Package 'DEET'

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Title Differential Expression Enrichment Tool

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Description RNA sequencing (RNA-seq) followed by differential gene expression analyses is a fundamental approach for making biological discoveries. Ongoing large-scale efforts to systematically process and normalize publicly available gene expression data facilitate rapid reanalyses of specific studies and the development of new methods for querying it. While there are several powerful tools for querying systematically processed publicly available RNAseq data at the individual sample level, there are fewer options for querying differentially expressed gene (DEG) lists generated from these experiments. Here, we present the Differential Expression Enrichment Tool (DEET), which allows users to interact with 3162 consistently processed DEG lists curated from 142 RNA-seq datasets obtained from recount2 database, which contains data from consortiums (GTex, TCGA) and individual labs (SRA). To establish DEET we integrated systematically processed human RNAseq data from recount2 with reported and predicted metadata from multiple sources and developed a CRAN R package and Shiny App where users can compare their genes, p-values, and coefficients against the DEG lists within DEET. Here we present DEET and demonstrate how it can facilitate hypothesis generation and provide biological insight from user-defined differential gene expression results. Reference: Sokolowski, D.J., Ahn J., Erdman, L., Hou, H., Ellis, K., Wang L., Goldenberg, A., and Wilson, M.D. (2022) Differential Expression Enrichment Tool (DEET): An interactive atlas of human differential gene expression. (In Preparation).

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DEET_data_download DEET_data_download

Description

Function to automatically download the files within the DEET database that are required for the DEET_enrich and DEET_feature_extractfunctions.

Usage

```
DEET_data_download(x = "enrich")
```

Arguments

x categorical variable containing options "ALL", "enrich", "metadata" or "feature_matrix".

Value

Named list with the neccesary data required to input into DEET_feature_extract or DEET_enrich. The metadata within DEET can also be downloaded.

- feature_matrix A gene by comparison matrix populated with the log2FC of gene expression for all genes, regardless of DE status.
- metadata a comparison by explanatory piece of data dataframe providing important details to contextualize each study. For every pairwise comparison, the study name, source (SRA, TCGA, GTEx and SRA-manual), description from the DRA compendium, the number of samples (total, up-condition, and down-condition), samples (total, up-condition, down-condition), tissue (including tumour from TCGA), number of DEs (total, up-condition, down-condition), age (mean +- sd), sex, top 15 DEGs up, top 15 DEGs down, top 5 enriched pathways, and top 5 enriched TFs. PMID are also available for studies selected from SRA. Lastly, each pairwise comparison was given an overall category based on those decided in Crow et al., 2019.
- DEET_enrich A named list of seven objects containing the data frames summarizing the DEGs from comparisons within DEET, GMT objects of comparisons within DEET for enrichment through ActivePathways, GMT objects for basic pathway and TF enrichment, and a dataframe for the metadata of each study. For more detail on each element of the list, please consult the vignette or "?DEET_example_data", as it is a subset of this object

Author(s)

Dustin Sokolowski, Jedid Ahn

References

Engebretsen, S., & Bohlin, J. (2019). Statistical predictions with glmnet. Clinical epigenetics, 11(1), 1-3.

```
# Download the metadata. Downloading other
# files within DEET are larger and take
# a bit more time.
downloaded <- DEET_data_download(x = "metadata")
# extract metadata from the list
metadata <- downloaded[["metadata"]]</pre>
```

DEET_enrich

Description

Core function of DEET where an input weighted human gene list will be queried to DEETs library of studies.

Usage

```
DEET_enrich(DEG_list, DEET_dataset, ordered = FALSE, background = NULL)
```

Arguments

DEG_list	Data frame or matrix of gene symbols with corresponding padj and log2FC values (3 columns in total). Can also be a character vector of gene symbols only. colnames of genes: c("gene_symbol", "padj", "coef") The rownames of the dataframe are also the gene symbols.
DEET_dataset	The databank of the differential expression enrichment tool. Appropriate inputs here are "DEET_example_data" stored within DEET, the "DEET_combined.rda" file from the DEET stable repository found at X, and the DEET database developmental repository found at Y. The DEET_dataset is a named list where details of it's structure can be found ?DEET_example_data.
ordered	Boolean value specifying whether DEG_list is a character vector of gene symbols that is ordered. Default value is FALSE.
background	Character vector of human gene symbols showing all possible genes. Default value is NULL.

Value

Named list where each element contains 6 objects. Each object will contain the results (enrichment or correlation) and corresponding metadata.

- AP_INPUT_BP_output Enriched BPs of input gene list.
- AP_INPUT_TF_output Enriched TFs of input gene list.
- AP_DEET_DE_output Enrichment of input gene list on DEETs studies.
- AP_DEET_BP_output Enrichment of BPs of input gene list on DEETs BPs of studies.
- AP_DEET_TF_output Enrichment of TFs of input gene list on DEETs TFs of studies.
- DE_correlations Correlation values of input gene list to DEETs studies (both Pearson and Spearman).

Author(s)

Dustin Sokolowski, Jedid Ahn

References

Paczkowska M, Barenboim J, Sintupisut N, et al. Integrative pathway enrichment analysis of multivariate omics data. Nat Commun. 2020;11(1):735. doi:10.1038/s41467-019-13983-9

Examples

```
data("example_DEET_enrich_input")
data("DEET_example_data")
DEET_out <- DEET_enrich(example_DEET_enrich_input, DEET_dataset = DEET_example_data)</pre>
```

Description

Generate barplots or dotplots from the output of DEET

Usage

```
DEET_enrichment_plot(
  enrich_list,
  outname,
  width = 8,
  text_angle = 0,
  horizontal = FALSE,
  topn = 5,
  ol_size = 1,
  exclude_domain = "",
  cluster_order = NULL,
  dot = FALSE,
  colors = "Set2",
  split_domain = FALSE
)
```

Arguments

enrich_list	A list of enrichments from DEET, with each element post-processed with the barplot enrichment function.
outname	A character giving the title of the barplot or dotplot.
width	The number of inches in the barplot or dotplot.
text_angle	The angle of the enriched studies.
horizontal	Whether the output barplot is vertical or horizontal
topn	the top number of studies (by p-value) to be plotted.

DEET_example_data

ol_size the minimum number of overlapping genes (or paths) in an enriched study.

exclude_domain Exclude studies enriched based on DEGs, Paths, or TF if the user happened to

aggregate the results into a single DF, generally unused.

cluster_order Factor to group studies based on the researchers custom annotation.

dot logical (T/F) of whether to produce a dotplot or a barplot

colors Type of color pallete to input into 'scale_fill_brewer' of ggplot.

split_domain logical (T/F) of whether to plot the "topn" studies for each "domain" (default

is source) or to plot the topn pathwys regardless of domain. default is set to

FALSE, meaning it plots the topn pathways regardless of domain.

Value

A ggplot2 object (barplot or dotplot) of enrichment identified within DEET.

Author(s)

Dustin Sokolowski, Hauyun Hou PhD

Examples

```
data("example_DEET_enrich_input")
data("DEET_example_data")
DEET_out <- DEET_enrich(example_DEET_enrich_input, DEET_dataset = DEET_example_data)

# converting output to format compatible with DEET_enrichment plot
DE_example <- DEET_out$AP_DEET_DE_output$results
DE_example$term.name <- DEET_out$AP_DEET_DE_output$metadata$DEET.Name
DE_example$domain <- "DE"
DE_example$overlap.size <- lengths(DE_example$overlap)
DE_example$p.value <- DE_example$adjusted.p.val</pre>
DE_example_plot <- DEET_enrichment_plot(list(DE_example = DE_example), "DE_example")</pre>
```

DEET_example_data

DEET_example_data

Description

Named list of gene-sets and representative metadata for studies associated with Alizada et al., 2021 This example data is the exact same as what is needed to run DEET enrich properly but subsetted to have 13 studies that are enriched by 'example_DEET_enrich_input'. This way, the example gives an output at all levels of enrichment and at the correlation level.

DEET_feature_extract

Usage

```
data(DEET_example_data)
```

Format

A named list of seven objects containing the data frames summarizing the DEGs from comparisons within DEET, GMT objects of comparisons within DEET for enrichment through ActivePathways, GMT objects for basic pathway and TF enrichment, and a dataframe for the metadata of each study.

#'

- **DEET_DE** A list of data frames containing the significant DE genes, mean expression, log2fold-change, and padj from DESeq (padj < 0.05).
- **DEET_gmt_BP** A list of class GMT, which is a list of studies where each study is populated by comparison id (internal DEET identifier), comparison name (interpretable comparison name), and a gene set. In this case the gene-set is the pathways that are enriched within that study.
- **DEET_gmt_TF** A list of class GMT, which is a list of studies where each study is populated by comparison id (internal DEET identifier), comparison name (interpretable comparison name), and a gene set. In this case the gene-set is the TFs that are enriched within that study.
- **DEET_gmt_DE** A list of class GMT, which is a list of studies where each study is populated by comparison id (internal DEET identifier), comparison name (interpretable comparison name), and a gene set. In this case the gene-set is the DEGs that are enriched within that study.
- gmt_BP A list of class GMT, which is a list of gene ontology gene-sets acquired from the bader lab 'http://download.baderlab.org/EM_Genesets/'#'
- gmt_TF A list of class GMT, which is a list of Transcription Factor gene-sets acquired from the bader lab 'http://download.baderlab.org/EM_Genesets/'
- **DEET_metadata** For every pairwise comparison, the study name, source (SRA, TCGA, GTEx and SRA-manual), description from the DRA compendium, the number of samples (total, upcondition, and down-condition), samples (total, up-condition, down-condition), tissue (including tumour from TCGA), number of DEs (total, up-condition, down-condition), age (mean +sd), sex, top 15 DEGs up, top 15 DEGs down, top 5 enriched pathways, and top 5 enriched TFs. PMID are also available for studies selected from SRA. Lastly, each pairwise comparison was given an overall category based on those decided in Crow et al., 2019.

Examples

```
data(DEET_example_data)
```

Description

Identify which genes are associated with pieces of metadata that a researcher queries.

Usage

```
DEET_feature_extract(mat, response, datatype)
```

Arguments

mat A gene-by-study matrix populated by the coefficients of that study. By default,

the coefficient is the log2Fold-change of genes as long as they are differentially

expressed (cutoff = padj < 0.05).

response A vector (binomial, categorical, or continuous) that is used to associated the

DEGs within the studies.

datatype indication of whether the response variable is binomial, categorical, or continu-

ous.

Value

Named list given the elastic net coefficients and the eleastic net regression between the response variable and the DEGs within DEET. It also outputs the correlation, ANOVA, and wilcoxon test of every gene against the response variable based on if it's continuous, categorical, or binomial in nature.

- elastic_net_coefficients Association that a gene has with the response variable based on the elastic net regression.
- elastic_net Output of the elastic net regression
- - basic_features gives the output of the correlation, ANOVA, and wilcoxon test of every gene against the response variable.

Author(s)

Dustin Sokolowski, Jedid Ahn

References

Engebretsen, S., & Bohlin, J. (2019). Statistical predictions with glmnet. Clinical epigenetics, 11(1), 1-3.

```
data(DEET_feature_extract_example_matrix)
data(DEET_feature_extract_example_response)
single1 <- DEET_feature_extract(DEET_feature_extract_example_matrix,
DEET_feature_extract_example_response, "categorical")</pre>
```

DEET_feature_extract_example_matrix

DEET_feature_extract_example_matrix

Description

An object of class data frame where rows are genes and columns are comparisons. The matrix is populated by the log2Fold-change of each gene within each study. If the gene is not DE within that study (padj < 0.05), it is populated with 0 instead of the log2Fold-change. This object is inputted into the 'mat' input variable for the 'DEET_feature_extract' function.

Usage

```
data(DEET_feature_extract_example_matrix)
```

Format

An object of class data frame where rows are genes and columns are comparisons (1000 randomly selected genes and 200 randomly selected studies).

Examples

```
data(DEET_feature_extract_example_matrix)
```

```
DEET_feature_extract_example_response

DEET_feature_extract_example_response
```

Description

Character vector giving the source (TCGA SRA, GTEx, SRA-manual) of 200 comparisons within DEET. Used as the input for the 'response' input of 'DEET_feature_extract' in the example. For this response variable to work, the 'datatype' input variable would also need to be set to "categorical".

Usage

```
data(DEET_feature_extract_example_response)
```

Format

Character vector giving the source (TCGA SRA, GTEx, SRA-manual) of 200 comparisons within DEET

```
data(DEET_feature_extract_example_response)
```

DEET_plot_correlation DEET_plot_correlation

Description

Take significant correlation outputs and generate scatterplots of the genes DE in one or the other.

Usage

```
DEET_plot_correlation(correlation_input)
```

Arguments

```
correlation_input
```

The "DE_correlations" element of the output of the DEET_enrich function. This function only works if there is at least one significantly correlated study.

Value

Named list of ggplot objects with the correlation between the input study and the study within DEET

Author(s)

Dustin Sokolowski, Jedid Ahn

```
data("example_DEET_enrich_input")
data("DEET_example_data")
DEET_out <- DEET_enrich(example_DEET_enrich_input, DEET_dataset = DEET_example_data)
correlation_input <- DEET_out$DE_correlations
correlation_plots <- DEET_plot_correlation(correlation_input)</pre>
```

Description

Exon-level DEGs of HAoEC after TNFa treatment for 45 mins from Alizada et al., 2021. Object is a data.frame with columns "gene_symbol" "padj" and "coef", which in this case is the log2Fold-change of differential expression.

Usage

```
data(example_DEET_enrich_input)
```

Format

A data frame with three columns. Rows are genes and it's populated by the gene symbol, padj of gene expression, and coef (log2Fold-change).

Examples

```
data(example_DEET_enrich_input)
```

```
proccess\_and\_plot\_DEET\_enrich \\ proccess\_and\_plot\_DEET\_enrich
```

Description

Generates barplots and dotplots based on the output of the DEET_enrich function.

Usage

```
proccess_and_plot_DEET_enrich(
   DEET_output,
   colour_barplot = "Source",
   width = 8,
   text_angle = 0,
   horizontal = F,
   topn = 5,
   ol_size = 1,
   exclude_domain = "",
   cluster_order = NULL,
   colors = "Set2"
)
```

Arguments

DEET_output Direct output of the DEET_enrich function. A list with all of the same names as

DEET_output.

colour_barplot Pick dotplot or barplot colours. It can be NULL, in which all bars are the same

or it can be a (case sensitive) column within the metadata. Defaults to "source".

width The number of inches in the barplot or dotplot.

text_angle The angle of the enriched studies.

horizontal Whether the output barplot is vertical or horizontal topn the top number of studies (by p-value) to be plotted.

ol_size the minimum number of overlapping genes (or paths) in an enriched study.

exclude_domain Exclude studies enriched based on DEGs, Paths, or TF if the user happened to

aggregate the results into a single DF, generally unused.

cluster_order Factor to group studies based on the researchers custom annotation.

colors Type of color pallete to input into 'scale_fill_brewer' of ggplot.

Value

Named list where each element is a ggplot object plotting the output of the enrichment tests within DEET. The final element is the output of ActivePathways (in DEET) that is directly compatible with the DEET_enrichment_barplot function.

- DEET_DotPlot ggplot object of Dotplot of enrichment of enrichment of DEET studies based on DE, BP, and TF information. Only plotted if 2/3 levels contain at least one significant study.
- Pathway_barplot ggplot object of Barplot of standard gene set enrichment based on gene ontology and TFs. Only plotted if there is at least one enriched significant pathway/TF.
- individual_barplot ggplot object of Barplot of the top enriched pathways or studies (depending on the input list).Barplot is only generated if each list has at least one pathway (or study) is enriched.
- DEET_output_forplotting output of Activepathways with "domain", "overlap.size", and "p.value" columns added to be compatible with the DEET_enrichment_barplot function.

Author(s)

Dustin Sokolowski, Hauyun Hou PhD

```
data("example_DEET_enrich_input")
data("DEET_example_data")
DEET_out <- DEET_enrich(example_DEET_enrich_input, DEET_dataset = DEET_example_data)
plotting_example <- proccess_and_plot_DEET_enrich(DEET_out, text_angle = 45,
horizontal = TRUE, topn=4)</pre>
```

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