RMF Investigation of GO-ONMF

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GO ONMF

Contact with Authors

As of Dec 1, 2015, a copy of this report was sent to Hwanjo Yu as the corresponding author listed on the manuscript for feedback, either in email, or ideally as issues or pull requests on this github repo.

Introduction

Kim, Sael and Yu, "A Mutation Profile for Top-k Patient Search Exploiting Gene-Ontology and Orthogonal Non-negative Matrix Factorization", Bioinformatics, 2015, http://dx.doi.org/10.1093/bioinformatics/btv409

Kim, Sael and Yu recently published the above paper in which they develop a method for creating a mutation-gene-ontology profile. As part of the overall procedure, they described (Section 2.3, *Gene Function Profile*):

To reduce the term correlation, we use only the most specific terms, i.e. the leaf node term after propagating the scores of the non-leaf terms down to the leaf node terms. This approach also resolves the problem of evaluating genes annotated with general term as the effect of the gene of function identification is spread out over several leaf node terms.

This sounds troubling to my ears, because the Gene Ontology (GO) is a directed acyclic graph (DAG), wherein there are specific relationships between the terms, and there is a directionality. There is also an expectation that when a gene product is annotated to a specific term, then that gene product is also automatically annotated to all of the less-specific parent terms in the DAG.

What the procedure quoted above sounds like it is doing is essentially **adding more specific** GO annotations to a gene product, which as far as I know is not allowed.

From The Gene Ontology Consortium, 2001:

The pathway from a child term to its top-level parent(s) must always be true.

This is why when considering gene-GO annotations, we consider the specific term annotation, and **all** parent terms as well, because of the **true path** rule. However, the converse is not true.

This document is a record of my investigation as to whether **more specific** GO annotations are being added to gene products during the creation of the mutation profiles.

Software

Kim and Yu made an Octave version of their software available at https://sites.google.com/site/postechdm/research/implementation/orgos, and a tarball of the Octave version can be downloaded from https://sites.google.com/site/postechdm/research/implementation/orgos/ONMF_octave-simple.zip

This code was used to fetch a copy of the software:

```
wget https://sites.google.com/site/postechdm/research/implementation/orgos/ONMF_octave-simple.zip
mkdir ONMF_source
unzip ONMF_octave-simple.zip -d ONMF_source
```

As part of the software package, a few data files are provided. The ones that are important for our purposes include:

- ONMF_source/brca/go_(merged).csv the index to GO term file
- ONMF_source/brca/gene_(merged).csv the index to gene file
- \bullet exp_onmf_brca.m the file that runs the *brca* analysis
- network_gene2go(merged).csv the gene 2 GO annotations

Running Code

After modifying $exp_onmf_brca.m$ to have the correct library path, and then adding some code that will save **gene - GO** indices before and after the optimization step (see below for the actual code run), I cd to the ONMF_source directory, and then start Octave, and then run:

```
cd brca
run exp_onmf_brca_mod.m
```

As this runs, it generates *gene2go.mat*, which is the propagated scores of gene to GO associations, and then saves indices (see file):

```
%% Orthogonal non-negative matrix factorization/Gene-Ontology-based stratification
library path = '/home/rmflight/Projects/personal/onmf/ONMF source/ONMF octave';
addpath(genpath(library path))
%% Convert somatic mutation cancer data to my data
patient2gene = load('network_patient2gene(merged).csv');
gene2go = load('network_gene2go(merged).csv');
go2go = load('network_allgo2allgo(merged).csv');
patient2gene = getSparse(patient2gene);
gene2go = getSparse(gene2go);
go2go = getSparse(go2go);
patient2gene(find(patient2gene)) = 1;
gene2go(find(gene2go)) = 1;
org_notzero = find(gene2go(1, :));
save -ascii org_notzero org_notzero;
[baseSMData, patient2gene, gene2go_new, go2go_new] = TuneData(patient2gene, gene2go, go2go);
isequal(go2go_new, go2go)
prop_notzero = find(gene2go_new(1, :));
save -ascii prop_notzero prop_notzero;
```

Prerequisites

To continue this analysis, you will need some other R packages installed.

```
install.packages(c("readr", "dplyr"))
library(BiocInstaller)
biocLite(c("GO.db", "graph", "org.Hs.eg.db"))
```

Load Packages

```
library(readr)
library(dplyr)
library(GO.db)
library(org.Hs.eg.db)
```

Check Actual Input Data

Lets check that the actual data being used as input looks right. Note that 1 gets added to the map indices so that we easily work with the data from the matrices in Octave. The input data includes the go_map, that maps GO terms to indices, gene_map mapping gene symbols to indices, gene2go mapping the GO annotations to genes, and go2go, giving the parent-child relationships of the GO terms.

And add the actual gene symbols and GO IDs to the gene2go to enable easy comparisons with any other data source we can find.

And then query org. Hs.eg.db for the gene-annotations as well.

Compare Gene-GO Annotations

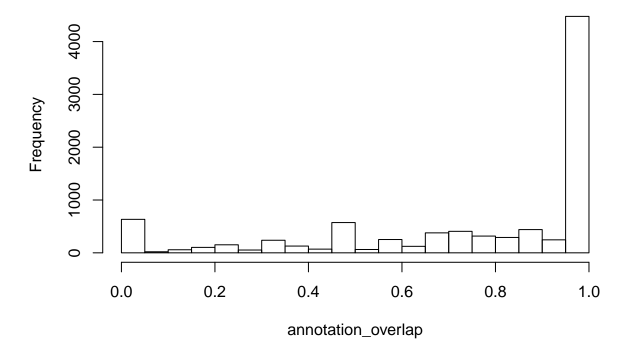
And compare the annotations for the genes between the data from Kim and from the Bioconductor database.

```
annotation_overlap <- vapply(unique(hs_gene2go$SYMBOL), function(in_loc){
  hs_data <- dplyr::filter(hs_gene2go, SYMBOL == in_loc) %>%
    dplyr::select(., GO) %>% unlist()
  onmf_data <- dplyr::filter(gene2go, geneid == in_loc) %>%
    dplyr::select(., goid) %>% unlist()
  sum(hs_data %in% onmf_data) / length(hs_data)
}, numeric(1))
```

And check the distribution of overlap:

```
hist(annotation_overlap)
```

Histogram of annotation_overlap



That is great, over 4000 of the genes have what appear to be very similar terms!

Compare GO-GO Relationships

Lets also check the ${\bf GO2GO}$ data.

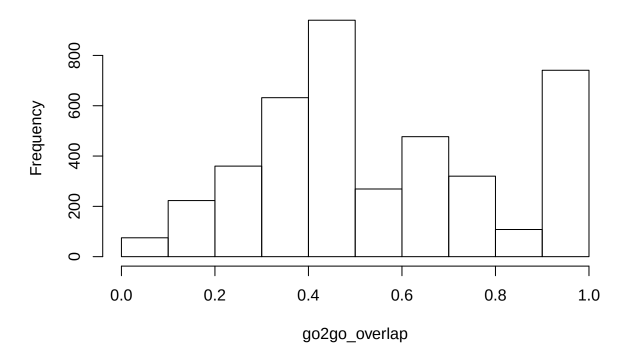
```
go_notself <- dplyr::filter(go2go, distance != 1)
go_notself$ID1 <- unlist(go_map[match(go_notself$G01, go_map$loc), "GO"], use.names = FALSE)
go_notself$ID2 <- unlist(go_map[match(go_notself$G02, go_map$loc), "GO"], use.names = FALSE)
onmf_pc <- split(go_notself$ID2, go_notself$ID1)
go_pc <- mget(names(onmf_pc), GOBPCHILDREN, ifnotfound = NA)</pre>
```

```
go2go_overlap <- vapply(names(onmf_pc), function(in_loc){
  go_data <- go_pc[[in_loc]]
  onmf_data <- onmf_pc[[in_loc]]
  sum(go_data %in% onmf_data) / length(go_data)
}, numeric(1))</pre>
```

And plot it:

```
hist(go2go_overlap)
```

Histogram of go2go_overlap



Work With Single Gene (A2M)

Now lets work with a single gene, and see what is going on. We'll keep it simple and play with the first one in the list, A2M.

Grab Starting and Propagated Terms

```
onmf_a2m_start <- dplyr::filter(gene2go, geneid == "A2M") %>%
   dplyr::select(goid) %>% unlist(., use.names = FALSE) %>% sort()
onmf_a2m_indices_org <- scan(file = "ONMF_source/brca/org_notzero", what = numeric(),</pre>
```

Compare To Database

Now, how many of the terms in the starting gene annotation actually in the database annotation?

```
hs_a2m_go <- dplyr::filter(hs_gene2go, SYMBOL == "A2M") %>%
    dplyr::select(GO) %>% unlist(., use.names = FALSE) %>% unique()
sum(onmf_a2m_matrix_org %in% hs_a2m_go) / length(onmf_a2m_matrix_org)
```

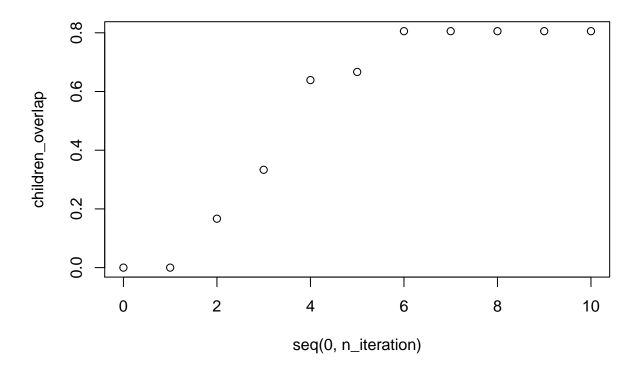
[1] 0.8

OK, seems we have most of them. Good.

Check if Propagated Are Children of Original

Next, are the propagated terms actually children of the starting terms? We will use multiple iterations of calls to GOBPCHILDREN and see if we get progressively more of the terms.

```
plot(seq(0, n_iteration), children_overlap)
```



FOUND THEM. The propagated terms have **NO** overlap with the original terms (see 0's at the start of the graph), and are children of the original terms. Just to be sure, I will also compare the propagated terms to the **GOALL** annotation, which would be all of the GO terms that can **technically** be considered as **annotated** to the gene (see *true path rule* from the Introduction), in that all **terms** and their **parent** terms are **annotated** to a gene product.

'select()' returned 1:many mapping between keys and columns

```
head(hs_a2m_all)
```

```
GOALL EVIDENCEALL ONTOLOGYALL
##
     SYMBOL
## 1
        A2M GO:0001775
                                  TAS
                                                ΒP
## 2
        A2M GO:0001867
                                  IDA
                                                ΒP
                                  IDA
## 3
        A2M GD:0001868
                                                ΒP
##
   4
        A2M GO:0001869
                                  IDA
                                                ΒP
## 5
        A2M GO:0002020
                                  IPI
                                                MF
## 6
        A2M GO:0002252
                                  IDA
                                                BP
```

```
sum(hs_a2m_all$GOALL %in% onmf_a2m_matrix_org) # the starting annotation
```

[1] 12

```
sum(hs_a2m_all$GOALL %in% onmf_a2m_matrix_prop) # propogated annotation
```

[1] 0

From this, none of the **propogated** terms can be properly considered as annotations to **A2M**. Effectively, the code has generated a **new set** of GO annotations to **A2M**. I am sure this is the situation for all of the genes.

Possible Solution

I wonder if a GOSlim would have fixed the problem of redundancy without introducing these new annotation relationships?