Identifying over-represented GO terms

In this notebook we will identify the over-represented GO terms in the proteins previously identified as changing RNA binding. The steps in this analysis are:

- 1. Model the relationship between differential RNA binding and an expected bias factor
- 2. Identify all the over-represented GO terms
- 3. Filter by FDR and degree of over-representation
- 4. Plot
- 5. Compare GO terms over-represented when using limma or lm

We need to source some functions which are stored in a script called "GO.R".

```
source("./GO.R")
```

Below, we load the result for differential RNA binding testing from the previous notebooks

```
limma_rna_binding_fit <- readRDS("../results/limma_rna_binding_fit.rds")
compare_methods <- readRDS("../results/compare_methods_rna_binding_results.rds")</pre>
```

To perform the differential RNA binding testing, we will use the R package goseq. This was orignally designed to account for length bias in RNA-Seq count-based differential expression testing but can be applied to any GO over-representation scenario where one expects there is a bias due to increased power to detect changes for some features. The basic idea is that the bias factor must have a monotonic relationship with the probability of a feature presenting as significantly changed. goseq estimates this relationship with a spline fit (we will see this later). goseq then takes this relationship into account when performing the GO over-representation testing. For more details about goseq, see here: https://genomebiology.biomedcentral.com/articles/10.1186/gb-2010-11-2-r14

In our case, we have reason to suspect that proteins with higher intensity will be more likely to be identified as having significantly altered RNA binding since the variance will be smaller so we will have more power to detect differences.

The first step is to make a probability weight function (PWF) linking the bias factor with the probability of significant difference using the nullp function.

For this, we need a boolean to indicate which proteins we identify as having a significant change and a vector of the same length which details our bias factor.

The limma output contains a "AveExpr" value for each protein which represents the mean intensity. This is precisely the bias factor we are after so we can use this directly.

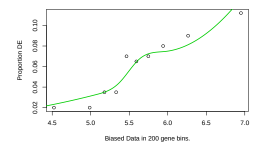
```
bias <- compare_methods$AveExpr</pre>
print(length(bias))
## [1] 1916
For the booleans, we can extract this from our results tables. We'll set our thresholds at 1% FDR and >=
1.4 fold change. Note that these are completely arbitrary thresholds of course!
limma_rna_binding_fit_treat <- treat(limma_rna_binding_fit, lfc=log2(1.4))</pre>
limma_rna_binding_changes <- topTreat(limma_rna_binding_fit_treat, coef = "conditionG1:type00PS", n = I</pre>
                                        p.value=0.01, adjust.method="BH", confint=0.95)
limma_rna_binding_changes <- limma_rna_binding_changes[limma_rna_binding_changes$logFC>log2(1.5),]
limma_sig_bool <- compare_methods$Row.names %in% rownames(limma_rna_binding_changes)
names(limma_sig_bool) <- compare_methods$Row.names</pre>
print(table(limma_sig_bool))
## limma_sig_bool
## FALSE TRUE
  1806
           110
lm_sig_bool <- (compare_methods$lm_BH < 0.01 & compare_methods$logFC>log2(1.4))
names(lm_sig_bool) <- compare_methods$Row.names</pre>
print(table(lm_sig_bool))
## lm_sig_bool
## FALSE TRUE
    1597
           319
##
```

5.1 Question: Why do we have fewer proteins with an increase in RNA binding according to limma?

By default, the function expects a genome name and it will go and fetch the bias data. However, we can also just provide this data ourselves. Below, we obtain the PWF and goseq outputs plots to show how closely these fit the observed relationship.

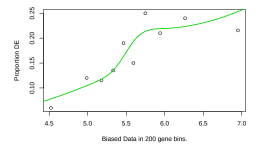
```
limma_pwf <- nullp(limma_sig_bool, bias.data=bias, plot.fit=TRUE)</pre>
```

Warning in pcls(G): initial point very close to some inequality constraints



```
lm_pwf <- nullp(lm_sig_bool, bias.data=bias, plot.fit=TRUE)</pre>
```

Warning in pcls(G): initial point very close to some inequality constraints



Now, we need those GO terms from the previous notebook

```
sapiens.go.full <- readRDS("../results/h_sapiens_go_full.rds")
head(sapiens.go.full)</pre>
```

##		UNIPROTKB	GO.ID			TERM	${\tt ONTOLOGY}$
##	1	P09874	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP
##	2	Q9UJZ1	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP
##	3	Q9Y243	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP
##	4	Q02078	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP
##	5	Q9BUK6	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP
##	6	Q96RR1	GD:0000002	${\tt mitochondrial}$	genome	${\tt maintenance}$	BP

And we're ready to run goseq. By default, goseq expects a genome and will obtain the GO terms automatically. However, if we provide our own to the gene2cat argument, it will use this instead. This also means we could use goseq to interrogate over-represented KEGG terms, interpro domains, etc by simply providing them at this point.

- ## Using manually entered categories.
- ## For 14 genes, we could not find any categories. These genes will be excluded.
- ## To force their use, please run with use genes without cat=TRUE (see documentation).
- ## This was the default behavior for version 1.15.1 and earlier.
- ## Calculating the p-values...
- ## 'select()' returned 1:1 mapping between keys and columns
- ## Using manually entered categories.
- ## For 14 genes, we could not find any categories. These genes will be excluded.
- ## To force their use, please run with use genes without cat=TRUE (see documentation).
- ## This was the default behavior for version 1.15.1 and earlier.
- ## Calculating the p-values...
- ## 'select()' returned 1:1 mapping between keys and columns

We need to adjust the p-values to account for multiple testing. Again, we'll use the Benjamini-Hochberg method.

```
limma_over_rep_go$BH <- p.adjust(limma_over_rep_go$over_represented_pvalue, method="BH")
lm_over_rep_go$BH <- p.adjust(lm_over_rep_go$over_represented_pvalue, method="BH")</pre>
```

```
5.2 Question: How many GO terms did we test for over-representation in the 'limma' results?
5.3 Question: How many would we expect to have a p-value < 0.05 by chance?</li>
5.4 Question: How many did have a p-value < 0.05?</li>
5.5 Question: What can you infer from this?
```

We can filter the output to only include the GO terms which are over-represented at 1% FDR (there are 112 for limma)

```
sig_terms <- limma_over_rep_go %>% filter(BH<0.01)</pre>
print(dim(sig_terms))
## [1] 112
print(head(sig_terms))
        category over_represented_pvalue under_represented_pvalue numDEInCat
                                4.028782e-24
## 1 GO:0044281
## 2 GO:0055114
                                1.958873e-19
                                                                             1
                                                                                         32
## 3 GD:0044429
                                3.100699e-19
                                                                             1
                                                                                         41
## 4 GO:0019752
                                4.011556e-18
                                                                             1
                                                                                         46
## 5 GO:0043436
                                4.942581e-18
                                                                             1
                                                                                         46
## 6 GD:0006082
                                6.292715e-18
                                                                                         46
     numInCat
                                                    term ontology
## 1
            310 small molecule metabolic process BP 4.769272e-20
## 2
            83
                  oxidation-reduction process
                                                               BP 1.159457e-15
           mitochondrial part CC 1.223536e-15

203 carboxylic acid metabolic process BP 1.170206e-14

204 oxoacid metabolic process BP 1.170206e-14

205 organic acid metabolic process BP 1.241553e-14
## 3
## 4
## 5
## 6
```

We would also like to know the effect size, e.g how over-represented are the terms. We could use the numDEInCat and numInCat columns and calculate the over-representation as:

(numInCat/ Number of proteins observed) / (numDEInCat/ Number of proteins with change in RNA binding).

5.6 Question: Why would it be sub-optimal to calculate the magnitude of the over-representation as indicated above?

The function below estimates the effect size, taking into account the bias (using the PWF)

```
term_features <- gene2cat[gene2cat[["GO.ID"]]==x[["category"]]], "UNIPROTKB"]</pre>
    term_weight <- mean(pwf[rownames(pwf) %in% term_features, "pwf"])</pre>
    non_term_weight <- mean(pwf[!rownames(pwf) %in% term_features, "pwf"])</pre>
    as.numeric(x[['numDEInCat']])/as.numeric(x[['numInCat']]) / (term_weight/non_term_weight) / (len_fo
  return(obj)
}
Now we can filter by the estimated over-representation too.
limma_over_rep_go %>% filter(BH<0.01) %>%
  addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
  filter(adj_over_rep>2) %>%
 head()
##
       category over_represented_pvalue under_represented_pvalue numDEInCat
## 1 GO:0055114
                            1.958873e-19
## 2 GO:0045333
                            4.456564e-11
                                                                            15
                                                                  1
## 3 GD:0015980
                            5.051024e-10
                                                                  1
                                                                            16
## 4 GD:0009060
                            7.197845e-09
                                                                  1
                                                                            11
## 5 GO:0004812
                            5.568684e-08
                                                                  1
                                                                            11
## 6 GD:0016875
                            5.568684e-08
                                                                  1
                                                                            11
     numInCat
##
                                                                term ontology
           83
## 1
                                        oxidation-reduction process
                                                                           BP
## 2
                                                                           ΒP
                                               cellular respiration
## 3
           46 energy derivation by oxidation of organic compounds
                                                                           BP
## 4
           22
                                                aerobic respiration
                                                                           BP
## 5
           24
                                    aminoacyl-tRNA ligase activity
                                                                           MF
## 6
           24
                      ligase activity, forming carbon-oxygen bonds
                                                                           MF
               BH adj_over_rep
                       2.195040
## 1 1.159457e-15
## 2 4.058216e-08
                       2.608945
                       2.102115
## 3 3.737126e-07
## 4 3.873095e-06
                      2.831174
## 5 2.354360e-05
                      2.400932
## 6 2.354360e-05
                      2.400932
lm_over_rep_go %>% filter(BH<0.01) %>%
  addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
  filter(adj_over_rep>2) %>%
       category over_represented_pvalue under_represented_pvalue numDEInCat
## 1 GO:0044281
                            4.514926e-33
                                                                           133
## 2 GD:0043436
                            9.437890e-27
                                                                  1
                                                                            97
## 3 GD:0006082
                            1.644155e-26
                                                                  1
                                                                            97
## 4 GO:0019752
                                                                            96
                            3.248454e-26
                                                                  1
## 5 GO:0055114
                            4.699760e-21
                                                                            52
                                                                  1
## 6 GD:0044429
                            2.148131e-19
                                                                            73
     numInCat
##
                                             term ontology
                                                                      BH adj_over_rep
## 1
          310 small molecule metabolic process
                                                        BP 5.344769e-29
                                                                             2.493771
## 2
          204
                       oxoacid metabolic process
                                                        BP 5.586287e-23
                                                                             2.715729
## 3
          205
                 organic acid metabolic process
                                                        BP 6.487837e-23
                                                                             2.699929
## 4
          203 carboxylic acid metabolic process
                                                        BP 9.613801e-23
                                                                             2.701236
## 5
                    oxidation-reduction process
                                                        BP 9.272627e-18
                                                                             3.566940
           83
```

Note that many of the above GO terms are essentially redundant and are simply the same set of proteins with each of the GO terms up the heirachy being over-represented. We can perform a simplification by taking each over-represented GO term, identifying all its offspring and ancesters and removing it if any of the terms above or below are more significantly over-represented.

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Below, I've wrapped this all up into a function so that we can pass a data frame of significantly over-represented GO terms and it will remove the redundant terms.

```
remove redundant GO terms <- function(go df){
  all_observed_go <- unique(go_df$category) # identify all the GO terms</pre>
  all_observed_go <- all_observed_go[!is.na(all_observed_go)] # Remove any NAs
  # Get the ontologies for the GO terms
  ontologies <- AnnotationDbi::select(GO.db, all_observed_go, columns = c('ONTOLOGY'), keytype='GOID')
  ontologies <- setNames(ontologies$ONTOLOGY, ontologies$GOID)</pre>
  # Get the mappings from GO term to parent GO terms using functions in GO.R
  go20ffspring <- getAllMappings(all_observed_go, ontologies, verbose=FALSE, direction="offspring")
  go2Ancesters <- getAllMappings(all_observed_go, ontologies, verbose=FALSE, direction="ancester")
  # start with all observed GO terms being retained
  retained <- all_observed_go
  # We want to keep track of the GO IDs we have processed
  processed <- NULL
  # If any GO term has no detected offspring or ancesters, mark them as already processed
  # This will also mean they are always retained
  no_anc_off <- setdiff(all_observed_go, union(names(go2Ancesters), names(go2Offspring)))</pre>
  if(length(no_anc_off)>0){
    cat(sprintf("No offspring or ancesters could be found for these terms: %s", no_anc_off))
   processed <- no_anc_off</pre>
  }
  # When all observed go terms are in processed, stop while loop
  while(length(setdiff(all_observed_go, processed))!=0){
    # take one of the GO terms
   go_id <- setdiff(all_observed_go, processed)[1]</pre>
    # Find all the ancesters and offspring = go_tree
    # (Only include those also observed as over-rep GO)
    go_tree <- union(go2Ancesters[[go_id]], go2Offspring[[go_id]]) %>%
      intersect(all_observed_go) %>% c(go_id)
   top_go <- go_df %>%
      filter(category %in% go_tree) %>% # subset to the terms in go_tree
      arrange(over_represented_pvalue) %>% # order by p-value (ascending by default)
      pull(category) %>%# pull out the category
      head(1) # keep the top GO term
    # We want to remove all ancester and offspring terms within the go_tree for the top GO term
```

```
terms_to_remove <- union(go2Ancesters[[top_go]], go2Offspring[[top_go]]) %>%
    intersect(go_tree)

processed <- union(processed, go_tree) # all terms in the tree are now considered "processed"
    retained <- setdiff(retained, terms_to_remove) # remove the unwanted terms from retained
}

go_df <- go_df %>% filter(category %in% retained) # subset to the retained terms
    return(go_df)
}

5.7 Task: Modify the remove_redundant_GO_terms function to keep an arbitrary number of top GO terms
within each sub tree of GO terms

limma_over_rep_go %>% filter(BH<0.01) %>%
    addAdjustedOverRep(limma_pwf, sapiens.go.full) %>%
```

```
filter(adj over rep>3) %>%
 remove_redundant_GO_terms() %>%
 head()
## 'select()' returned 1:1 mapping between keys and columns
       category over represented pvalue under represented pvalue numDEInCat
## 1 GO:0044281
                           4.028782e-24
                                                                 1
                                                                           63
## 2 GO:0055114
                           1.958873e-19
                                                                 1
                                                                           32
## 3 GD:0044429
                           3.100699e-19
                                                                 1
                                                                           41
## 4 GD:0006091
                           1.414456e-13
                                                                 1
                                                                           27
## 5 GO:0017144
                           5.526715e-12
                                                                           30
                                                                 1
## 6 GD:0016491
                           6.280439e-11
                                                                 1
                                                                           25
     numInCat
                                                                                  BH
##
                                                         term ontology
## 1
          310
                            small molecule metabolic process
                                                                     BP 4.769272e-20
## 2
          83
                                  oxidation-reduction process
                                                                     BP 1.159457e-15
## 3
          159
                                           mitochondrial part
                                                                    CC 1.223536e-15
## 4
           92 generation of precursor metabolites and energy
                                                                     BP 1.860482e-10
## 5
          128
                                       drug metabolic process
                                                                    BP 5.947750e-09
## 6
           95
                                      oxidoreductase activity
                                                                    MF 5.310560e-08
##
     adj_over_rep
## 1
         3.494409
## 2
         6.204881
## 3
         4.640499
         5.018165
## 4
## 5
         3.918863
## 6
         4.245300
lm_over_rep_go %>% filter(BH<0.01) %>%
 addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
  filter(adj_over_rep>3) %>%
 remove_redundant_GO_terms() %>%
 head()
```

'select()' returned 1:1 mapping between keys and columns
category over_represented_pvalue under_represented_pvalue numDEInCat

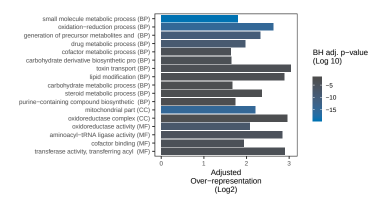
```
## 1 GO:0055114
                            4.699760e-21
                                                                             52
                                                                  1
                            2.495342e-14
## 2 GD:0044282
                                                                             37
                                                                  1
## 3 GD:0004812
                            1.039095e-11
                                                                  1
                                                                             20
## 4 GD:0006418
                            2.455402e-10
                                                                             20
                                                                  1
## 5 GO:0005743
                            2.561231e-09
                                                                  1
                                                                             26
## 6 GD:0006734
                            6.737147e-08
                                                                             12
                                                                  1
     numInCat
                                                        term ontology
## 1
           83
                               oxidation-reduction process
                                                                   BP 9.272627e-18
## 2
           62
                          small molecule catabolic process
                                                                   BP 1.969324e-11
           24
## 3
                            aminoacyl-tRNA ligase activity
                                                                   MF 6.150401e-09
           26 tRNA aminoacylation for protein translation
                                                                   BP 7.855960e-08
                                                                   CC 6.890877e-07
## 5
           48
                              mitochondrial inner membrane
## 6
           13
                                    NADH metabolic process
                                                                   BP 1.035771e-05
##
     adj_over_rep
## 1
         3.566940
## 2
         3.440306
## 3
         4.365332
## 4
         3.928257
## 5
         3.221653
## 6
         4.298834
```

OK so now we're getting a more useful set of terms. The next thing we might want to do is some basic plot to show the result. Again, I've wrapped this up into a a function below so we can pass the GO dataframe straight through the above pipe to filter the results and then plot the filtered results.

```
plotTerms <- function(go_df,</pre>
                      horizontal=FALSE, # make plot hoizontal
                      plot top=10, # plot the top n most significant GO terms
                      shorten_term=FALSE){ # shorten the term to max 30 char
  # re-order data frame by p-value
  if(horizontal){
    go_df <- go_df %>% arrange(ontology, over_represented_pvalue)
    go_df <- go_df %% head(plot_top) # subset to top n most significant terms</pre>
  else{
    go_df <- go_df %>% arrange(desc(ontology), desc(over_represented_pvalue))
    go_df <- go_df %>% tail(plot_top) # subset to top n most significant terms
  if(shorten term){
    go_df$term_for_plot <- substr(go_df$term, 1, 40) # cut at character 40</pre>
  }
 else{
    go_df$term_for_plot <- go_df$term</pre>
  # add the ontology (BP, MF, CC) to the end of the term
  go_df$term_for_plot <- paste0(go_df$term_for_plot, " (", go_df$ontology, ")")</pre>
  # re-level factor make keep plotting order in order of dataframe (ontology, p-value)
  go_df$term_for_plot <- factor(go_df$term_for_plot, levels=rev(go_df$term_for_plot))</pre>
  p <- go_df %>%
    ggplot(aes(x=term_for_plot, y=log(adj_over_rep,2), fill=log(BH,10))) +
```

```
geom_bar(stat="identity") + # When geom_bar is plotting a single data point, need to set stat="iden
    xlab("") +
    ylab("Adjusted\nOver-representation\n(Log2)") +
    scale_fill_continuous(name="BH adj. p-value\n(Log 10)\n", low=cbPalette[5], high="grey30", limits=c
    theme(text=element_text(size=15),
          plot.title=element_text(hjust=0.5))
  if(horizontal){
    p <- p + coord_flip() # Flip the coordinates</pre>
  else{
    # If vertical bards, set the x-axis text at an angle so it fits better
    p <- p + theme(axis.text.x=element_text(size=12, angle=30, vjust=1, hjust=1))
  return(list("p"=p, "data"=go_df))
}
limma_over_rep_go %>% filter(BH<0.01) %>%
  addAdjustedOverRep(limma_pwf, sapiens.go.full) %>%
  filter(adj_over_rep>3) %>%
  remove_redundant_GO_terms() %>%
  plotTerms(horizontal=TRUE, shorten_term=TRUE, plot_top=20)
```

'select()' returned 1:1 mapping between keys and columns
\$p

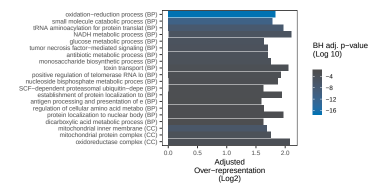


```
##
## $data
##
        category over_represented_pvalue under_represented_pvalue numDEInCat
     GO:0044281
                            4.028782e-24
                                                         1.0000000
                                                                            63
## 1
                                                                            32
## 2
     GO:0055114
                            1.958873e-19
                                                         1.0000000
## 3
     GD:0006091
                            1.414456e-13
                                                         1.0000000
                                                                            27
## 4 GO:0017144
                            5.526715e-12
                                                         1.0000000
                                                                            30
## 5 GO:0051186
                            1.245389e-06
                                                         0.9999997
                                                                            21
## 6
     GO:1901137
                            1.910455e-06
                                                         0.999996
                                                                            20
                            2.425164e-06
## 7 GO:1901998
                                                         0.999999
                                                                             7
```

```
## 8 GD:0030258
                             4.666279e-06
                                                          0.9999997
                                                                              8
## 9 GD:0005975
                             1.073440e-05
                                                          0.9999976
                                                                             17
## 10 GO:0008202
                             1.109125e-05
                                                          0.9999987
                                                                             10
## 11 GO:0072522
                             1.876277e-05
                                                          0.9999961
                                                                             15
## 12 GO:0044429
                             3.100699e-19
                                                          1.0000000
                                                                             41
## 13 GO:1990204
                             1.136036e-05
                                                          0.999993
                                                                              7
## 14 GO:0016491
                             6.280439e-11
                                                          1.0000000
                                                                             25
## 15 GO:0004812
                             5.568684e-08
                                                          1.0000000
                                                                             11
## 16 GD:0048037
                             3.662326e-07
                                                          0.9999999
                                                                             18
## 17 GD:0016747
                             1.009188e-06
                                                          0.9999999
                                                                              9
      numInCat
## 1
           310
## 2
            83
## 3
            92
## 4
           128
## 5
           115
## 6
           107
## 7
            10
## 8
            21
## 9
            89
## 10
            35
## 11
            71
## 12
           159
## 13
            16
## 14
            95
## 15
            24
## 16
            85
## 17
            23
##
                                                                                 term
                                                   small molecule metabolic process
## 1
## 2
                                                         oxidation-reduction process
## 3
                                    generation of precursor metabolites and energy
## 4
                                                              drug metabolic process
## 5
                                                         cofactor metabolic process
## 6
                                      carbohydrate derivative biosynthetic process
## 7
                                                                     toxin transport
## 8
                                                                  lipid modification
## 9
                                                     carbohydrate metabolic process
## 10
                                                           steroid metabolic process
## 11
                                   purine-containing compound biosynthetic process
## 12
                                                                  mitochondrial part
## 13
                                                             oxidoreductase complex
## 14
                                                             oxidoreductase activity
## 15
                                                     aminoacyl-tRNA ligase activity
## 16
                                                                    cofactor binding
   17 transferase activity, transferring acyl groups other than amino-acyl groups
##
      ontology
                          BH adj_over_rep
## 1
            BP 4.769272e-20
                                 3.494409
## 2
            BP 1.159457e-15
                                 6.204881
## 3
            BP 1.860482e-10
                                 5.018165
## 4
            BP 5.947750e-09
                                 3.918863
## 5
            BP 2.541882e-04
                                 3.113399
## 6
            BP 3.627390e-04
                                 3.145541
## 7
            BP 4.284939e-04
                                 8.233090
```

```
## 8
            BP 7.672140e-04
                                 7.426975
## 9
            BP 1.588422e-03
                                 3.177366
## 10
            BP 1.606653e-03
                                 5.153704
            BP 2.524019e-03
## 11
                                 3.335070
## 12
            CC 1.223536e-15
                                 4.640499
            CC 1.620289e-03
                                 7.800007
## 13
            MF 5.310560e-08
                                 4.245300
## 14
            MF 2.354360e-05
## 15
                                 7.172311
## 16
            MF 9.224386e-05
                                 3.832178
            MF 2.203678e-04
## 17
                                 7.483026
##
                                       term_for_plot
              small molecule metabolic process (BP)
##
  1
##
  2
                   oxidation-reduction process (BP)
      generation of precursor metabolites and
##
  3
## 4
                         drug metabolic process (BP)
## 5
                    cofactor metabolic process (BP)
## 6
      carbohydrate derivative biosynthetic pro (BP)
## 7
                                toxin transport (BP)
## 8
                             lipid modification (BP)
## 9
                carbohydrate metabolic process (BP)
## 10
                      steroid metabolic process (BP)
      purine-containing compound biosynthetic
## 11
                             mitochondrial part (CC)
## 12
## 13
                         oxidoreductase complex (CC)
## 14
                        oxidoreductase activity (MF)
## 15
                aminoacyl-tRNA ligase activity (MF)
## 16
                               cofactor binding (MF)
## 17 transferase activity, transferring acyl
                                                (MF)
lm_over_rep_go %>% filter(BH<0.01) %>%
  addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
  filter(adj_over_rep>3) %>%
  remove_redundant_GO_terms() %>%
  plotTerms(horizontal=TRUE, shorten_term=TRUE, plot_top=20)
```

'select()' returned 1:1 mapping between keys and columns
\$p



```
##
## $data
##
        category over_represented_pvalue under_represented_pvalue numDEInCat
     GO:0055114
                             4.699760e-21
                                                           1.0000000
## 1
   2
      GO:0044282
                             2.495342e-14
                                                           1.0000000
                                                                              37
## 3
      GO:0006418
                             2.455402e-10
                                                           1.0000000
                                                                              20
      GD:0006734
                             6.737147e-08
                                                           1.0000000
                                                                              12
                                                                              21
      GD:0006006
## 5
                             1.455207e-07
                                                           1.0000000
## 6
      GD:0033209
                             4.392061e-07
                                                           0.999999
                                                                              18
## 7
                                                                              18
      GD:0016999
                             5.618775e-07
                                                           0.999999
## 8
      GD:0046364
                             6.939182e-07
                                                           0.999999
                                                                              16
                                                                               9
## 9
      GO:1901998
                             6.388143e-06
                                                           0.999998
## 10 GO:1904874
                             7.699857e-06
                                                           0.9999996
                                                                              10
## 11 GO:0033865
                             1.516983e-05
                                                           0.9999981
                                                                              12
## 12 GO:0031146
                             2.457018e-05
                                                           0.9999961
                                                                              14
## 13 GD:0070199
                             2.482724e-05
                                                           0.9999988
                                                                               9
## 14 GD:0002479
                             2.993548e-05
                                                                              14
                                                           0.9999951
## 15 GO:0006521
                             3.676311e-05
                                                           0.9999945
                                                                              13
## 16 GO:1903405
                             3.785542e-05
                                                           0.9999989
                                                                               8
## 17 GD:0043648
                             5.548052e-05
                                                           0.9999909
                                                                              13
## 18 GO:0005743
                             2.561231e-09
                                                           1.0000000
                                                                              26
## 19 GO:0098798
                             1.044075e-06
                                                           0.999998
                                                                              17
## 20 GO:1990204
                             3.925597e-06
                                                           0.999997
                                                                              11
      numInCat
## 1
            83
## 2
            62
## 3
            26
## 4
            13
## 5
            38
## 6
            31
## 7
            32
## 8
            25
## 9
            10
## 10
            12
## 11
            21
## 12
            26
## 13
            11
## 14
            26
## 15
            23
             9
## 16
## 17
            25
## 18
            48
## 19
            32
## 20
            16
##
                                                                                                       term
## 1
                                                                              oxidation-reduction process
## 2
                                                                         small molecule catabolic process
## 3
                                                             tRNA aminoacylation for protein translation
## 4
                                                                                   NADH metabolic process
## 5
                                                                                 glucose metabolic process
## 6
                                                        tumor necrosis factor-mediated signaling pathway
## 7
                                                                             antibiotic metabolic process
## 8
                                                                      monosaccharide biosynthetic process
## 9
                                                                                           toxin transport
```

```
## 10
                                      positive regulation of telomerase RNA localization to Cajal body
## 11
                                                              nucleoside bisphosphate metabolic process
## 12
                               SCF-dependent proteasomal ubiquitin-dependent protein catabolic process
## 13
                                                    establishment of protein localization to chromosome
  14
      antigen processing and presentation of exogenous peptide antigen via MHC class I, TAP-dependent
                                                    regulation of cellular amino acid metabolic process
## 15
## 16
                                                                   protein localization to nuclear body
## 17
                                                                    dicarboxylic acid metabolic process
## 18
                                                                            mitochondrial inner membrane
## 19
                                                                           mitochondrial protein complex
## 20
                                                                                  oxidoreductase complex
##
      ontology
                          BH adj_over_rep
## 1
            BP 9.272627e-18
                                 3.566940
## 2
            BP 1.969324e-11
                                 3.440306
## 3
            BP 7.855960e-08
                                 3.928257
## 4
            BP 1.035771e-05
                                 4.298834
## 5
            BP 1.980085e-05
                                 3.111141
## 6
            BP 5.147843e-05
                                 3.268283
## 7
            BP 6.521084e-05
                                 3.260468
## 8
            BP 7.975343e-05
                                 3.377015
                                 4.160712
## 9
            BP 5.817141e-04
## 10
            BP 6.858492e-04
                                 3.813483
## 11
            BP 1.221635e-03
                                 3.662603
            BP 1.825496e-03
## 12
                                 3.083310
                                 3.853160
## 13
            BP 1.825496e-03
## 14
            BP 2.134797e-03
                                 3.026172
## 15
            BP 2.441537e-03
                                 3.105898
            BP 2.441537e-03
## 16
                                 3.927636
            BP 3.333900e-03
## 17
                                 3.086860
## 18
            CC 6.890877e-07
                                 3.221653
## 19
            CC 1.166015e-04
                                 3.376489
## 20
            CC 3.809116e-04
                                 4.232450
##
                                       term_for_plot
## 1
                   oxidation-reduction process (BP)
##
              small molecule catabolic process (BP)
## 3
      tRNA aminoacylation for protein translat (BP)
## 4
                         NADH metabolic process (BP)
## 5
                     glucose metabolic process (BP)
      tumor necrosis factor-mediated signaling (BP)
## 7
                  antibiotic metabolic process (BP)
## 8
           monosaccharide biosynthetic process (BP)
## 9
                                toxin transport (BP)
## 10 positive regulation of telomerase RNA lo (BP)
## 11 nucleoside bisphosphate metabolic proces (BP)
## 12 SCF-dependent proteasomal ubiquitin-depe (BP)
## 13 establishment of protein localization to (BP)
      antigen processing and presentation of e (BP)
## 15
      regulation of cellular amino acid metabo (BP)
## 16
          protein localization to nuclear body (BP)
## 17
           dicarboxylic acid metabolic process (BP)
```

mitochondrial inner membrane (CC)

oxidoreductase complex (CC)

mitochondrial protein complex (CC)

18

19

20

Ok, so now we have a reasonable list of over-represented GO terms and some pretty plots to show off our results...

5.8 Task: Modify the plotTerms function so that it makes a separate plot for each ontology

One thing you may have noticed in the above is that the over-represented terms are slightly different for 1m and limma above. For example, "tRNA aminoacylation for protein translation" is one of the most significantly overrepresented MF GO term in the 1m proteins but not present in the 1imma over-represented GO terms (although "aminoacyl-tRNA ligase activity" is there in the MF terms)

Below, we take the GO term "tRNA aminoacylation for protein translation". Note that very few of the proteins annotated with "tRNA aminoacylation for protein translation" are detected as having a significant change in RNA binding according to limma - 11/26 vs 20/26 for lm!!

```
lm_over_rep_go %% filter(term=="tRNA aminoacylation for protein translation") %%
  addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
 head(10)
##
       category over_represented_pvalue under_represented_pvalue numDEInCat
## 1 GD:0006418
                            2.455402e-10
##
     numInCat
                                                                               BH
                                                       term ontology
## 1
           26 tRNA aminoacylation for protein translation
                                                                  BP 7.85596e-08
##
     adj_over_rep
## 1
         3.928257
limma_over_rep_go %>% filter(term=="tRNA aminoacylation for protein translation") %>%
  addAdjustedOverRep(lm_pwf, sapiens.go.full) %>%
 head(10)
##
       category over_represented_pvalue under_represented_pvalue numDEInCat
## 1 GO:0006418
                            2.046793e-07
     numInCat
##
                                                       term ontology
                                                                                BH
## 1
           26 tRNA aminoacylation for protein translation
                                                                  BP 6.376299e-05
##
     adj_over_rep
         2.160541
## 1
We want to take a look at the intensity values, so let's make that plotIntensities() function again
combined_intensities <- readRDS("../results/combined_intensities.rds")</pre>
plotIntensities <- function(obj){</pre>
  p <- tidy(obj, addPheno=TRUE) %>%
    ggplot(aes(Condition, value, colour=Type, group=Type)) +
    geom_point() +
    stat_summary(geom="line", fun.y=mean) +
    facet_wrap(~gene, scales='free') +
    ylab("Intensity (log2)")
  invisible(p)
}
```

If we take a look at the intensity values for these 26 proteins annotated with "tRNA aminoacylation for protein translation", most do seem to have a clear increase in RNA binding. Note that not all of these are cytosolic tRNA ligases. Q5JPH6 + Q5JTZ9 are mitchondrial tRNA ligase. Q15181=Inorganic pyrophosphatase and Q12904 & Q13155 interact with the Aminoacyl tRNA synthase complex.

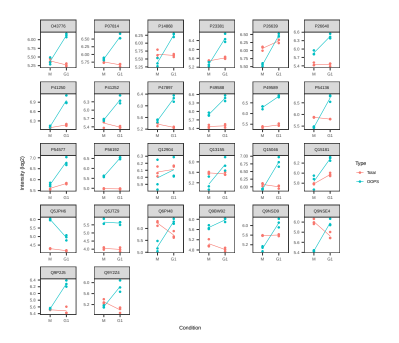
So, why does limma only detect 11/26 of these as increasing RNA binding given the apparent significant

changes...

```
tRNA_aa <- sapiens.go.full %>%
  filter(TERM=="tRNA aminoacylation for protein translation") %>%
  pull(UNIPROTKB)

tRNA_aa_intensities <- combined_intensities[intersect(rownames(combined_intensities), tRNA_aa),] %>%
  plotIntensities()

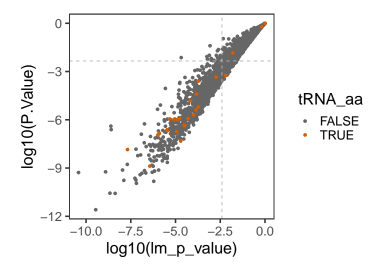
print(tRNA_aa_intensities + theme(text=element_text(size=10)))
```



Next, we compare the p-values again. Note that the four proteins not in the bottom left quadrant (e.g <1% FDR with both 1m and 1imma) are the two Mt tRNA-ligases and the two aminoacyl tRNA synthase complex interactors.

```
max_p_sig_lm <- compare_methods %>% filter(lm_BH<0.01) %>% pull(lm_p_value) %>% max()
max_p_sig_limma <- compare_methods %>% filter(adj.P.Val<0.01) %>% pull(P.Value) %>% max()

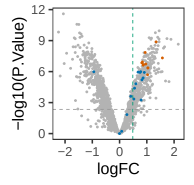
compare_methods %>%
  mutate(tRNA_aa=Row.names %in% tRNA_aa) %>%
  arrange(tRNA_aa) %>%
  ggplot(aes(x=log10(lm_p_value), y=log10(P.Value), colour=tRNA_aa)) +
  geom_point() +
  scale_colour_manual(values=c("grey40", cbPalette[6])) +
  geom_vline(xintercept=log10(max_p_sig_lm), linetype=2, colour="grey70") +
  geom_hline(yintercept=log10(max_p_sig_limma), linetype=2,colour="grey70")
```



Finally, we look at the point estimates for the fold change. The green dashed lines represent our 1.4-fold threshold. The dashed grey-line is the threshold for the BH adjusted p-value being < 0.01.

```
max_p_sig_limma <- compare_methods %>% filter(adj.P.Val<0.01) %>% pull(P.Value) %>% max()
compare methods %>%
  mutate(tRNA_aa=Row.names %in% tRNA_aa) %>%
  mutate(limma_sig=Row.names %in% rownames(limma_rna_binding_changes)) %>%
  mutate(tRNA_aa_limma=recode(interaction(tRNA_aa, limma_sig),
                              "TRUE.TRUE"="tRNA AA limma sig",
                              "TRUE.FALSE"="tRNA AA limma not sig",
                              "FALSE.TRUE"="Other",
                              "FALSE.FALSE"="Other")) %>%
  arrange(tRNA_aa) %>%
  ggplot(aes(logFC, -log10(P.Value), colour=tRNA_aa_limma)) +
  geom_point(size=1) +
  scale_colour_manual(values=c("grey70", cbPalette[5], cbPalette[6]), name="") +
  geom_vline(xintercept=log2(1.4), linetype=2, colour=cbPalette[3]) +
  geom_hline(yintercept=-log10(max_p_sig_limma), linetype=2, colour="grey60") +
  theme(legend.position="top", legend.direction=2)
```

- Other
- · tRNA AA limma not sig
- tRNA AA limma sig



5.9 Question: Why are there proteins with p-values less than the FDR threshold (above the grey line) and >1.4 fold change but not identified as significant with limma?

This demonstrates the downsides of applying a thresholds to the confidence interval for the estimated fold change when we don't know what fold change is biological relevant. These proteins were subsequently shown to have a consistent (if still slight) change in RNA binding in a Thymidine + Nocadazole experiment suggesting this is probably a real change in RNA binding. So, while a threshold on the log fold change is a very sensible approach, be careful about what threshold you use!

And that's the end of the workshop!