

Task 3: RNA-seq Differential Gene Expression & RNA-seq Functional Profiling

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1 Introduction

2 Raw Reads and Mapping QC

2.1 FastQC:

- a) According to FastQC: What was the minimum and the maximum number of read pairs sequenced per sample?

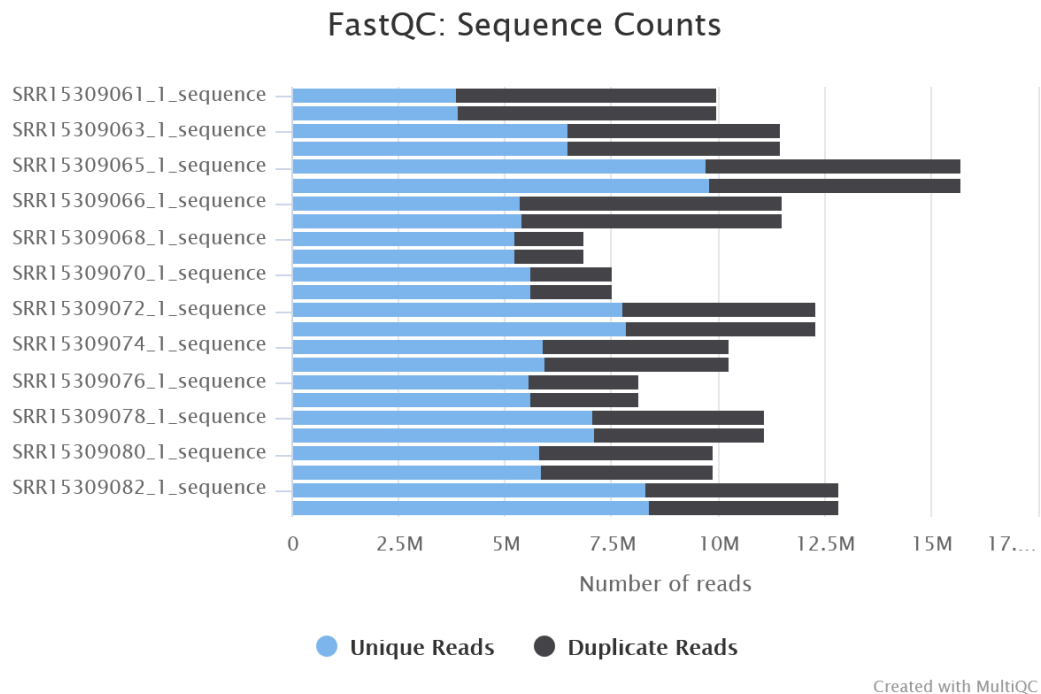


Figure 1: Barplot of the number of read pairs per sample.

Answer: As can be seen in Figure 1 the minimum number of reads per sample is around 6.8 million and the maximum number of reads per sample is around 15.7 million.

- b) What is the most overrepresented sequence (string of nucleotides) that was found by FastQC?

Answer: According to the MultiQC report the most overrepresented sequence was:

"AA"

- c) What might be the reason there is so much of this specific sequence/homopolymer ?

Answer: This could be from A tailed adapter dimers and PCR slippage products that outcompete genuine RNA fragments in the library.

2.2 HTSeq-count:

- d) According to HTSeq Count: What was the minimum and the maximum number of read pairs reported per sample?

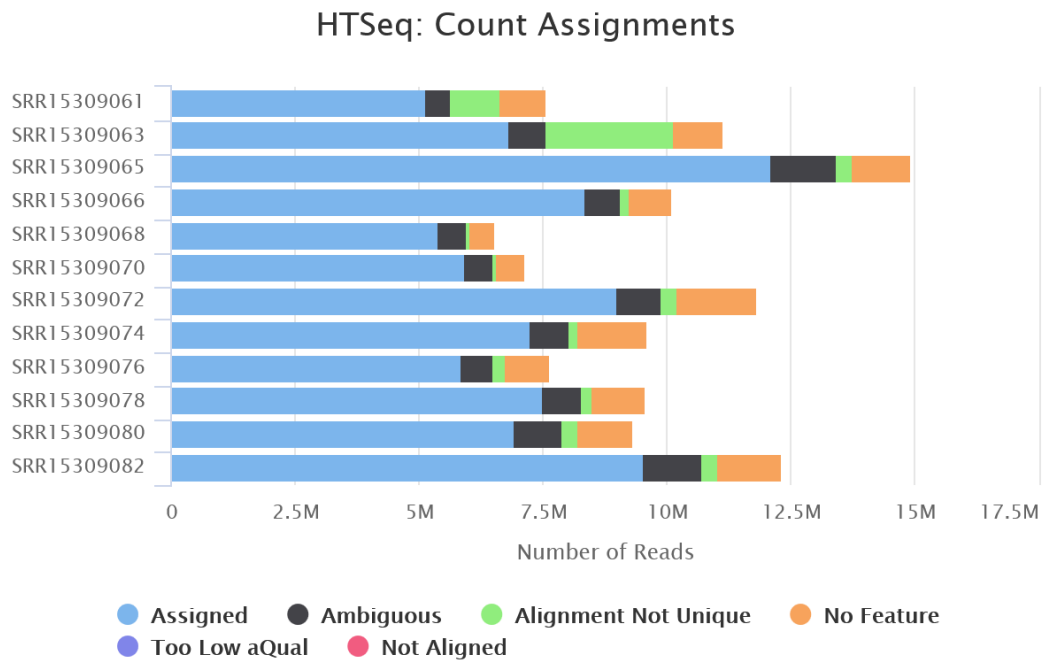


Figure 2: Barplot of the number of read pairs, per sample HTSeq.

Answer: As can be seen in Figure 2 the minimum number of reads per sample is around 5.9 million and the maximum number of reads per sample is around 14.8 million.

- f) Which was the minimum and maximum percentage of reads uniquely assigned to a gene, as reported by HTSeq-count?

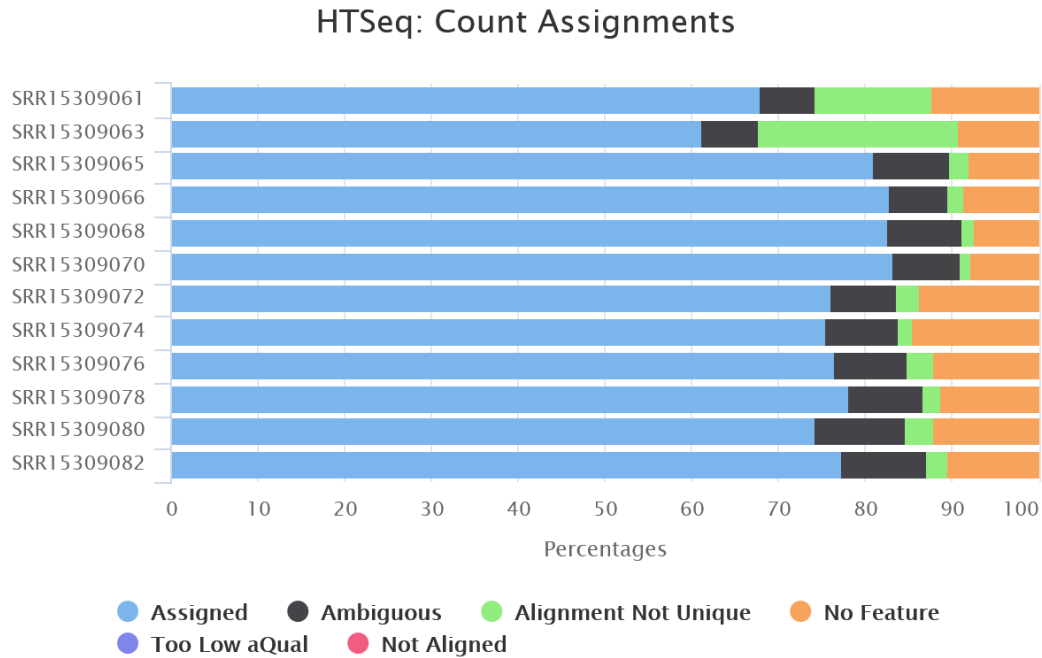


Figure 3: HTSeq assignment plot in percentages.

Answer: As can be seen in Figure 3 the minimum percentage of reads uniquely assigned to a gene is 61.3% and the maximum percentage of reads uniquely assigned to a gene is around 83.3%.

3 Dataset

3.1 Raw read counts

3.2 Sample information

- a) Which columns define the used base strain and substrain (WT or TGT mutant), respectively? Can you spot an error in one of those columns?

Answer: The genotype column shows if the wild type or tgt mutant was used. The strain column shows which *E.coli* strain was used. The error is in the strain column suggesting that for every experiment a different strain was used. But comparing this with the strains and method subsection of the methods section of the paper shows that only “*Escherichia coli* K-12 MG1655 was used as the WT strain.”.[1]

- b) Which column defines if nickel was added to the media?

Answer: The treatment column defines if nickel was added to the media.

- c) Find the following information on the SRA Study page:

Which type of Illumina machine was used for sequencing?

Answer: Illumina HiSeq 2500

What was the library layout?

Answer: PAIRED

When was the data released?

Answer: 2022-05-16

4 Preprocessing of the data

4.1 Filtering of the data

- a) For how many genes did we originally retrieve count data?

```
dim(rawCounts)
#[1] 4295 12
```

Answer: Originally count data were retrieved for 4295 genes

- b) How many will be left after applying the filter?

```
dim(rawCounts[rowSums(rawCounts) > 10, ])
#[2] 4221 12
```

Answer: After applying the filter 4221 genes will be left.

5 DGE Analysis

5.1 Differential Expression Analysis

5.2 Extracting results

Interpreting the summary:

- a) How many genes are significantly up-regulated and how many are significantly down-regulated in the nickel treated WT strain as compared to the untreated WT strain, using the default cutoff for the adjusted p-value?

```

# Extract results with default alpha (0.1)
DESeq2Results_WT_nickel <- results(DESeq2Data,
                                   contrast = c("group", "WT.Nickel", "WT.none"))

# View summary
summary(DESeq2Results_WT_nickel)

# out of 4221 with nonzero total read count
# adjusted p-value < 0.1
# LFC > 0 (up)      : 1069, 25%
# LFC < 0 (down)    : 1063, 25%
# outliers [1]      : 6, 0.14%
# low counts [2]     : 0, 0%
# (mean count < 1)
# [1] see 'cooksCutoff' argument of ?results
# [2] see 'independentFiltering' argument of ?results

```

Answer: 1069 genes are significantly up-regulated and 1063 genes are significantly down-regulated in the nickel treated WT strain as compared to the untreated WT strain, using the default cutoff for the adjusted p-value of 10%.

- b) What is the standard cutoff used for the significance level (adjusted p-value), if we don't change it?

Answer: The standard cutoff used for the significance level (adjusted p-value), if we don't change it is 10% (p-value < 0.1).

- c) How many significantly differentially expressed genes does that make in total?

```

# Extract TGT.Nickel vs TGT.none with alpha = 0.1
DESeq2Results_TGT_nickel <- results(DESeq2Data,
                                   contrast = c("group", "TGT.Nickel", "TGT.none"),
                                   alpha = 0.1)

# View summary
summary(DESeq2Results_TGT_nickel)

# out of 4221 with nonzero total read count
# adjusted p-value < 0.1
# LFC > 0 (up)      : 986, 23%
# LFC < 0 (down)    : 877, 21%
# outliers [1]      : 6, 0.14%
# low counts [2]     : 164, 3.9%
# (mean count < 5)

```

```
# [1] see 'cooksCutoff' argument of ?results
# [2] see 'independentFiltering' argument of ?results
```

Answer: If we add up the up and down regulated genes we get the total amount of significantly differentially expressed genes. Looking at the summary this would be 1863 genes.

Changing the alpha factor:

- d) For the comparison of the genotypes under standard conditions. How many significantly differentially expressed genes in total are reported for a significance level of 0.05? (Go to the RStudio Help and search for “results” function, to identify the attribute you have to change.)

```
# Extract genotype comparison (TGT vs WT, no nickel) with alpha = 0.05
DESeq2Results_genotype <- results(DESeq2Data,
                                   contrast = c("group", "TGT.none", "WT.none"),
                                   alpha = 0.05)

# View summary (this shows the numbers you need)
summary(DESeq2Results_genotype)

# out of 4221 with nonzero total read count
# adjusted p-value < 0.05
# LFC > 0 (up)      : 187, 4.4%
# LFC < 0 (down)    : 275, 6.5%
# outliers [1]      : 6, 0.14%
# low counts [2]     : 0, 0%
# (mean count < 1)
# [1] see 'cooksCutoff' argument of ?results
# [2] see 'independentFiltering' argument of ?results
```

Answer: If we add up the up and down regulated genes we get the total amount of significantly differentially expressed genes. Looking at the summary this would be 462 genes at a p-value < 0.05.

Comparing the nickel treatment to no treatment in the TGT-mutant:

- e) Repeat the steps above for the comparison of the TGT-mutant strain treated with nickel to the TGT-mutant strain not treated with nickel. How many significantly differentially expressed genes in total are reported for a significance level of 0.05?

```
# Extract TGT mutant nickel effect with alpha = 0.05
DESeq2Results_TGT_nickel <- results(DESeq2Data,
                                     contrast = c("group", "TGT.Nickel", "TGT.none"),
                                     alpha = 0.05)
```



```
# View summary (this shows the numbers you need)
summary(DESeq2Results_TGT_nickel)

# out of 4221 with nonzero total read count
# adjusted p-value < 0.05
# LFC > 0 (up)      : 826, 20%
# LFC < 0 (down)    : 752, 18%
# outliers [1]      : 6, 0.14%
# low counts [2]    : 82, 1.9%
# (mean count < 3)
# [1] see 'cooksCutoff' argument of ?results
# [2] see 'independentFiltering' argument of ?results
```

Answer: If we add up the up and down regulated genes we get the total amount of significantly differentially expressed genes. Looking at the summary this would be 1578 genes at a p-value < 0.05.

6 Vizualising data

6.1 Experimental QC - Clustering of samples (PCA)

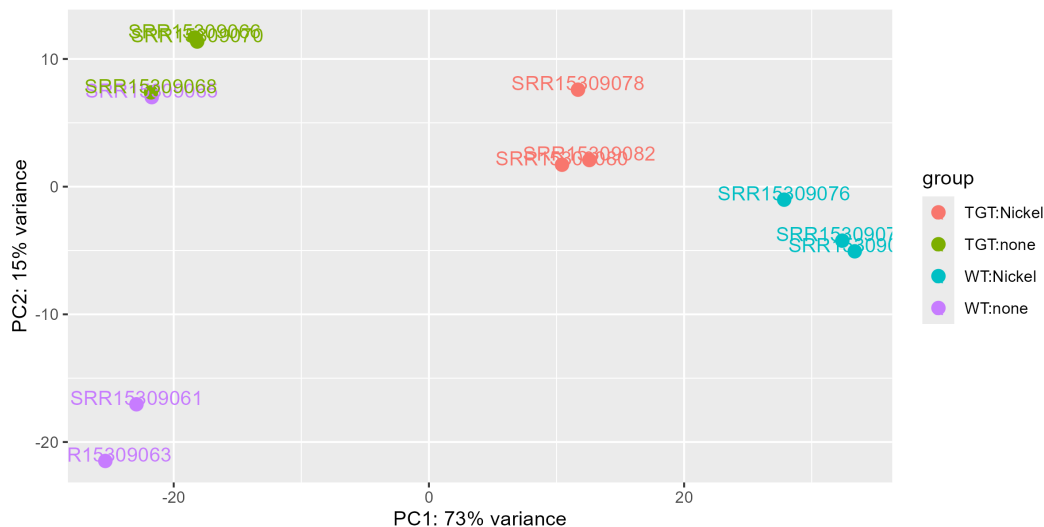


Figure 4: Clustering of samples PCA dot plot.

a) Do the groups of replicates behave as expected?

Answer: Looking at the plot in figure 4 we can observe that the individual groups are clearly separated from each other. It also makes sense that there is a big distance between untreated and Nickel treated strains (WT as well as TGT strains).

b) Which sample would you identify as an outlier?

Answer: Looking at figure 4 we would identify the WT strain untreated as an outlier as it does not completely cluster with its biological replicates but one point also clusters with the TGT untreated cluster.

6.2 Viewing counts for a single geneID

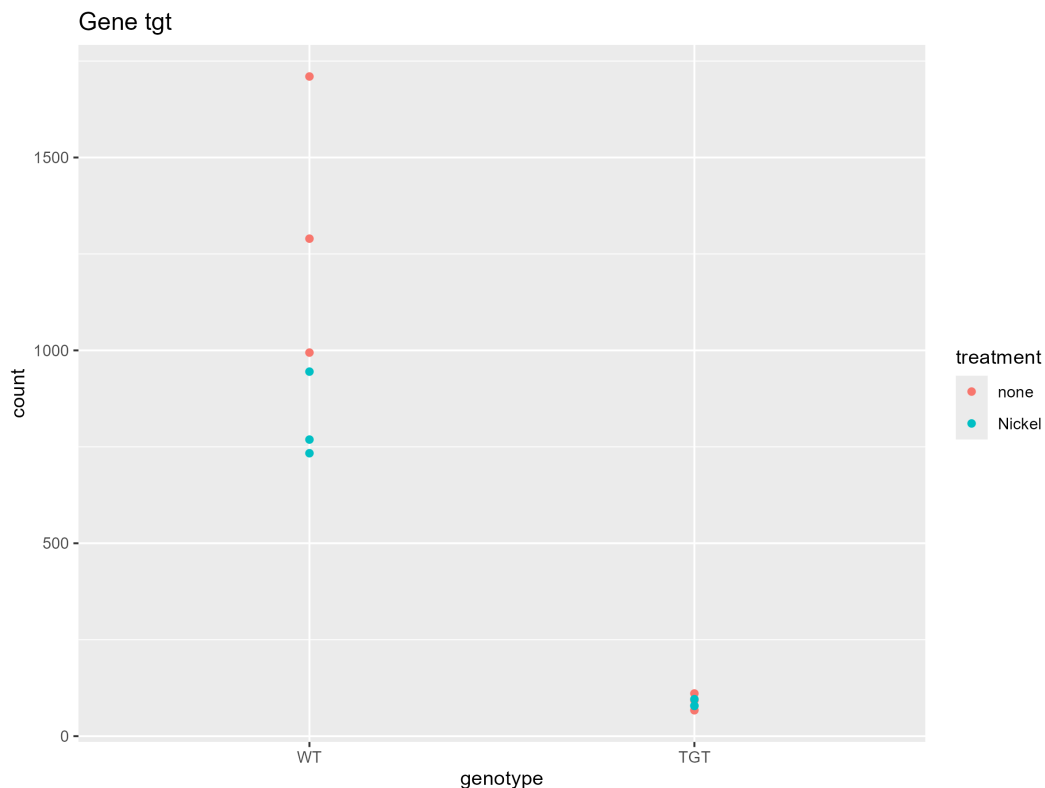


Figure 5: Viewing counts for a single geneID in a dot plot.

c) Think about the mutations in the E. coli strains and how that influences the transcripts of a gene. Are the read counts for the tgt gene in the wild type and the knockout strain what you expected? Explain why.

Answer:

Wild type + none: tgt encodes tRNA guanine transglycosylase, an essential enzyme for queuosine synthesis. It is expressed under normal growth conditions to modify tRNAs. Expected Counts are going to be high.

Wild type + Nickel: tgt is transcriptionally repressed by nickel stress. Expected Counts are going to be lower than that of Wt + none but still higher than the expected counts of the knock out mutants.

tgt + none: The tgt gene is deleted so No functional tgt should be transcribed. Expected counts are going to be very low to near zero.

tgt + Nickel: The tgt knock out stais the same but now Nickel is added. This should not have a significant affect on the already very low to near zero counts.

7 Part 2: Functional Analysis and Vizualisation

7.1 Setup

```
BiocManager::install(c("clusterProfiler"))
install.packages("tidyverse")

install.packages("devtools")
devtools::install_github('kevinblighe/EnhancedVolcano')
```

```
library(EnhancedVolcano)
library(clusterProfiler)
library(tidyverse)
library(ggplot2)
library(dplyr)
library(DESeq2)
```

```
### load the data with read delim, because it is tab seperated
annotatedRawCounts <- read_delim("Counts_raw.tsv")

head(annotatedRawCounts)
```

```
## # A tibble: 6 x 15
##   ID      product      gene SRR15309076 SRR15309078 SRR15309074 SRR15309066
##   <chr> <chr>      <chr>      <dbl>      <dbl>      <dbl>      <dbl>
## 1 b0001 thr operon leader~ thrL          113          432          119          107
## 2 b0002 fused aspartate k~ thrA         2910         27851         3646         2378
## 3 b0003 homoserine kinase  thrB         1274          6956         1497         1476
## 4 b0004 threonine synthase thrC         1814         10638         2256         2462
## 5 b0005 DUF2502 domain-co~ yaaX          124           216          171          124
## 6 b0006 peroxide stress r~ yaaA          405           443          530          741
```

```
## # i 8 more variables: SRR15309082 <dbl>, SRR15309080 <dbl>, SRR15309070 <dbl>,
## #   SRR15309065 <dbl>, SRR15309072 <dbl>, SRR15309068 <dbl>, SRR15309063 <dbl>,
## #   SRR15309061 <dbl>
```

```
annotatedRawCounts <- annotatedRawCounts %>%
  column_to_rownames(var = "ID")

### split data into rawCounts and Annotations
rawCounts <- annotatedRawCounts[,3:14]
annotations <- annotatedRawCounts[,1:2]

DESeq2ResultsDF <- read_delim("DESeq2Result_treatment.tsv")
```

8 Gene annotation (databases)

- a) If you look closely at the 21 OrgDB packages, there are 2 different packages for E. coli. Which one should we use?

The screenshot shows the Bioconductor website interface. On the left, there's a sidebar with 'Bioconductor version 3.22 (Release)' and a 'Go to 3.23 (Devel)' button. Below this is a search bar and a list of categories like 'Software (2361)', 'AnnotationData (928)', etc. The main content area is titled 'Packages found under Escherichia_coli:' and shows a table of search results. The table has columns for 'Package', 'Maintainer', 'Title', and 'Rank'. The first row, 'org.EcK12.eg.db', is highlighted with a red box. Other rows include 'BSgenome.Ecoli.NCBI.20080805', 'org.EcSakai.eg.db', 'ecoliK12.db', and 'ecoliSakai.db'.

Package	Maintainer	Title	Rank
org.EcK12.eg.db	Bioconductor Package Maintainer	Genome wide annotation for E coli strain K12	59
BSgenome.Ecoli.NCBI.20080805	Bioconductor Package Maintainer	Escherichia coli full genomes	93
org.EcSakai.eg.db	Bioconductor Package Maintainer	Genome wide annotation for E coli strain Sakai	165
ecoliK12.db	Bioconductor Package Maintainer	Base Level Annotation databases for E coli K12 Strain	296
ecoliSakai.db	Bioconductor Package Maintainer	Base Level Annotation databases for E coli Sakai Strain	284

Figure 6: Website of the used database for E.coli K12.

```
BiocManager::install("org.EcK12.eg.db")

library(org.EcK12.eg.db)
```

Answer: Since we are working with E. coli K12 MG1655 it would make sense to use the genome wide annotation package for E.coli K12 (org.EcK12.eg.db)

- c) List the different available keytypes/identifiers.

```
#gene names/identifiers
genes <- annotations$gene
head(genes,10)
```

```
## [1] "thrL" "thrA" "thrB" "thrC" "yaaX" "yaaA" "yaaJ" "talB" "mog" "satP"
```

```
#View all available keytypes
keytypes(org.EcK12.eg.db)
```

```
## [1] "ACCNUM" "ALIAS" "ENTREZID" "ENZYME" "EVIDENCE"
## [6] "EVIDENCEALL" "GENENAME" "GO" "GOALL" "ONTOLOGY"
## [11] "ONTOLOGYALL" "PATH" "PMID" "REFSEQ" "SYMBOL"
```

Answer: The different keytypes and identifiers are shown above.

d) Which of the keytypes/identifiers include the gene names given in the annotation?

```
keys <- keys(org.EcK12.eg.db, keytype = "SYMBOL")
head(keys)
```

```
## [1] "yjhR" "nfrA" "thrL" "insB1" "sspA" "yaaJ"
```

```
#make sure our IDs are contained in the keys
#prints the number of genes that are found within the keys
sum(genes %in% keys)
```

```
## [1] 4274
```

Answer: After iterating through all different keytypes, SYMBOL was found to contain the genes.

9 Running functional analysis

9.1 Overrepresentation analysis

a) How many significantly up- and downregulated genes are left, after applying the LFC cutoff?

```
# filter our results by padj and log2FoldChange
res_up <- dplyr::filter(DESeq2ResultsDF, padj < 0.05 & log2FoldChange > 1)
res_down <- dplyr::filter(DESeq2ResultsDF, padj < 0.05 & log2FoldChange < -1)

# extract gene IDs for upregulated genes
genes_up_id <- res_up$ID
```

```
genes_up <- annotations[genes_up_id,"gene"]

# extract gene names for downregulated genes
genes_down_id <- res_down$ID
genes_down <- annotations[genes_down_id,"gene"]

# append both lists to get all deregulated genes
genes_de <- c(genes_up,genes_down)

length(genes_up)
```

```
## [1] 660
```

```
length(genes_down)
```

```
## [1] 582
```

```
length(genes_de)
```

```
## [1] 1242
```

Answer: With the *length* command we can show the number of genes up- or downregulated. In total, 660 genes were up- and 582 genes were downregulated. In sum, 1242 genes are deregulated.

- b) How many significantly over-represented biological processes (GO terms) are there, per subset of genes (all differentially expressed genes, up-regulated genes only, down-regulated genes only.)

```
EC <- "org.EcK12.eg.db"
EC_KEY <- "SYMBOL"

orBP <- enrichGO(genes_de,
                  EC,
                  ont="BP",
                  keyType = EC_KEY,
                  pvalueCutoff=0.05)

orUpBP <- enrichGO(genes_up,
                   EC,
                   ont="BP",
                   keyType = EC_KEY,
                   pvalueCutoff=0.05)
```

Table 1: Overrepresented BP terms - All DE genes

	Description	p.adjust	Count
GO:0006935	chemotaxis	0	34
GO:0042330	taxis	0	34
GO:0040011	locomotion	0	42
GO:0001539	cilium or flagellum-dependent cell motility	0	49
GO:0071973	bacterial-type flagellum-dependent cell motility	0	49

```
orDownBP <- enrichGO(genes_down,
                      EC,
                      ont="BP",
                      keyType = EC_KEY,
                      pvalueCutoff=0.05)

#To show the number of statistically significant GO terms
n_BP_all <- nrow(orBP)
n_BP_up <- nrow(orUpBP)
n_BP_down <- nrow(orDownBP)

print(n_BP_all)
```

```
## [1] 100
```

```
print(n_BP_up)
```

```
## [1] 59
```

```
print(n_BP_down)
```

```
## [1] 93
```

Answer: There are in summary 100 GO terms deregulated, 59 up and 93 down. For a quick look into the first few results, take a look at the tables below.

10 Vizualization

10.1 Vulcano Plot

- Which is the most down-regulated gene and which is the most up-regulated gene, in terms of LFC?

Table 2: Overrepresented BP terms - Upregulated genes

	Description	p.adjust	Count
GO:0046377	colanic acid metabolic process	0.00e+00	18
GO:0009242	colanic acid biosynthetic process	2.00e-07	15
GO:0072348	sulfur compound transport	4.00e-07	23
GO:0006857	oligopeptide transport	2.20e-06	21
GO:0042938	dipeptide transport	3.42e-05	15

Table 3: Overrepresented BP terms - Downregulated genes

	Description	p.adjust	Count
GO:0006935	chemotaxis	0	32
GO:0042330	taxis	0	32
GO:0001539	cilium or flagellum-dependent cell motility	0	42
GO:0071973	bacterial-type flagellum-dependent cell motility	0	42
GO:0097588	archaeal or bacterial-type flagellum-dependent cell motility	0	42

```

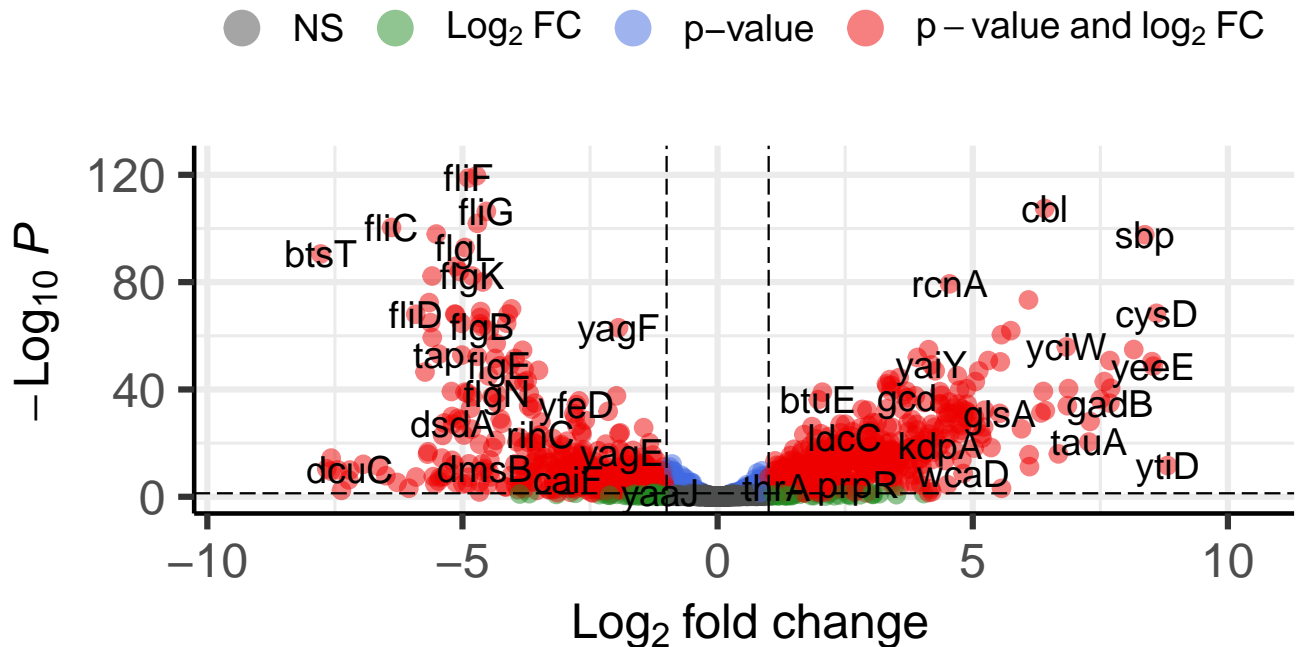
# Set a boolean column for significance
DESeq2ResultsDF$significant <- ifelse(!is.na(DESeq2ResultsDF$padj) & DESeq2ResultsDF$padj < .05, TRUE, FALSE)

## Add gene annotations
DESeq2ResultsDF$gene <- annotations[DESeq2ResultsDF$ID, "gene"]

EnhancedVolcano(DESeq2ResultsDF,
  lab = DESeq2ResultsDF$gene,
  x = 'log2FoldChange',
  y = 'padj',
  pCutoff = 0.05) # + xlim(0,2.5) + ylim(0,10) you can further restrict the plot

```


Enhanced Volcano



total = 4221 variables

```
#show the entry with the highest log2FoldChange
DESeq2ResultsDF[DESeq2ResultsDF$log2FoldChange == max(DESeq2ResultsDF$log2FoldChange),]
```

```
## # A tibble: 1 x 9
##   baseMean log2FoldChange lfcSE  stat    pvalue    padj ID      significant gene
##   <dbl>         <dbl> <dbl> <dbl>    <dbl>    <dbl> <chr> <lgl>      <chr>
## 1      41.6           8.82  1.20  7.33  2.25e-13  2.33e-12 b4721 TRUE      ytid
```

```
#show the entry with the lowest log2FoldChange
DESeq2ResultsDF[DESeq2ResultsDF$log2FoldChange == min(DESeq2ResultsDF$log2FoldChange),]
```

```
## # A tibble: 1 x 9
##   baseMean log2FoldChange lfcSE  stat    pvalue      padj ID      significant gene
##   <dbl>         <dbl> <dbl> <dbl>    <dbl>    <dbl> <chr> <lgl>      <chr>
## 1   38216.         -7.78 0.378 -20.6  7.41e-94  3.13e-91 b4354 TRUE      btsT
```

```
ggsave('figs/volcano_plot.png', width = 12, height = 8)
```

Answer: The most upregulated gene is *ytiD* with a L2FC of 8.82. The most downregulated gene is *btsT* with a L2FC of -7.78.

- b) How many times are those genes higher or lower expressed in treated compared to untreated WT strain? (Calculate the fold change from the LFC)

```
DESeq2ResultsDF <- read_delim("DESeq2Result_treatment.tsv")
res_df <- as.data.frame(DESeq2ResultsDF)

most_up_idx <- which.max(res_df$log2FoldChange)
most_up <- res_df[most_up_idx, ]

most_down_idx <- which.min(res_df$log2FoldChange)
most_down <- res_df[most_down_idx, ]

gene_name_up <- annotations[most_up$ID, "gene"]
gene_name_down <- annotations[most_down$ID, "gene"]

most_up$foldChange <- 2^most_up$log2FoldChange
most_down$foldChange <- 2^most_down$log2FoldChange

most_up$gene <- gene_name_up
most_down$gene <- gene_name_down

print(c(most_up$log2foldChange, most_up$foldChange ,most_up$gene ))
```

```
## [1] "451.110505802606" "ytiD"
```

```
print(c(most_down$log2foldChange, most_down$foldChange, most_down$gene))
```

```
## [1] "0.00455492673103271" "btsT"
```

Answer: For a comprehensive comparison, we can do this manually by thinking of how the LFC is defined. Since it's a logarithmic function with base 2, we can calculate the FoldChange: $FC = 2^{LFC}$ and compare the results manually. For faster and automated calculation, one may use the code above, which takes the data from the DESeq dataset for treated and untreated WT and calculates the FC. Gene *ytiD* is 451.1 times higher transcribed than in untreated WT. Gene *btsT* is 217.4 times lower transcribed than in untreated WT.

- c) Which of the genes shows the most significant deregulation (the lowest p-value)?

```
min_pvalue_idx <- which.min(res_df$pvalue)
gene_min_pvalue <- res_df[min_pvalue_idx, ]
```

```

gene_id_sign <- gene_min_pvalue$ID
gene_name_sign <- annotations[gene_id_sign, "gene"]

most_sign <- gene_min_pvalue
most_sign$gene <- gene_name_sign

print(c(gene_min_pvalue$pvalue, most_sign$gene))

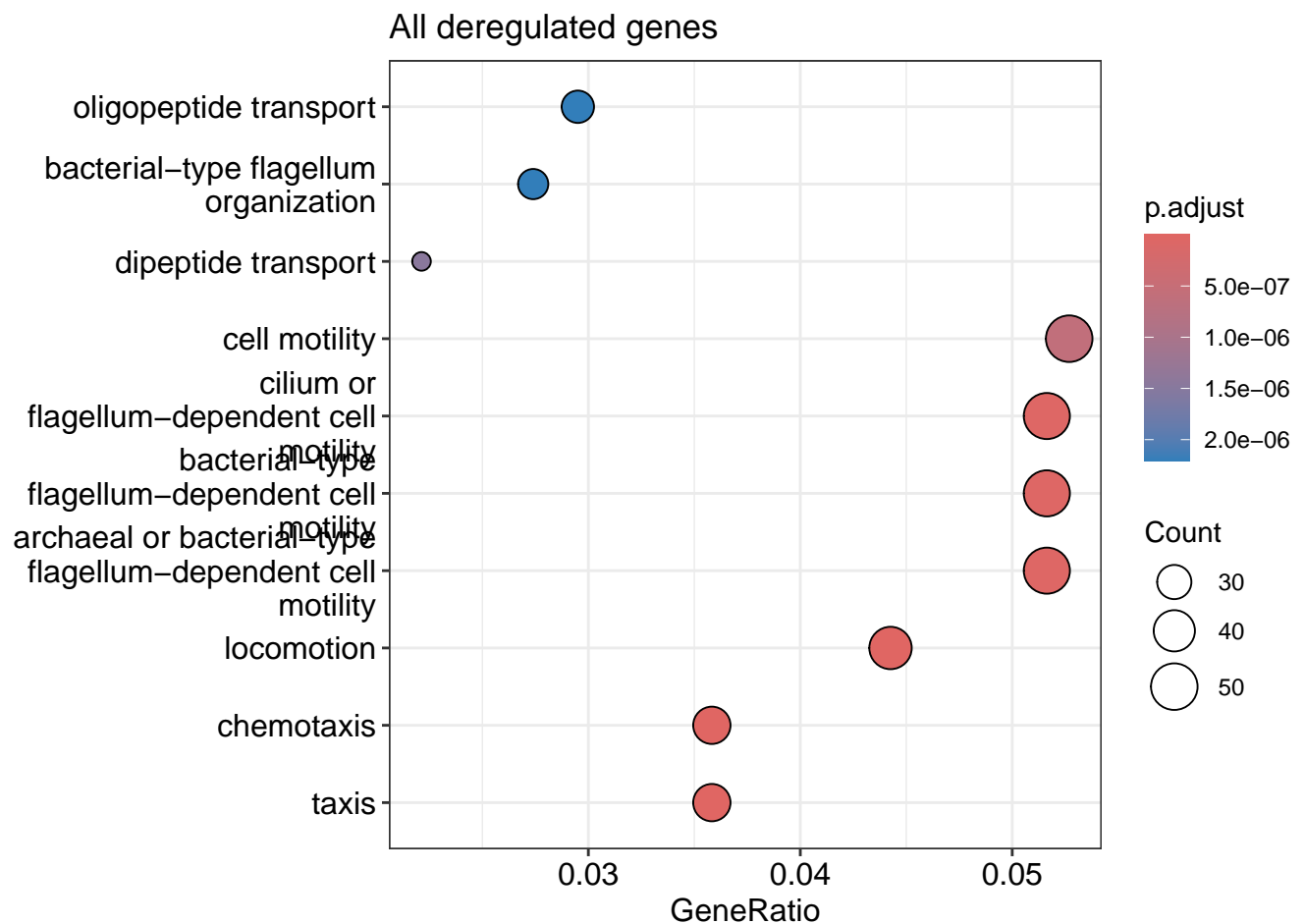
```

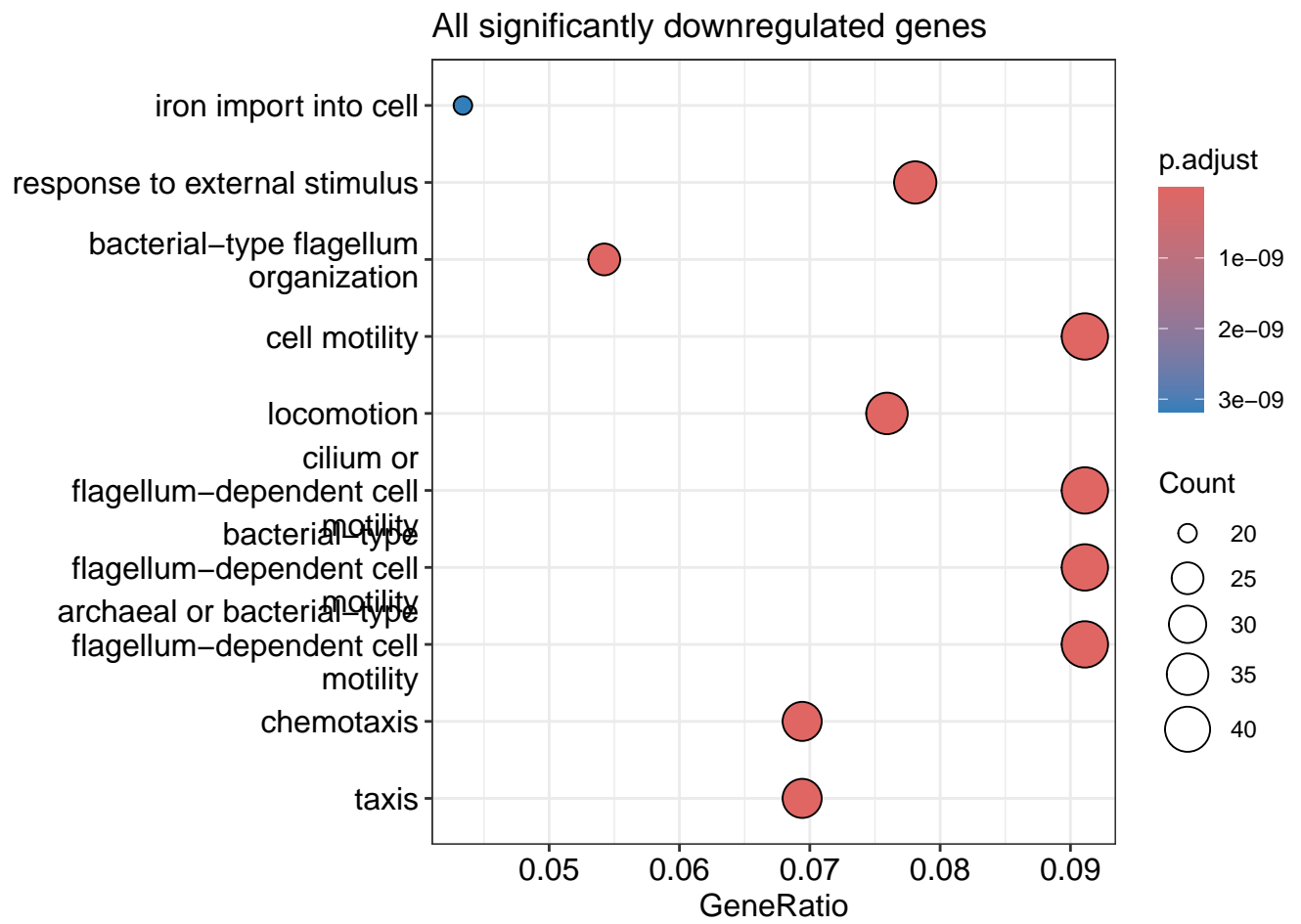
```
## [1] "5.66136528139081e-124" "fliM"
```

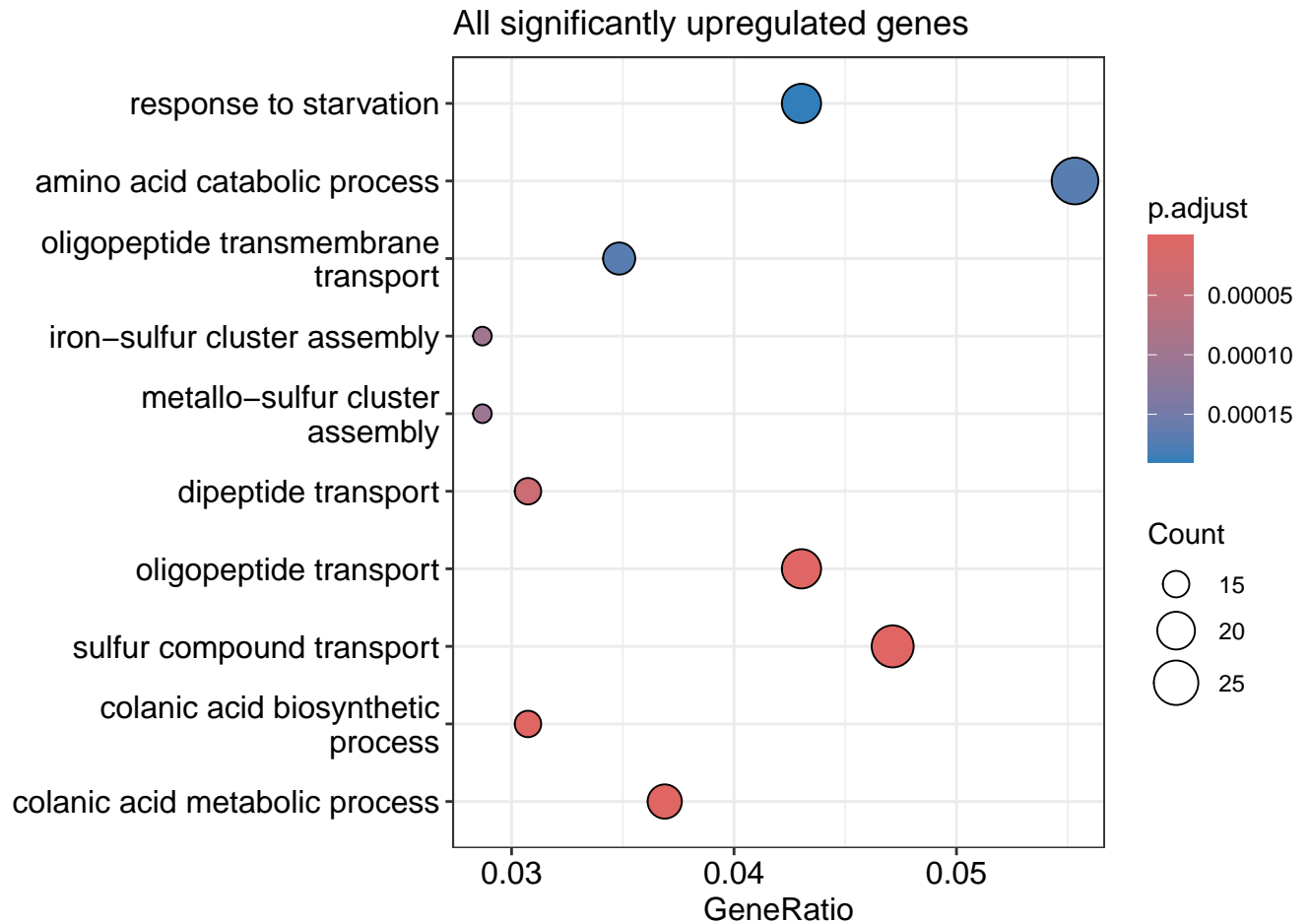
Answer: The gene with the lowest p-value (the most significant deregulation): *fliM* with a **p-value** = **5.66e-124**.

10.2 Dotplot

a) Which is the most significantly enriched GO term ?







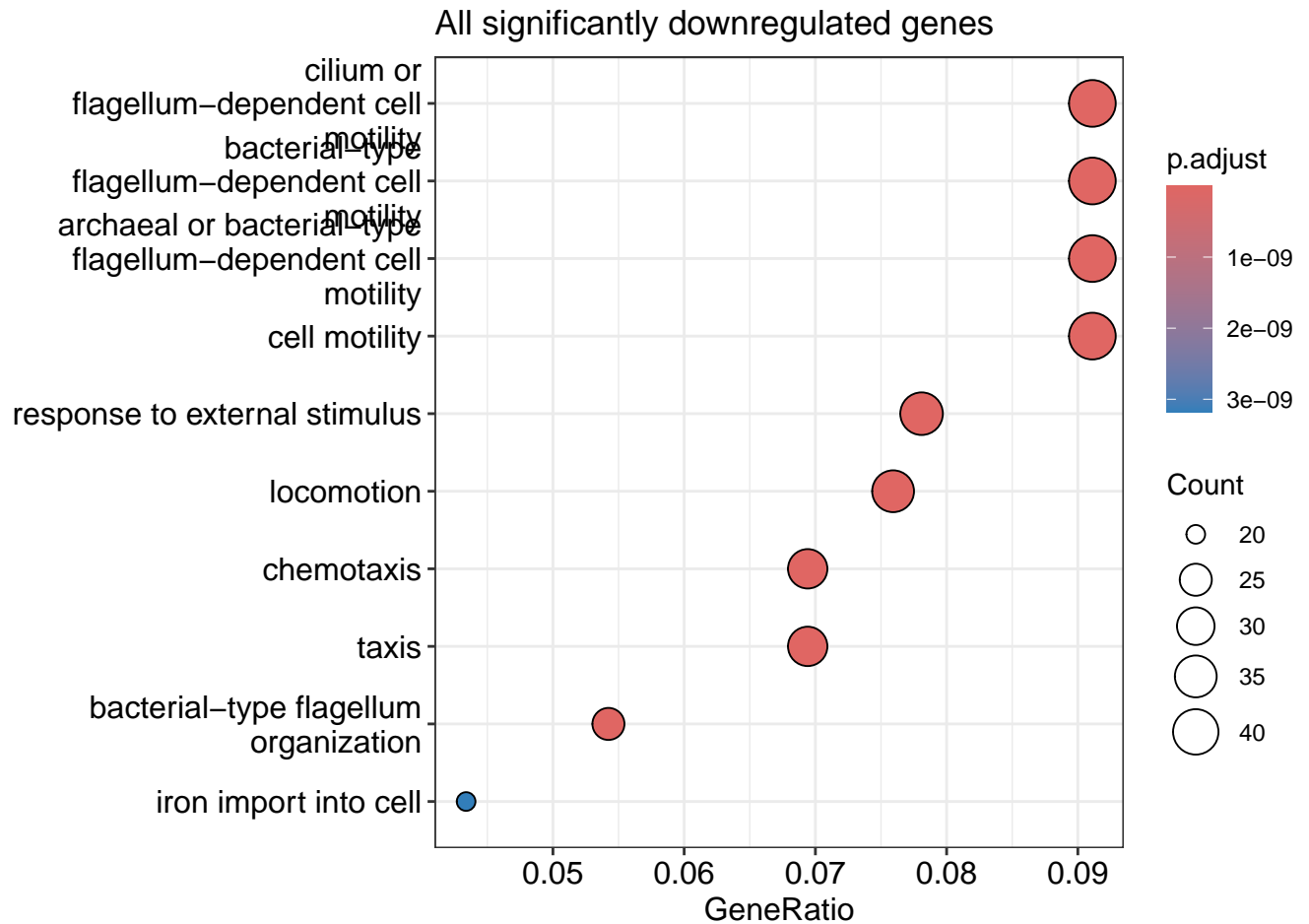
a) Which is the most significantly enriched GO term in terms of upregulated genes?

Answer: By looking at the graphs, which are sorted by adjusted p-values, we can see that the **genes for response to starvation** have the lowest p-value, therefore are the most significant upregulated GO term.

b) Which is the most significantly enriched GO term in terms of downregulated genes?

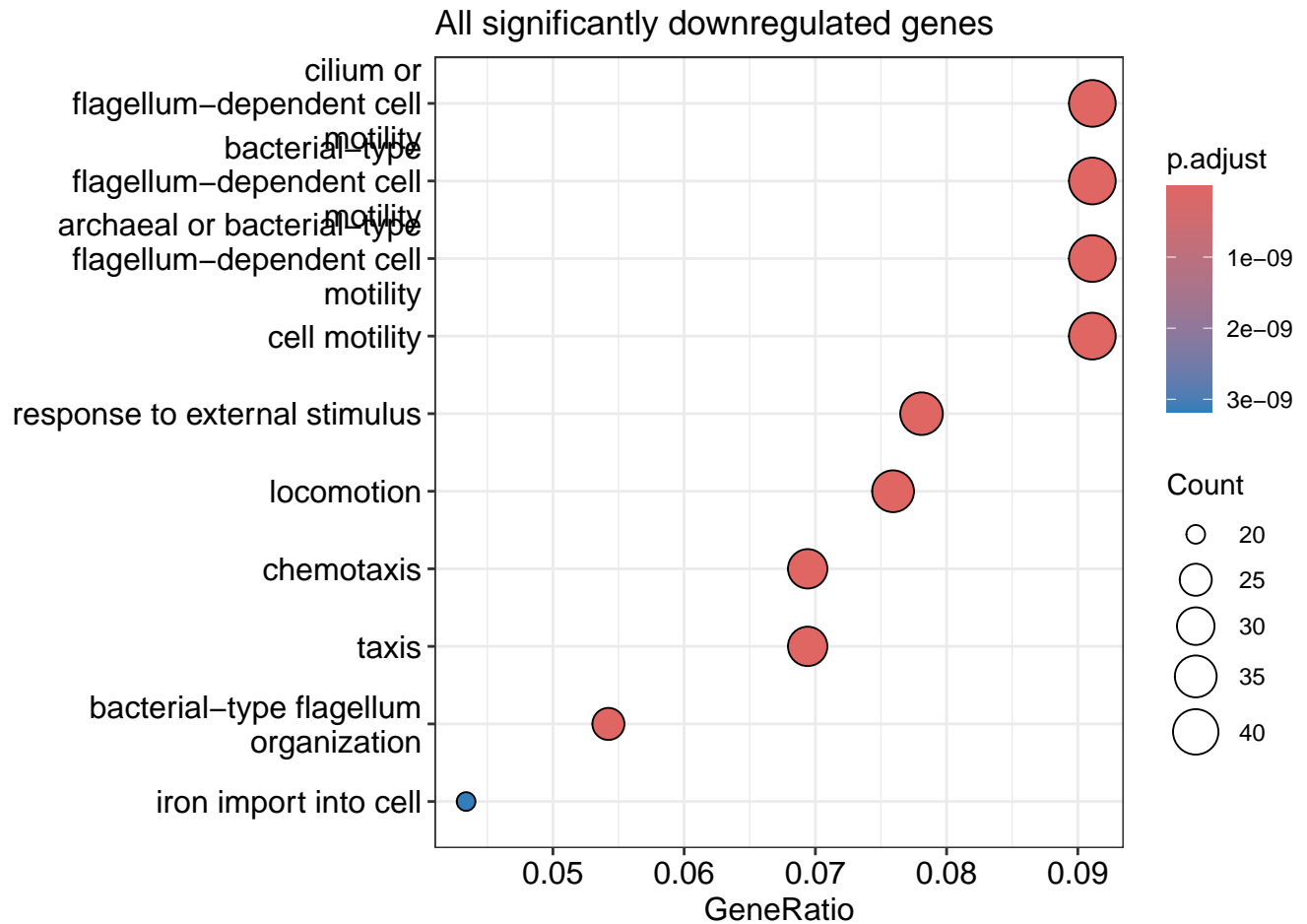
Answer: By looking at the graphs, which are sorted by adjusted p-values, we can see that the **genes for iron import into the cell** have the lowest p-value, therefore are the most significant downregulated GO term.

c) Which of the shown GO terms has the highest GeneRatio ?



Answer: By looking at the graphs, which are sorted by GeneRatio, we can see that a gene cluster with mobility functions like cell motility or flagellum-dependent cell motility rank highest in terms of GeneRatio.

d) Which of the shown GO terms has the highest Count ?

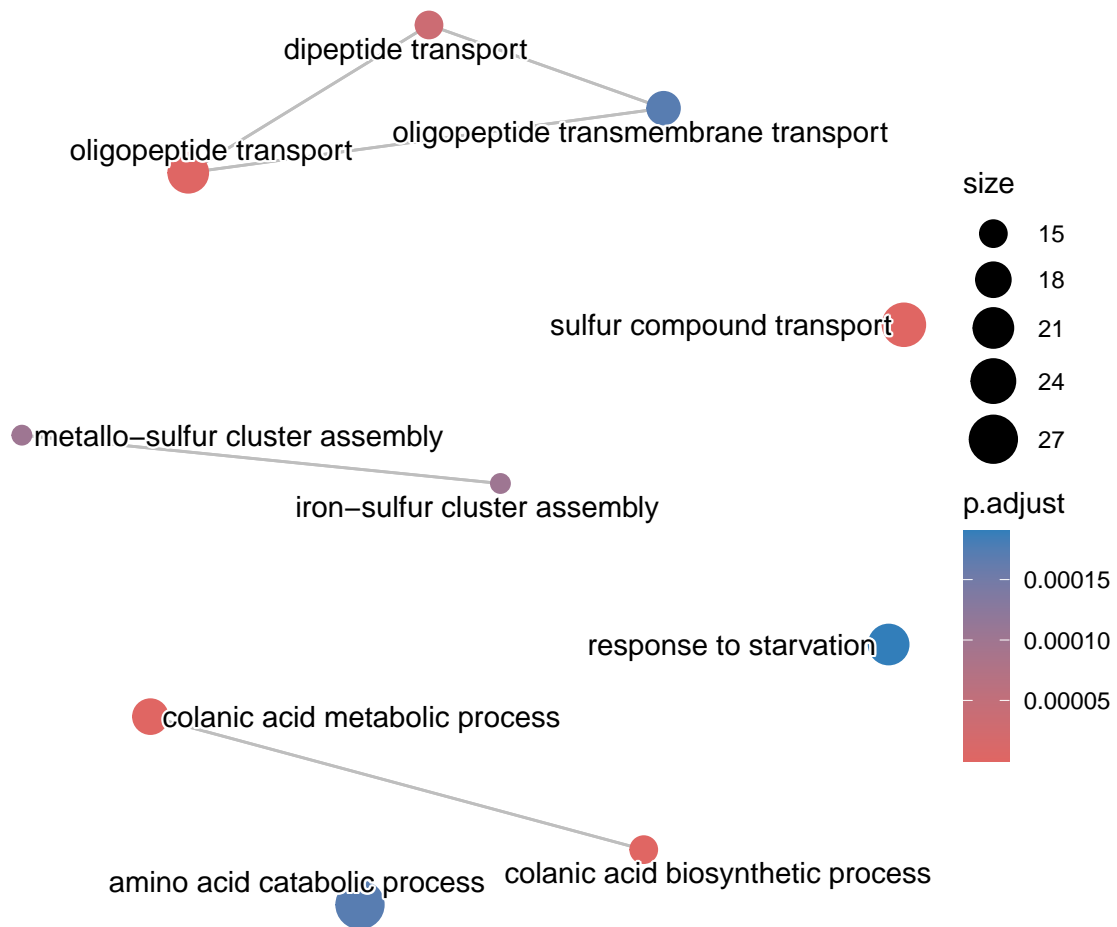


Answer: By looking at the graphs, which are sorted by Counts, we can see that **the same gene cluster with mobility functions like cell motility or flagellum-dependent cell motility** rank also highest in terms of Counts.

10.3 Clustering by GO relations

a) How many clusters can you identify when showing 10, 20, and 30 categories

```
enrichedBP <- enrichplot::pairwise_termsim(orUpBP)
emapplot(enrichedBP, showCategory = 10)
```



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