

Package ‘GSEAlens’

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Type Package

Title Gene Set Enrichment Analysis Interactive Explorer

Version 0.99.35

Description GSEAlens provides an interactive exploration layer on top of standard Bioconductor RNA-seq workflows. It consumes fitted model objects from limma (MArrayLM) or DESeq2 (DESeqDataSet) as input and accepts expression matrices and sample metadata as SummarizedExperiment objects, ensuring interoperability with core Bioconductor data containers. For core computation, GSEAlens wraps clusterProfiler::GSEA() as its statistical engine (thereby inheriting the fgsea fast GSEA methodology) and draws on MSigDB gene set collections via the msigdbR package from CRAN; multi-contrast parallel computation is handled by future (future::multisession). Visualization output relies on Bioconductor graphics packages including enrichplot, ComplexHeatmap, and circlize, producing figures suitable for publication pipelines. The package also includes a built-in Shiny application for interactive exploration of enrichment results after DEG analysis, with the ability to export self-contained reproducible R scripts.

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URL <https://github.com/DDI095/GSEAlens>

BugReports <https://github.com/DDI095/GSEAlens/issues>

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GSEAlens-package

GSEAlens: Gene Set Enrichment Analysis Interactive Explorer

Description

GSEAlens provides an interactive exploration layer on top of standard Bioconductor RNA-seq workflows. It consumes fitted model objects from limma (MAArrayLM) or DESeq2 (DESeqDataSet) as input and accepts expression matrices and sample metadata as SummarizedExperiment objects, ensuring interoperability with core Bioconductor data containers.

For core computation, GSEAlens wraps [clusterProfiler::GSEA()] as its statistical engine (thereby inheriting the fgsea fast GSEA methodology) and draws on MSigDB gene set collections via the msigdb package from CRAN. Multi-contrast parallel computation is handled by future (future::multisession).

Visualization output relies on Bioconductor graphics packages including enrichplot, ComplexHeatmap, and circlize, producing figures suitable for publication pipelines. The package also includes a built-in Shiny application for interactive exploration of enrichment results after DEG analysis, with the ability to export self-contained reproducible R scripts.

Details

****Main entry points:****

- [setup_gsea_env()]: Initialize a GSEA analysis environment from limma or DESeq2 fitted objects.
- [batch_calc_gsea()]: Run GSEA across multiple contrasts in parallel.
- [build_gsea_pathways()]: Build MSigDB pathway database (interactive or automated selection).
- [extract_gsea_task()]: Extract a single contrast for downstream visualization and reporting.

****Built-in Shiny application:****

Launch the interactive explorer via [launch_gsea_app()] after DEG analysis. The app provides pathway-level drill-down, multi-contrast joint canvases, quadrant plots, hub-gene networks, and one-click export of self-contained reproducible R scripts.

****Vignettes:****

- 'vignette("GSEAlens", package = "GSEAlens")': Main workflow
- 'vignette("GSEAlens-preprocessing", package = "GSEAlens")': Data preprocessing

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References

<<https://github.com/DDL095/GSEALens>>

See Also

Useful links:

- <https://github.com/DDL095/GSEALens>
- Report bugs at <https://github.com/DDL095/GSEALens/issues>

Examples

```
library(GSEALens)
packageDescription("GSEALens")$Version
```

```
.auto_detect_addition_data
```

Auto-detect Addition Data

Description

Search working directory for default addition_data files. Checks for 'addition_data_gsealens.rds' or 'addition_data_gsealens.csv' in the current working directory.

Usage

```
.auto_detect_addition_data()
```

Value

A validated data.frame or NULL if no file found

```
.build_expr_bundle
```

Build Expression Data Bundle

Description

Uniformly encapsulate expression matrix and metadata

Usage

```
.build_expr_bundle(obj, backend)
```

<code>.check_gsea_env</code>	<i>Validate GseaEnv Object Integrity</i>
------------------------------	--

Description

Internal function to ensure object structure conforms to specification.

Usage

```
.check_gsea_env(env_obj)
```

Arguments

<code>env_obj</code>	GseaEnv object to validate
----------------------	----------------------------

Value

TRUE if validation passes, otherwise stops with an error

<code>.check_gsea_res</code>	<i>Validate GseaRes Object Integrity</i>
------------------------------	--

Description

Internal function to ensure GseaRes object contains required components.

Usage

```
.check_gsea_res(res_obj)
```

Arguments

<code>res_obj</code>	GseaRes object to validate
----------------------	----------------------------

Value

TRUE if validation passes, otherwise stops with an error

`.convert_expr_rownames`*Convert Expression Matrix Row Names to Gene Symbols*

Description

Automatically detect and convert expression matrix row names, supporting Ensembl ID to gene symbol conversion.

Usage

```
.convert_expr_rownames(expr_mat, gene_meta, gsea_res)
```

Arguments

<code>expr_mat</code>	An expression matrix.
<code>gene_meta</code>	A gene metadata data frame containing both ID and symbol columns.
<code>gsea_res</code>	A GseaRes object (used to access backend information).

Value

The expression matrix with updated row names.

`.derive_task_seed` *Derive Deterministic Per-Task Seed*

Description

Derive Deterministic Per-Task Seed

Usage

```
.derive_task_seed(seed, task_id, all_task_ids)
```

Arguments

<code>seed</code>	Integer or NULL. Base seed chosen by the user.
<code>task_id</code>	Character. Contrast task identifier.
<code>all_task_ids</code>	Character vector. All task ids in the current run.

Value

Integer seed derived from the base seed and the task's rank in the alphabetically sorted task list, or NULL when 'seed' is NULL, NA or non-numeric. Ranking against the sorted full task list (not the chunk-local order) makes the derived seed independent of 'workers' and 'chunk_size'.

`.detect_gene_symbol_column`

Detect Gene Symbol Column

Description

Automatically detect the gene symbol column in gene metadata. Uses housekeeping gene probe matching for robust cross-species detection.

Usage

```
.detect_gene_symbol_column(gene_meta)
```

Arguments

gene_meta

A data frame containing gene metadata.

Value

The detected column name if successful, or NULL if detection fails.

`.enrich_gsea_result`

GSEA Result Metadata Injection

Description

GSEA Result Metadata Injection

Usage

```
.enrich_gsea_result(result_df, meta_dict)
```

Arguments

result_df

GSEA result dataframe

meta_dict

Metadata dictionary

Value

Enriched dataframe

`.extract_deseq2_data` *Extract DESeq2 Data*

Description

Extract contrast groups and differential analysis results from DESeqDataSet object.

Usage

```
.extract_deseq2_data(dds, target_factor = NULL)
```

Arguments

<code>dds</code>	DESeqDataSet object (must have been processed with DESeq())
<code>target_factor</code>	Character. Target factor. If NULL, automatically inferred as the last term in the design formula.

Value

A list containing `contrast_registry`, `de_store`, and `expr_bundle`

`.extract_limma_data` *Extract Limma-Voom Data*

Description

Extract contrast groups and differential analysis results from MArrayLM object.

Usage

```
.extract_limma_data(fit, expr_data = NULL)
```

Arguments

<code>fit</code>	MArrayLM object (must have been processed with eBayes)
<code>expr_data</code>	Optional. DGEList or expression matrix.

Value

A list containing `contrast_registry`, `de_store`, and `expr_bundle`

<code>.get_display_symbol</code>	<i>Get Display Gene Symbol</i>
----------------------------------	--------------------------------

Description

Retrieve the original case gene symbol based on the case mapping.

Usage

```
.get_display_symbol(gene, symbol_map)
```

Arguments

<code>gene</code>	A gene identifier (case-insensitive).
<code>symbol_map</code>	A case mapping vector generated by <code>.rebuild_symbol_map</code> .

Value

The original case gene symbol, or the input if no mapping exists.

<code>.get_display_symbols</code>	<i>Batch Get Display Gene Symbols</i>
-----------------------------------	---------------------------------------

Description

Batch Get Display Gene Symbols

Usage

```
.get_display_symbols(genes, symbol_map)
```

Arguments

<code>genes</code>	A character vector of gene identifiers.
<code>symbol_map</code>	A case mapping vector generated by <code>.rebuild_symbol_map</code> .

Value

A character vector of gene symbols in their original case.

<code>.get_expr_internal</code>	<i>Internal Expression Matrix Extraction Logic</i>
---------------------------------	--

Description

Core logic for extracting and computing expression matrices. Handles backend-specific computations (DESeq2, limma-voom) and normalizes sample ordering between expression matrices and metadata.

Usage

```
.get_expr_internal(expr_bundle, backend_info, type = "default", ...)
```

<code>.gsea_nb_core</code>	<i>Internal GSEA Native Plotting Engine</i>
----------------------------	---

Description

Core plotting logic migrated from `GseaVis::gseaNb`, stripped of unused features (`newGsea`, `filePath`, `KEGG`, `geneExpHt`, etc.). Returns a list of `ggplot` objects for downstream assembly via `patchwork`.

Usage

```
.gsea_nb_core(  
  object,  
  geneSetID,  
  subPlot = 3,  
  curveCol = c("#76BA99", "#EB4747", "#996699"),  
  htCol = c("#08519C", "#A50F15"),  
  lineSize = 0.8,  
  addPval = FALSE,  
  pvalX = 0.9,  
  pvalY = 0.9,  
  pvalSize = 4,  
  pCol = "grey0",  
  nesDigit = 2,  
  pDigit = 2,  
  show_contrast_in_axis = FALSE,  
  left_group = "Left",  
  right_group = "Right",  
  base_size = 14,  
  subRatio = c(0.5, 0.2, 0.3),  
  rankSeq = 5000,  
  htHeight = 0.3,  
  htAlpha = 0.8  
)
```

Arguments

object	gseaResult object.
geneSetID	Character vector of pathway IDs.
subPlot	Integer, 1/2/3.
curveCol	Curve color vector.
htCol	Heatmap gradient colors.
lineSize	Line width for enrichment curve.
addPval	Logical, add NES/Pvalue text annotation (single pathway only).
pvalX	X position for pval text (relative fraction).
pvalY	Y position for pval text (relative fraction).
pvalSize	Text size for pval annotation.
pCol	Color for pval text.
nesDigit	NES rounding digits.
pDigit	P-value rounding digits.
show_contrast_in_axis	Logical, if TRUE shows "Left vs Right" on X-axis.
left_group	Left group label.
right_group	Right group label.
base_size	Theme base size.
subRatio	Relative heights for 3 panels.
rankSeq	X-axis break interval for rank plot.
htHeight	Relative height of heatmap strip inside Panel 2.
htAlpha	Alpha for heatmap rectangles.

Value

Named list: p1 (curve), p2 (heatmap strip), p3 (rank distribution), data_ga, gsdata, glist.

.gs_info	<i>Internal GSEA Data Extractor</i>
----------	-------------------------------------

Description

Extract running score data from gseaResult object. Adapted from GseaVis::gsInfo, using enrichit::gseaScores internally.

Usage

```
.gs_info(object, geneSetID)
```

Arguments

object	A gseaResult object.
geneSetID	Character, gene set ID.

Value

data.frame with runningScore, position, geneList, etc.

<code>.launch_gsea_shiny</code>	<i>Launch GSEAlens Shiny App (Internal)</i>
---------------------------------	---

Description

Core Shiny application launcher. Builds the UI layout and initializes all server modules for interactive GSEA analysis.

Usage

```
.launch_gsea_shiny(gsea_res, addition_data)
```

Arguments

<code>gsea_res</code>	A valid GseaRes object
<code>addition_data</code>	Optional data.frame with pathway annotations

Value

A Shiny app object

<code>.prepare_rank_vector_fast</code>	<i>Fast Rank Vector Preparation</i>
--	-------------------------------------

Description

Fast Rank Vector Preparation

Usage

```
.prepare_rank_vector_fast(de_table, flip = FALSE)
```

Arguments

<code>de_table</code>	DE result table
<code>flip</code>	Whether to flip sign

Value

Named numeric vector

`.process_addition_data`*Process Addition Data*

Description

Internal function to handle various `addition_data` input types. Supports `data.frame` objects or file paths (`.csv`, `.rds` formats).

Usage

```
.process_addition_data(addition_data)
```

Arguments

`addition_data` A `data.frame` or character vector (file path)

Value

A validated `data.frame` with addition data

`.process_sample_meta` *Internal Sample Metadata Processing*

Description

Automatically infers `target_factor` from `dds_obj@design` for DESeq2 workflows, handles `DFrame` conversion, and creates a unified 'group' column for consistent downstream processing.

Usage

```
.process_sample_meta(sample_meta, expr_bundle)
```

Arguments

`sample_meta` Raw sample metadata (possibly `DFrame`).

`expr_bundle` Expression bundle containing `raw_counts`, `dds_obj`, and other components for row name recovery.

<code>.rebuild_symbol_map</code>	<i>Rebuild Gene Symbol Case Mapping</i>
----------------------------------	---

Description

Reconstruct gene symbol case mapping from differential expression results for display purposes.

Usage

```
.rebuild_symbol_map(gsea_res, contrast_id)
```

Arguments

<code>gsea_res</code>	A GseaRes object.
<code>contrast_id</code>	Character string specifying the contrast ID.

Value

A named character vector where names are uppercase gene symbols and values are the original case. Returns NULL if mapping cannot be constructed.

<code>.serialize_df_tsv</code>	<i>Serialize a Data Frame to TSV</i>
--------------------------------	--------------------------------------

Description

Internal helper used by the Quadrant module's "Export Boxplot Data" modal. Writes a `data.frame` as tab-separated values with a header row and NA rendered as empty strings. Values are quoted only when they contain tab / newline / double-quote characters, so clean sample IDs stay unquoted.

Usage

```
.serialize_df_tsv(df)
```

Arguments

<code>df</code>	A <code>data.frame</code> .
-----------------	-----------------------------

Value

A character scalar containing the TSV (no trailing newline).

```
.standardize_de_columns
```

Standardize Differential Expression Table Column Names

Description

Unify column names from different backends to: gene_symbol, logFC, stat, pvalue, padj

Usage

```
.standardize_de_columns(df, backend)
```

```
.validate_addition_data
```

Validate Addition Data

Description

Internal function to validate addition_data structure. Checks that input is a data.frame with required 'ID' column and removes duplicate entries if present.

Usage

```
.validate_addition_data(data)
```

Arguments

data	A data.frame to validate
------	--------------------------

Value

Validated data.frame with character ID column

```
.validate_deseq2_design
```

Validate DESeq2 Target Factor

Description

Check if target_factor exists in colData and is properly formatted.

Usage

```
.validate_deseq2_design(dds, target_factor)
```

Arguments

dds	DESeqDataSet object
target_factor	String, the target factor to validate

Value

TRUE if validation passes, otherwise stops with an error

`.validate_limma_design`*Validate Limma Design Matrix*

Description

Enforce no-intercept design ($\sim 0 + \text{group}$) to ensure accurate contrast parsing.

Usage

```
.validate_limma_design(fit)
```

Arguments

`fit` MArrayLM object from limma

Value

TRUE if validation passes, otherwise stops with an error

`backends`*Backend Data Extractor*

Description

Extract standardized data structures from limma or DESeq2 objects.

`batch_calc_gsea`*Batch Parallel GSEA Calculation*

Description

Consumes a standard GseaEnv object and performs efficient parallel GSEA calculation.

Usage

```
batch_calc_gsea(
  gsea_env,
  custom_series_name = "Auto_Analysis",
  output_dir = tempdir(),
  workers = 2,
  bidirectional = TRUE,
  minGSSize = 10,
  maxGSSize = 500,
  pvalueCutoff = 1,
  seed = 123,
  force = FALSE,
  use_progress = FALSE,
  chunk_size = NULL
)
```

Arguments

<code>gsea_env</code>	GseaEnv object
<code>custom_series_name</code>	String. Analysis series name
<code>output_dir</code>	String. Output directory for results, default <code>"/GSEA_Output"</code>
<code>workers</code>	Number of parallel cores. Default 4
<code>bidirectional</code>	Logical. Whether to automatically generate reverse contrasts, default TRUE
<code>minGSSize</code>	Minimum gene set size, default 10
<code>maxGSSize</code>	Maximum gene set size, default 500
<code>pvalueCutoff</code>	P-value threshold, default 1
<code>seed</code>	Integer or NULL. Base random seed for per-contrast GSEA permutation, default 123. Each contrast derives its own deterministic seed from the base seed and the contrast id, so results are reproducible across runs and independent of <code>'workers'/'chunk_size'</code> . On <code>clusterProfiler <= 4.20.x</code> (where the <code>'seed'</code> argument of <code>'GSEA()'</code> is silently dropped) reproducibility is guaranteed by seeding the worker R RNG via <code>'withr::local_seed()'</code> ; on <code>clusterProfiler >= 4.21.x</code> the derived seed is also forwarded to <code>'clusterProfiler::GSEA(seed = ...)'</code> . Set to NULL to restore native (non-reproducible) engine sampling.
<code>force</code>	Logical. Whether to force recalculation, default FALSE
<code>use_progress</code>	Logical. Whether to show progress bar, default TRUE
<code>chunk_size</code>	Integer. Number of tasks per worker per chunk, default NULL (auto)

Value

GseaRes object

Examples

```
# Build a GseaEnv from pre-computed inputs shipped with the package
data(preprocessed_limma, package = "GSEAlens")

precomp <- preprocessed_limma
data(gsea_pathwaysets_toy, package = "GSEAlens")
```

```

pathways <- gsea_pathwaysets_toy
gsea_env <- setup_gsea_env(
  fit = precomp$fit,
  pathway_obj = pathways,
  expr_data = precomp$gsea_limma_voom_data
)
# Run batch GSEA (writes to a temporary directory; ~20s on the toy set)
gsea_res <- batch_calc_gsea(gsea_env, workers = 2, output_dir = tempdir())

```

build_edge_list_safely

Build Pathway Similarity Network Edges (Safe Version)

Description

Safely construct edge list for pathway network based on Jaccard similarity of core genes. Includes defensive checks for edge cases. Now includes Overlap Coefficient and Dice Coefficient for comprehensive similarity assessment.

Usage

```
build_edge_list_safely(core_genes_list, min_shared_genes = 2)
```

Arguments

core_genes_list

Named list: pathway_id -> core_gene_vector

min_shared_genes

Minimum number of shared genes between pathways to form an edge

Value

Data frame with columns: from, to, shared, weight (Jaccard index), overlap_coef, dice_coef, shared_genes

Examples

```

# Build edges from two pathways sharing 2 core genes
core_genes <- list(
  PATHWAY_A = c("GENE_A", "GENE_B", "GENE_C"),
  PATHWAY_B = c("GENE_B", "GENE_C", "GENE_D")
)
edges <- build_edge_list_safely(core_genes, min_shared_genes = 2)
print(edges)

```

build_gsea_pathways	<i>Build GSEA Pathway Database with Interactive Selection</i>
---------------------	---

Description

Provides an interactive or automated gene set selection interface based on msigdb, generating standardized pathway objects with intelligent semantic tagging. Supports both human (*Homo sapiens*) and mouse (*Mus musculus*) species.

Usage

```
build_gsea_pathways(species = "HS", auto_select = NULL)
```

Arguments

species	Character string specifying the species. Options: "HS": <i>Homo sapiens</i> (human, default) "MM": <i>Mus musculus</i> (mouse)
auto_select	Optional parameter for automated selection. Can be: - NULL: Launch interactive menu (default) "ALL": Load entire MSigDB database for the specified species - Character vector: Collection names (e.g., c("H", "C5:GO:BP") for human)

Value

A list object containing:

- TERM2GENE: Data frame mapping pathway IDs to gene symbols
- meta_dict: Metadata dictionary with descriptions, URLs, and collection info
- SuperTag: Intelligent batch identifier for the selected collections
- collections_used: Data frame of selected collection metadata
- species: Species identifier ("HS" or "MM")

Examples

```
# Automated selection by collection name -- safe for non-interactive use
pathways <- build_gsea_pathways(species = "HS",
                               auto_select = c("H", "C5:GO:BP"))

# MM uses M-prefixed collection codes (MH, M5:GO:BP, ...)
pathways_mm <- build_gsea_pathways(species = "MM",
                                   auto_select = c("MH", "M5:GO:BP"))

## Not run:
# Interactive menu -- only runs when user is at a terminal
if (interactive()) {
  pathways <- build_gsea_pathways(species = "HS")
}
```

```
# Load ALL collections -- downloads the entire MSigDB (slow, needs network)
pathways <- build_gsea_pathways(species = "HS", auto_select = "ALL")

## End(Not run)
```

build_hubgene_network	<i>Build HubGene Network Data Structure (Enhanced with Leading Edge)</i>
-----------------------	--

Description

Build complete network data with leading edge information for edges. Now properly passes `de_df` to `extract_hub_genes` for stat extraction.

Usage

```
build_hubgene_network(
  gsea_task,
  pathway_ids,
  min_hub_degree = 2,
  de_df = NULL,
  res_df = NULL,
  seed = 123
)
```

Arguments

<code>gsea_task</code>	GseaTask object
<code>pathway_ids</code>	Character vector of selected pathway IDs
<code>min_hub_degree</code>	Minimum degree threshold for hub genes (default: 2)
<code>de_df</code>	Optional DE data frame containing stat column
<code>res_df</code>	Optional pathway result data frame
<code>seed</code>	Random seed for reproducibility (default: 123)

Value

A list with three components:

<code>nodes</code>	A list containing pathway and gene data frames
<code>edges</code>	Edge data frame with source, target, and leading edge info
<code>hub_df</code>	Hub gene data frame from extraction

Examples

```
# Load pre-computed GseaRes shipped with the package and pick a task
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")

# Pick top 3 enriched pathway IDs from the task result
```

```
res_df <- as.data.frame(task$gsea_res)
top_ids <- utils::head(res_df$ID, 3)

# Build the hub-gene network from these pathways
network <- build_hubgene_network(task, top_ids)
names(network)
```

```
calculate_overlap_ratio
```

Calculate ORA Ratio (for DotPlot)

Description

Calculate overlap ratio between pathway genes and differentially expressed genes.

Usage

```
calculate_overlap_ratio(
  pathway_genes,
  de_genes,
  ratio_mode = c("ora", "leading")
)
```

Arguments

pathway_genes	Gene set of pathway (defined by TERM2GENE)
de_genes	Differentially expressed genes (filtered by p-value threshold)
ratio_mode	Character, "ora" (intersection/pathway size) or "leading" (intersection/DE size)

Value

Numeric, ratio value

Examples

```
ratio <- calculate_overlap_ratio(c("GAPDH", "TP53", "MYC"), c("TP53", "MYC", "EGFR"))
```

```
class_validations
```

Core Class Definitions and Validation Functions

Description

Define GseaEnv, GseaRes, GseaTask structures and provide internal validation logic.

color_by_direction	<i>Color Nodes by Direction</i>
--------------------	---------------------------------

Description

Assign colors based on NES (pathways) or log2FC (genes).

Usage

```
color_by_direction(  
  node_data,  
  color_mode = "logFC",  
  left_group = "A",  
  right_group = "B"  
)
```

Arguments

- node_data Node data frame with type and direction columns
- color_mode Color mode: "logFC", "pathway", or "uniform"
- left_group Left group name for legend label
- right_group Right group name for legend label

Value

Node data with added color and color_label columns

Examples

```
# Build a minimal node data frame (pathway + gene nodes)  
nodes <- data.frame(  
  name = c("P_POS", "P_NEG", "G_UP", "G_DOWN"),  
  type = c("pathway", "pathway", "gene", "gene"),  
  NES = c( 2.5, -2.0, NA, NA),  
  log2FC = c( NA, NA, 1.5, -1.2),  
  stringsAsFactors = FALSE  
)  
colored <- color_by_direction(nodes, color_mode = "logFC")  
head(colored$color)
```

create_addition_template	<i>Create Additional Data Template</i>
--------------------------	--

Description

Generate a template CSV file containing all pathway IDs from the GseaRes object. Users can edit this template to add custom annotations such as Brief_Description, Custom_Category, Custom_Tag, and User_Notes.

Usage

```
create_addition_template(
  gsea_res,
  output_path = "addition_template_gsealens.csv",
  include_cols = c("Brief_Description", "Custom_Category", "Custom_Tag", "User_Notes")
)
```

Arguments

<code>gsea_res</code>	A GseaRes object (generated by <code>batch_calc_gsea</code>).
<code>output_path</code>	Character string specifying the output template file path. Default: "addition_template_gsealens.csv".
<code>include_cols</code>	Character vector specifying additional columns to include besides ID. Default includes standard annotation columns.

Value

Invisible TRUE on success.

Examples

```
# Load pre-computed GseaRes shipped with the package and write a template
# CSV to a temporary file (no side effects on the user's filesystem).
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
out_csv <- file.path(tempdir(), "addition_template_demo.csv")
create_addition_template(gsea_res, output_path = out_csv)

# The template contains one row per pathway ID, ready to edit.
template <- utils::read.csv(out_csv)
utils::head(template)
```

<code>create_gsea_env</code>	<i>Define GseaEnv Class</i>
------------------------------	-----------------------------

Description

Standardized GSEA input environment object.

Usage

```
create_gsea_env(
  backend_info,
  contrast_registry,
  de_store,
  expr_bundle,
  geneset,
  raw_obj
)
```

Arguments

backend_info	Backend information object
contrast_registry	Contrast registry
de_store	Differential expression store
expr_bundle	Expression data bundle
geneset	Gene set collection
raw_obj	Raw data object

Value

A GseaEnv object

Examples

```
gsea_env <- create_gsea_env(
  backend_info = list(),
  contrast_registry = data.frame(),
  de_store = data.frame(),
  expr_bundle = list(),
  geneset = character(),
  raw_obj = NULL
)
```

create_gsea_res	<i>Define GseaRes Class</i>
-----------------	-----------------------------

Description

GSEA computation result capsule.

Usage

```
create_gsea_res(
  metadata,
  backend_info,
  contrast_registry,
  de_store,
  expr_bundle,
  geneset_info,
  results
)
```

Arguments

metadata	Metadata information
backend_info	Backend information object
contrast_registry	Contrast registry

de_store	Differential expression store
expr_bundle	Expression data bundle
geneset_info	Gene set information
results	GSEA results

Value

A GseaRes object

Examples

```
gsea_res <- create_gsea_res(  
  metadata = list(),  
  backend_info = list(),  
  contrast_registry = data.frame(),  
  de_store = data.frame(),  
  expr_bundle = list(),  
  geneset_info = list(),  
  results = list()  
)
```

create_gsea_task	<i>Define GseaTask Class</i>
------------------	------------------------------

Description

Single contrast extraction result object.

Usage

```
create_gsea_task(gsea_res, meta)
```

Arguments

gsea_res	GseaRes object
meta	Metadata for the task

Value

A GseaTask object

Examples

```
gsea_task <- create_gsea_task(  
  gsea_res = NULL,  
  meta = data.frame()  
)
```

`creat_addition_data_rdsfile`*Create Additional Data RDS File*

Description

Convert a CSV file to standardized gzip-compressed RDS format with validation. The output file will be saved to the specified directory with a standardized naming convention.

Usage

```
creat_addition_data_rdsfile(  
  csv_path,  
  output_dir = getwd(),  
  output_name = "addition_data_gsealens.rds",  
  overwrite = FALSE  
)
```

Arguments

<code>csv_path</code>	Character string specifying the input CSV file path.
<code>output_dir</code>	Character string specifying the output directory path. Default: current working directory.
<code>output_name</code>	Character string specifying the output filename. Default: "addition_data_gsealens.rds"
<code>overwrite</code>	Logical scalar indicating whether to overwrite an existing file. Default: FALSE.

Value

Invisible TRUE on success. The RDS file path is printed to the console.

Examples

```
# Build a small CSV in a temp directory, then convert to RDS.  
tmp_csv <- file.path(tempdir(), "annotations_demo.csv")  
utils::write.csv(  
  data.frame(  
    ID = c("HALLMARK_HYPOXIA", "HALLMARK_INFLAMMATORY_RESPONSE"),  
    Brief_Description = c("Hypoxia", "Inflammatory response"),  
    stringsAsFactors = FALSE  
  ),  
  tmp_csv, row.names = FALSE  
)  
  
out_rds <- file.path(tempdir(), "annotations_demo.rds")  
creat_addition_data_rdsfile(  
  tmp_csv,  
  output_dir = tempdir(),  
  output_name = "annotations_demo.rds",  
  overwrite = TRUE  
)  
file.exists(out_rds)
```

extract_boxplot_data	<i>Extract Boxplot Data (Long and Wide)</i>
----------------------	---

Description

Internal helper used by the Quadrant module's "Export Boxplot Data" button. Computes the exact per-sample expression data frame that the Shiny boxplot (panel 4) is built on, and returns it in two formats:

long Tidy data frame with columns Sample, Group, Expression. One row per sample. Suitable for GraphPad Prism ("Grouped" data table) and Excel.

wide A one-row data frame. The first column holds the gene symbol (column name Gene); the remaining columns hold the expression value of that gene in each sample (column names are the sample IDs, in the expression matrix's original column order). Suitable for direct copy-paste into Prism ("Column" data table) or any spreadsheet.

Usage

```
extract_boxplot_data(
  GSEAlens_res,
  contrast_id,
  gene_symbol,
  expr_type = "logcpm"
)
```

Arguments

GSEAlens_res	A GseaRes object.
contrast_id	Character. Contrast ID (e.g. "Treat_vs_Control").
gene_symbol	Character. Gene symbol as shown in the boxplot title.
expr_type	Character. Expression matrix type passed to get_expr_matrix(). Default "logcpm".

Value

A named list with elements long, wide, gene, contrast_id, expr_type, left_group, right_group, sample_order.

extract_gsea_task	<i>Extract GSEA Results for Specific Contrast</i>
-------------------	---

Description

Extract results for a specified contrast from a GseaRes capsule. Supports subsetting by gene set collection and automatically recalculates FDR.

Usage

```
extract_gsea_task(gsea_res, contrast_id, target_collection = "ALL")
```

Arguments

gsea_res GseaRes object (generated by batch_calc_gsea)
 contrast_id String. Contrast group ID (e.g., "A_vs_B").
 target_collection String vector. Gene set collection to extract (e.g., "H", "C2:CP:KEGG_LEGACY").
 Default "ALL" means extract all.

Value

GseaTask object.

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")
class(task)
```

extract_hub_genes	<i>Extract Hub Genes from Selected Pathways (Enhanced with Leading Edge)</i>
-------------------	--

Description

Extract hub genes with their leading edge status for each pathway. A gene is considered a hub if it appears in at least min_degree pathways.

Usage

```
extract_hub_genes(gsea_task, pathway_ids, min_degree = 2, de_df = NULL)
```

Arguments

gsea_task GseaTask object
 pathway_ids Character vector of pathway IDs to analyze
 min_degree Minimum number of pathways a gene must appear in (default: 2)
 de_df Optional differential expression data frame with stat column

Value

A data frame with columns:

gene Gene symbol
 degree Number of pathways the gene appears in
 pathways Comma-separated list of pathway IDs
 is_hub Logical indicating if gene is a hub (degree >= min_degree)
 stat Statistic value from de_df or geneList
 pathway_leading_edges List column with per-pathway leading edge status

Examples

```
# Load pre-computed GseaRes shipped with the package and pick a task
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")

# Pick top 3 enriched pathway IDs from the task result
res_df <- as.data.frame(task$gsea_res)
top_ids <- utils::head(res_df$ID, 3)

# Extract hub genes appearing in >= 2 of these pathways
hubs <- extract_hub_genes(task, top_ids, min_degree = 2)
utils::head(hubs)
```

```
generate_boxplot_data_code
```

Generate Boxplot Expression-Data Extraction Code

Description

Produce a self-contained, reproducible R script that extracts the per-sample expression values (Sample / Group / Expression) of a single gene from a GseaRes object. The returned data frame matches exactly what is shown in the Quadrant module's expression boxplot (panel 4), so the user can paste it into other software (e.g. GraphPad Prism, Excel, R) for downstream plotting.

Usage

```
generate_boxplot_data_code(
  GSEAlens_res,
  contrast_id,
  gene_symbol,
  expr_type = "logcpm",
  custom_order = NULL
)
```

Arguments

GSEAlens_res	A GseaRes object.
contrast_id	Character. Contrast ID (e.g. "Treat_vs_Control").
gene_symbol	Character. Gene symbol as shown in the boxplot title.
expr_type	Character. Expression matrix type passed to get_expr_matrix() (e.g. "logcpm", "vst", "count"). Default "logcpm".
custom_order	Character or NULL. Custom group ordering string (e.g. "GroupA -> GroupB") or "default" / NULL to use natural order. When non-default, the generated code will re-order the Group factor accordingly.

Value

A character string of executable R code.

Examples

```
data(precomputed_gseares, package = "GSEAlens")
code <- generate_boxplot_data_code(
  GSEAlens_res = precomputed_gseares,
  contrast_id = "untrt_vs_trt",
  gene_symbol = "GAPDH",
  expr_type = "logcpm"
)
cat(code)
```

```
generate_boxplot_image_code
```

Generate Publication Boxplot Image Code

Description

Return an executable ggplot2 script reproducing the Quadrant module's per-gene expression boxplot (panel 4) as a static, publication-ready figure. Mirrors the interactive plotly: boxplot + jittered points, group colors, optional zero baseline, optional custom group order.

Usage

```
generate_boxplot_image_code(
  box_data_long,
  gene_symbol = "Gene",
  expr_type = "Expression",
  left_group = NA_character_,
  right_group = NA_character_,
  use_zero_baseline = TRUE,
  custom_order = "default",
  base_size = 12,
  margin_top = 15,
  margin_bottom = 15,
  margin_left = 18,
  margin_right = 18,
  fig_width = 5,
  fig_height = 5,
  fig_dpi = 300
)
```

Arguments

box_data_long	data.frame with columns: Sample, Group, Expression. (Same long format returned by <code>extract_boxplot_data()</code> \$long.)
gene_symbol	Character. Gene name for the plot title.
expr_type	Character. Y-axis label (e.g. "logcpm", "log2FC").
left_group	Character. First group name (gets red fill).
right_group	Character. Second group name (gets blue fill).

use_zero_baseline	Logical. Add dashed red line at $y = 0$.
custom_order	Character scalar. Custom group ordering (e.g. "A->B,C") or "default".
base_size	Numeric.
margin_top	Numeric. Top plot margin (mm).
margin_bottom	Numeric. Bottom plot margin (mm).
margin_left	Numeric. Left plot margin (mm).
margin_right	Numeric. Right plot margin (mm).
fig_width	Numeric. Figure width in inches for ggsave.
fig_height	Numeric. Figure height in inches for ggsave.
fig_dpi	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```
# Build a minimal long-format expression data frame (self-contained)
box_long <- data.frame(
  Sample = c("S1", "S2", "S3", "S4"),
  Group = factor(c("untrt", "untrt", "trt", "trt"),
    levels = c("untrt", "trt")),
  Expression = c(8.1, 8.3, 9.5, 9.7)
)
code <- generate_boxplot_image_code(
  box_data_long = box_long,
  gene_symbol = "GAPDH",
  expr_type = "logcpm",
  left_group = "untrt",
  right_group = "trt"
)
cat(code)
```

generate_combined_plot_code

Generate Combined Pathway Plotting Code

Description

Return an executable R script reproducing the Combined Pathway Plotting module (mod_multi_plot / module 12) as a static figure. Mirrors the interactive plot_directional_gsea() call the app performs when the user selects pathways in the main table's "Combined Display" column.

Usage

```
generate_combined_plot_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  target_pathways,
  subPlot = 3,
  curve_colors = NULL,
  left_group = "Left",
  right_group = "Right"
)
```

Arguments

<code>gsea_res_var</code>	Character. Name of the GseaRes variable the generated code should reference (default "gsea_res"). The user replaces the placeholder assignment with their own object.
<code>contrast_id</code>	Character. Contrast ID passed to <code>extract_gsea_task()</code> .
<code>target_pathways</code>	Character vector. Pathway IDs to plot.
<code>subPlot</code>	Integer (1-3). <code>plot_directional_gsea</code> subPlot argument (1 = running score, 2 = rank position, 3 = both). Default 3.
<code>curve_colors</code>	Character vector of color hex codes for pathway curves. If NULL, falls back to a sensible default palette.
<code>left_group</code>	Character. Left group label for axis annotation.
<code>right_group</code>	Character. Right group label.

Value

Character scalar of executable R code.

Examples

```
code <- generate_combined_plot_code(
  contrast_id = "untrt_vs_trt",
  target_pathways = c("HALLMARK_TNFA_SIGNALING_VIA_NFKB")
)
cat(code)
```

generate_de_volcano_code

Generate Publication DE Volcano Code

Description

Return an executable ggplot2 script reproducing the Quadrant module's gene-level Differential Expression Volcano as a static figure. Assumes a GseaRes object is already loaded. Six-color classification (both / user / pathway / up / down / ns) mirrors the interactive view.

Usage

```
generate_de_volcano_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  user_genes = character(0),
  pathway_genes = character(0),
  logfc_thresh = 1,
  pval_thresh = 0.05,
  left_group = "Left",
  right_group = "Right",
  show_annotations = FALSE,
  base_size = 12,
  margin_top = 15,
  margin_bottom = 15,
  margin_left = 18,
  margin_right = 18,
  fig_width = 8,
  fig_height = 6,
  fig_dpi = 300
)
```

Arguments

<code>gsea_res_var</code>	Character. Name of the GseaRes variable.
<code>contrast_id</code>	Character. Contrast ID for DE table extraction.
<code>user_genes</code>	Character vector of user-selected genes (upper-cased).
<code>pathway_genes</code>	Character vector of pathway genes (upper-cased).
<code>logfc_thresh</code>	Numeric. Default 1.
<code>pval_thresh</code>	Numeric. Default 0.05.
<code>left_group</code>	Character.
<code>right_group</code>	Character.
<code>show_annotations</code>	Logical. Add ggrepel labels.
<code>base_size</code>	Numeric.
<code>margin_top</code>	Numeric. Top plot margin (mm).
<code>margin_bottom</code>	Numeric. Bottom plot margin (mm).
<code>margin_left</code>	Numeric. Left plot margin (mm).
<code>margin_right</code>	Numeric. Right plot margin (mm).
<code>fig_width</code>	Numeric. Figure width in inches for ggsave.
<code>fig_height</code>	Numeric. Figure height in inches for ggsave.
<code>fig_dpi</code>	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```
code <- generate_de_volcano_code(
  gsea_res_var = "gsea_res", contrast_id = "untrt_vs_trt",
  left_group = "Ctrl", right_group = "Trt")
cat(code)
```

generate_dotplot_code *Generate Publication DotPlot Code*

Description

Return an executable ggplot2 script reproducing the Pathway Relationship module's dotplot as a static, publication-ready figure. The generated script assumes a GseaRes object is already loaded in the R environment and derives plot data via `extract_gsea_task()` + `get_core_genes_list()`, producing concise reproducible code without embedding raw data.

Usage

```
generate_dotplot_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  pathways,
  color_mode = "padj",
  size_mode = "core_size",
  left_group = "Left",
  right_group = "Right",
  palette = "D",
  point_range = c(3, 8),
  target_collection = "ALL",
  base_size = 12,
  margin_top = 18,
  margin_bottom = 18,
  margin_left = 18,
  margin_right = 18,
  fig_width = 9,
  fig_height = 7,
  fig_dpi = 300
)
```

Arguments

<code>gsea_res_var</code>	Character. Name of the GseaRes variable in the user's environment (default "gsea_res").
<code>contrast_id</code>	Character. Contrast ID to extract.
<code>pathways</code>	Character vector. Pathway IDs to display.
<code>color_mode</code>	Character. One of "padj", "pval", "nes".
<code>size_mode</code>	Character. One of "core_size", "setsize".
<code>left_group</code>	Character. Left contrast label.
<code>right_group</code>	Character. Right contrast label.

palette	Character. Viridis option letter ("A"-"H").
point_range	Numeric length 2. Min/max point size in mm.
target_collection	Character. Gene set collection(s) to extract (e.g. "H", c("C2", "C2:CP")). Default "ALL".
base_size	Numeric. theme_bw base font size.
margin_top	Numeric. Top plot margin (mm).
margin_bottom	Numeric. Bottom plot margin (mm).
margin_left	Numeric. Left plot margin (mm).
margin_right	Numeric. Right plot margin (mm).
fig_width	Numeric. Figure width in inches for ggsave.
fig_height	Numeric. Figure height in inches for ggsave.
fig_dpi	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```
code <- generate_dotplot_code(
  gsea_res_var = "gsea_res", contrast_id = "untrt_vs_trt",
  pathways = c("HALLMARK_TNFA_SIGNALING_VIA_NFKB"))
cat(code)
```

```
generate_gsea_html_report
```

Generate GSEA HTML Report

Description

Automatically avoids column name contamination, intelligently extracts expression matrices from objects, and outputs comfortable web reports with dual p-value display and heatmaps. Uses the ComplexHeatmap engine to draw high-quality heatmaps, perfectly replicating pheatmap aesthetics.

Usage

```
generate_gsea_html_report(
  res_obj,
  output_base_dir = NULL,
  p_adjust_cutoff = 1,
  top_plots = c(15, 15),
  dpi = 200
)
```

Arguments

res_obj	GseaTask object
output_base_dir	Output folder path. If left empty (NULL), it will automatically follow the original project location of the calculation capsule.
p_adjust_cutoff	FDR filtering threshold, default 1
top_plots	Number of pathways to plot detailed sub-pages, format c(positive_count, negative_count)
dpi	Resolution for GSEA enrichment plots, default 200

Value

Path to the generated HTML report (invisible)

Examples

```
data(precomputed_gseares, package = "GSEALens")
task <- extract_gsea_task(precomputed_gseares, contrast_id = "untrt_vs_trt")
report_path <- generate_gsea_html_report(task,
                                         output_base_dir = tempdir())
```

generate_hubgene_code *Generate Publication HubGene Network Code*

Description

Return an executable R script reproducing the HubGene module's visNetwork view as a static ggplot2 figure. Assumes a GseaRes object is already loaded. Node size is controlled by the same three-mode logic (fixed / fdr / setsize) as the interactive view.

Usage

```
generate_hubgene_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  pathways,
  min_hub_degree = 2,
  pw_size_mode = "setsize",
  seed = 42L,
  target_collection = "ALL",
  base_size = 12,
  margin_top = 18,
  margin_bottom = 18,
  margin_left = 18,
  margin_right = 18,
  fig_width = 10,
  fig_height = 8,
  fig_dpi = 300
)
```

Arguments

gsea_res_var	Character. Name of the GseaRes variable.
contrast_id	Character. Contrast ID to extract.
pathways	Character vector. Pathway IDs to include.
min_hub_degree	Integer. Minimum hub degree for gene inclusion.
pw_size_mode	Character. One of "fixed", "fdr", "setsize".
seed	Integer.
target_collection	Character. Gene set collection(s) to extract. Default "ALL".
base_size	Numeric.
margin_top	Numeric. Top plot margin (mm).
margin_bottom	Numeric. Bottom plot margin (mm).
margin_left	Numeric. Left plot margin (mm).
margin_right	Numeric. Right plot margin (mm).
fig_width	Numeric. Figure width in inches for ggsave.
fig_height	Numeric. Figure height in inches for ggsave.
fig_dpi	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```
code <- generate_hubgene_code(
  gsea_res_var = "gsea_res", contrast_id = "untrt_vs_trt",
  pathways = c("HALLMARK_TNFA_SIGNALING_VIA_NFKB"))
cat(code)
```

```
generate_hubgene_hover_text
```

Generate HubGene Hover Text

Description

Generate HTML text for plotly hover tooltips.

Usage

```
generate_hubgene_hover_text(
  node,
  node_type,
  left_group = "A",
  right_group = "B"
)
```

Arguments

node	Node data row (as list with named elements)
node_type	Node type: "pathway" or "gene"
left_group	Left group name
right_group	Right group name

Value

Character string with HTML content for hover tooltip

Examples

```
# Example with a pathway node
pathway_node <- list(
  id = "HALLMARK_INFLAMMATORY_RESPONSE",
  NES = 2.3, FDR = 0.001, pvalue = 0.0005,
  n_core = 15, n_total = 50
)
hover_pathway <- generate_hubgene_hover_text(pathway_node, "pathway")
cat(hover_pathway)

# Example with a gene node
gene_node <- list(id = "GENE_A", log2FC = 1.8)
hover_gene <- generate_hubgene_hover_text(gene_node, "gene")
cat(hover_gene)
```

```
generate_joint_canvas_code
```

Generate Joint Canvas Code

Description

Generate Joint Canvas Code

Usage

```
generate_joint_canvas_code(
  GSEAlens_res,
  contrast_ids,
  target_pathways,
  ncol = 2
)
```

Arguments

GSEAlens_res	GseaRes object
contrast_ids	Character vector. Multiple contrast IDs
target_pathways	Character vector. Pathway IDs
ncol	Integer. Number of columns

Value

A character string containing executable R code

Examples

```
data(precomputed_gseares, package = "GSEALens")
task <- extract_gsea_task(precomputed_gseares, contrast_id = "untrt_vs_trt")
target <- as.data.frame(task$gsea_res)$ID[1]
code <- generate_joint_canvas_code(GSEALens_res = precomputed_gseares,
                                  contrast_ids = "untrt_vs_trt",
                                  target_pathways = target)

cat(code)
```

generate_network_code *Generate Publication Pathway Network Code*

Description

Return an executable R script reproducing the Pathway Relationship module's network view as a static ggplot2 figure. Assumes a GseaRes object is already loaded. Uses igraph layout + geom_segment edges + geom_point nodes (no ggraph dependency).

Usage

```
generate_network_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  pathways,
  min_shared_genes = 2,
  width_mode = "weight",
  seed = 42L,
  target_collection = "ALL",
  base_size = 12,
  margin_top = 18,
  margin_bottom = 18,
  margin_left = 18,
  margin_right = 18,
  fig_width = 10,
  fig_height = 8,
  fig_dpi = 300
)
```

Arguments

gsea_res_var	Character. Name of the GseaRes variable.
contrast_id	Character. Contrast ID to extract.
pathways	Character vector. Pathway IDs to include in the network.
min_shared_genes	Integer. Minimum shared core genes for an edge.

width_mode	Character. "rank" or "weight".
seed	Integer. Layout seed.
target_collection	Character. Gene set collection(s) to extract. Default "ALL".
base_size	Numeric.
margin_top	Numeric. Top plot margin (mm).
margin_bottom	Numeric. Bottom plot margin (mm).
margin_left	Numeric. Left plot margin (mm).
margin_right	Numeric. Right plot margin (mm).
fig_width	Numeric. Figure width in inches for ggsave.
fig_height	Numeric. Figure height in inches for ggsave.
fig_dpi	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```
code <- generate_network_code(
  gsea_res_var = "gsea_res", contrast_id = "untrt_vs_trt",
  pathways = c("HALLMARK_TNFA_SIGNALING_VIA_NFKB", "HALLMARK_HYPOXIA"))
cat(code)
```

```
generate_pathway_plot_code
```

Generate Pathway Plotting Code

Description

Generate Pathway Plotting Code

Usage

```
generate_pathway_plot_code(
  GSEAlens_res,
  contrast_id,
  target_pathways,
  user_genes = character(0),
  pathway_genes = character(0),
  expr_type = "logcpm"
)
```

Arguments

GSEAlens_res	GseaRes object
contrast_id	Character. Contrast ID
target_pathways	Character vector. Pathway IDs
user_genes	Character vector. Genes of interest
pathway_genes	Character vector. Pathway genes (for coloring)
expr_type	Character. Expression type

Value

A character string containing executable R code

Examples

```
# Generate plotting code for the top enriched pathway in a pre-computed task
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")
res_df <- as.data.frame(task$gsea_res)
top_pathway <- res_df$ID[1]

code <- generate_pathway_plot_code(
  GSEAlens_res = gsea_res,
  contrast_id = "untrt_vs_trt",
  target_pathways = top_pathway
)
cat(code)
```

generate_volcano_code *Generate Publication Volcano Code*

Description

Return an executable ggplot2 script reproducing the Quadrant module's pathway volcano plot as a static, publication-ready figure. Assumes a GseaRes object is already loaded in the R environment.

Usage

```
generate_volcano_code(
  gsea_res_var = "gsea_res",
  contrast_id,
  left_group = "Left",
  right_group = "Right",
  n_selected = 0L,
  selected_ids = character(0),
  show_annotations = FALSE,
  target_collection = "ALL",
  base_size = 12,
  margin_top = 15,
```

```

margin_bottom = 15,
margin_left = 18,
margin_right = 18,
fig_width = 8,
fig_height = 6,
fig_dpi = 300
)

```

Arguments

<code>gsea_res_var</code>	Character. Name of the GseaRes variable.
<code>contrast_id</code>	Character. Contrast ID to extract.
<code>left_group</code>	Character.
<code>right_group</code>	Character.
<code>n_selected</code>	Integer. Number of selected pathways (subtitle).
<code>selected_ids</code>	Character vector. Pathway IDs for ggrepel annotation.
<code>show_annotations</code>	Logical. Add ggrepel labels for selected pathways.
<code>target_collection</code>	Character. Gene set collection(s) to extract. Default "ALL".
<code>base_size</code>	Numeric.
<code>margin_top</code>	Numeric. Top plot margin (mm).
<code>margin_bottom</code>	Numeric. Bottom plot margin (mm).
<code>margin_left</code>	Numeric. Left plot margin (mm).
<code>margin_right</code>	Numeric. Right plot margin (mm).
<code>fig_width</code>	Numeric. Figure width in inches for ggsave.
<code>fig_height</code>	Numeric. Figure height in inches for ggsave.
<code>fig_dpi</code>	Numeric. DPI for PNG raster output (default 300).

Value

Character scalar of executable R code.

Examples

```

code <- generate_volcano_code(
  gsea_res_var = "gsea_res", contrast_id = "untrt_vs_trt",
  left_group = "Ctrl", right_group = "Trt")
cat(code)

```

get_contrast_registry *Get Contrast Registry*

Description

Retrieve the registered contrasts used for differential expression analysis, including contrast definitions and metadata.

Usage

```
get_contrast_registry(obj)
```

Arguments

obj A GseaEnv or GseaRes object.

Value

The contrast registry object stored in the input object.

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
registry <- get_contrast_registry(gsea_res)
print(registry)
```

get_core_genes_for_pathway
Extract Pathway Core Genes (Leading Edge)

Description

Parse the core_enrichment field from GSEA results to extract the gene list. This is the fundamental data interface for all visualization modules in 03/04.

Usage

```
get_core_genes_for_pathway(gsea_res_obj, pathway_id)
```

Arguments

gsea_res_obj GseaRes object or single GSEA result (list(status=, data=, ...))
pathway_id Character, pathway ID

Value

Character vector, core genes (uppercase normalized)

Examples

```
# Load a pre-computed GseaRes and extract a task
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")
res_df <- as.data.frame(task$gsea_res)
pw_id <- res_df$ID[1]
core_genes <- get_core_genes_for_pathway(task, pw_id)
head(core_genes)
```

get_core_genes_list	<i>Batch Extract Core Genes for Multiple Pathways</i>
---------------------	---

Description

Batch data preparation for 03 module Network/UpSet/Chord visualizations.

Usage

```
get_core_genes_list(gsea_task_obj, pathway_ids)
```

Arguments

gsea_task_obj	GseaTask object (single contrast)
pathway_ids	Character vector, pathway ID list

Value

Named list, names = pathway_id, values = core_genes vector

Examples

```
# Load a pre-computed GseaRes and extract a task
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")
res_df <- as.data.frame(task$gsea_res)
pw_ids <- head(res_df$ID, 3)
genes_list <- get_core_genes_list(task, pw_ids)
length(genes_list)
```

get_de_table	<i>Get Differential Expression Table</i>
--------------	--

Description

Extract differential expression results for a specific contrast from the stored results. Automatically handles reverse contrasts (e.g., B_vs_A) by retrieving the forward contrast and flipping the sign of logFC.

Usage

```
get_de_table(obj, contrast_id)
```

Arguments

obj	A GseaRes object containing differential expression results.
contrast_id	Character string specifying the contrast ID (e.g., "A_vs_B"). If the reverse contrast exists in storage, it will be retrieved and logFC/stat signs will be inverted.

Value

A data frame containing at minimum: gene_symbol, logFC, and p-value columns. Additional columns (stat, pvalue, padj, t) may be present depending on the analysis backend.

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
de_result <- get_de_table(gsea_res, contrast_id = "untrt_vs_trt")
head(de_result)
```

get_expr_matrix	<i>Get Expression Matrix</i>
-----------------	------------------------------

Description

Extract expression matrix from GseaEnv or GseaRes object, supporting multiple normalization methods including raw counts, CPM, logCPM, VST, FPKM, TPM, and log-normalized counts.

Usage

```
get_expr_matrix(obj, type = "default", ...)
```

Arguments

obj	A GseaEnv or GseaRes object.
type	Character string specifying the type of expression values to return. Options include: "default" (default display matrix), "raw" (raw counts), "cpm" (counts per million), "logcpm" (log2-transformed CPM), "vst" (variance stabilizing transformation, DESeq2 only), "fpkm" (fragments per kilobase per million), "tpm" (transcripts per million), and "lognorm" (log2 of normalized counts).
...	Additional arguments passed to backend-specific methods.

Value

A numeric matrix with genes as rows and samples as columns.

Examples

```
# Load pre-computed inputs shipped with the package
data(preprocessed_limma, package = "GSEAlens")

precomp <- preprocessed_limma
data(gsea_pathwaysets_toy, package = "GSEAlens")

pathways <- gsea_pathwaysets_toy
gsea_env <- setup_gsea_env(
  fit = precomp$fit,
  pathway_obj = pathways,
  expr_data = precomp$gsea_limma_voom_data
)
expr <- get_expr_matrix(gsea_env)
dim(expr)
```

get_geneset_info

Get Gene Set Information

Description

Retrieve gene set collection metadata including pathway definitions and annotation dictionary.

Usage

```
get_geneset_info(obj)
```

Arguments

obj	A GseaEnv or GseaRes object.
-----	------------------------------

Value

An object containing geneset_info with pathway definitions and metadata dictionary.

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
gs_info <- get_geneset_info(gsea_res)
names(gs_info)
```

get_hubgene_legend	<i>Get Color Legend Configuration</i>
--------------------	---------------------------------------

Description

Generate color legend data for HubGene network plot.

Usage

```
get_hubgene_legend(left_group = "A", right_group = "B")
```

Arguments

left_group	Left group name (e.g., treatment or case)
right_group	Right group name (e.g., control or reference)

Value

A named list containing:

pathway_up	Color for up-regulated pathways
pathway_down	Color for down-regulated pathways
gene_up	Color for up-regulated genes
gene_down	Color for down-regulated genes
labels	Named list of legend labels

Examples

```
legend <- get_hubgene_legend("Control", "Treatment")
```

get_sample_meta	<i>Get Sample Metadata</i>
-----------------	----------------------------

Description

Extract and process sample metadata, handling DFrame conversion from DESeq2 and unifying group column naming conventions across backends.

Usage

```
get_sample_meta(obj)
```

Arguments

obj	GseaEnv or GseaRes object
-----	---------------------------

Value

A data frame with sample metadata. Includes a standardized 'group' column derived from the target factor (inferred from dds_obj@design for DESeq2 workflows). Sample row names are preserved for downstream matching.

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
meta <- get_sample_meta(gsea_res)
head(meta)
```

get_term_genes	<i>Get Full Pathway Gene Set (TERM2GENE)</i>
----------------	--

Description

Extract complete gene list for a pathway from geneset_info.

Usage

```
get_term_genes(gsea_res, pathway_id)
```

Arguments

gsea_res	GseaRes object
pathway_id	Pathway ID

Value

Character vector

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
pw_id <- gsea_res$geneset_info$term2gene$gs_name[1]
genes <- get_term_genes(gsea_res, pw_id)
head(genes)
```

`gsea_pathwaysets_toy` *Toy MSigDB pathway collection (Hallmark + KEGG_LEGACY)*

Description

A lightweight pathway object for examples and vignettes, containing 236 pathways (Hallmark + KEGG_LEGACY) pre-assembled by [GSEALens::build_gsea_pathways()]. See ‘inst/scripts/make_gsea_pathwaysets_to’ for regeneration.

Format

A list with elements ‘TERM2GENE’, ‘meta_dict’, ‘SuperTag’, ‘collections_used’, ‘species’.

Source

Assembled from msigdb (Hallmark + KEGG_LEGACY collections).

Examples

```
data(gsea_pathwaysets_toy, package = "GSEALens")
names(gsea_pathwaysets_toy)
```

`gsea_pathwaysets_toy_hallmark`
Toy MSigDB pathway collection (Hallmark only)

Description

A minimal Hallmark-only pathway object for fast examples and tests. See ‘inst/scripts/make_gsea_pathwaysets_toy.R’ for regeneration.

Format

A list with elements ‘TERM2GENE’, ‘meta_dict’, ‘SuperTag’, ‘collections_used’, ‘species’.

Source

Assembled from msigdb (Hallmark collection).

Examples

```
data(gsea_pathwaysets_toy_hallmark, package = "GSEALens")
names(gsea_pathwaysets_toy_hallmark)
```

import_gsea_capsule *Load and Smart-Relocate GSEA Computation Capsule*

Description

Safely load a computation capsule. If the file is moved from its original project path (e.g., copied to desktop), the engine will automatically restore the standard folder structure in the current working directory and relocate the capsule. Fully compatible with GseaEnv and GseaRes objects.

Usage

```
import_gsea_capsule(file_path, auto_relocate = TRUE, inspect = TRUE)
```

Arguments

file_path	Character. Absolute or relative path to the capsule rds file.
auto_relocate	Logical. Whether to automatically fix the folder structure and relocate based on the built-in lineage in the current directory? Default TRUE.
inspect	Logical. Whether to automatically print capsule overview after loading? Default TRUE.

Value

Returns the parsed GseaRes or GseaEnv capsule object.

Examples

```
# Load the pre-computed GseaRes capsule shipped with the package, write it
# to a temp file, then import it (auto_relocate = FALSE to avoid side
# effects).
data(precomputed_gseares, package = "GSEAlens")
tmp_capsule <- file.path(tempdir(), "import_demo.rds")
saveRDS(precomputed_gseares, tmp_capsule)

gsea_res <- import_gsea_capsule(
  tmp_capsule,
  auto_relocate = FALSE,
  inspect = FALSE
)
class(gsea_res)
```

inspect_gsea_env *View GSEA Environment Object Overview*

Description

Print detailed information about the GseaEnv object to the console, including backend type, contrast list, and gene set status.

Usage

```
inspect_gsea_env(env_obj)
```

Arguments

env_obj GseaEnv object

Value

Invisibly returns the input object (for chaining).

Examples

```
# Build a GseaEnv from pre-computed inputs shipped with the package
data(preprocessed_limma, package = "GSEAlens")

precomp <- preprocessed_limma
data(gsea_pathwaysets_toy, package = "GSEAlens")

pathways <- gsea_pathwaysets_toy
env <- setup_gsea_env(
  fit      = precomp$fit,
  pathway_obj = pathways,
  expr_data = precomp$gsea_limma_voom_data
)
inspect_gsea_env(env)
```

inspect_gsea_res

View GSEA Result Overview

Description

Print detailed summary of GseaRes object to console.

Usage

```
inspect_gsea_res(gsea_res)
```

Arguments

gsea_res GseaRes object

Value

Invisibly returns the input object (for chaining).

Examples

```
# Load a pre-computed GseaRes shipped with the package
data(precomputed_gseares, package = "GSEAlens")

gsea_res <- precomputed_gseares
inspect_gsea_res(gsea_res)
```

launch_gsea_app

GSEALens Interactive Analysis Workstation

Description

Modular Shiny application based on GseaRes object, supporting Limma and DESeq2 dual back-ends. Provides interactive visualization for GSEA pathway enrichment analysis including volcano plots, heatmaps, expression boxplots, and pathway relationship networks.

Usage

```
launch_gsea_app(gsea_res, addition_data = NULL)
```

Arguments

gsea_res	GseaRes object (generated by batch_calc_gsea())
addition_data	Optional. A data frame or file path containing pathway annotations to merge into the main table. Can also be: • A data.frame with 'ID' column as primary key • A character vector: file path to .csv, .csv.gz, .rds, or .rds.gz • NULL: automatically searches for "addition_data_gsealens.rds" or "addition_data_gsealens.csv" in the working directory

Additional columns will be merged to the main pathway table.

Value

Shiny application object

Examples

```
if(interactive()){
# Basic usage
gsea_res <- batch_calc_gsea(gsea_env)
launch_gsea_app(gsea_res)

# With addition data from file path
launch_gsea_app(gsea_res, "pathway_annotations.csv")

# With pre-loaded data frame
add_data <- read.csv("annotations.csv")
launch_gsea_app(gsea_res, add_data)
}
```

mod_ai_abs_page_server	<i>AI Interpretation Page Server</i>
------------------------	--------------------------------------

Description

AI Interpretation Page Server. Called for side effects within a Shiny module.

Usage

```
mod_ai_abs_page_server(id, gsea_res, data_prep_list, table_controller)
```

Arguments

id	Module ID
gsea_res	GseaRes object
data_prep_list	Data preprocessing list
table_controller	Table controller

Value

No return value; called for side effects within a Shiny module. Extract Leading Edge genes and format as comma-separated Determine confidence level

mod_ai_abs_page_ui	<i>AI Interpretation Page UI</i>
--------------------	----------------------------------

Description

AI Interpretation Page UI

Usage

```
mod_ai_abs_page_ui(id)
```

Arguments

id	Module ID
----	-----------

mod_data_prep_server	<i>Data Preprocessing Module Server</i>
----------------------	---

Description

Server logic for the data preprocessing module. Handles:

- Contrast and gene set selection
- Data processing and sorting
- Gene marker management (add, confirm, clear)
- Boxplot ordering configuration
- Joint canvas parameter passing
- Expression data type selection based on backend

Usage

```
mod_data_prep_server(id, gsea_res)
```

Arguments

id	Module namespace ID
gsea_res	GSEA results object containing contrast registry and backend info

Value

A list of reactive functions and values:

- data: Reactive data object with processed GSEA results
- highlight_genes: Reactive character vector of confirmed gene markers
- boxplot_order: Reactive value for boxplot group ordering
- joint_contrasts: Reactive vector of contrasts for joint canvas
- joint_ncol: Reactive number of columns for joint canvas
- joint_generate: Event reactive for canvas regeneration trigger
- update_pending_genes: Function to update pending genes from modal
- get_pending_genes: Function to retrieve current pending genes

mod_data_prep_ui	<i>Data Preprocessing Module UI</i>
------------------	-------------------------------------

Description

User interface for the data preprocessing module, providing controls for:

- Contrast selection and gene set subgroup filtering
- Joint GSEA canvas for multi-pathway visualization
- Differential expression gene marker selection
- Group display order customization
- Expression metrics configuration

Usage

```
mod_data_prep_ui(id)
```

Arguments

id	Module namespace ID
----	---------------------

mod_hubgene_vis_server	<i>HubGene Network (visNetwork) Module Server</i>
------------------------	---

Description

Shiny module server for HubGene network visualization using visNetwork. Handles pathway selection, network building, rendering with physics simulation, and interactive node/edge display.

Usage

```
mod_hubgene_vis_server(id, data_prep_list, table_controller, gsea_res)
```

Arguments

id	Character string used to namespace the module.
data_prep_list	A reactive expression returning a list containing GSEA results, data frame, contrast groups, and metadata.
table_controller	A list containing reactive expressions for table interactions, including selected pathways.
gsea_res	A GSEA result object used for symbol mapping.

Value

A list containing reactive expressions:

final_pathways	Reactive character vector of selected pathway IDs
----------------	---

mod_hubgene_vis_ui	<i>HubGene Network Module UI</i>
--------------------	----------------------------------

Description

Shiny module UI for visualizing HubGene networks using visNetwork. Provides interactive controls for pathway selection modes, physics simulation parameters, node sizing, and network statistics display.

Usage

```
mod_hubgene_vis_ui(id)
```

Arguments

id	Character string used to namespace the module.
----	--

Value

A Shiny tagList containing the module's UI elements.

mod_joint_canvas_server	<i>Joint GSEA Fill Canvas Module Server (Sidebar Control Version)</i>
-------------------------	---

Description

Server-side logic for the joint GSEA canvas display module. This module:

1. Receives sidebar control parameters (arrangement mode)
2. Applies no maximum row limit
3. Maintains aesthetic consistency with multi_plot
4. Generates combined visualization with multiple contrasts and pathways
5. Provides R code export functionality for reproducibility

Usage

```
mod_joint_canvas_server(id, gsea_res, data_prep_list, table_result)
```

Arguments

id	Character string identifying the module namespace.
gsea_res	Reactive expression containing GSEA results.
data_prep_list	Reactive list containing data preparation parameters including contrasts, ncol, and custom colors.
table_result	Reactive list containing pathway selection results.

mod_joint_canvas_ui	<i>Joint GSEA Fill Canvas Module UI (Display-Only Version)</i>
---------------------	--

Description

All controls have been moved to the sidebar; only canvas results are displayed here. This module provides a display interface for joint GSEA visualization results with export functionality for reproducible R code generation.

Usage

```
mod_joint_canvas_ui(id)
```

mod_master_table_server	<i>Master Workspace Table Server</i>
-------------------------	--------------------------------------

Description

Provides data display, column display control with decoupled checkbox state. Supports merging addition_data columns into the main table. Includes external selection update capabilities for bidirectional sync.

Usage

```
mod_master_table_server(id, data_prep, addition_data = NULL)
```

Arguments

id	Module ID
data_prep	Reactive data from the data preprocessing module
addition_data	Optional. A data frame with pathway annotations to merge.

Value

List containing selected_pathways, show_modal, and selection control functions

mod_master_table_ui	<i>Master Workspace Table UI</i>
---------------------	----------------------------------

Description

Interactive data table module for displaying GSEA pathway results with checkbox selection and modal integration.

Usage

```
mod_master_table_ui(id)
```

Arguments

id	Module ID
----	-----------

Value

Shiny UI tagList

mod_multi_plot_server	<i>Combined Pathway Plotting Server</i>
-----------------------	---

Description

Combined Pathway Plotting Server

Usage

```
mod_multi_plot_server(id, data_prep, table_controller, gsea_res = NULL)
```

mod_multi_plot_ui	<i>Combined Pathway Plotting UI</i>
-------------------	-------------------------------------

Description

Combined Pathway Plotting UI

Usage

```
mod_multi_plot_ui(id)
```

mod_pathway_modal_server

Pathway Detail Modal Server

Description

Modal with checkbox selection using standard DT approach.

Usage

```
mod_pathway_modal_server(  
  id,  
  data_prep,  
  trigger_event,  
  gsea_res,  
  pending_genes_reactive,  
  update_pending_genes  
)
```

Arguments

- | | |
|------------------------|---------------------------------------|
| id | Module ID |
| data_prep | Reactive data from data prep module |
| trigger_event | Event that triggers modal opening |
| gsea_res | GseaRes object |
| pending_genes_reactive | Function to get current pending genes |
| update_pending_genes | Function to update pending genes list |

mod_pathway_modal_ui

Pathway Detail Modal UI

Description

Pathway Detail Modal UI

Usage

```
mod_pathway_modal_ui(id)
```

mod_pathway_relation_server
<i>Pathway Relationship Exploration Module Server</i>

Description

Dual-mode pathway network visualization with defensive rendering

Usage

```
mod_pathway_relation_server(id, data_prep_list, gsea_res, table_result = NULL)
```

mod_pathway_relation_ui
<i>Pathway Relationship Exploration Module UI</i>

Description

Network visualization module with dual-mode pathway selection

Usage

```
mod_pathway_relation_ui(id)
```

mod_quadrant_server	<i>Quadrant Linkage Server</i>
---------------------	--------------------------------

Description

Server logic with unified gene pool and bidirectional table-volcano sync. Now properly responds to table selection changes.

Usage

```
mod_quadrant_server(id, data_prep_list, gsea_res, table_controller)
```

Arguments

- | | |
|------------------|---|
| id | Module ID |
| data_prep_list | List from data prep module |
| gsea_res | GseaRes object |
| table_controller | Table controller with update_selection method |

mod_quadrant_ui	<i>Quadrant Linkage Module</i>
-----------------	--------------------------------

Description

Four-quadrant interactive visualization module for pathway and gene analysis. Provides synchronized pathway selection, gene ranking, differential expression volcano, and expression boxplot visualizations with bidirectional table-volcano synchronization.

UI components for the four-quadrant interactive visualization

Usage

```
mod_quadrant_ui(id)
```

Arguments

id	Module ID
----	-----------

Value

Shiny UI tagList

plot_directional_gsea	<i>Plot Directional GSEA</i>
-----------------------	------------------------------

Description

Professional-grade GSEA plotting engine with seamless support for single or multiple pathway visualization. Automatically generates elegant legends and native red-blue gene distribution track.

Usage

```
plot_directional_gsea(  
  directional_gsea_obj,  
  target_pathways,  
  main_title = NULL,  
  subPlot = 3,  
  curveCol = NULL,  
  add_pval = FALSE,  
  show_contrast_in_axis = FALSE,  
  ...  
)
```

Arguments

directional_gsea_obj	A wrapped GseaTask object (returned by extract_gsea_task)
target_pathways	Vector of pathway IDs to plot (supports single or multiple)
main_title	Main title name
subPlot	Controls the number of subplots generated (1: enrichment plot only; 2: enrichment + heatmap track; 3: full rank track)
curveCol	Custom curve color vector
add_pval	Whether to add statistical annotations on the plot. Default FALSE.
show_contrast_in_axis	Logical. Whether to display contrast group information on the x-axis of the Rank plot. Default FALSE; recommended to set TRUE only in combined canvas mode.
...	Additional parameters passed to internal engine

Value

ggplot2 composite object

Examples

```
# Load pre-computed GseaRes and pick a task + an enriched pathway ID
data(precomputed_gseares, package = "GSEALens")

gsea_res <- precomputed_gseares
task <- extract_gsea_task(gsea_res, contrast_id = "untrt_vs_trt")
res_df <- as.data.frame(task$gsea_res)
top_pathway <- res_df$ID[1]

# Render the directional GSEA plot (enrichment + rank + heatmap tracks)
p <- plot_directional_gsea(task, target_pathways = top_pathway)
class(p)
```

plot_gsea_memory	<i>Plot GSEA Benchmark Memory Curve</i>
------------------	---

Description

Generates a standard memory usage curve plot for publication.

Usage

```
plot_gsea_memory(aligned_data, highlight_phases = TRUE)
```

Arguments

aligned_data	Aligned data frame returned by align_benchmark_data
highlight_phases	Whether to color by phase, default TRUE

Value

A ggplot object

Examples

```
# Build a small synthetic aligned-data frame (mimics the output of
# align_benchmark_data() without needing an external CSV or GseaRes object).
syn <- data.frame(
  timestamp_ms = seq(0, 20000, by = 1000),
  rss_mb = c(120, 150, 220, 280, 310, 330, 340, 335, 320, 305,
             290, 270, 255, 245, 240, 235, 230, 225, 220, 215, 210),
  relative_sec = seq(0, 20, by = 1),
  phase = c(rep("startup", 4), rep("core_compute", 12), rep("cleanup", 5))
)
attr(syn, "gsea_info") <- list(duration_sec = 20, workers = 2)
class(syn) <- c("GseaBenchmarkAligned", "data.frame")
plot_gsea_memory(syn)
plot_gsea_memory(syn, highlight_phases = FALSE)
```

precomputed_gseares	<i>Pre-computed GSEA results from the airway dataset</i>
---------------------	--

Description

A pre-computed ‘GseaRes’ object built from the airway dataset (DESeq2 + clusterProfiler::GSEA). Used in examples and vignettes so that users can try visualization and code-generation functions without having to re-run the full GSEA pipeline.

Format

A ‘GseaRes’ object (list-like).

Source

Built from the ‘airway’ Bioconductor dataset via DESeq2 + clusterProfiler::GSEA.

Examples

```
data(precomputed_gseares, package = "GSEALens")
class(precomputed_gseares)
```

```
prepare_hubgene_nodes Prepare HubGene Nodes for Plotting
```

Description

Prepare node data with layout positions for plotly visualization.

Usage

```
prepare_hubgene_nodes(network_data, layout = "fr", seed = 42)
```

Arguments

network_data	Output from build_hubgene_network
layout	Layout algorithm: "fr" (Fruchterman-Reingold), "kk" (Kamada-Kawai), or "circle"
seed	Random seed for reproducibility (default: 42)

Value

A named list with three components:

pathway	Pathway node data frame with coordinates
gene	Gene node data frame with coordinates
all	Combined node data frame

Examples

```
# Build a minimal network_data object (2 pathways + 2 shared genes)
network_data <- list(
  nodes = list(
    pathway = data.frame(
      id = c("PATHWAY_A", "PATHWAY_B"), type = "pathway",
      NES = c(2.0, -1.5), direction = c("up", "down"),
      stringsAsFactors = FALSE
    ),
    gene = data.frame(
      id = c("GENE_X", "GENE_Y"), type = "gene",
      stringsAsFactors = FALSE
    )
  ),
  edges = data.frame(
    from = c("PATHWAY_A", "PATHWAY_A", "PATHWAY_B", "PATHWAY_B"),
    to = c("GENE_X", "GENE_Y", "GENE_X", "GENE_Y"),
    stringsAsFactors = FALSE
  ),
  hub_df = data.frame(gene = c("GENE_X", "GENE_Y"), degree = c(2, 2),
    stringsAsFactors = FALSE)
)
nodes <- prepare_hubgene_nodes(network_data, layout = "fr", seed = 42)
if (!is.null(nodes)) head(nodes$all[, c("id", "type")])
```

preprocessed_dds	<i>Pre-computed DESeq2 DESeqDataSet from the airway dataset</i>
------------------	---

Description

A slimmed ‘DESeqDataSet’ object fitted with ‘DESeq()’, suitable as input to [GSEAlens::setup_gsea_env()].

Format

A ‘DESeqDataSet’ object.

Source

Built from the ‘airway’ Bioconductor dataset via DESeq2.

Examples

```
data(preprocessed_dds, package = "GSEAlens")
class(preprocessed_dds)
```

preprocessed_dds_se	<i>Pre-computed DESeqDataSet (SummarizedExperiment form)</i>
---------------------	--

Description

A ‘DESeqDataSet’ based on a ‘SummarizedExperiment’ input, suitable for demonstrating the SE-based entry point of [GSEAlens::setup_gsea_env()].

Format

A ‘DESeqDataSet’ object.

Source

Built from the ‘airway’ Bioconductor dataset via DESeq2.

Examples

```
data(preprocessed_dds_se, package = "GSEAlens")
class(preprocessed_dds_se)
```

preprocessed_limma	<i>Pre-computed limma-voom fit and DGEList from the airway dataset</i>
--------------------	--

Description

A list with elements 'fit' ('MArrayLM') and 'gsea_limma_voom_data' (filtered 'DGEList'), suitable as input to [GSEALens::setup_gsea_env()] for the limma-voom workflow.

Format

A list with elements 'fit' and 'gsea_limma_voom_data'.

Source

Built from the 'airway' Bioconductor dataset via limma-voom.

Examples

```
data(preprocessed_limma, package = "GSEALens")
names(preprocessed_limma)
```

read_addition_data	<i>Read Additional Data</i>
--------------------	-----------------------------

Description

Read and validate additional data from CSV or RDS file. Supports gzip-compressed formats (.gz). The data frame must contain an 'ID' column as the primary key for joining with pathway results.

Usage

```
read_addition_data(file_path)
```

Arguments

file_path	Character string specifying the file path. Supported formats: • .csv / .csv.gz - Comma-separated values (optionally gzip compressed) • .rds / .rds.gz - R serialized data (optionally gzip compressed)
-----------	--

Value

A data frame with validated ID column. Rows are deduplicated by ID, keeping the first occurrence of each unique ID.

Examples

```
# Build a small CSV in a temp directory and read it back.
tmp_csv <- file.path(tempdir(), "pathway_annotations_demo.csv")
utils::write.csv(
  data.frame(
    ID = c("HALLMARK_HYPOXIA", "HALLMARK_INFLAMMATORY_RESPONSE"),
    Brief_Description = c("Hypoxia", "Inflammatory response"),
    stringsAsFactors = FALSE
  ),
  tmp_csv, row.names = FALSE
)

add_data <- read_addition_data(tmp_csv)
add_data
```

setup_gsea_env

Initialize GSEA Environment Object

Description

Unified entry point supporting limma-voom (no intercept design) and DESeq2 objects. Automatically extracts contrasts, differential analysis results, and expression matrices to generate a standardized GseaEnv object.

Usage

```
setup_gsea_env(
  fit,
  pathway_obj,
  expr_data = NULL,
  target_factor = NULL,
  verbose = TRUE
)
```

Arguments

fit	Differential analysis object. Supports MArrayLM (limma) or DESeqDataSet (DESeq2).
pathway_obj	Gene set object. Typically generated by build_gsea_pathways().
expr_data	Optional. Expression matrix or DGEList. If NULL, extraction from fit object will be attempted (partial support only).
target_factor	Only for DESeq2. Specifies the target factor of interest. If NULL, automatically inferred as the last term in the design formula.
verbose	Logical. If TRUE (default), print progress messages. Set to FALSE for silent batch-mode use.

Value

Returns a standardized object of class GseaEnv.

Examples

```
# Limma-voom workflow using pre-computed inputs shipped with the package
data(preprocessed_limma, package = "GSEAlens")

precomp <- preprocessed_limma
data(gsea_pathwaysets_toy, package = "GSEAlens")

pathways <- gsea_pathwaysets_toy
env <- setup_gsea_env(
  fit      = precomp$fit,
  pathway_obj = pathways,
  expr_data = precomp$gsea_limma_voom_data
)
class(env)
```

utils-accessors	<i>Data Accessor Utilities</i>
-----------------	--------------------------------

Description

Provides unified data extraction interface, isolating underlying object structure differences. Fixes DESeq2 sample name matching issues.

utils-imports	<i>GSEAlens Global Imports</i>
---------------	--------------------------------

Description

Centralized management of all external dependency imports to avoid redundant definitions

validate_param	<i>Safe Parameter Validation with Fallback (Global Strategy)</i>
----------------	--

Description

Unified handling of invalid parameters (e.g., ncol <= 0) to ensure 03/04 module stability.

Usage

```
validate_param(
  value,
  default,
  min_val = 1,
  max_val = NULL,
  param_name = "parameter"
)
```

Arguments

value	Input value
default	Default value
min_val	Minimum value (inclusive)
max_val	Maximum value (inclusive, optional)
param_name	Parameter name (for warning messages)

Value

Validated value

Examples

```
workers <- validate_param(4, default = 2, min_val = 1, max_val = 8, param_name = "workers")
```

visualization

GSEA Visualization and Report Generation

Description

Provides static plotting functions and interactive HTML report generation capabilities. This is a documentation landing page for the visualization family of functions in this file.

Value

NULL (this is a documentation landing page; individual functions return their own objects, documented separately).

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