Package 'oncoEnrichR'

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Type Package
```

Title Functional annotation and enrichment analysis of genesets in the context of cancer

Version 0.7.1

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URL https://github.com/sigven/oncoEnrichR

BugReports https://github.com/sigven/oncoEnrichR/issues

Description oncoEnrichR performs annotation and prioritization of genesets discovered from high-throughput experiments within

cancer biology. The tool queries a number of high-

quality resources, including Project Score (cellular fitness in genome-scale

drop-out screens), Open Targets Platform (comprehensive target-drug and target-

disease associations), TCGA (genetic aberrations

and gene expression patterns in human tumor samples), STRING (protein-

protein interaction networks), CORUM (curated protein complexes),

COMPPI (subcellular compartments), and Gene Ontology/MSigDB/REACTOME/KEGG (functional enrichment/over-representation analysis

with respect to gene function and cellular pathways). The package outputs an interac-

tive HTML report in which the user

can interrogate the the various types of annotations and analyses performed with the geneset.

Depends R (>= 3.5)

biocViews

Imports tidyverse (>= 1.3.0), visNetwork (>= 2.0.9), plotly (>= 4.9.1), pals (>= 1.5), org.Hs.eg.db, clusterProfiler, igraph, reshape2 (>= 1.4.4), rlogging, gganatogram, openxlsx, speed2

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Encoding UTF-8

Remotes mjkallen/rlogging, jespermaag/gganatogram, molsysbio/speed2

LazyData true

RoxygenNote 7.1.1

Roxygen list(markdown = TRUE)

onco_enrich

R topics documented:

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onco_enrich		Function that interrogates and analyzes a list of human protein-coding genes for cancer relevance			

Description

Function that interrogates and analyzes a list of human protein-coding genes for cancer relevance

Usage

```
onco_enrich(
  query,
  query_source = "symbol",
  ignore_unknown = FALSE,
  project_title = "Project title",
  project_owner = "Project owner",
  project_description = "Project description",
  background_enrichment = NULL,
  background_enrichment_source = "symbol",
  background_enrichment_description = "All protein-coding genes",
  p_value_cutoff_enrichment = 0.05,
  p_value_adjustment_method = "BH",
  q_value_cutoff_enrichment = 0.2,
  min_geneset_size = 10,
  max_geneset_size = 500,
  simplify_go = F,
  ppi_add_nodes = 50,
  ppi_score_threshold = 900,
  show_gene_summary = F,
  show_ppi = T,
  show_drugs_in_ppi = F,
  show_disease = T,
  show_drug = T,
  show_enrichment = T,
  show_tcga_aberration = T,
  show_tcga_coexpression = T,
  show\_subcell\_comp = T,
  show_crispr_lof = T,
  show\_complex = T
)
```

Arguments

query

character vector with gene/query identifiers

character indicating source of query (one of "uniprot acc", "symbol", "entrezquery_source gene", or "ensembl_gene_id") ignore_unknown logical indicating if analysis should continue when uknown query identifiers are encountered project_title project title (title of report) project_owner name of project owner project_description project background information background_enrichment character vector with gene identifiers, used as reference/background for enrichment/overrepresentation analysis background_enrichment_source character indicating source of background ("uniprot_acc", "symbol", "entrezgene", "ensembl_gene_id" background_enrichment_description character indicating type of background (e.g. "All lipid-binding proteins (n = 200)") p_value_cutoff_enrichment cutoff p-value for enrichment/over-representation analysis p_value_adjustment_method one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none" q_value_cutoff_enrichment cutoff q-value for enrichment analysis min_geneset_size minimal size of geneset annotated by term for testing in enrichment/over-representation analysis max_geneset_size maximal size of geneset annotated by term for testing in enrichment/over-representation analysis remove highly similar GO terms in results from GO enrichment/over-representation simplify_go analysis number of nodes to add to query set when computing the protein-protein interppi_add_nodes action network (STRING) ppi_score_threshold minimum score (0-1000) for retrieval of protein-protein interactions (STRING) show_gene_summary logical indicating if report should fetch summary of gene function (from Ref-Seq/mygene.info, will increase processing time) logical indicating if report should contain protein-protein interaction data (STRING) show_ppi show_drugs_in_ppi logical indicating if targeted drugs (> phase 3) should be displayed in proteinprotein interaction network (Open Targets Platform) logical indicating if report should contain disease associations (Open Targets show_disease Platform) show enrichment logical indicating if report should contain functional enrichment/over-representation analysis (MSigDB, GO, KEGG, REACTOME etc.) show_tcga_aberration logical indicating if report should contain TCGA aberration plots (amplifica-

tions/deletions)

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show_tcga_coexpression

logical indicating if report should contain TCGA co-expression data (RNAseq)

of queryset with oncogenes/tumor suppressor genes

show_subcell_comp

logical indicating if report should list subcellular compartment annotations (ComPPI)

show_crispr_lof

logical indicating if report should list results from CRISPR/Cas9 loss-of-fitness

screens (Project Score)

show_complex logical indicating if report should list proteins in known protein complexes (CO-

RUM)

write

Function that writes the contents in the oncoEnrichR report object to an interactive HTML report

Description

Function that writes the contents in the oncoEnrichR report object to an interactive HTML report

Usage

```
write(report, project_directory, report_name, format = "html")
```

Arguments

report object with oncoEnrichR report data (returned by oncoEnrichR::onco_enrich)

project_directory

working directory

report_name prefix filename for report output format file format of output (html/json)

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