

# Package: OmicNetR (via r-universe)

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**Type** Package

**Title** Network-Based Integration of Multi-Omics Data Using Sparse CCA

**Version** 0.1.1

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**Description** Provides an end-to-end workflow for integrative analysis of two omics layers using sparse canonical correlation analysis (sCCA), including sample alignment, feature selection, network edge construction, and visualization of gene-metabolite relationships. The underlying methods are based on penalized matrix decomposition and sparse CCA (Witten, Tibshirani and Hastie (2009) <doi:10.1093/biostatistics/kxp008>), with design principles inspired by multivariate integrative frameworks such as mixOmics (Rohart et al. (2017) <doi:10.1371/journal.pcbi.1005752>).

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**Encoding** UTF-8

**LazyData** true

**Imports** stats, graphics, grDevices, ggplot2, igraph

**Suggests** knitr, rmarkdown, testthat (>= 3.0.0), mixOmics

**VignetteBuilder** knitr

**RoxygenNote** 7.3.2

**NeedsCompilation** no

**Repository** https://cran.r-universe.dev

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align_omics	<i>Align Multi-Omic Datasets</i>
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### Description

Ensures that X and Y matrices have matching samples in the exact same order.

### Usage

```
align_omics(X, Y)
```

### Arguments

X	Matrix or data frame (Samples x Features).
Y	Matrix or data frame (Samples x Features).

### Value

A list containing the aligned X and Y matrices.

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generate_dummy_omics	<i>Generate Dummy Multi-Omics Data</i>
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### Description

Generates synthetic, linked RNA-seq and Metabolomics datasets.

### Usage

```
generate_dummy_omics(
  n_samples = 50,
  n_genes = 1000,
  n_metabolites = 200,
  n_linked = 10
)
```

**Arguments**

n_samples	Number of samples (rows).
n_genes	Number of genes (columns in X).
n_metabolites	Number of metabolites (columns in Y).
n_linked	Number of features linked by a hidden variable.

**Value**

A list containing X (RNA-seq matrix), Y (Metabolomics matrix), and metadata.

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omic_scca	<i>Perform Sparse Canonical Correlation Analysis (sCCA)</i>
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**Description**

Fits a sparse PLS model in canonical mode to identify shared variation.

**Usage**

```
omic_scca(X, Y, n_components = 2, penalty_X = 0.9, penalty_Y = 0.9)
```

**Arguments**

X	Normalized RNA-seq matrix (Samples x Features).
Y	Normalized Metabolomics matrix (Samples x Features).
n_components	Number of components.
penalty_X	Sparsity for X (0 to 1, where 1 is most sparse).
penalty_Y	Sparsity for Y (0 to 1, where 1 is most sparse).

**Value**

An object of class "OmicNetR\_sCCA" (a named list) with:

- canonical\_correlations: numeric vector of per-component correlations/variance explained from the fitted model.
- loadings: list with matrices X and Y (feature weights) for each component.
- variates: list with matrices X and Y (sample scores) for each component.
- penalties: list with penalty\_X and penalty\_Y used to set sparsity.

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`omics_example`*Example Multi-Omics Dataset*

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**Description**

A simulated multi-omics dataset included in the OmicNetR package. This dataset is intended for demonstrating data alignment, sparse CCA analysis, and network visualization functions.

**Usage**`omics_example`**Format**

A list with the following components:

**X** A numeric matrix of gene expression values (samples in rows, genes in columns).

**Y** A numeric matrix of metabolite abundances (samples in rows, metabolites in columns).

**Details**

The dataset is small by design and should not be used for biological inference. It is provided solely for examples, vignettes, and unit testing.

**Source**

Simulated data generated within the OmicNetR package.

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`plot_bipartite_network`*Plot Bi-partite sCCA Weight Network*

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**Description**

Optimized version using Base-R igraph engine to prevent memory exhaustion.

**Usage**

```
plot_bipartite_network(  
  net_data,  
  gene_color = "#1F77B4",  
  metabolite_color = "#FF7F0E",  
  layout_type = "fr"  
)
```

**Arguments**

- net\_data            The edge list data frame from `scca_to_network()`.
- gene\_color        Color for gene nodes.
- metabolite\_color            Color for metabolite nodes.
- layout\_type       igraph layout to use (default "fr").

**Value**

A graph object (invisibly).

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plot\_correlation\_heatmap

*Global Gene-Metabolite Correlation Heatmap*

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**Description**

Visualizes the correlation matrix with a gradient color scale.

**Usage**

```
plot_correlation_heatmap(scca_model, X, Y, top_n = 20)
```

**Arguments**

- scca\_model        The result object from `omic_scca()`.
- X                Aligned RNA-seq matrix.
- Y                Aligned Metabolomics matrix.
- top\_n            Number of top features from each omic to include.

**Value**

(Invisible) A numeric matrix of correlations between the selected features in X and Y.

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`plot_pathway_circle`     *Canonical Loading Pathway Circle Plot*

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**Description**

Visualizes top feature importance in a radial layout.

**Usage**

```
plot_pathway_circle(scca_model, top_features = 40, pathway_db = "KEGG")
```

**Arguments**

<code>scca_model</code>	The result object from <code>omic_scca()</code> .
<code>top_features</code>	Number of most weighted features to map.
<code>pathway_db</code>	Conceptual database name for labeling.

**Value**

A `ggplot2` object.

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`scca_to_network`     *Convert sCCA Loadings to Network Edges*

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**Description**

Generates an edge list for network plotting from sCCA loadings.

**Usage**

```
scca_to_network(scca_model, comp_select = 1, weight_threshold = 0.05)
```

**Arguments**

<code>scca_model</code>	The result object from <code>omic_scca()</code> .
<code>comp_select</code>	Which canonical component to use.
<code>weight_threshold</code>	Minimum absolute product of weights to include an edge.

**Value**

A data frame of edges with one row per gene-metabolite pair passing the threshold, containing:

- `Gene`: character, feature name from X.
- `Metabolite`: character, feature name from Y.
- `Weight_Product`: numeric, product of the selected loadings (edge weight).
- `Interaction_Type`: character, "Positive" or "Negative" based on the sign of `Weight_Product`.

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