## Package 'oncoEnrichR'

August 11, 2020

```
Type Package
Title Functional annotation and enrichment analysis of genesets in the context of cancer
Version 0.7.2
Author Sigve Nakken
Maintainer Sigve Nakken <sigven@ifi.uio.no>
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 (func-
      tional 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
License MIT
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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