## main

August 21, 2021

## 1 GO analysis using GOATOOLS

```
[1]: import functools
  import pandas as pd
  import collections as cx
  from pybiomart import Dataset
  # GO analysis
  from goatools.base import download_go_basic_obo
  from goatools.base import download_ncbi_associations
  from goatools.obo_parser import GODag
  from goatools.anno.genetogo_reader import Gene2GoReader
  from goatools.goea.go_enrichment_ns import GOEnrichmentStudyNS
```

```
[2]: Ofunctools.lru_cache()
     def get_database():
         dataset = Dataset(name="hsapiens_gene_ensembl",
                           host="http://www.ensembl.org",
                           use_cache=True)
         db = dataset.query(attributes=["ensembl_gene_id",
                                        "external_gene_name",
                                         "entrezgene_id"],
                            use_attr_names=True).dropna(subset=['entrezgene_id'])
         return db
     @functools.lru_cache()
     def get_deg():
         fn = '../../_m/genes/diffExpr_EAvsAA_FDR05.txt'
         return pd.read_csv(fn, sep='\t')
     @functools.lru_cache()
     def convert2entrez():
         df = get_deg()
         if 'EntrezID' in df.columns:
             return df.rename(columns={'EntrezID': 'entrezgene_id'})
         else:
```

```
[3]: def obo_annotation(alpha=0.05):
         # database annotation
         fn_obo = download_go_basic_obo()
         fn_gene2go = download_ncbi_associations() # must be qunzip to work
         obodag = GODag(fn_obo) # downloads most up-to-date
         anno_hs = Gene2GoReader(fn_gene2go, taxids=[9606])
         # get associations
         ns2assoc = anno_hs.get_ns2assc()
         for nspc, id2gos in ns2assoc.items():
             print("{NS} {N:,} annotated human genes".format(NS=nspc, N=len(id2gos)))
         goeaobj = GOEnrichmentStudyNS(
             get_database()['entrezgene_id'], # List of human genes with entrez IDs
             ns2assoc, # geneid/GO associations
             obodag, # Ontologies
             propagate_counts = False,
             alpha = alpha, # default significance cut-off
             methods = ['fdr_bh'])
         return goeaobj
     def run_goea(direction):
         if direction == "Up":
             df = get_upregulated()
         elif direction == "Down":
             df = get_downregulated()
         else:
             df = convert2entrez()
         geneids_study = {z[0]:z[1] for z in zip(df['entrezgene_id'], df['Symbol'])}
         goeaobj = obo_annotation()
         goea_results_all = goeaobj.run_study(geneids_study)
         goea_results_sig = [r for r in goea_results_all if r.p_fdr_bh < 0.05]</pre>
```

```
ctr = cx.Counter([r.NS for r in goea_results_sig])
print('Significant results[{TOTAL}] = {BP} BP + {MF} MF + {CC} CC'.format(
    TOTAL=len(goea_results_sig),
    BP=ctr['BP'], # biological_process
    MF=ctr['MF'], # molecular_function
    CC=ctr['CC'])) # cellular_component

if direction == "Up":
    label = "upregulated"
elif direction == "Down":
    label = "downregulated"
else:
    label = "allDEG"
goeaobj.wr_xlsx("GO_analysis_%s.xlsx" % label, goea_results_sig)
goeaobj.wr_txt("GO_analysis_%s.txt" % label, goea_results_sig)
```

## 1.1 Gene ontology

```
[4]: for direction in ["All", "Up", "Down"]:
         run_goea(direction)
    requests.get(http://purl.obolibrary.org/obo/go/go-basic.obo, stream=True)
      WROTE: go-basic.obo
    FTP RETR ftp.ncbi.nlm.nih.gov gene/DATA gene2go.gz -> gene2go.gz
      gunzip gene2go.gz
    go-basic.obo: fmt(1.2) rel(2021-08-18) 47,217 GO Terms
    HMS:0:00:04.253571 330,313 annotations, 20,685 genes, 18,684 GOs, 1 taxids READ:
    CC 19,422 annotated human genes
    BP 18,505 annotated human genes
    MF 18,190 annotated human genes
    Load BP Gene Ontology Analysis ...
     70% 20,236 of 29,107 population items found in association
    Load CC Gene Ontology Analysis ...
     74% 21,428 of 29,107 population items found in association
    Load MF Gene Ontology Analysis ...
     70\% 20,354 of 29,107 population items found in association
    Run BP Gene Ontology Analysis: current study set of 2759 IDs ... 81% 1,719 of
    2,111 study items found in association
     77% 2,111 of 2,759 study items found in population(29107)
    Calculating 12,429 uncorrected p-values using fisher
      12,429 GO terms are associated with 17,848 of 29,107 population items
       4,694 GO terms are associated with 1,719 of 2,759 study items
```

```
METHOD fdr_bh:
      11 GO terms found significant (< 0.05=alpha) ( 8 enriched +
purified): statsmodels fdr_bh
     159 study items associated with significant GO IDs (enriched)
      37 study items associated with significant GO IDs (purified)
Run CC Gene Ontology Analysis: current study set of 2759 IDs ... 86% 1,820 of
2,111 study items found in association
77% 2,111 of 2,759 study items found in population(29107)
Calculating 1,753 uncorrected p-values using fisher
   1,753 GO terms are associated with 18,711 of 29,107 population items
     838 GO terms are associated with 1,820 of 2,759 study items
 METHOD fdr_bh:
      39 GO terms found significant (< 0.05=alpha) ( 39 enriched +
purified): statsmodels fdr_bh
   1,696 study items associated with significant GO IDs (enriched)
       O study items associated with significant GO IDs (purified)
Run MF Gene Ontology Analysis: current study set of 2759 IDs ... 84% 1,768 of
2,111 study items found in association
77% 2,111 of 2,759 study items found in population(29107)
Calculating 4,420 uncorrected p-values using fisher
   4,420 GO terms are associated with 17,838 of 29,107 population items
   1,596 GO terms are associated with 1,768 of 2,759 study items
 METHOD fdr_bh:
      15 GO terms found significant (< 0.05=alpha) ( 13 enriched +
                                                                     2
purified): statsmodels fdr_bh
   1,452 study items associated with significant GO IDs (enriched)
       3 study items associated with significant GO IDs (purified)
Significant results[65] = 11 BP + 15 MF + 39 CC
     65 items WROTE: GO_analysis_allDEG.xlsx
     65 GOEA results for 1826 study items. WROTE: GO_analysis_allDEG.txt
 EXISTS: go-basic.obo
 EXISTS: gene2go
go-basic.obo: fmt(1.2) rel(2021-08-18) 47,217 GO Terms
HMS:0:00:04.541304 330,313 annotations, 20,685 genes, 18,684 GOs, 1 taxids READ:
gene2go
CC 19,422 annotated human genes
BP 18,505 annotated human genes
MF 18,190 annotated human genes
Load BP Gene Ontology Analysis ...
70% 20,236 of 29,107 population items found in association
Load CC Gene Ontology Analysis ...
74% 21,428 of 29,107 population items found in association
```

Load MF Gene Ontology Analysis ...

```
70% 20,354 of 29,107 population items found in association
```

```
Run BP Gene Ontology Analysis: current study set of 1337 IDs ... 79%
                                                                      760 of
967 study items found in association
        967 of 1,337 study items found in population(29107)
72%
Calculating 12,429 uncorrected p-values using fisher
  12,429 GO terms are associated with 17,848 of 29,107 population items
  2,805 GO terms are associated with 760 of 1,337 study items
 METHOD fdr bh:
       5 GO terms found significant (< 0.05=alpha) ( 4 enriched +
                                                                     1
purified): statsmodels fdr_bh
      48 study items associated with significant GO IDs (enriched)
       2 study items associated with significant GO IDs (purified)
Run CC Gene Ontology Analysis: current study set of 1337 IDs ... 84%
                                                                      813 of
967 study items found in association
        967 of 1,337 study items found in population(29107)
Calculating 1,753 uncorrected p-values using fisher
   1,753 GO terms are associated with 18,711 of 29,107 population items
     530 GO terms are associated with
                                         813 of 1,337 study items
 METHOD fdr bh:
      16 GO terms found significant (< 0.05=alpha) ( 16 enriched +
purified): statsmodels fdr_bh
     602 study items associated with significant GO IDs (enriched)
       O study items associated with significant GO IDs (purified)
Run MF Gene Ontology Analysis: current study set of 1337 IDs ... 82%
                                                                      793 of
967 study items found in association
        967 of 1,337 study items found in population(29107)
Calculating 4,420 uncorrected p-values using fisher
   4,420 GO terms are associated with 17,838 of 29,107 population items
     942 GO terms are associated with
                                      793 of 1,337 study items
 METHOD fdr_bh:
       8 GO terms found significant (< 0.05=alpha) ( 7 enriched +
purified): statsmodels fdr bh
     630 study items associated with significant GO IDs (enriched)
       1 study items associated with significant GO IDs (purified)
Significant results[29] = 5 BP + 8 MF + 16 CC
     29 items WROTE: GO_analysis_upregulated.xlsx
                          770 study items. WROTE: GO_analysis_upregulated.txt
     29 GOEA results for
 EXISTS: go-basic.obo
 EXISTS: gene2go
go-basic.obo: fmt(1.2) rel(2021-08-18) 47,217 GO Terms
HMS:0:00:04.466198 330,313 annotations, 20,685 genes, 18,684 GOs, 1 taxids READ:
gene2go
CC 19,422 annotated human genes
BP 18,505 annotated human genes
MF 18,190 annotated human genes
```

```
Load BP Gene Ontology Analysis ...
70% 20,236 of 29,107 population items found in association
Load CC Gene Ontology Analysis ...
74\% 21,428 of 29,107 population items found in association
Load MF Gene Ontology Analysis ...
 70% 20,354 of 29,107 population items found in association
Run BP Gene Ontology Analysis: current study set of 1423 IDs ... 84%
                                                                      960 of
1,145 study items found in association
80% 1,145 of 1,423 study items found in population(29107)
Calculating 12,429 uncorrected p-values using fisher
  12,429 GO terms are associated with 17,848 of 29,107 population items
   3,212 GO terms are associated with 960 of 1,423 study items
 METHOD fdr_bh:
      10 GO terms found significant (< 0.05=alpha) ( 7 enriched +
                                                                     3
purified): statsmodels fdr_bh
      81 study items associated with significant GO IDs (enriched)
      15 study items associated with significant GO IDs (purified)
Run CC Gene Ontology Analysis: current study set of 1423 IDs ... 88% 1,008 of
1,145 study items found in association
80% 1,145 of 1,423 study items found in population(29107)
Calculating 1,753 uncorrected p-values using fisher
   1,753 GO terms are associated with 18,711 of 29,107 population items
     657 GO terms are associated with 1,008 of 1,423 study items
 METHOD fdr bh:
      28 GO terms found significant (< 0.05=alpha) ( 28 enriched +
purified): statsmodels fdr_bh
     910 study items associated with significant GO IDs (enriched)
       O study items associated with significant GO IDs (purified)
Run MF Gene Ontology Analysis: current study set of 1423 IDs ... 85%
                                                                      976 of
1,145 study items found in association
80% 1,145 of 1,423 study items found in population(29107)
Calculating 4,420 uncorrected p-values using fisher
  4,420 GO terms are associated with 17,838 of 29,107 population items
   1,101 GO terms are associated with 976 of 1,423 study items
 METHOD fdr_bh:
      14 GO terms found significant (< 0.05=alpha) ( 13 enriched +
purified): statsmodels fdr_bh
     768 study items associated with significant GO IDs (enriched)
       O study items associated with significant GO IDs (purified)
Significant results[52] = 10 BP + 14 MF + 28 CC
     52 items WROTE: GO_analysis_downregulated.xlsx
     52 GOEA results for 1007 study items. WROTE: GO_analysis_downregulated.txt
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