisa Monzón-Casanova [ctb, fnd] (ORCID: <https://orcid.org/0000-0001-6617-6138>),
 Prof. Dr. Eduard Sabidó-Aguade [res, ctb, dtc, fnd] (ORCID: <https://orcid.org/0000-0001-6506-7714>)

Maintainer Aitor Moruno-Cuenca <morunoaitor@gmail.com>

Contents

ProteinBatcher-package	2
.pp_export_volcano_tables	3
.pp_infer_interaction_factor	5
.pp_organize_effects	5
filter_se_missing	7
impute_se	9
plot_deregulogram	11
plot_volcano_customized	13
run_proteomics_pipeline	15
test_limma_customized	20

Index	24
--------------	-----------

ProteinBatcher-package

ProteinBatcher: An end-to-end proteomics workflow with condition-aware imputation, flexible statistical modelling and interactive visualization

Description

ProteinBatcher provides utilities to read, validate, and harmonize quantitative proteomics result tables generated by DIA-NN from data-independent acquisition mass spectrometry experiments. The package enables batch processing of multiple DIA-NN output files, consistent handling of protein identifiers and associated metadata, and transformation of tabular data into analysis-ready formats. It facilitates downstream differential abundance analysis using linear modeling frameworks based on limma, with explicit support for multivariable experimental designs including batch effects and other confounding covariates, as well as correlation-aware designs via duplicateCorrelation.

Author(s)

Maintainer: Aitor Moruno-Cuenca <morunoaitor@gmail.com> ([ORCID](#))

Authors:

- Aitor Moruno-Cuenca <morunoaitor@gmail.com> ([ORCID](#))

Other contributors:

- Dr. Sergi Sayols-Puig <sergi.sayolspuig@almirall.com> ([ORCID](#)) [contributor]
- Dr. Bruna Oriol-Tordera <bruna.orioltordera@almirall.com> ([ORCID](#)) [contributor]
- Dr. Hiba Salim <hiba.salim@crg.eu> ([ORCID](#)) [data contributor]
- Dr. Francesc Fernández-Albert <francesc.fernandez@almirall.com> ([ORCID](#)) [thesis advisor]
- Prof. Dr. Alexandre Perera-Lluna <alexandre.perera@upc.edu> ([ORCID](#)) [thesis advisor]
- Dr. Guadalupe Espadas-García <guadalupe.espadas@crg.eu> ([ORCID](#)) [contributor, data contributor]
- Dr. Elisa Monzón-Casanova <elisa.monzoncasanova@almirall.com> ([ORCID](#)) [contributor, funder]
- Prof. Dr. Eduard Sabidó-Aguade <eduard.sabido@upf.edu> ([ORCID](#)) [researcher, contributor, data contributor, funder]

See Also

Useful links:

- <https://github.com/DataScienceRD-Almirall/ProteinBatcher>
- Report bugs at <https://github.com/DataScienceRD-Almirall/ProteinBatcher/issues>

.pp_export_volcano_tables

Export volcano plots and result tables from organized limma effects

Description

Generates volcano plots (PDF) and corresponding result tables for a given effect type (main, common, or interaction) produced by .pp_organize_effects().

Usage

```
.pp_export_volcano_tables(  
  effects,  
  tests,  
  path_output,  
  experiment,  
  effect_slot = c("all_common_effect", "common_effect", "interaction_effect"),  
  file_tag = c("MainEffect_AllProteins", "CommonEffect", "InteractionEffect"),  
  plot_contrasts = NULL,  
  name_col = "Genes",  
  name_imputed = "imputed",  
  alpha = 0.05,  
  lfc = 1  
)
```

Arguments

<code>effects</code>	Named list as returned by <code>.pp_organize_effects()</code> . Each element corresponds to a main contrast and contains <code>SummarizedExperiment</code> objects for different effect types.
<code>tests</code>	Character vector of main contrast names (e.g. "IL13_vs_NoTreated").
<code>path_output</code>	Character. Output directory where PDF files are written.
<code>experiment</code>	Character. Experiment identifier used in output filenames.
<code>effect_slot</code>	Character. Which effect to export. One of "all_common_effect", "common_effect", or "interaction_effect".
<code>file_tag</code>	Character. Tag used in output filenames describing the exported effect (e.g. "MainEffect_AllProteins", "CommonEffect", "InteractionEffect").
<code>plot_contrasts</code>	Character vector of contrasts to be plotted. By default, this is set to <code>tests</code> . For <code>effect_slot = "interaction_effect"</code> , this should typically be the corresponding interaction contrasts (e.g. <code>tests_interaction</code>), since interaction result objects do not contain statistics for the main contrasts.
<code>name_col</code>	Character. Column in <code>rowData()</code> used for labeling significant points in volcano plots (default: "Genes").
<code>name_imputed</code>	Character. Column in <code>rowData()</code> indicating imputed features.
<code>alpha</code>	Numeric. Significance threshold used in volcano plots.
<code>lfc</code>	Numeric. Absolute log2 fold-change threshold used in volcano plots.

Details

For each contrast, the function extracts the requested effect slot, generates a volcano plot using [plot_volcano_customized](#), and returns the corresponding `rowData()` as a named list.

If the selected effect object is `NULL` or contains zero rows, the corresponding contrast is skipped silently.

Output PDF filenames follow the pattern:

```
<experiment>_<file_tag>_<plot_contrast>.pdf
```

This function does not perform additional filtering beyond what is already encoded in the provided effects object.

Value

A list (one element per main contrast) containing named lists of `rowData()` tables. Contrasts that are skipped return `NULL`.

See Also

[.pp_organize_effects](#), [plot_volcano_customized](#)

`.pp_infer_interaction_factor`*Infer a 2-level interaction factor from colData() and tests_interaction*

Description

Internal helper to infer the interaction factor name and its two levels by scanning colData(se) for variables with exactly two levels and matching the non-reference level against tests_interaction (e.g. "sexmale", "batchday2"). Returns NULL if no suitable factor is found.

Usage

```
.pp_infer_interaction_factor(se, tests_interaction)
```

Arguments

`se` SummarizedExperiment.
`tests_interaction` Character vector of interaction contrasts/coefficients.

Value

A list with name and levels (length 2) or NULL.

`.pp_organize_effects` *Organize limma outputs into common vs interaction effects (plot-ready)*

Description

Organize limma outputs into common vs interaction effects (plot-ready)

Usage

```
.pp_organize_effects(  
  se_limma,  
  tests,  
  tests_interaction = "NA",  
  alpha = 0.05,  
  base_cols = c("Protein.Group", "Protein.Names", "Genes", "First.Protein.Description",  
    "name", "ID", "Index", "imputed")  
)
```

Arguments

<code>se_limma</code>	SummarizedExperiment returned by <code>test_limma_customized()</code> .
<code>tests</code>	Character vector of main contrasts (prefixes in <code>rowData</code>), e.g. "IL13_vs_NoTreated".
<code>tests_interaction</code>	Character vector of interaction tests (prefixes in <code>rowData</code>) or "NA". If length 1, it will be recycled to match <code>length(tests)</code> .
<code>alpha</code>	FDR threshold for interaction significance (default 0.05).
<code>base_cols</code>	Annotation columns to keep (if present).

Value

Named list (by tests). Each element is a list with: `common_effect`, `interaction_effect`, `all_common_effect` (SummarizedExperiment objects).

Examples

```
if (requireNamespace("limma", quietly = TRUE)) {
  library(SummarizedExperiment)
  library(S4Vectors)

  ## Minimal imputed SE-like object
  mat <- matrix(
    c(10,11, 9,10,
      20,21,19,20,
      5, NA, 6, 7),
    nrow = 3, byrow = TRUE,
    dimnames = list(paste0("p",1:3), paste0("s",1:4))
  )

  cd <- DataFrame(
    condition = c("A", "A", "B", "B"),
    batch      = c("d1", "d2", "d1", "d2"),
    replicate  = c("r1", "r2", "r1", "r2"),
    donor_id   = c("1", "1", "1", "1"),
    label      = colnames(mat),
    row.names  = colnames(mat)
  )

  rd <- DataFrame(
    name  = rownames(mat),
    ID    = rownames(mat),
    Genes = paste0("G",1:3),
    row.names = rownames(mat)
  )

  se_imp <- SummarizedExperiment(
    assays = list(intensity = mat),
    colData = cd,
    rowData = rd
  )

  ## minimal imputation map required by your limma wrapper
  metadata(se_imp)$imputation_map <- matrix(
    "none", nrow = nrow(mat), ncol = ncol(mat),
    dimnames = dimnames(mat)
  )
}
```

```

)

## Run limma wrapper -> this is se_limma
se_limma <- test_limma_customized(
  se = se_imp,
  test = "B_vs_A",
  test_interaction = "NA",
  design_formula = ~ 0 + condition + batch,
  ref_condition = "A"
)

## Organize effects (no interaction)
eff <- ProteinBatcher:::pp_organize_effects(
  se_limma = se_limma,
  tests = "B_vs_A",
  tests_interaction = "NA"
)

eff[["B_vs_A"]][all_common_effect
}

```

filter_se_missing	<i>Filter proteins by missingness within conditions (SummarizedExperiment)</i>
-------------------	--

Description

filter_se_missing() removes proteins (rows) that fail a per-condition missingness threshold. For each condition in colData(se)\$condition, the function computes the **proportion of non-missing intensities per protein** and keeps a protein if it meets the threshold in **at least one** condition.

This is useful in label-free proteomics pipelines prior to imputation and differential analysis, ensuring that proteins are not uniformly sparse across all conditions.

Usage

```
filter_se_missing(se, percentage = 50)
```

Arguments

se	A SummarizedExperiment (or compatible) object with: - an assay matrix accessible via assay(se) (numeric, may contain NA) - colData(se)\$condition (character or factor) giving the experimental condition per sample. (optional) rowData(se)\$Protein.Names and rowData(se)\$Genes used to annotate the returned table of removed proteins. If missing, they will be safely filled with NA_character_.
percentage	Numeric scalar in [0, 100]. Minimum proportion of non-missing values required within a condition for a protein to be retained. Default: 50 (i.e., at least half of samples observed within any one condition).

Details

Let X be the assay matrix (`assay(se)`), rows = proteins and columns = samples. For every distinct condition in `colData(se)$condition`, we compute `rowMeans(!is.na(X[, condition == cn]))`. A protein is **kept** if the per-condition proportion of observed values is $\geq \text{percentage} / 100$ for **any** condition. Otherwise, it is **removed**.

Value

A named list with two elements:

`se_filt` SummarizedExperiment containing only retained proteins.

`removed` A data.frame with metadata for proteins that were filtered out, columns: `protein_id`, `gene_name`, and `reason`.

Notes

- Samples with NA in `colData(se)$condition` are **excluded** from per-condition computations and a message is emitted.
- If `percentage = 0`, all proteins with at least one observed value in any condition are kept. If `percentage = 100`, only proteins fully observed within at least one condition are kept.

See Also

- `SummarizedExperiment::SummarizedExperiment`
- Downstream steps like `impute_se()` for imputation.

Examples

```
library(SummarizedExperiment)

## Toy expression matrix: 4 proteins x 4 samples
mat <- matrix(
  c(
    1, NA, 2, 3, # p1: OK in cond B
    NA, NA, NA, NA, # p2: always missing -> removed
    5, 6, NA, NA, # p3: OK in cond A
    NA, NA, 1, NA # p4: too sparse everywhere
  ),
  nrow = 4, byrow = TRUE,
  dimnames = list(
    paste0("p", 1:4),
    paste0("s", 1:4)
  )
)

## Sample conditions
cd <- DataFrame(
  condition = c("A", "A", "B", "B"),
  row.names = colnames(mat)
)

## Protein annotations (optional)
rd <- DataFrame(
  Protein.Names = paste0("Prot", 1:4),
```

```

    Genes = paste0("Gene", 1:4),
    row.names = rownames(mat)
  )

  se <- SummarizedExperiment(
    assays = list(intensity = mat),
    colData = cd,
    rowData = rd
  )

  ## Keep proteins with >= 50% observed values in at least one condition
  out <- filter_se_missing(se, percentage = 50)

  ## Filtered SummarizedExperiment
  out$se_filt

  ## Table of removed proteins with reasons
  out$removed

```

impute_se	<i>Impute missing proteomics intensities within conditions (Summarized-Experiment)</i>
-----------	--

Description

`impute_se()` imputes missing values using a two-step, condition-aware strategy: (1) within-condition mean when missingness < threshold; (2) left-censored draws (LDV) for remaining NAs, using a low-intensity reference and per-protein SD.

Usage

```

impute_se(
  se,
  threshold = 0.5,
  ldv_source = c("global", "per_condition"),
  sd_fallback = NULL,
  lower_bound = NULL
)

```

Arguments

<code>se</code>	A <code>SummarizedExperiment</code> with numeric assay and <code>colData(se)\$condition</code> .
<code>threshold</code>	Numeric in $(0, 1]$, default 0.5.
<code>ldv_source</code>	<code>c("global", "per_condition")</code> , default "global".
<code>sd_fallback</code>	Optional numeric fallback SD; default is median non-zero SD.
<code>lower_bound</code>	Optional minimum clip for imputed values (avoid on log scale).

Details

Let `assay(se)` be a numeric matrix (rows = proteins, cols = samples); `colData(se)$condition` provides the condition for each sample. Step (1) replaces NAs in a condition with the mean of observed values for that protein in the same condition when the NA proportion is below threshold. Step (2) replaces remaining NAs with `rnorm(1, mean = LDV, sd = sd_protein)`, where LDV is the global (or per-condition) minimum observed intensity; per-protein SDs are computed from available values with a robust fall-back when needed.

The per-cell imputation method is recorded in `metadata(se)$imputation_map("none"|"mean"|"ldv")`. A row-level flag is added at `rowData(se)$imputed`.

Value

A `SummarizedExperiment` with imputed assay, updated `metadata/rowData`.

See Also

`SummarizedExperiment::SummarizedExperiment`

Examples

```
library(SummarizedExperiment)

## Toy matrix with missing values
mat <- matrix(
  c(
    10, 11, NA, NA, # p1: partial missing
    NA, NA, NA, NA, # p2: fully missing
    5, NA, 6, 7 # p3: mixed
  ),
  nrow = 3, byrow = TRUE,
  dimnames = list(
    paste0("p", 1:3),
    paste0("s", 1:4)
  )
)

## Conditions for samples
cd <- DataFrame(
  condition = c("A", "A", "B", "B"),
  row.names = colnames(mat)
)

## Build SummarizedExperiment
se <- SummarizedExperiment(
  assays = list(intensity = mat),
  colData = cd
)

## Impute with default strategy:
## - mean within condition if missingness < threshold
## - LDV otherwise
set.seed(123)
se_imp <- impute_se(se, threshold = 0.5, ldv_source = "global")

## Imputed assay
```

```

assay(se_imp)

## Per-cell imputation method
metadata(se_imp)$imputation_map

## Row-level flag: was any value imputed?
rowData(se_imp)$imputed

```

plot_deregulogram	<i>Deregulogram plot for interaction models (2-level factors)</i>
-------------------	---

Description

Creates a deregulogram: a scatter of full effects in two levels of an interaction factor (e.g. sex female vs male, batch day1 vs day2), colored by whether the interaction is significant. This visualization complements volcano plots by showing effect concordance and sex-/batch-bias in magnitude and direction, as described in the proteomics workflow documentation.

Usage

```

plot_deregulogram(
  se_limma,
  tests,
  tests_interaction,
  factor_name,
  factor_levels,
  path_output,
  experiment,
  alpha = 0.05,
  lfc = 1,
  label_col = "Genes"
)

```

Arguments

se_limma	A SummarizedExperiment containing limma results in rowData() (e.g., output of test_limma_customized()).
tests	Character vector of main contrasts (e.g. "IL13_vs_NoTreated").
tests_interaction	Character vector of interaction contrasts/coefficients aligned with tests (e.g. "IL13.sexmale" or "IL13.batchday2").
factor_name	Character scalar, name of the interaction factor (e.g. "sex", "batch"). Used only for titles/labels.
factor_levels	Character vector of length 2 giving the factor levels in order c(ref, other) (e.g. c("female", "male")).
path_output	Output directory where PDFs are written.
experiment	Experiment identifier used in output filenames.
alpha	Numeric. Significance threshold (default 0.05).
lfc	Numeric. Absolute log2FC threshold used for highlighting (default 1).
label_col	Column in rowData(se_limma) used for point labels (default "Genes").

Details

The function assumes that differential testing has already been performed and stored in `rowData(se_limma)` using the "<stat>_<contrast>" naming convention (e.g., "log2FC_IL13_vs_NoTreated", "p.adj_IL13_vs_NoTreated", "log2FC_IL13.batchday2", "p.adj_IL13.batchday2").

Deregulogram axes:

- y-axis: full effect in the reference level (level 1)
- x-axis: full effect in the non-reference level (level 2), computed as $\log_2FC_{\text{main}} + \log_2FC_{\text{interaction}}$

This plot is meaningful only when the interaction factor has exactly two levels. Use `.pp_infer_interaction_factor()` (internal) to infer the factor from `colData()` and `tests_interaction`, or pass factor name and levels explicitly.

Important: gene sets do not have to match the interaction volcano plot. The interaction volcano plot (see `plot_volcano_customized` on the `interaction_effect` object) identifies proteins with a statistically significant interaction term (e.g. `p.adj_interaction < alpha` and `|log2FC_interaction| >= 1fc`). In contrast, the deregulogram is designed to highlight a more interpretable subset where the interaction is not only statistically significant, but also occurs in the context of a relevant condition effect (i.e., the main/full effects reach the chosen alpha and 1fc thresholds). Therefore, the deregulogram typically shows a subset of the interaction-volcano hits and is not expected to reproduce the same labeled proteins one-to-one.

Deregulogram axes compare full effects between two levels of the interaction factor. For a main contrast C and a 2-level factor (ref vs level2), the full effect in the reference level is $\log_2FC_C$, while the full effect in the second level is $\log_2FC_C + \log_2FC_{\{C: \text{factor}\}}$. Points are colored by whether the interaction is significant and optionally labeled to emphasize proteins whose condition effect is factor-dependent.

Value

Invisibly returns NULL. Side effect: writes one PDF per contrast.

Examples

```
library(SummarizedExperiment)
library(S4Vectors)

## -----
## Minimal SummarizedExperiment with main + interaction statistics
## -----

mat <- matrix(
  rnorm(12),
  nrow = 3,
  dimnames = list(
    paste0("p", 1:3),
    paste0("s", 1:4)
  )
)

## Sample metadata (interaction factor has 2 levels)
cd <- DataFrame(
  condition = c("A", "A", "B", "B"),
  sex       = c("female", "male", "female", "male"),
  row.names = colnames(mat)
)
```

```

## Main and interaction contrasts
main_contrast <- "B_vs_A"
inter_contrast <- "B_vs_A.sexmale"

## rowData with columns expected by plot_deregulogram()
rd <- DataFrame(
  Genes = c("G1", "G2", "G3"),
  row.names = rownames(mat)
)

## Main effect
rd[[paste0("log2FC_", main_contrast)]] <- c( 1.5, 0.2, -1.3)
rd[[paste0("p.adj_", main_contrast)]] <- c( 0.01, 0.8, 0.03)

## Interaction effect
rd[[paste0("log2FC_", inter_contrast)]] <- c( 0.7, -0.1, 1.1)
rd[[paste0("p.adj_", inter_contrast)]] <- c( 0.02, 0.9, 0.01)

se_limma <- SummarizedExperiment(
  assays = list(intensity = mat),
  colData = cd,
  rowData = rd
)

## -----
## Create deregulogram (PDF written to tempdir())
## -----

plot_deregulogram(
  se_limma = se_limma,
  tests = main_contrast,
  tests_interaction = inter_contrast,
  factor_name = "sex",
  factor_levels = c("female", "male"),
  path_output = tempdir(),
  experiment = "EXAMPLE",
  alpha = 0.05,
  lfc = 1,
  label_col = "Genes"
)

```

plot_volcano_customized

Volcano plot for differential abundance results

Description

Generates a volcano plot from differential testing results stored in a SummarizedExperiment. The function expects contrast-specific columns in rowData(dep) following the naming convention "<contrast>_diff" and either "<contrast>_p.adj" or "<contrast>_p.val".

Usage

```
plot_volcano_customized(
  dep,
  contrast,
  label_size = 3,
  name_col = "ID",
  name_imputed = "imputed",
  add_names = TRUE,
  adjusted = TRUE,
  plot = TRUE,
  alpha = 0.05,
  lfc = 1
)
```

Arguments

dep	A SummarizedExperiment containing differential testing results in <code>rowData()</code> .
contrast	Character scalar specifying the contrast to plot. Must match the prefix used in <code>rowData(dep)</code> (e.g. "IL13_vs_NoTreated").
label_size	Numeric. Text size for feature labels in the plot.
name_col	Character. Column in <code>rowData(dep)</code> used for labeling significant points (default: "ID").
name_imputed	Character or NULL. Column in <code>rowData(dep)</code> indicating imputed features (logical). Used to color points. If NULL or not present, all features are treated as non-imputed.
add_names	Logical. If TRUE, labels significant features using ggrepel .
adjusted	Logical. If TRUE, uses adjusted p-values (" <code><contrast>_p.adj</code> "); otherwise uses raw p-values (" <code><contrast>_p.val</code> ").
plot	Logical. If TRUE, returns a ggplot object; otherwise returns a data.frame with volcano statistics.
alpha	Numeric. Significance threshold for p-values.
lfc	Numeric. Absolute log2 fold-change threshold.

Details

Significance is determined internally using a log2 fold-change threshold (`lfc`) and a p-value or adjusted p-value threshold (`alpha`). Optionally, significant features can be labeled using a column from `rowData(dep)` (e.g. gene symbols).

Points are colored according to significance and imputation status: imputed & significant, non-imputed & significant, or non-significant. The plot title and axis annotations are derived from the contrast name.

This function does not rely on precomputed `*_significant` columns; significance is evaluated on the fly from `diff`, `p.adj/p.val`, `alpha`, and `lfc`.

Value

If `plot = TRUE`, a **ggplot** volcano plot. If `plot = FALSE`, a data.frame with log2 fold change, $-\log_{10}(\text{p-value})$, and significance flag.

Examples

```

library(SummarizedExperiment)
library(S4Vectors)

## Minimal SummarizedExperiment with differential results in rowData
mat <- matrix(
  rnorm(12),
  nrow = 3,
  dimnames = list(
    paste0("p", 1:3),
    paste0("s", 1:4)
  )
)

cd <- DataFrame(
  condition = c("A", "A", "B", "B"),
  row.names = colnames(mat)
)

rd <- DataFrame(
  ID = c("Prot1", "Prot2", "Prot3"),
  imputed = c(FALSE, TRUE, FALSE),
  row.names = rownames(mat)
)

contrast <- "B_vs_A"

## Columns expected by plot_volcano_customized()
rd[[paste0("log2FC_", contrast)]] <- c(2.0, 0.3, -1.5)
rd[[paste0("p.adj_", contrast)]] <- c(0.01, 0.9, 0.03)

se <- SummarizedExperiment(
  assays = list(intensity = mat),
  colData = cd,
  rowData = rd
)

## Return volcano statistics as a data.frame (no plotting)
df <- plot_volcano_customized(
  dep = se,
  contrast = contrast,
  plot = FALSE
)

df

```

run_proteomics_pipeline

Run a the downstream proteomics pipeline (filter, impute, limma, reporting)

Description

This workflow performs a reproducible downstream analysis for quantitative proteomics: (i) imports quantification and sample annotation into a SummarizedExperiment, (ii) filters features by

missingness, (iii) imputes remaining missing values using a mean vs LDV (left-censored) strategy, (iv) runs differential abundance testing with limma (optionally including an interaction term), (v) organizes results into plot-ready effect objects, and (vi) optionally exports volcano plots, deregulograms and results to disk.

Usage

```
run_proteomics_pipeline(
  path_pgmatrix,
  path_annotation,
  path_output,
  level = "protein",
  type = "DIA",
  experiment,
  percent_missing,
  ldv_source = c("global", "per-condition"),
  threshold = 0.3,
  tests,
  tests_interaction,
  formula,
  reference_condition,
  paired = FALSE,
  block_effect = FALSE,
  block_var = "donor_id",
  annotation_format = c("standard", "sdrf"),
  sdrf_map = NULL,
  plots = FALSE,
  ...
)
```

Arguments

<code>path_pgmatrix</code>	Character scalar. Path to the quantitative matrix file.
<code>path_annotation</code>	Character scalar. Path to the sample annotation TSV.
<code>path_output</code>	Character scalar. Output directory where files may be written.
<code>level</code>	Character. Quantification level (e.g. "protein").
<code>type</code>	Character. Quantification type (e.g. "DIA").
<code>experiment</code>	Character scalar. Experiment identifier used in output filenames.
<code>percent_missing</code>	Numeric. Missingness threshold used for filtering (percentage, e.g. 50).
<code>ldv_source</code>	Character. One of "global" or "per-condition" controlling LDV definition.
<code>threshold</code>	Numeric in (0, 1]. Forwarded to <code>impute_se()</code> . Missingness proportion below which within-condition mean imputation is used; at or above it, LDV (left-censored) imputation is used instead.
<code>tests</code>	Character vector of main contrasts to test. Each element must correspond to a valid condition contrast present in the design matrix (e.g. "IL13_vs_NoTreated").
<code>tests_interaction</code>	Character vector defining interaction contrasts or interaction coefficients aligned with tests. Use "NA" to disable interaction testing for a given contrast. Typical values encode condition-by-factor interactions (e.g. "IL13.batchday2").

formula	A model formula specifying the design matrix for limma. Common examples include <code>~ 0 + condition + batch + condition:batch</code> . All variables referenced in the formula must be present in <code>colData()</code> .
reference_condition	Character scalar specifying the reference level for the condition variable. This level is used to relevel the design prior to fitting the limma model.
paired	Logical. Forwarded to <code>test_limma_customized()</code> . If TRUE, treats the design as paired/repeated-measures (see "Paired designs and blocking" section below).
block_effect	Logical. Forwarded to <code>test_limma_customized()</code> . If TRUE, models correlation between repeated observations via <code>limma::duplicateCorrelation</code> (see "Paired designs and blocking" section below); the blocking variable is specified by <code>block_var</code> .
block_var	Character. Forwarded to <code>test_limma_customized()</code> . Name of the <code>colData</code> column to use as the blocking variable when <code>block_effect = TRUE</code> . Defaults to <code>"donor_id"</code> . Ignored when <code>block_effect = FALSE</code> .
annotation_format	Character. One of <code>"standard"</code> (default) or <code>"sdrf"</code> . Use <code>"sdrf"</code> when <code>path_annotation</code> is an SDRF-Proteomics file rather than a plain annotation TSV with <code>file/sample_name/condition</code> columns; see <code>sdrf_map</code> .
sdrf_map	Named list, only used when <code>annotation_format = "sdrf"</code> . Maps SDRF column names (which are not fixed by the standard) to the roles the pipeline needs, e.g. <code>list(condition = "factor value[treatment]", batch = "comment[batch]", donor_id = "characteristics[individual]")</code> . At minimum <code>condition</code> must be supplied. Ignored when <code>annotation_format = "standard"</code> .
plots	Logical. If TRUE, exports plots/tables to <code>path_output</code> .
...	Additional arguments forwarded only to <code>test_limma_customized()</code> (beyond <code>test</code> , <code>test_interaction</code> , <code>design_formula</code> , <code>ref_condition</code> , <code>paired</code> and <code>block_effect</code> , which are already explicit parameters above).

Value

A named list with:

`se_raw` Raw SummarizedExperiment imported from files.

`se_filt` Filtered SummarizedExperiment.

`removed` A data.frame describing filtered-out features.

`se_imp` Imputed SummarizedExperiment.

`effects` A named list keyed by main contrast (the values of tests). Each element is itself a list of three SummarizedExperiment objects: `all_common_effect`, `common_effect` and `interaction_effect`. Per-contrast statistics are stored in their `rowData()` using the `"<stat>_<contrast>"` naming convention (e.g. `log2FC_IL13_vs_NoTreated`). This element is always returned, independently of `plots`, so a contrast can be accessed with `$`, e.g. `res$effects$IL13_vs_NoTreated$all_common_`

Input files

`path_pgmatrix` Path to a quantitative matrix file produced by the upstream quantification tool (e.g., a TSV). Rows represent quantified features (e.g. protein groups) and columns represent samples.

path_annotation Path to a tab-delimited sample annotation file describing the experimental design. At minimum it must encode sample-to-condition mapping. Additional columns are required (batch, replicate, block). Accepts either a standard annotation TSV (default, `annotation_format = "standard"`) or an SDRF-Proteomics file (`annotation_format = "sdrf"`, with column roles supplied via `sdrf_map`).

Main steps

1. **Import:** creates a raw SummarizedExperiment (`se_raw`).
2. **Missingness filter:** removes features not sufficiently observed across samples/conditions according to `percent_missing`. Filtered-out features are returned and may be written to disk.
3. **Imputation:** imputes remaining missing values using a two-step strategy: within-condition mean imputation when missingness is below a threshold and LDV (left-censored) imputation when missingness is above the threshold. The imputation method is controlled by `ldv_source` and `threshold`.
4. **Differential testing:** calls `test_limma_customized` to fit a limma model and evaluate user-specified contrasts. Interaction testing is optional.
5. **Organize outputs:** splits the limma results into plot-ready effect objects using `.pp_organize_effects`:
 - `all_common_effect`: main-effect statistics for all proteins (no interaction filtering).
 - `common_effect`: proteins not significant for the interaction term (main effect stable).
 - `interaction_effect`: proteins significant for the interaction term.
6. **Optional reporting** (`plots = TRUE`): exports volcano plots and tables for the three effect types above, and generates deregulograms when an interaction factor with exactly two levels is detected (e.g., female vs male, day1 vs day2, ...).

Differential testing inputs

The limma step requires the following explicit parameters:

tests Character vector of main contrasts to test, using a consistent naming scheme (e.g. "A_vs_B"). These define the primary condition effects.

tests_interaction Character vector aligned with `tests` defining interaction contrasts or interaction coefficients (or "NA" to disable interaction testing). These represent factor-dependent differences in the condition effect (e.g. condition-by-batch).

formula A model formula describing the design matrix (e.g. `~ 0 + condition + batch + condition:batch`). Variables referenced in the formula must be present in `colData(se)`.

reference_condition Character scalar indicating the reference level for condition. The function will `relevel colData(se)$condition` to this value before model fitting.

Paired designs and blocking

paired Logical (forwarded to `test_limma_customized`). If TRUE, the model is treated as a paired / repeated-measures design by ensuring `replicate` is included in the design formula. In the current implementation, if `replicate` is not already present in `formula`, it is added automatically. This preserves any additional covariates already present in the formula.

block_effect Logical (forwarded to `test_limma_customized`). If TRUE, correlation between repeated observations is modeled using `limma::duplicateCorrelation` and `lmFit(..., block=..., correlation=...)`. The blocking variable is specified by `block_var` (default: "donor_id"). Use this when there is a known blocking factor (e.g., donor/subject) inducing correlation across samples.

`block_var` Character (forwarded to `test_limma_customized`). Name of the `colData` column to use as the blocking variable for `duplicateCorrelation` when `block_effect = TRUE`. Defaults to `"donor_id"`. Any column present in the annotation file can be used (e.g., `"batch"`, `"patient_id"`). Ignored when `block_effect = FALSE`.

Volcano vs deregulogram (interpretation note)

Interaction volcano plots highlight proteins with significant interaction coefficients (i.e., where the condition effect depends on a second factor). Deregulograms are more restrictive: they visualize full effects across the two levels of the interaction factor and emphasize proteins where interaction occurs in the context of a relevant condition effect (effect-size and FDR thresholds). Therefore, the labeled proteins in the deregulogram are not expected to match one-to-one with those in the interaction volcano plot.

See Also

[test_limma_customized](#), [plot_volcano_customized](#), [plot_deregulogram](#)

Examples

```
## Load package
library(ProteinBatcher)

## -----
## Input files shipped with the package
## -----

path_annotation <- system.file(
  "extdata", "annotation_HaCaT.tsv",
  package = "ProteinBatcher"
)

path_pgmatrix <- system.file(
  "extdata",
  "2024MK017_HaCaT_Stimulation_1to24_Astral_report.pg_matrix.tsv",
  package = "ProteinBatcher"
)

## Output directory
path_output <- tempdir()

## -----
## Workflow parameters (single contrast example)
## -----

experiment <- "HaCaT"
percent_missing <- 50

formula <- ~ 0 + condition + batch + condition:batch

tests <- c("IL13_vs_NoTreated")
tests_interaction <- c("IL13.batchday2")

reference_condition <- "NoTreated"

## -----
```

```

## Run downstream proteomics workflow
## -----

res <- run_proteomics_pipeline(
  path_pgmatrix      = path_pgmatrix,
  path_annotation    = path_annotation,
  path_output        = path_output,
  tests              = tests,
  tests_interaction   = tests_interaction,
  formula            = formula,
  reference_condition = reference_condition,
  percent_missing    = percent_missing,
  ldv_source          = "per-condition",
  experiment         = experiment,
  plots              = FALSE
)

## Inspect outputs
names(res)
res$se_imp

## `effects` is keyed by contrast, so it can be accessed with `$`:
names(res$effects)
res$effects$IL13_vs_NoTreated$all_common_effect

```

test_limma_customized *Differential abundance testing with limma (imputation-aware FDR)*

Description

Performs differential testing on an imputed SummarizedExperiment using **limma**. The function fits a linear model defined by `design_formula`, applies user-specified contrasts (`test`) and, optionally, interaction-related contrasts (`test_interaction`), and stores the resulting statistics in `rowData(se)`.

Usage

```

test_limma_customized(
  se,
  type = c("manual"),
  test = NULL,
  test_interaction = "NA",
  design_formula = stats::formula(~0 + condition),
  ref_condition = NULL,
  paired = FALSE,
  block_effect = FALSE,
  block_var = "donor_id"
)

```

Arguments

<code>se</code>	A SummarizedExperiment. The assay is taken from <code>SummarizedExperiment::assay(se)</code> .
<code>type</code>	Character. Currently only "manual" is supported (kept for API compatibility).

test	Character vector of contrasts to test for the main effect. Contrasts must be supplied in the "A_vs_B" format (internally translated to "A - B" for limma).
test_interaction	Character vector defining interaction-related tests. Use "NA" (default) to skip interaction testing. If provided (not "NA"), results for these contrasts will also be added to rowData(se).
design_formula	A model formula describing the linear model. Default is $\sim 0 + \text{condition}$. The function expects condition to be present in colData(se) and will relevel it to ref_condition. If paired = TRUE and replicate is not present in the formula, replicate is added.
ref_condition	Character scalar. Reference level for condition. Must be a level present in colData(se)\$condition.
paired	Logical. If TRUE, replicate is added to the design (if absent) to support paired/repeated-measure designs for technical replicates.
block_effect	Logical. If TRUE, fits limma with a blocking factor using limma::duplicateCorrelation() and the colData column specified by block_var.
block_var	Character. Name of the colData(se) column to use as the blocking variable for duplicateCorrelation when block_effect = TRUE. Defaults to "donor_id". The column must exist in colData(se) and have at least two levels. Ignored when block_effect = FALSE.

Details

A key feature of this customized implementation is that adjusted p-values (BH-FDR) are recomputed after excluding features that were imputed as *LDV* in both groups of a tested contrast, using an imputation_map stored in metadata(se). This prevents artificially significant results driven by LDV/LDV imputations.

When paired = TRUE, the function ensures the design includes replicate by updating the right-hand side of design_formula (it does not overwrite other covariates such as batch). When block_effect = TRUE, the function fits the model using limma::duplicateCorrelation() with the column specified by block_var (default: donor_id) as the blocking variable.

Results are written into rowData(se) as wide columns per contrast, using prefixes corresponding to each tested comparison. For each contrast, the following statistics are stored (naming depends on internal helpers):

- diff: log2 fold-change estimate (logFC)
- CI.L, CI.R: confidence interval bounds (if computed)
- p.val: raw p-value
- p.adj: BH-adjusted p-value, recomputed after excluding LDV/LDV

Interaction handling is optional and depends on test_interaction. In the associated proteomics workflow, interaction tests are used to distinguish "common" effects from "interaction" effects (e.g., condition-by-sex or condition-by-batch interactions).

Value

The input SummarizedExperiment with additional columns appended to rowData(se) containing differential testing results.

See Also

[lmFit](#), [makeContrasts](#), [contrasts.fit](#), [eBayes](#), [topTable](#), [duplicateCorrelation](#)

Examples

```

library(SummarizedExperiment)

## Toy proteomics matrix: 4 proteins x 4 samples
mat <- matrix(
  c(
    10, 11, 9, 10,
    20, 21, 19, 20,
    5, NA, 6, 7,
    NA, NA, NA, NA
  ),
  nrow = 4, byrow = TRUE,
  dimnames = list(
    paste0("p", 1:4),
    paste0("s", 1:4)
  )
)

## Sample metadata
cd <- DataFrame(
  condition = c("A", "A", "B", "B"),
  batch      = c("d1", "d1", "d2", "d2"),
  replicate  = c("r1", "r2", "r1", "r2"),
  donor_id   = c("1", "1", "1", "1"),
  label      = colnames(mat),
  row.names  = colnames(mat)
)

## Feature metadata
rd <- DataFrame(
  name      = rownames(mat),
  Genes     = paste0("G", 1:4),
  row.names = rownames(mat)
)

se <- SummarizedExperiment(
  assays = list(intensity = mat),
  colData = cd,
  rowData = rd
)

## Imputation map required by the FDR logic
imp_map <- matrix(
  "none", nrow = nrow(mat), ncol = ncol(mat),
  dimnames = dimnames(mat)
)
metadata(se)$imputation_map <- imp_map

## Run differential testing (main effect only)
se_out <- test_limma_customized(
  se = se,
  test = "B_vs_A",
  test_interaction = "NA",
  design_formula = ~ 0 + condition + batch,
  ref_condition = "A"
)

```

```
## Results are appended to rowData(se_out)  
rowData(se_out)
```

Index

- * **internal**
 - .pp_export_volcano_tables, [3](#)
 - .pp_infer_interaction_factor, [5](#)
 - .pp_organize_effects, [5](#)
 - ProteinBatcher-package, [2](#)
 - .pp_export_volcano_tables, [3](#)
 - .pp_infer_interaction_factor, [5](#)
 - .pp_organize_effects, [4](#), [5](#)
- contrasts.fit, [21](#)
- duplicateCorrelation, [21](#)
- eBayes, [21](#)
- filter_se_missing, [7](#)
- ggplot, [14](#)
- impute_se, [9](#)
- lmFit, [21](#)
- makeContrasts, [21](#)
- plot_deregulogram, [11](#), [19](#)
- plot_volcano_customized, [4](#), [13](#), [19](#)
- ProteinBatcher
 - (ProteinBatcher-package), [2](#)
- ProteinBatcher-package, [2](#)
- run_proteomics_pipeline, [15](#)
- test_limma_customized, [18](#), [19](#), [20](#)
- topTable, [21](#)