## Functional analysis of gene lists

### Ferran Briansó and Alex Sánchez-Pla. Statistics department. UB & Statistics and Bioinformatics Unit (UEB). VHIR.

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### 1 Introduction

This document provides information on how to extract subsets of genes from previously available gene lists by setting different filtering conditions such as the fold change, the p-value or the availability of Entrez identifier.

#### 1.1 From gene lists to Functional Analysis

The main, but not the only, goal of creating a gene list is to use it as input for some type of functional analysis such as Enrichment Analysis (ORA) or Gene set Enrichment Analysis.

Functional analysis can be made, on a first approach on

- A list of genes selected by being differentially expressed in a given experimental setting.
- The whole list of genes -or even the whole expression matrix- that has been used in the analysis.

Most tools require that gene list consist of gene identifiers in some standard notation such as Entrez, ENSEMBL or other related to these.

These gene lists can be easily extracted from output tables provided by microarrays or RNA-seq data analysis tools.

The analysis below is applied on a set of three gene lists obtained from a renal cancer study, but it can be easily extended to more lists or other studies.

```
x1<- AvsB <- read.table(file.path(resultsDir, "ExpressAndTop_AvsB.csv2"), head=T, sep=";", or a sep=";", or a
x2<- AvsL <- read.table(file.path(resultsDir, "ExpressAndTop_AvsL.csv2"), head=T, sep=";", or read.table(file.path(resultsDir, "ExpressAndTop_AvsL.csv2"))</pre>
x3<- BvsL <- read.table(file.path(resultsDir, "ExpressAndTop_BvsL.csv2"), head=T, sep=";", or sep=";",
dim(x1);
## [1] 6221
                                                18
cat("\nHeader of top Table for comparison AvsB\n")
## Header of top Table for comparison AvsB
cat("----\n")
## -----
head(x1[1:10, 1:8])
                                                    SymbolsA EntrezsA logFC AveExpr t
                                                        FOXA1 3169 -3.038344 8.651157 -14.362164 5.741793e-11
## 204667_at
## 215729_s_at VGLL1 51442 3.452290 6.137595 12.814829 3.439769e-10
## 220192_x_at SPDEF 25803 -3.016315 9.521883 -10.859194 4.337504e-09
## 214451_at TFAP2B
## 217528_at CLCA2
                                                                                              7021 -5.665059 7.432823 -10.829548 4.519412e-09
                                                                                        9635 -5.622086 6.763101 -9.666128 2.431610e-08
## 217284_x_at SERHL2 253190 -4.313116 9.133307 -9.528373 2.996253e-08
##
                                                           adj.P.Val
                                                                                                                                R
## 204667_at 3.571969e-07 14.648730
## 215729_s_at 1.069940e-06 13.148992
## 220192_x_at 7.028816e-06 10.928314
## 214451_at 7.028816e-06 10.891489
## 217528_at 3.025409e-05 9.363419
## 217284_x_at 3.106615e-05 9.171294
cat("\nHeader of top Table for comparison AvsL\n")
## Header of top Table for comparison AvsL
cat("----\n")
## -----
dim(x2); head(x2[1:10, 1:8])
## [1] 6221 18
                                                 SymbolsA EntrezsA logFC AveExpr
                                                                                                                                                                                                   t
## 205009_at TFF1 7031 4.735050 8.692478 10.564670 6.548729e-09
```

```
## 205862_at
             GREB1
                      9687 3.958563 6.082835 9.983993 1.513906e-08
## 205225_at
                 ESR1
                          2099 3.964939 9.300546 9.849787 1.846739e-08
## 209443_at
             SERPINA5
                          5104 2.198392 7.586226 8.531873 1.448630e-07
## 217528_at
              CLCA2
                          9635 -4.429254 6.763101 -7.615275 6.877151e-07
                          2674 2.333785 6.239876 7.600491 7.058428e-07
## 205696_s_at
                GFRA1
##
                adj.P.Val
                                  В
## 205009_at
             3.829521e-05 10.133204
## 205862_at
             3.829521e-05 9.434088
## 205225_at
             3.829521e-05 9.266376
## 209443_at
              2.252982e-04
                          7.488507
## 217528_at
             7.318414e-04 6.101859
## 205696_s_at 7.318414e-04 6.078427
cat("\nHeader of top Table for comparison BvsL\n")
##
## Header of top Table for comparison BvsL
## -----
dim(x3); head(x3[1:10, 1:8])
## [1] 6221
             18
##
              SymbolsA EntrezsA
                                 logFC
                                         AveExpr
                                                                 P. Value
                                                         t
## 204667_at
              FOXA1
                         3169 2.961042 8.651157 13.996760 8.630583e-11
## 215729_s_at VGLL1 51442 -3.744599 6.137595 -13.899875 9.630024e-11
## 205009_at
                TFF1
                         7031 5.729322 8.692478 12.783054 3.575181e-10
## 205225_at
                ESR1
                          2099 3.939276
                                        9.300546
                                                  9.786035 2.030957e-08
## 205862_at
                GREB1
                         9687 3.774303 6.082835
                                                  9.519268 3.038123e-08
## 218211_s_at
                MLPH
                         79083 2.808408 10.932769
                                                  8.813968 9.162548e-08
##
                adj.P.Val
## 204667_at
            2.995419e-07 14.040383
## 215729_s_at 2.995419e-07 13.953517
## 205009 at 7.413733e-07 12.893342
             3.158646e-05 9.422443
## 205225_at
## 205862_at
              3.780032e-05 9.061947
## 218211_s_at 8.534352e-05 8.062668
```

## 2 Input data preprocessing

Sometimes lists may need some preprocessing (e.g. in this example the gene list has multiple transcripts per gene identifier that have to be unitized previous to the analysis).

source("https://raw.githubusercontent.com/alexsanchezpla/scripts/master/usefulFunctions/gene source("https://raw.githubusercontent.com/alexsanchezpla/scripts/master/usefulFunctions/extrapressions)

We can use the available functions to extract only the gene lists

```
geneList1 <- genesFromTopTable (x1, entrezOnly = TRUE, uniqueIds=TRUE,</pre>
                                  adjOrrawP = "adj", Pcutoff = 0.1, FCcutoff = .75,
                                  id2Select = "EntrezsA" , cols2Select =3)
length(geneList1)
## [1] 874
geneList1Up <- genesFromTopTable (x1, entrezOnly = TRUE, uniqueIds=TRUE,</pre>
                                  adjOrrawP = "adj", Pcutoff = 0.1, FCcutoff = .75, updown="t
                                  id2Select = "EntrezsA" , cols2Select =3)
length(geneList1Up)
## [1] 534
geneList1Down <- genesFromTopTable (x1, entrezOnly = TRUE, uniqueIds=TRUE,</pre>
                                  adjOrrawP = "adj", Pcutoff = 0.1, FCcutoff = .75, updown="c
                                  id2Select = "EntrezsA" , cols2Select =3)
length(geneList1Down)
## [1] 340
geneList2 <- genesFromTopTable (x2, entrezOnly = TRUE, uniqueIds=TRUE,</pre>
                                  adjOrrawP = "adj", Pcutoff = 0.1, FCcutoff = .75,
                                  id2Select = "EntrezsA" , cols2Select =3)
geneList3 <- genesFromTopTable (x3, entrezOnly = TRUE, uniqueIds=TRUE,</pre>
                                  adjOrrawP = "adj", Pcutoff = 0.1, FCcutoff = .75,
                                  id2Select = "EntrezsA" , cols2Select =3)
```

Another possibility is to use function extractInfo do a "batch extraction"

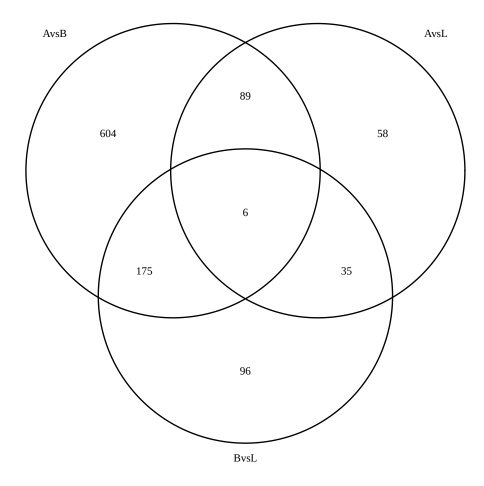
```
universeList2 <-List2[[2]]; geneList2<- List2[[1]];
cat("\nNumber of genes selectable (AvsL) with adjusted p-value < 0.1 and logFC > 0.75", leng
##
## Number of genes selectable (AvsL) with adjusted p-value < 0.1 and logFC > 0.75 188

List3 <- extractInfo(x3, "BvsL", "B|L", resultsDir, adjOrraw="adj", pCutOff=0.1, fcCutoff=...
universeList3 <-List3[[2]]; geneList3<- List3[[1]];
cat("\nNumber of genes selectable (BvsL) with adjusted p-value < 0.1 and logFC > 0.75", leng
##
## Number of genes selectable (BvsL) with adjusted p-value < 0.1 and logFC > 0.75 312

# test
# pattern <- "WL/PS"; cols2select<- grep(pattern, colnames(x1)); colnames(x1)[cols2select]
# pattern <- "WL\\.M/PS\\.M"; cols2select<- grep(pattern, colnames(x1M)); colnames(x1M)[cols2select]
# pattern <- "WL\\.F/PS\\.F"; cols2select<- grep(pattern, colnames(x1F)); colnames(x1F)[cols2select]</pre>
```

The following diagram shows which genes there are in common (or not) between the three lists.

```
require(VennDiagram)
vd2<- venn.diagram(list(AvsB=geneList1, AvsL=geneList2, BvsL=geneList3), filename=NULL)
grid.draw(vd2)</pre>
```



```
dev.off()
## null device
## 1
```

# 3 Case study

Imagine a user wants to do the following analysis:

1. Select three lists from my study (In this example we choose AvsB, AvsL, BvsL) We can do a preliminar optional filtering to keep only genes with Entrez Identifier and remove duplicates keeping only the most variable one.

```
AvsB0 <- genesFromTopTable (AvsB, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj" AvsL0 <- genesFromTopTable (AvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj" BvsL0 <- genesFromTopTable (BvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj"
```

2. Filter lists with adjusted-p-value less than 0.05

```
AvsB1 <- genesFromTopTable (AvsB, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj' AvsL1 <- genesFromTopTable (AvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj' BvsL1 <- genesFromTopTable (BvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "adj' cat("AvsB: ", length(AvsB0), "-->", length(AvsB1), "\n")

## AvsB: 708 --> 434

cat("AvsL: ", length(AvsL0), "-->", length(AvsL1), "\n")

## AvsL: 336 --> 80

cat("BvsL: ", length(BvsL0), "-->", length(BvsL1), "\n")

## BvsL: 412 --> 132
```

3. Create separate lists with up and down regulated genes

```
AvsB1Up <- genesFromTopTable (AvsB, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "ad AvsL1Up <- genesFromTopTable (AvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "ad BvsL1Up <- genesFromTopTable (BvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "ad Cat("AvsB: ", length(AvsB1), "-->", length(AvsB1Up), "\n")

## AvsB: 434 --> 243

cat("AvsL: ", length(AvsL1), "-->", length(AvsL1Up), "\n")

## AvsL: 80 --> 44

cat("BvsL: ", length(BvsL1), "-->", length(BvsL1Up), "\n")

## BvsL: 132 --> 77

AvsB1Down <- genesFromTopTable (AvsB, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "AvsL1Down <- genesFromTopTable (AvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "BvsL1Down <- genesFromTopTable (BvsL, entrezOnly = TRUE, uniqueIds=TRUE, adjOrrawP = "cat("AvsB: ", length(AvsB1), "-->", length(AvsB1Down), "\n")
```

```
## AvsB: 434 --> 191

cat("AvsL: ", length(AvsL1), "-->", length(AvsL1Down), "\n")

## AvsL: 80 --> 36

cat("BvsL: ", length(BvsL1), "-->", length(BvsL1Down), "\n")

## BvsL: 132 --> 55
```

4. Create a gene list with genes shared by AvsL and BvsL

```
commonAvsLandBvsL <- intersect(AvsL0, BvsL0)
length(commonAvsLandBvsL)
## [1] 104</pre>
```