## Package 'DeeDeeExperiment'

December 1, 2025

Title DeeDeeExperiment: An S4 Class for managing and exploring omics

```
analysis results
Version 1.0.0
Description DeeDeeExperiment is an S4 class extending the SingleCellExperiment
      class, designed to integrate and manage omics analysis results. It
      introduces two dedicated slots to store Differential Expression Analysis
      (DEA) results and Functional Enrichment Analysis (FEA) results, providing a
      structured approach for downstream analysis.
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clusterPro\_res

clusterPro\_res

### Description

A list of FE results generated with clusterProfiler::enrichGO()

### **Format**

A list of enrichResult objects

### **Details**

A list of FE result for the macrophage data (salmonella vs naive and IFNg vs naive)

### Value

A list

### See Also

Other enrich\_resus: enrichr\_res, fgseaRes, gost\_res, gsea\_res, topGO\_results\_list

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dd\_list\_original

dd\_list\_original

### **Description**

A list of deedee\_prepared DE results.

### **Format**

A list object

### **Details**

documented creation in ... script

### Value

A list of DE results

DeeDeeExperiment-class

The DeeDeeExperiment class

### **Description**

The DeeDeeExperiment class is integrate and manage omics analysis results. It inherits from the SingleCellExperiment class, and additionally stores DE-related/functional enrichment information via dedicated slots and rowData.

### Usage

```
DeeDeeExperiment(
   sce = SingleCellExperiment(),
   de_results = NULL,
   enrich_results = NULL
)
```

### **Arguments**

sce A SingleCellExperiment object, that will be used as a scaffold to store the DE

related information.

de\_results A named list of DE results, in any of the formats supported by the package

(currently: results from DESeq2, edgeR, limma).

enrich\_results A named list of functional enrichment results. Each element can be either a

data.frame (currently supports results from topGO, enrichR, gProfiler, fgsea, gsea, DAVID, and output of GeneTonic shakers), or an enrichResult/gseaResult

objects (currently supports clusterProfiler)

#### **Details**

The sce parameter can be optionally left unspecified. If this is the case, the resulting DeeDeeExperiment object will contain as features the ones specified by the provided components of the object supplied via the de\_results parameter.

The conversion of the components of the de\_results list will be handled via conversion functions to uniform the names and set of information which will be stored in the returned DeeDeeExperiment object. The names of the list will be used to define the contrasts for the different DE analyses included, which will determine the way to access the information stored in the dea slot of the DeeDeeExperiment object.

The content of the enrich\_results provided by the user will be validated to ensure that it is properly formatted and correctly named. The FE tool can be automatically detected, and based on that, the appropriate shaking method is used to return a standardized format of the FEA results. The names of the list will be used to attempt to associate each enrichment result with a corresponding DE contrast stored in the DeeDeeExperiment object, but it also can be defined by the user.

Since a DeeDeeExperiment is also a SummarizedExperiment object, it can be seamlessly provided downstream for visualization and in-depth exploration to packages such as iSEE or similar.

#### Value

A DeeDeeExperiment object.

### **Slots**

dea This slot is designed to hold the DE-related information. This is internally being created upon importing from the list of DE results objects, provided when instantiating the DeeDeeExperiment

fea This slot is designed to hold Functional Enrichment related information.

#### Author(s)

Najla Abassi, Lea Rothörl, and Federico Marini

#### **Examples**

```
data("de_named_list", package = "DeeDeeExperiment")

dde_onlyde <- DeeDeeExperiment(
    de_results = de_named_list
)

# or, with a SE object as support - even without assay data available
library("SummarizedExperiment")

rd_macrophage <- DataFrame(
    gene_id = rownames(de_named_list$ifng_vs_naive)
)

rownames(rd_macrophage) <- rownames(de_named_list$ifng_vs_naive)

se_macrophage_noassays <- SummarizedExperiment(
    assays = SimpleList(),
    rowData = rd_macrophage
)

dde <- DeeDeeExperiment(</pre>
```

```
se_macrophage_noassays,
de_results = de_named_list
)
```

DeeDeeExperiment-methods

Methods for DeeDeeExperiment objects

### Description

The DeeDeeExperiment() class provides a family of methods to get and set DE-related information and functional enrichment results in DeeDeeExperiment objects.

### Usage

```
## S4 method for signature 'DeeDeeExperiment'
getDEAInfo(x)
## S4 replacement method for signature 'DeeDeeExperiment'
getDEAInfo(x) \leftarrow value
## S4 method for signature 'DeeDeeExperiment'
getDEANames(x)
## S4 method for signature 'DeeDeeExperiment'
renameDEA(x, old_name, new_name)
## S4 method for signature 'DeeDeeExperiment'
addDEA(x, dea, force = FALSE)
## S4 method for signature 'DeeDeeExperiment'
removeDEA(x, dea_name, remove_linked_fea = FALSE)
## S4 method for signature 'DeeDeeExperiment'
getDEA(
  Χ,
  dea_name = NULL,
  format = c("minimal", "original"),
  extra_rd = NULL,
  type = c("DFrame", "data.frame"),
  verbose = FALSE
## S4 method for signature 'DeeDeeExperiment'
getDEAList(x, format = c("minimal", "original"), verbose = FALSE)
## S4 method for signature 'DeeDeeExperiment'
addScenarioInfo(x, dea_name, info = NULL, force = FALSE)
## S4 method for signature 'DeeDeeExperiment'
getFEAInfo(x)
```

```
## S4 replacement method for signature 'DeeDeeExperiment'
getFEAInfo(x) <- value</pre>
## S4 method for signature 'DeeDeeExperiment'
getFEANames(x)
## S4 method for signature 'DeeDeeExperiment'
renameFEA(x, old_name, new_name)
## S4 method for signature 'DeeDeeExperiment'
addFEA(
  х,
  fea,
  de_name = NA_character_,
  fe_name = NULL,
  fea_tool = "auto",
  force = FALSE,
  verbose = FALSE
## S4 method for signature 'DeeDeeExperiment'
removeFEA(x, fea_name)
## S4 method for signature 'DeeDeeExperiment'
getFEA(x, fea_name = NULL, format = c("minimal", "original"), verbose = FALSE)
## S4 method for signature 'DeeDeeExperiment'
getFEAList(
  х,
  dea_name = NULL,
  format = c("minimal", "original"),
  verbose = FALSE
)
## S4 method for signature 'DeeDeeExperiment'
linkDEAandFEA(x, dea_name, fea_name, force = FALSE)
```

### Arguments

force

X	A DeeDeeExperiment() object
value	Replacement value for replacement methods.
old_name	A character vector of existing DEA names to be renamed in a DeeDeeExperiment object
new_name	A character vector with new names to assign to existing DEA names in a DeeDeeExperiment object. It must be the same length of old_name, and contains unique values that don't overlap with existing DEA names.
dea	A named list of DE results, in any of the formats supported by the package (currently: results from DESeq2, edgeR, limma).

A logical, indicating whether to overwrite results when introducing the same

results name. It defaults to FALSE.

dea\_name Character value, specifying the name of the DE analysis to get or remove, or

match against (e.g., to fetch associated FEA results), or to which additional

context and information can be attached

remove\_linked\_fea

A logical, specifying whether to remove or not the linked FEA when a DEA

results is removed

format A character string, specifying the DEA/FEAs output format. It takes either

"minimal" to return only essential columns (e.g. log2FC, p-value, adjusted p-value for DEAs, or gs\_id, gs\_description, gs\_pvalue, gs\_genes... for FEAs), or

"original" to return the full result object. It defaults to "minimal"

extra\_rd A character vector of additional columns from rowData(x) to include. It defaults

to c("gene\_id", "SYMBOL").

type A character string referring to the type of object returned by getDEA(). It de-

faults to DFrame, but can also take the value of data.frame

verbose Logical, whether or not to display warnings. If TRUE, warnings/messages will

be displayed. If FALSE, the function runs silently

info A character vector, containing contextual information about the specified DE

analysis. It defaults to NULL

fea A named list of Functional Enrichment results. Each element can be either a

 $data. frame \ (currently \ supports \ results \ from \ top GO, \ enrich R, \ gProfiler, \ fgsea, \ gsea, \ DAVID, \ and \ output \ of \ Gene Tonic \ shakers), \ or \ an \ enrich Result/gsea Result$ 

objects (currently supports clusterProfiler)

de\_name A character string to explicitly specify the name of the de result this fea should

be linked to. If not provided, the function will attempt to match fea names to de

results automatically.

fe\_name A character string giving a name to the FE results.

fea\_tool A character string indicating the FEA tool used. It can take any of the following

values: "topGO", "clusterProfiler", "GeneTonic", "DAVID", "gsea", "fgsea", "enrichr", "gProfiler". When not specified, it defaults to "auto" and the tool is

inferred automatically based on the input.

fea\_name Character value, specifying the name of the functional enrichment result to add

or remove

#### Details

#### **DEAs**

- getDEAInfo and getDEAInfo<- are the methods to get and set the dea information as a whole. These methods return DeeDeeExperiment objects.
- getDEANames returns the names of the available DE contrasts in DeeDeeExperiment objects.
- renameDEA is the method to rename one or multiple DEAs stored in a DeeDeeExperiment object.
- addDEA and removeDEA are used to respectively add or remove DE-results items. These methods also return DeeDeeExperiment objects, with updated content in the dea slot.
- dea and getDEAList retrieve the DEA information, as well as some extra rowData information and provide this as a DataFrame object (for a specific analysis) or as a list, with one element for each reported analysis.

• addScenarioInfo is the method to add user defined contextual information for a specific DE analysis. It allows users to attach free-text notes to a specific DEA results that stored in a DeeDeeExperiment object. This information can include any other relevant information to help document that DEA scenario. This context is stored in the dea slot under the name scenario\_info, which is not a default element in dea.

#### **FEAs**

- getFEAInfo and getFEAInfo<- are the methods to get and set the fea information as a whole. These methods return DeeDeeExperiment objects.
- getFEANames returns the names of the available enrichment results in DeeDeeExperiment objects.
- renameFEA is the method to rename one or multiple FEAs stored in a DeeDeeExperiment object.
- addFEA and removeFEA are used to respectively add or remove functional enrichment results items. These methods also return DeeDeeExperiment objects, with updated content in the fea slot.
- fea is the method to retrieve FE results stored in a DeeDeeExperiment object for a specific contrast, as a standardized format similar to the output of GeneTonic shakers.
- getFEAList is the method that retrieves FEA results as a list. if the dea\_name is indicated, the method will return only FEAs linked to that dea\_name, otherwise it returns all FEAs in the fea slot.
- linkDEAandFEA is the method that allows the user to manually link a FEA result to a specific DEA result.
- show is the method to nicely print out the information of a DeeDeeExperiment object.
- summary is the method to print a summary of the available DE and FE results in a DeeDeeExperiment object.

### Value

Return value varies depending on the individual methods, as described below.

### **Examples**

```
data("de_named_list", package = "DeeDeeExperiment")
data("topGO_results_list", package = "DeeDeeExperiment")
library("SummarizedExperiment")

rd_macrophage <- DataFrame(
    gene_id = rownames(de_named_list$ifng_vs_naive)
)
rownames(rd_macrophage) <- rownames(de_named_list$ifng_vs_naive)
se_macrophage_noassays <- SummarizedExperiment(
    assays = SimpleList(),
    rowData = rd_macrophage
)

# creating a `DeeDeeExperiment`
dde <- DeeDeeExperiment(
    se_macrophage_noassays,
    de_results = de_named_list
)
dde</pre>
```

```
new_del <- list(</pre>
  ifng2 = de_named_list$ifng_vs_naive,
  ifngsalmo2 = de_named_list$ifngsalmo_vs_naive
# add a new (set of) DE result(s)
dde_new <- addDEA(dde, new_del)</pre>
dde_new
# removing DEAs
dde_removed <- removeDEA(dde, "ifng_vs_naive")</pre>
dde_removed
# add a new (set of) FE result(s)
dde_new <- addFEA(dde, fea = topGO_results_list)</pre>
# removing FEAs
dde_rem <- removeFEA(dde_new, "ifng_vs_naive")</pre>
# display available DEAs
getDEANames(dde)
# display available FEAs
getFEANames(dde)
# print a summary of the available DEAs and FEAs
summary(dde, FDR = 0.01)
# rename DEA
dde_new <- renameDEA(dde_new,</pre>
  old_name = "salmonella_vs_naive",
  new_name = "Salmo_vs_Naive_renamed"
# assign DEA to FEA
dde_new <- linkDEAandFEA(dde_new,</pre>
  dea_name = "ifngsalmo_vs_naive",
  fea_name = "ifngsalmo_vs_naive"
```

DeeDeeExperiment-misc Miscellaneous DeeDeeExperiment methods

### **Description**

Miscellaneous methods for the DeeDeeExperiment class and its descendants that do not fit into any other documentation category such as, for example, show and summary methods.

### Usage

```
## S4 method for signature 'DeeDeeExperiment'
```

```
show(object)
## S4 method for signature 'DeeDeeExperiment'
summary(object, FDR = 0.05, show_scenario_info = FALSE, ...)
```

### **Arguments**

object a DeeDeeExperiment object

FDR Numeric, sets the significance threshold for subsetting differentially expressed

genes based on adjusted p-values. Defaults to 0.05

show\_scenario\_info

Logical; if TRUE, displays the associated scenario info for each DE contrast, if available. Defaults to FALSE

. . additional argument passed to the summary method. Currently supports:

- FDR: Numeric, sets the significance threshold for subsetting differentially expressed genes based on adjusted p-values. Defaults to 0.05
- show\_scenario\_info: Logical; if TRUE, displays the associated scenario info for each DE contrast, if available.

#### Value

Returns NULL

DeeDeeExperiment-pkg DeeDeeExperiment

### Description

DeeDeeExperiment is a an S4 class that allows integrating and managing transcriptomic analysis results.

### **Details**

DeeDeeExpeirment is an S4 class extending the SummarizedExperiment framework to facilitate the integration and management of transcriptomic analysis results. It introduces two dedicated slots to store Differential Expression (DE) analysis results and Functional Enrichment analysis outcomes, providing a structured approach for downstream analysis.

### Author(s)

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- Federico Marini <marinif@uni-mainz.de> (ORCID)

#### See Also

Useful links:

- https://github.com/imbeimainz/DeeDeeExperiment
- Report bugs at https://github.com/imbeimainz/DeeDeeExperiment/issues

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de\_limma

A sample MArrayLM object

### **Description**

A sample MArrayLM object, generated with limma

#### **Format**

A MArrayLM object

### **Details**

This MArrayLM object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with different stimulation conditions, including IFNg treatment, Salmonella infection, and their combined effects.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

### Value

A sample MArrayLM object, generated with limma

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

de\_named\_list

de\_named\_list

### **Description**

A named list of DE results, in their original format (from DESeq2, edgeR or limma)

### **Format**

A list object

### **Details**

documented creation in the create\_dataset\_example.R script in the scripts package folder

#### Value

A named list of DE results, in their original format

### Description

A sample DGEExact object, generated with edgeR

### **Format**

A DGEExact object

### **Details**

This DGEExact object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with IFNg to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

#### Value

A sample DGEExact object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive, dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_both, dge\_lrt\_Salm\_naive

dge\_exact\_IFNg\_naive A sample DGEExact object

#### **Description**

A sample DGEExact object, generated with edgeR

### **Format**

A DGEExact object

### **Details**

This DGEExact object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from naive macrophage to those associated with IFNg.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

dge\_exact\_Salm\_both 13

#### Value

A sample DGEExact object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

#### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive, dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_both, dge\_lrt\_Salm\_naive

### Description

A sample DGEExact object, generated with edgeR

#### **Format**

A DGEExact object

### **Details**

This DGEExact object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with Salmonella to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

### Value

A sample DGEExact object, generated with edgeR

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_naive, dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_naive

14 dge\_lrt\_IFNg\_both

dge\_exact\_Salm\_naive A sample DGEExact object

### Description

A sample DGEExact object, generated with edgeR

### **Format**

A DGEExact object

### **Details**

This DGEExact object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from naive macrophage to those associated with Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

#### Value

A sample DGEExact object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_both, dge\_lrt\_Salm\_naive

dge\_lrt\_IFNg\_both

A sample DGELRT object

#### **Description**

A sample DGELRT object, generated with edgeR

### **Format**

A DGELRT object

#### **Details**

This DGELRT object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with IFNg to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

dge\_lrt\_IFNg\_naive 15

#### Value

A sample DGELRT object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

#### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_both, dge\_lrt\_Salm\_naive

dge\_lrt\_IFNg\_naive

A sample DGELRT object

### Description

A sample DGELRT object, generated with edgeR

#### **Format**

A DGELRT object

### **Details**

This DGELRT object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with IFNg to those associated with IFNg and naive.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

### Value

A sample DGELRT object, generated with edgeR

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive dge\_lrt\_IFNg\_both, dge\_lrt\_Salm\_both, dge\_lrt\_Salm\_naive

dge\_lrt\_Salm\_naive

dge\_lrt\_Salm\_both

A sample DGELRT object

### Description

A sample DGELRT object, generated with edgeR

### **Format**

A DGELRT object

### **Details**

This DGELRT object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with Salmonella to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

#### Value

A sample DGELRT object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_naive

dge\_lrt\_Salm\_naive

A sample DGELRT object

#### **Description**

A sample DGELRT object, generated with edgeR

### **Format**

A DGELRT object

### **Details**

This DGELRT object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from naive macrophage to those associated with Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

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### Value

A sample DGELRT object, generated with edgeR

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other edgeRresus: dge\_exact\_IFNg\_both, dge\_exact\_IFNg\_naive, dge\_exact\_Salm\_both, dge\_exact\_Salm\_naive dge\_lrt\_IFNg\_both, dge\_lrt\_IFNg\_naive, dge\_lrt\_Salm\_both

enrichr\_res

enrichr\_res

### **Description**

A list of FE results generated with enrichr::enrichr()

#### **Format**

A list of data.frame objects

#### **Details**

A list of FE result for the macrophage data (salmonella vs naive)

### Value

A list

### See Also

Other enrich\_resus: clusterPro\_res, fgseaRes, gost\_res, gsea\_res, topGO\_results\_list

fgseaRes

fgseaRes

### Description

A data frame of FE results generated with fgsea::fgsea()

### **Format**

A data.table/data.frame object

### **Details**

Tabular representation of FE result for the macrophage data (IFNg vs naive)

18 gsea\_res

### Value

A data.table/data.frame

### See Also

Other enrich\_resus: clusterPro\_res, enrichr\_res, gost\_res, gsea\_res, topGO\_results\_list

gost\_res

 ${\sf gost\_res}$ 

### Description

A list of FE results generated with gprofiler2::gost()

### **Format**

A list, as returned by gprofiler2

### **Details**

A list of FE result for the macrophage data (salmonella vs naive)

### Value

A list

### See Also

Other enrich\_resus: clusterPro\_res, enrichr\_res, fgseaRes, gsea\_res, topGO\_results\_list

gsea\_res

gsea\_res

### Description

An individual set of FE results generated with clusterProfiler::gseGO()

### **Format**

A gseaResult object

### **Details**

A set of FE results for the macrophage data (IFNg vs naive)

### Value

A gseaResult object

### See Also

Other enrich\_resus: clusterPro\_res, enrichr\_res, fgseaRes, gost\_res, topGO\_results\_list

IFNg\_both

IFNg\_both

A sample DESeqResults object

### **Description**

A sample DESeqResults object, generated with DESeq2

#### **Format**

A DESeqResults object

### **Details**

This DESeqResults object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with IFNg to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

#### Value

A sample DESeqResults object, generated with DESeq2

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

#### See Also

Other DEresus: IFNg\_naive, Salm\_both, Salm\_naive

IFNg\_naive

A sample DESeqResults object

### **Description**

A sample DESeqResults object, generated with DESeq2

### **Format**

A DESeqResults object

### **Details**

This DESeqResults object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from naive macrophage to those associated with IFNg.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

20 Salm\_both

#### Value

A sample DESeqResults object, generated with DESeq2

#### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

#### See Also

Other DEresus: IFNg\_both, Salm\_both, Salm\_naive

Salm\_both

A sample DESeqResults object

### **Description**

A sample DESeqResults object, generated with DESeq2

#### **Format**

A DESeqResults object

### **Details**

This DESeqResults object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from macrophage associated with Salmonella to those associated with IFNg and Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

### Value

A sample DESeqResults object, generated with DESeq2

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other DEresus: IFNg\_both, IFNg\_naive, Salm\_naive

Salm\_naive 21

Salm\_naive

A sample DESeqResults object

### **Description**

A sample DESeqResults object, generated with DESeq2

#### **Format**

A DESeqResults object

#### **Details**

This DESeqResults object contains the results of a Differential Expression Analysis performed on data from the macrophage package, more precisely contrasting the counts from naive macrophage to those associated with Salmonella.

The code to create said object can be found in the folder /inst/scripts in the DeeDeeExperiment package, the file is called generate\_data.R.

### Value

A sample DESeqResults object, generated with DESeq2

### References

Alasoo, et al. "Shared genetic effects on chromatin and gene expression indicate a role for enhancer priming in immune response", Nature Genetics, January 2018 doi: 10.1038/s41588-018-0046-7.

### See Also

Other DEresus: IFNg\_both, IFNg\_naive, Salm\_both

 $supported_fea_formats$  Display available FEA formats

### Description

Display available FEA formats

### Usage

```
supported_fea_formats()
```

#### Value

a data.frame of possible FEA input formats

### **Examples**

```
supported_fea_formats()
```

22 topGO\_results\_list

topGO\_results\_list topGO\_results\_list

### Description

A list of FE results generated with mosdef::topGOtable()

### **Format**

A list of data.frame objects

### **Details**

A list of FE results for the macrophage data

### Value

A list

### See Also

Other enrich\_resus: clusterPro\_res, enrichr\_res, fgseaRes, gost\_res, gsea\_res

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