Ageing, Dementia and TBI Data Analysis

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Here is a brief introduction of analyzing the results of our proposed approach on Ageing, Dementia and TBI dataset.

Preparations

```
suppressPackageStartupMessages({
  require(ggplot2)
  require(formatR)
  require(knitr)
  require(cluster)
  require(factoextra)
  require(dplyr)
  require(RColorBrewer)
  require(clusterProfiler)
  require(org.Hs.eg.db)
  require(enrichplot)
  require(stringr)
  require(forcats)
  require(ggplot2)
  require(ggdendro)
  require(graphics)
  require(gridExtra)
  require(extrafont)
  require(viridis)
  require(hrbrthemes)
})
truncated var <- function(x){</pre>
    remove_idx <- c(which.max(x), which.min(x))</pre>
    var(x[-remove_idx])
}
wrap_labal <- function(x, width = 60){</pre>
    str_wrap(x, width=60)
}
simes.test <- function(x, returnstat = FALSE){</pre>
    r = rank(x, ties.method = "random")
    t = min(length(x) * x / r)
    if (returnstat) c(t, t) else t
```

```
setwd("/Users/zhaokai/data/Results/ageing")
load("ageing_dataset_annotated_with_phenotypes_filtered.RData")
pheno <- dataset[,2]</pre>
tissue <- dataset[,3]</pre>
dataset <- dataset[, -(1:4)]</pre>
# our fitted model
load("insider ageing R23 fitted object.RData")
# load("gene_expression_iMF_L1_penalty_25v2.RData")
# attach(fitted obj)
attach(object)
disease_factor <- cfd_matrices[[1]]</pre>
tissue_factor <- cfd_matrices[[2]]</pre>
donor_factor <- cfd_matrices[[3]]</pre>
interactions <- cfd_matrices[[4]]</pre>
# read meta information to facilitate our analyis
meta <- read.csv("meta_info.csv", header = TRUE, stringsAsFactors = F)</pre>
gene_info <- read.csv("rows-genes.csv", header = TRUE, stringsAsFactors = F)</pre>
structure_info <- unique(read.csv("structure_id_mapping.csv", stringsAsFactors = F)[, c(14, 15)])
structure_info$snames[c(4,7)] <- c("hippocampus_right", "hippocampus_left")
# match gene_info with genes included in the study
gene_included <- data.frame(gene_id = as.numeric(gsub("X", "", colnames(data))))</pre>
gene_info_inc <- inner_join(gene_included, gene_info, by = "gene_id")</pre>
row.names(tissue_factor) <- structure_info$snames</pre>
row.names(disease_factor) <- c("ctrl", "case")</pre>
# head to get a sense of our results and meta information
# str(fitted_obj)
head(meta)
      donor id
                     name age sex apo_e4_allele education_years age_at_first_tbi
## 1 326765665 H14.09.078 87
                                 М
                                                N
                                                                16
                                                                                   0
## 2 326765656 H14.09.069
                                 М
                                                N
                                                                17
                                                                                  12
## 3 326765654 H14.09.067 85 M
                                                Y
                                                                                  72
                                                                10
## 4 467056391 H15.09.103 92
                                 F
                                                                                  87
                                                N
                                                                11
                                                Y
## 5 309335447 H14.09.010 100
                                 М
                                                                16
                                                                                   0
## 6 309335457 H14.09.020 97
                                                N
                                                                18
                                                                                   0
     longest_loc_duration cerad num_tbi_w_loc dsm_iv_clinical_diagnosis
## 1
           Unknown or N/A
                               0
                                              0
                                                               No Dementia
                               2
## 2
                  1-2 min
                                              1
                                                               No Dementia
## 3
                  < 10 sec
                               3
                                                                  Vascular
                                              1
## 4
                 < 10 sec
                               0
                                              1
                                                               No Dementia
## 5
           Unknown or N/A
                               3
                                              O Alzheimer's Disease Type
## 6
           Unknown or N/A
                               2
                                              0
                                                               No Dementia
##
     control_set
                        nincds_arda_diagnosis ever_tbi_w_loc
                                                                     race
## 1
                                   No Dementia
                                                                    White
                                                              N
## 2
                                                              Υ
              26
                                   No Dementia
                                                                    White
## 3
              25
                        Dementia, Type Unknown
                                                             Y
                                                                    White
                                                             Y
## 4
              52
                                                                    White
                                   No Dementia
## 5
              28 Possible Alzheimer'S Disease
                                                             N
                                                                    White
## 6
                                                             N Non-white
                                   No Dementia
               1
```

```
hispanic act_demented braak nia_reagan
## 1 Not Hispanic
                   No Dementia
                                    1
                                                1
## 2 Not Hispanic
                   No Dementia
                                    5
                                                2
                                               2
## 3 Not Hispanic
                                    4
                      Dementia
## 4 Not Hispanic
                   No Dementia
                                    4
                                                0
                                                2
## 5 Not Hispanic
                      Dementia
                                    4
## 6 Not Hispanic No Dementia
                                                2
head(gene_info)
##
       gene id chromosome gene entrez id
                                           gene symbol
## 1 499304660
                         1
                                100287102
                                               DDX11L1
## 2 499304661
                                   653635
                                                 WASH7P
                         1
## 3 499304662
                                             MIR6859-1
                                102466751
                         1
## 4 499304663
                                100302278
                                              MIR1302-2
                         1
## 5 499304664
                         1
                                   645520
                                                FAM138A
## 6 499304665
                                105379212 LOC105379212
##
                                                 gene_name
## 1 DEAD/H (Asp-Glu-Ala-Asp/His) box helicase 11 like 1
## 2
                 WAS protein family homolog 7 pseudogene
## 3
                                          microRNA 6859-1
## 4
                                          microRNA 1302-2
## 5
           family with sequence similarity 138, member A
## 6
                             uncharacterized LOC105379212
```

Cluster analysis of donor_factor

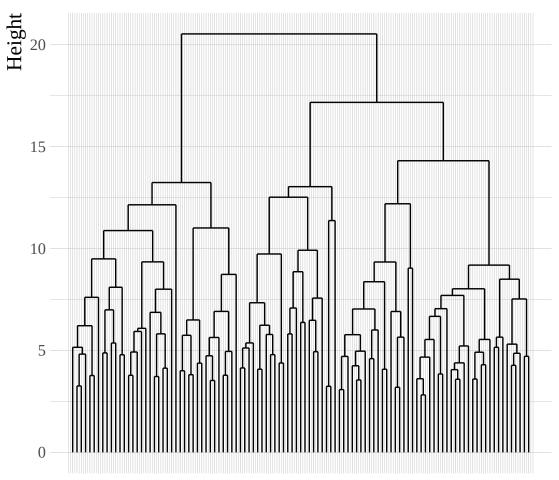
The donor_factor is a low rank representation of genetic information from the dementia dataset, which is a matrix of N rows and K columns. N is the number of donors and K is the number of latent features representing gene expression information in low dimensions. In this section, the cluster analysis basically follows this tutorial.

We exclude a number of metagenes that is irrelevant to synapse function, cognition, and memory. The selection of metagenes is somewhat subjective. However, we observe that this strategy shows some interesting results.

```
# for detail, see https://uc-r.github.io/hc_clustering methods to assess
m <- c( "average", "single", "complete", "ward")</pre>
names(m) <- c( "average", "single", "complete", "ward")</pre>
# function to compute coefficient
ac <- function(x) {
  agnes(donor_factor, method = x)$ac
ac_vec <- sapply(m, function(x) ac(x))</pre>
ac_vec
     average
                single complete
                                       ward
## 0.5436436 0.4432775 0.6453565 0.7564181
# carry out hierarchical cluster analysis using the method with the greatest coefficient
hc3 <- agnes(donor_factor[,-c(14)], method = unname(m[which(ac_vec == max(ac_vec))]))
hc <- as.hclust(hc3)
ggdendrogram(hc, theme_dendro = FALSE) +
  ggtitle("Dendrogram of donor clustering") + ylab("Height") +
 theme_ipsum(base_family = "Times New Roman", base_size= 12, plot_title_face = "bold", axis_title_size
```

```
theme(plot.title = element_text(hjust = 0.5, size=16,face="bold"),
    text=element_text(family="Times New Roman"),
    axis.title.x=element_blank(),
    axis.text.x=element_blank(),
    axis.ticks.x=element_blank())
```

Dendrogram of donor clustering



```
# select cluster number using Gap statistics
# gap_stat <- clusGap(donor_factor, FUN = hcut, nstart = 25, K.max = 10, B = 200)
# fviz_gap_stat(gap_stat)

# choose 3 clusters by investigating Dendrogram and Gap statistics plots
sub_grp <- cutree(hc3, k = 2)
result <- cbind(meta, cluster_id = sub_grp)
aov_res <- aov(age ~ cluster_id, data = result)
unname(unlist(summary(aov_res)))[9]</pre>
```

```
## [1] 0.02517354
kruskal.test(age ~ cluster_id, data = result)
```

```
##
## Kruskal-Wallis rank sum test
##
```

```
## data: age by cluster_id
## Kruskal-Wallis chi-squared = 5.5526, df = 1, p-value = 0.01845
fisher.test(table(result$cluster_id, result$braak), simulate.p.value=TRUE)

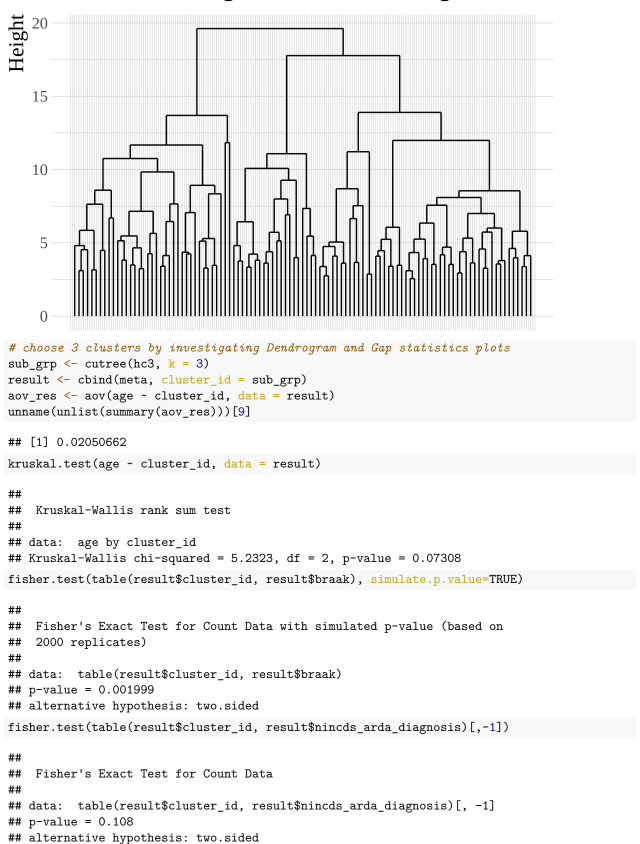
##
## Fisher's Exact Test for Count Data with simulated p-value (based on
## 2000 replicates)
##
## data: table(result$cluster_id, result$braak)
## p-value = 0.3243
## alternative hypothesis: two.sided
```

explore the relavance of the clustering

In this section, we only examined the association between the age of donors and their clusters. Explorations of the relavance of the clustering to other clinical variables can also be carried out. Pie charts and histograms can be drawn to visualize the association. Furthermore, some statistical tests can also be used to check the significance. The result below shows that there is a statistical significant association between the age and clustering.

```
exc_idx <- c(4, 5, 8, 19)
# carry out hierarchical cluster analysis using the method with the greatest coefficient
hc3 <- agnes(donor_factor[,-exc_idx], method = unname(m[which(ac_vec == max(ac_vec))]))
hc <- as.hclust(hc3)
ggdendrogram(hc, theme_dendro = FALSE) +
    ggtitle("Dendrogram of donor clustering") + ylab("Height") +
    theme_ipsum(base_family = "Times New Roman", base_size= 12, plot_title_face = "bold", axis_title_size
    theme(plot.title = element_text(hjust = 0.5, size=16,face="bold"),
        text=element_text(family="Times New Roman"),
        axis.title.x=element_blank(),
        axis.text.x=element_blank())</pre>
```

Dendrogram of donor clustering



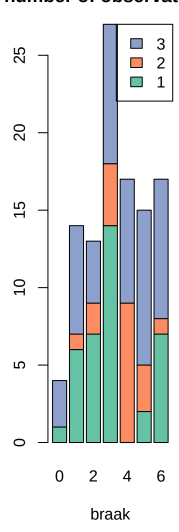
```
fisher.test(table(result$cluster_id, result$dsm_iv_clinical_diagnosis)[,-c(4,5,6)])
```

```
##
## Fisher's Exact Test for Count Data
##
## data: table(result$cluster_id, result$dsm_iv_clinical_diagnosis)[, -c(4, 5, 6)]
## p-value = 0.5234
## alternative hypothesis: two.sided
```

The below histogram show the dribution of clusters cross different braak stages, which are clinical diagnoses of stage of dementia.

```
cluster_tbi <- table(result$cluster_id, result$braak)
par(mar=c(5, 13, 3, 13))
barplot(cluster_tbi, main="number of observations",
    xlab="braak", col= brewer.pal(3, "Set2") ,
    legend = rownames(cluster_tbi), space = 0.2, width = 0.2,
    args.legend = list(x = "toprigh"))</pre>
```

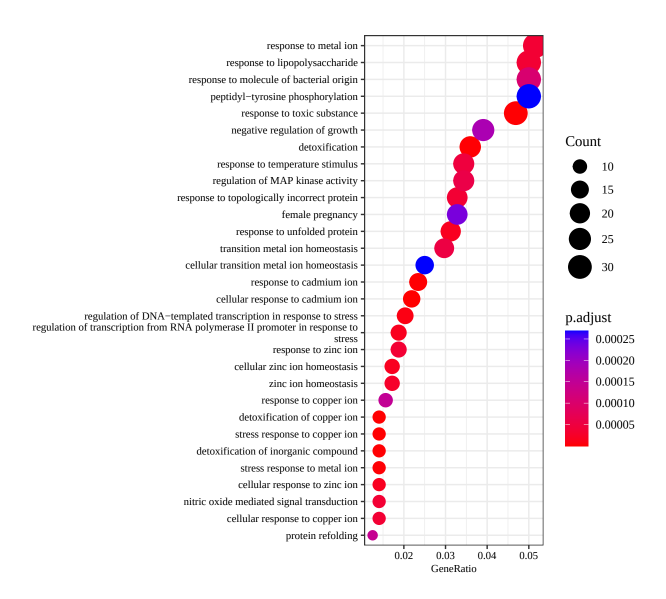
number of observations



Enrichment analysis of biological processes involved

We can also investigate biological processes we are interested in with our results. For example, we explored the mechanism of dementia with the disease and gene factors. We obtained the expression profiles for the dementia and control, extracted the genes with greatest postive difference between them, and examine the biological processes enriched by those genes.

```
idx \leftarrow order(apply(disease factor, 2, function(x) abs(x[1] - x[2])), decreasing = T)[1:3]
disease_matrix <- disease_factor[,idx, drop = F] %*% column_factor[idx, , drop = F]</pre>
diff <- disease_matrix[2,] - disease_matrix[1,]</pre>
cutoffs \leftarrow quantile(diff, probs = seq(0, 1, 0.025))
selected <- (diff <= cutoffs[2])</pre>
# selected <- (column_factor[idx,] >= cutoffs[length(cutoffs)-1]) # greatest negative difference
# cutoffs <- quantile(column_factor[idx,], probs = seq(0, 1, 0.025))</pre>
# selected <- (column_factor[idx,] <= cutoffs[2]) # greatest negative difference
                                = unique(gene_info_inc[selected,3]),
down_reg <- enrichGO(gene</pre>
                      OrgDb
                                = 'org.Hs.eg.db',
                      ont = "BP",
                      readable = TRUE)
dotplot(down_reg, font.size = 8, showCategory=30, label_format = 80) +
 theme(text=element_text(family="Times New Roman"))
```



compare expression level across different brain structures for selected genes

```
loadfonts(quiet = T)

structure_names <- c("TC left", "FWM right", "FWM left", "HPC right", "PC right", "TC right", "HPC left

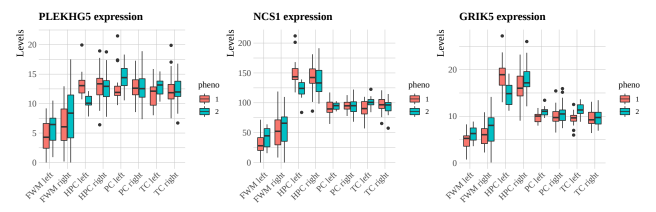
# result <- as.data.frame(upreg)
# gene_names <- unlist(strsplit(result[1,8], split = "/"))

selected <- c("PLEKHG5", "NCS1", "GRIK5", "NRGN")

row_ids <- unlist(sapply(selected, function(x) which(gene_info_inc[[4]] == x)))
data <- cbind(pheno, structure_names[tissue], dataset[,row_ids])
colnames(data) <- c("pheno", "tissue", names(row_ids))
data$pheno <- as.factor(data$pheno)

# boxplots and t-tests for the 4 variables at once
test_results <- sapply(3:ncol(data), function(j){</pre>
```

```
pvalues <- sapply(structure_names, function(i) t.test(data[data$tissue == i, j] ~ data$pheno[data$t</pre>
})
simes_pvalues <- apply(test_results, 2, function(x)simes.test(x))</pre>
names(simes_pvalues) <- names(row_ids)</pre>
colnames(test_results) <- names(row_ids)</pre>
rownames(test results) <- structure names</pre>
p1 <- ggplot(data, aes(x=tissue, y=PLEKHG5, fill=pheno)) +
    scale_color_viridis(discrete = TRUE) +
    geom_boxplot() +
    ggtitle("PLEKHG5 expression") +
    xlab("") + ylab("Levels") +
    theme_ipsum(base_family = "Times New Roman", base_size= 12, axis_title_size = 14, plot_title_size =
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1), plot.margin = unit(c(5, 2, 5, 2),
p2 <- ggplot(data, aes(x=tissue, y=NCS1, fill=pheno)) +
    scale_color_viridis(discrete = TRUE) +
    geom_boxplot() +
    ggtitle("NCS1 expression") +
    xlab("") + ylab("Levels") +
    theme_ipsum(base_family = "Times New Roman", base_size= 12, axis_title_size = 14, plot_title_size =
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1), plot.margin = unit(c(5, 2, 5, 2),
p3 <- ggplot(data, aes(x=tissue, y=GRIK5, fill=pheno)) +
    scale_color_viridis(discrete = TRUE) +
    geom_boxplot() +
    ggtitle("GRIK5 expression") +
    xlab("") + ylab("Levels") +
    theme_ipsum(base_family = "Times New Roman", base_size= 12, axis_title_size = 14, plot_title_size =
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1), plot.margin = unit(c(5, 2, 5, 2),
\# p4 <- ggplot(data, aes(x=tissue, y=NRGN, fill=pheno)) +
      scale_color_viridis(discrete = TRUE) +
      geom_boxplot() +
#
      qqtitle("NRGN expression") +
      xlab("") + ylab("Levels") +
      theme_ipsum(base_family = "Times New Roman", base_size= 12, plot_title_face = "bold", axis_title_
      theme(axis.text.x = element\_text(angle = 45, vjust = 1, hjust=1), plot.margin = unit(c(5, 2, 5, 2))
grid.arrange(p1, p2, p3, nrow=1, ncol=3)
```

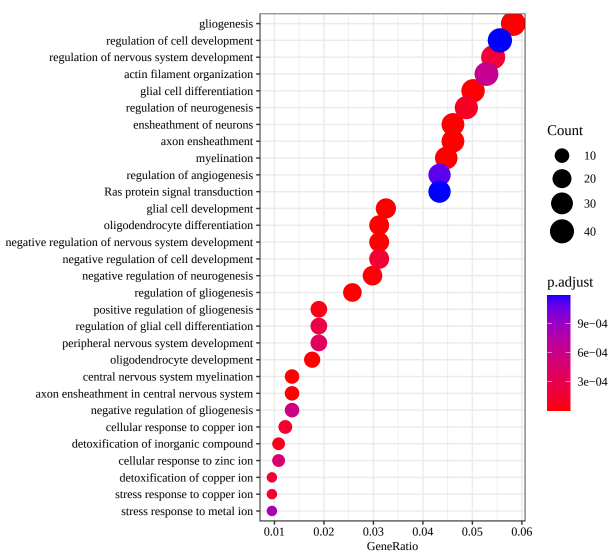


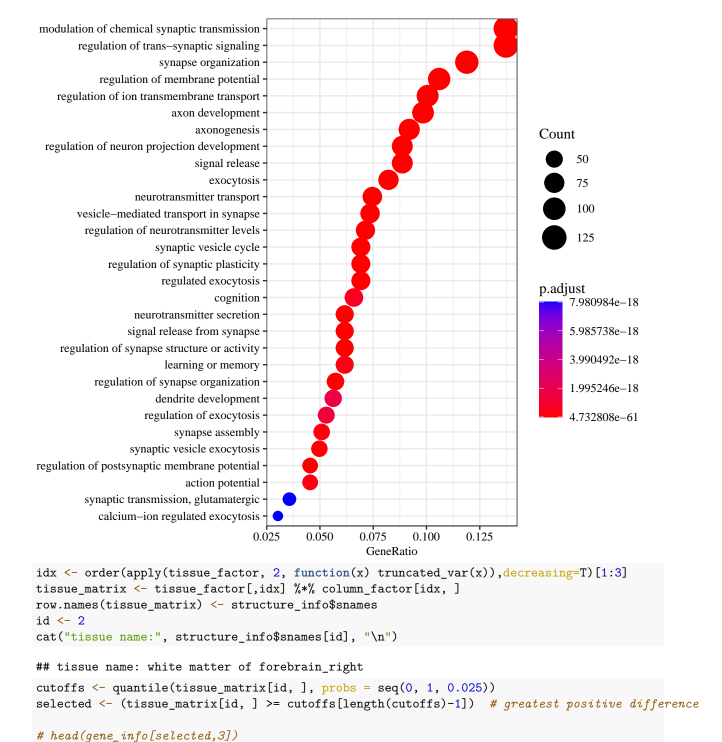
Here is a another pieces of code to demenstrate molecular functions of genes with the largest effects in different brain structures.

In this part of analysis, I only demonstrate with the second tissue. In order to expand, analysis of other metagenes with a single for loop is fine to generate all results.

```
idx <- order(apply(tissue_factor, 2, function(x) truncated_var(x)),decreasing=T)[1:3]</pre>
tissue_matrix <- tissue_factor[,idx] %*% column_factor[idx, ]</pre>
row.names(tissue_matrix) <- structure_info$snames</pre>
id <- 3
cat("tissue name:", structure_info$snames[id], "\n")
## tissue name: white matter of forebrain_left
cutoffs <- quantile(tissue_matrix[id, ], probs = seq(0, 1, 0.025))</pre>
selected <- (tissue_matrix[id, ] >= cutoffs[length(cutoffs)-1]) # greatest positive difference
# head(gene_info[selected,3])
up_reg <- enrichGO(gene</pre>
                              = unique(gene_info_inc[selected,3]),
                              = 'org.Hs.eg.db',
                    OrgDb
                              = "BP",
                    readable = TRUE)
dotplot(up_reg, font.size = 9, showCategory=30, label_format = 60) +
```

theme(text=element_text(family="Times New Roman"))





= unique(gene_info_inc[selected,3]),

= 'org.Hs.eg.db',

= "BP",

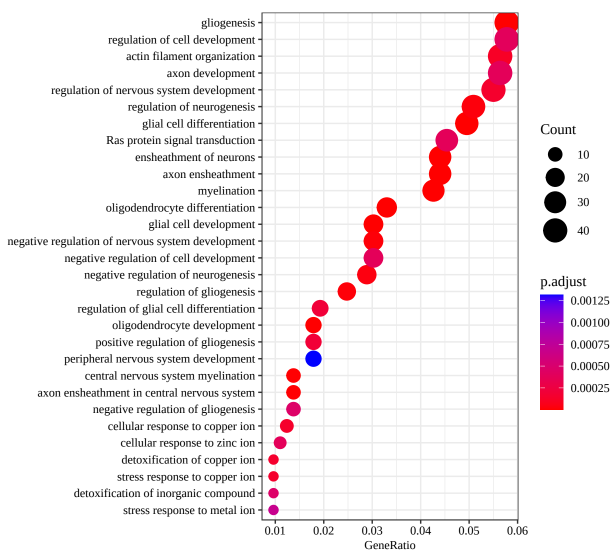
dotplot(up_reg, font.size = 9, showCategory=30, label_format = 60) +

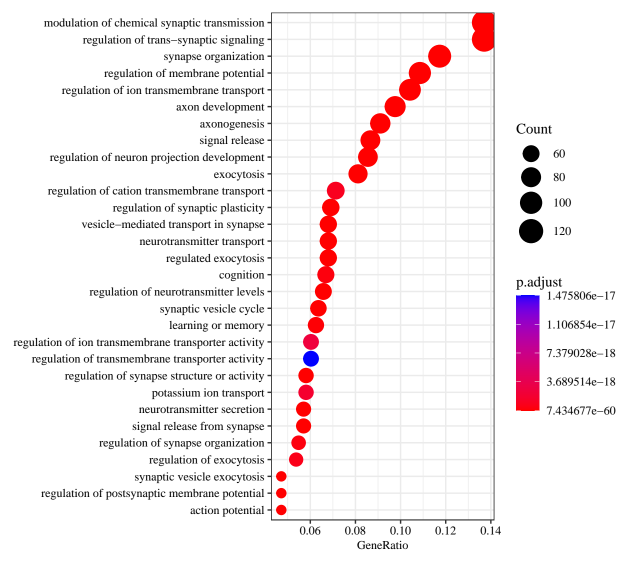
readable = TRUE)

theme(text=element_text(family="Times New Roman"))

up_reg <- enrichGO(gene</pre>

OrgDb





Furthermore, we could also examine the interaction between tissue and disease factors. The code below explores the interaction between different tissues and dementia. Then, similar techniques can be employed to examine the contribution of underlying biological processes to the interaction.

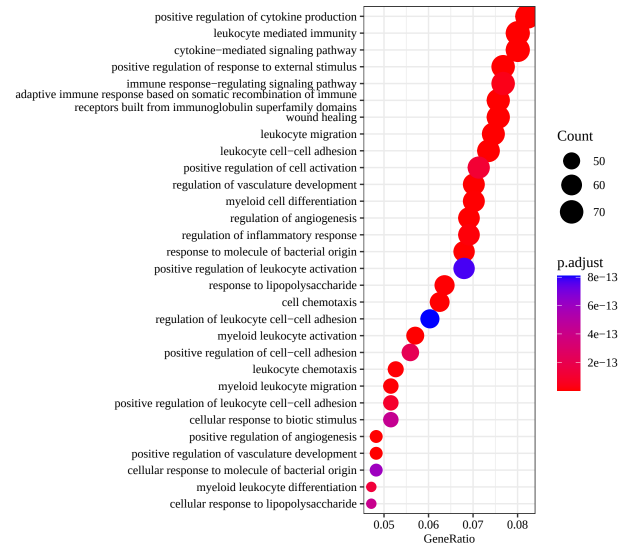
```
# since all latent vectors are restricted in the same space, we can compute the correlation between dis
confounder <- object[['confounder']][,-3]
unique_cfd <- unique(confounder)

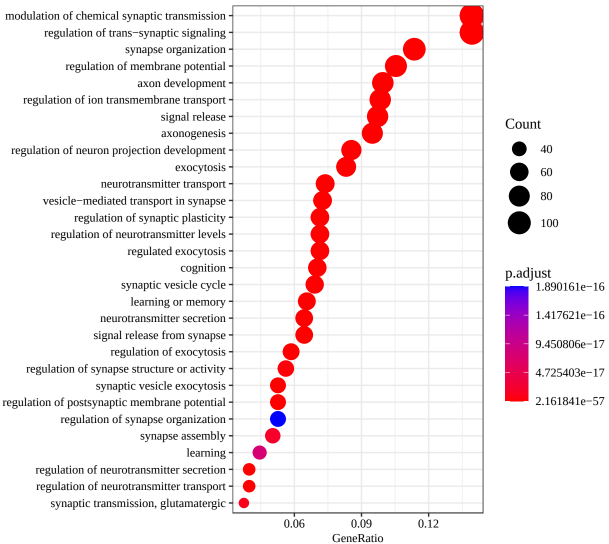
distances <- sapply(seq(8), function(i){
   idx <- which(unique_cfd[,2] == i)
   mean((interactions[idx,1] - interactions[idx,2])^2)
   # t.test(interactions[idx,1], interactions[idx,2], paired = T)$p.value
})

# then we can examine the tissue with the largest distance between dementia and control
# tissue_id <- which.min(distances)
tissue_id <- 4

cat("Tissue name:", structure_info$snames[tissue_id], "\n")</pre>
```

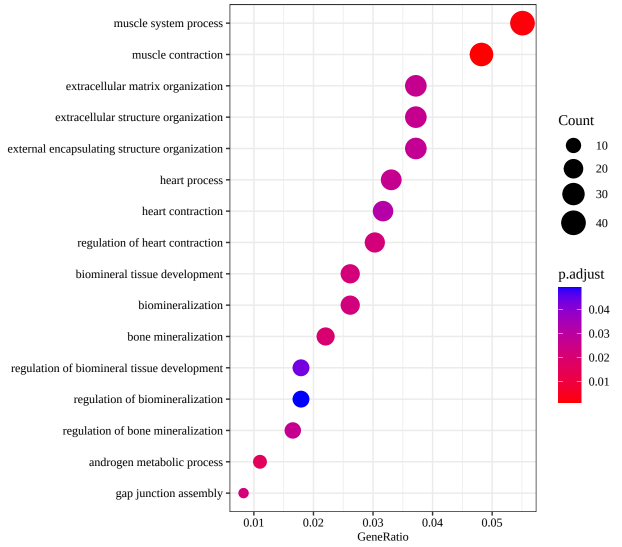
Tissue name: hippocampus_right



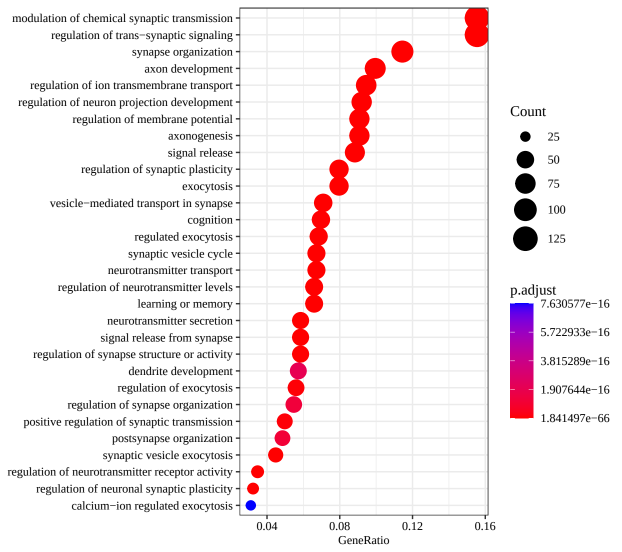


```
# then we can examine the tissue with the largest distance between dementia and control
tissue_id <- 7
# tissue_id <- which.min(distances)
cat("Tissue name:", structure_info$snames[tissue_id], "\n")</pre>
```

```
## Tissue name: hippocampus_left
cfd_id <- which(unique_cfd[,2] == tissue_id)</pre>
```



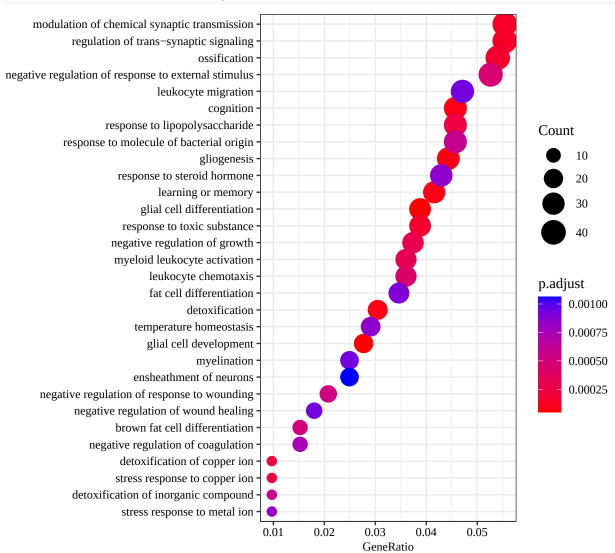
down-regulation, greatest negative difference
selected <- (diff <= cutoffs[2])</pre>

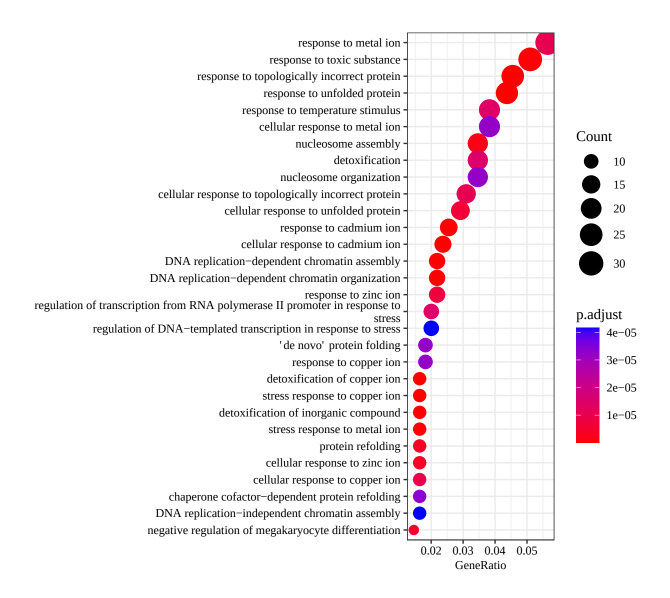


metagene links cognitive dysfunction

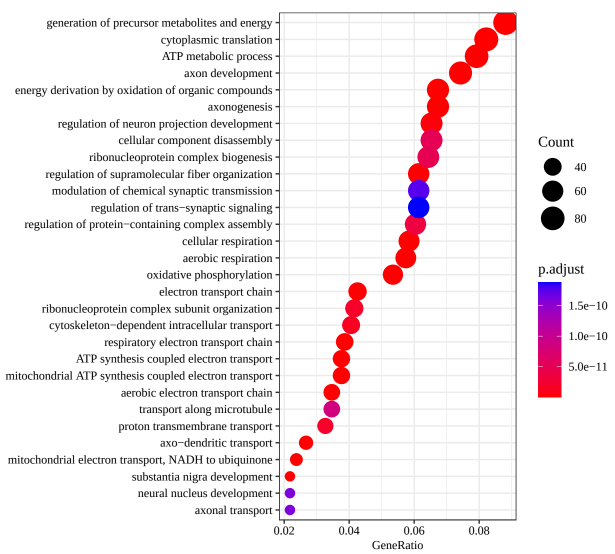
```
readable = TRUE)

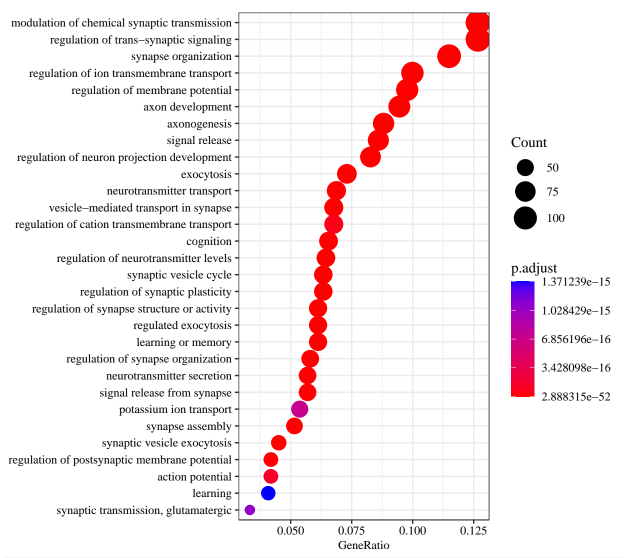
dotplot(down_reg, font.size = 9, showCategory=30, label_format = 80) +
    theme(text=element_text(family="Times New Roman"))
```

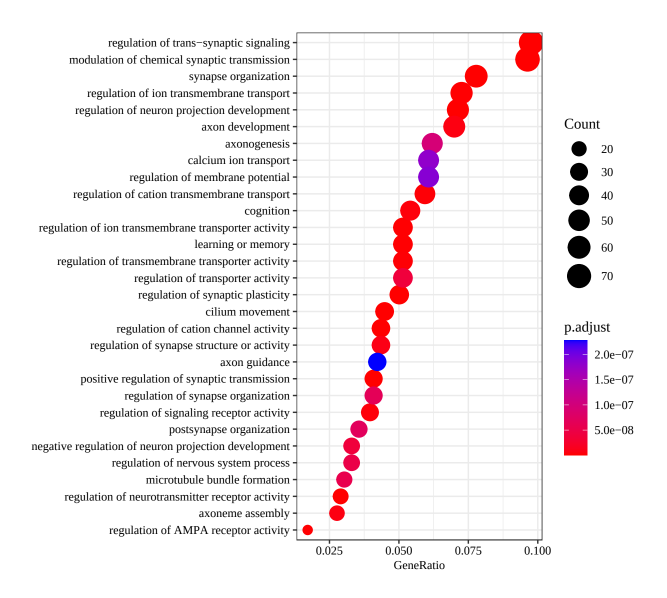




metagene characterize the funtion of brain regions

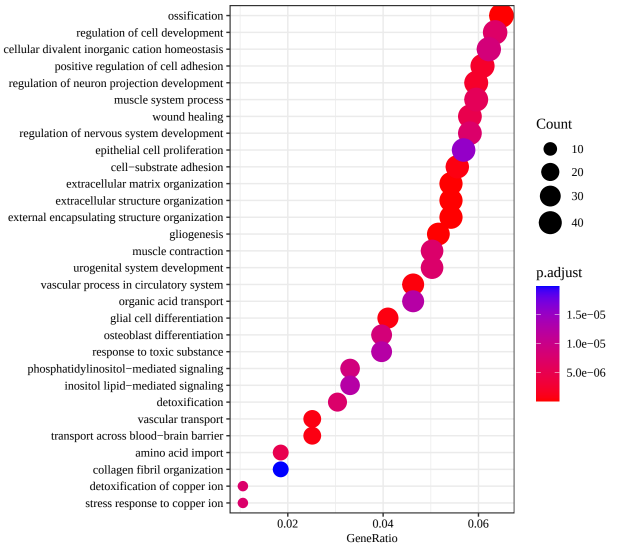






metagene characterize donors

```
result <- data.frame(meta[,c(4,14)], donor_factor)</pre>
t.test(X5 ~ sex, data = result)
##
##
    Welch Two Sample t-test
##
## data: X5 by sex
## t = -2.4849, df = 71.336, p-value = 0.01531
## alternative hypothesis: true difference in means between group F and group M is not equal to 0
## 95 percent confidence interval:
   -0.87454203 -0.09589633
## sample estimates:
  mean in group F mean in group M
                         0.2086001
        -0.2766190
##
```



```
idx <- 12
cutoffs <- quantile(column_factor[idx, ], probs = seq(0, 1, 0.025))
# selected <- (column_factor[idx, ] <= cutoffs[2])
selected <- (column_factor[idx,] >= cutoffs[length(cutoffs)-1]) # greatest negative difference
```

