

# Package ‘DeSciDe’

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**Type** Package

**Title** Tool for Unbiased Literature Searching and Gene List Curation

**Version** 1.0.3

**Description** Designed for genomic and proteomic data analysis, enabling unbiased PubMed searching, protein interaction network visualization, and comprehensive data summarization. This package aims to help users identify novel targets within their data sets based on protein network interactions and publication precedence of target's association with research context based on literature precedence. Methods in this package are described in detail in: Douglas et al. (2025) <[doi:10.1039/D5MO00160A](https://doi.org/10.1039/D5MO00160A)>. Key functionalities of this package also leverage methodologies from previous works, such as:

- Szklarczyk et al. (2023) <[doi:10.1093/nar/gkac1000](https://doi.org/10.1093/nar/gkac1000)>
- Winter (2017) <[doi:10.32614/RJ-2017-066](https://doi.org/10.32614/RJ-2017-066)>.

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**Imports** rentrez, ComplexHeatmap, circlize, STRINGdb, data.table, igraph, ggplot2, openxlsx, dplyr, tidyr, magrittr, tibble, ggrepel

**RoxygenNote** 7.3.3

**URL** <https://github.com/camdouglas/DeSciDe>

**BugReports** <https://github.com/camdouglas/DeSciDe/issues>

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

**VignetteBuilder** knitr

**Config/testthat/edition** 3

**Depends** R (>= 4.0.0)

**NeedsCompilation** no

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combine_summary	<i>Combine PubMed and STRING Metrics</i>
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## Description

Combine PubMed search summary and STRING gene metrics.

## Usage

```
combine_summary(
  pubmed_search_results,
  string_results,
  file_directory = NULL,
  export_format = "csv",
  export = FALSE,
  threshold_percentage = 20
)
```

## Arguments

`pubmed_search_results` Data frame with PubMed search results.

`string_results` Data frame with STRING metrics.

`file_directory` Directory for saving the output summary. Defaults to NULL.

`export_format` Format for export, either "csv", "tsv", or "excel".

`export` Logical indicating whether to export the summary. Defaults to FALSE.

`threshold_percentage` Percentage threshold for ranking (default is 20%).

## Value

A data frame with combined summary including connectivity, precedence, and category.

## Examples

```
pubmed_data <- data.frame(Gene = c("Gene1", "Gene2"), PubMed_Rank = c(1, 2))
string_data <- data.frame(Gene = c("Gene1", "Gene2"), Connectivity_Rank = c(2, 1))
combined <- combine_summary(pubmed_data, string_data, export = FALSE)
print(combined)
```

---

descide

*Run DeSciDe pipeline*

---

## Description

Run the entire analysis pipeline including PubMed search, STRING database search, and plotting.

## Usage

```
descide(
  genes_list,
  terms_list,
  rank_method = "weighted",
  species = 9606,
  network_type = "full",
  score_threshold = 400,
  threshold_percentage = 20,
  export = FALSE,
  file_directory = NULL,
  export_format = "csv"
)
```

## Arguments

<code>genes_list</code>	A list of gene IDs.
<code>terms_list</code>	A list of search terms.
<code>rank_method</code>	The method to rank pubmed results, either "weighted" or "total". Weighted ranks results based on order of terms inputted. Total ranks results on total sum of publications across all search term combinations. Defaults to "weighted".
<code>species</code>	The NCBI taxon ID of the species. Defaults to 9606 (Homo sapiens).
<code>network_type</code>	The type of string network to use, either "full" or "physical". Defaults to "full".
<code>score_threshold</code>	The minimum score threshold for string interactions. Defaults to 400.
<code>threshold_percentage</code>	Percentage threshold for ranking (default is 20%).
<code>export</code>	Logical indicating whether to export the results. Defaults to FALSE.
<code>file_directory</code>	Directory for saving the output files. Defaults to NULL.
<code>export_format</code>	Format for export, either "csv", "tsv", or "excel".

**Value**

A list containing the PubMed search results, STRING results, and summary results.

**Examples**

```
genes <- c("TP53", "BRCA1")
terms <- c("cancer", "tumor")
results <- descide(genes, terms, export = FALSE)
str(results)
```

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plot_clustering	<i>Plot STRING Interactions</i>
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**Description**

Plot STRING interactions degree vs. clustering.

**Usage**

```
plot_clustering(string_results, file_directory = NULL, export = FALSE)
```

**Arguments**

`string_results` Data frame with STRING metrics.  
`file_directory` Directory for saving the output plot. Defaults to NULL.  
`export` Logical indicating whether to export the plot. Defaults to FALSE.

**Value**

Invisibly returns the ggplot object.

**Examples**

```
# Example data frame
string_results <- data.frame(Degree = c(10, 5), Clustering_Coefficient_Percent = c(20, 10))
plot_clustering(string_results, file_directory = tempdir(), export = FALSE)
```

---

`plot_connectivity_precedence`*Plot Connectivity vs. Precedence*

---

**Description**

Create a scatter plot of Connectivity Rank vs. PubMed Rank.

**Usage**

```
plot_connectivity_precedence(  
  combined_summary,  
  file_directory = NULL,  
  export = FALSE  
)
```

**Arguments**

`combined_summary` Data frame with combined summary including categories.

`file_directory` Directory for saving the output plot. Defaults to NULL.

`export` Logical indicating whether to export the plot. Defaults to FALSE.

**Value**

Invisibly returns a ggplot object.

**Examples**

```
combined_data <- data.frame(Gene = c("Gene1", "Gene2"), Connectivity_Rank = c(1, 2),  
  PubMed_Rank = c(2, 1),  
  Category = c("High Connectivity - High Precedence", "Other"))  
plot_connectivity_precedence(combined_data, export = FALSE)
```

---

`plot_heatmap`*Plot Heatmap*

---

**Description**

Create and optionally save a heatmap of the PubMed search results.

**Usage**

```
plot_heatmap(pubmed_search_results, file_directory = NULL, export = FALSE)
```

**Arguments**

pubmed\_search\_results      A data frame containing raw search results with genes and terms.

file\_directory      Directory for saving the output plot. Defaults to NULL.

export      Logical indicating whether to export the plot. Defaults to FALSE.

**Value**

Invisibly returns a HeatmapList object.

**Examples**

```
# Example data frame
data <- data.frame(Gene = c("Gene1", "Gene2"),
                  Term1 = c(10, 20),
                  Term2 = c(5, 15),
                  Total = c(15, 35),
                  PubMed_Rank = c(1, 2))
plot_heatmap(data, file_directory = tempdir(), export = FALSE)
```

---

plot\_string\_network      *Plot STRING Network*

---

**Description**

Plot STRING network interactions using STRINGdb.

**Usage**

```
plot_string_network(
  string_db,
  string_ids,
  file_directory = NULL,
  export = FALSE
)
```

**Arguments**

string\_db      A STRINGdb object.

string\_ids      A list of STRING IDs.

file\_directory      Directory for saving the output plot. Defaults to NULL.

export      Logical indicating whether to export the plot. Defaults to FALSE.

**Value**

Invisibly returns NULL.

## Examples

```
## Not run:
library(STRINGdb)
string_db <- STRINGdb$new(species = 9606)
string_ids <- c("9606.ENSP00000269305", "9606.ENSP00000357940")
plot_string_network(string_db, string_ids, file_directory = tempdir(), export = FALSE)

## End(Not run)
```

---

rank\_search\_results     *Rank Search Results*

---

## Description

Rank search results based on a chosen method.

## Usage

```
rank_search_results(data, terms_list, rank_method = "weighted")
```

## Arguments

data	A data frame containing search results.
terms_list	A list of search terms.
rank_method	The method to rank pubmed results, either "weighted" or "total". Weighted ranks results based on order of terms inputted. Total ranks results on total sum of publications across all search term combinations. Defaults to "weighted".

## Value

A data frame with ranked search results, which includes the genes and their corresponding ranks based on the search method.

## Examples

```
# Example data frame
data <- data.frame(Gene = c("Gene1", "Gene2"),
                  Term1 = c(10, 20),
                  Term2 = c(5, 15))
terms_list <- c("Term1", "Term2")
ranked_results <- rank_search_results(data, terms_list, rank_method = "weighted")
print(ranked_results)
```

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search_pubmed	<i>Search PubMed with Multiple Genes and Terms</i>
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**Description**

Perform a PubMed search for multiple genes and terms.

**Usage**

```
search_pubmed(genes_list, terms_list, rank_method = "weighted", verbose = TRUE)
```

**Arguments**

genes_list	A list of gene IDs.
terms_list	A list of search terms.
rank_method	The method to rank results, either "weighted" or "total". Defaults to "weighted".
verbose	Logical flag indicating whether to display messages. Default is TRUE.

**Value**

A data frame with search results, including genes, terms, and their corresponding publication counts and ranks.

**Examples**

```
genes <- c("TP53", "BRCA1")
terms <- c("cancer", "tumor")
search_results <- search_pubmed(genes, terms, rank_method = "weighted", verbose = FALSE)
print(search_results)
```

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search_string_db	<i>Search STRING Database</i>
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**Description**

Search the STRING database for protein interactions.

**Usage**

```
search_string_db(
  genes_list,
  species = 9606,
  network_type = "full",
  score_threshold = 400
)
```

**Arguments**

genes_list	A list of gene IDs.
species	The NCBI taxon ID of the species. Defaults to 9606 (Homo sapiens).
network_type	The type of network to use, either "full" or "physical". Defaults to "full".
score_threshold	The minimum score threshold for string interactions. Defaults to 400.

**Value**

A list containing the following elements:

**string\_results** A data frame with STRING interaction metrics.

**string\_db** The STRINGdb object used.

**string\_ids** The STRING IDs for the input genes.

**Examples**

```
## Not run:  
library(STRINGdb)  
genes <- c("TP53", "BRCA1")  
results <- search_string_db(genes)  
print(results)  
  
## End(Not run)
```

---

single\_pubmed\_search *Search PubMed*

---

**Description**

Perform a PubMed search for a given gene and term.

**Usage**

```
single_pubmed_search(gene, term)
```

**Arguments**

gene	A character string representing the gene symbol.
term	A character string representing the search term.

**Value**

An integer representing the number of PubMed articles found from the search query in PubMed.

**Examples**

```
# Perform a PubMed search for gene 'TP53' with term 'cancer'  
result <- single_pubmed_search("TP53", "cancer")  
print(result)
```

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