

Exploratory Data Analysis

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Introduction

The aim of this project is to find links between certain gene expressions and familial Alzheimer's disease, using machine learning. To be more specific for the sake of data analysis: The mutation being observed is a presenilin 2 mutation, using patient-specific induced pluripotent stem cells (iPSC) to facilitate expression of the mutant type. Four different expression profiles were collected, using the Affymetrix Human Genome U133 Plus 2.0 Array. When looking at the names of columns, genes and differing values of expression, it's important to consider those are all Affymetrix standards, which may need to be converted to further down the line. For example: Converting the gene IDs to ensembl IDs.

Initial Data and Variables

Let's first take a look at the provided .csv file, it's structure and first entries.

```
raw.df = read.csv("../data/GSE28379.csv")
head(raw.df, 5)
```

```
##      ID_REF GSM701542 GSM701543 GSM701544 GSM701545 no.mutation mutation
## 1 1007_s_at 615.52540 739.77800 720.90040 735.84750   677.65170 728.3740
## 2 1053_at 319.87120 654.39166 319.87140 319.87150   487.13143 319.8714
## 3 117_at 20.04304 32.15144 14.41752 24.94408    26.09724 19.6808
## 4 121_at 239.84415 171.02960 137.31161 176.75978   205.43687 157.0357
## 5 1255_g_at 155.14342 335.75186 177.99786 128.04279   245.44764 153.0203
##      log.2.fold.change fold.change
## 1      0.1041354    1.0748500
## 2     -0.6068188    0.6566430
## 3     -0.4071086    0.7541332
## 4     -0.3876026    0.7643988
## 5     -0.6816920    0.6234337
```

ID_Ref.

This column indicates the probe ID's, as sequenced by the Affymetrix Human Genome U133 Plus 2.0 Array. This is the result of the sequencing technique. These are probe ID's, which don't represent a lot by themselves. They can, however, be used to find the ensembl ID's and gene symbols, which will be attempted below with Bioconductor:

```
genes <- select(hgu133plus2.db, c(raw.df[,1]), c("SYMBOL", "ENTREZID", "GENENAME", "ONTOLOGY", "PATH"))
```

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

```
na.list <- genes[is.na(genes$SYMBOL),] # Storing the genes which were not detected for whatever reason.
colnames(genes)[1] <- "ID_REF" # Renaming the column so the following merge works.
main <- merge(raw.df, genes, by=c("ID_REF")) # Merging the two dataframes together into a new one.
main <- main[!(main$ID_REF == "!series_matrix_table_end"),] # Removing an indicator row.
head(main[,c(1,10,11,12)], 5)
```

```
##      ID_REF SYMBOL ENTREZID GENENAME
## 2 1007_s_at  DDR1      780 discoidin domain receptor tyrosine kinase 1
## 3 1007_s_at  DDR1      780 discoidin domain receptor tyrosine kinase 1
## 4 1007_s_at  DDR1      780 discoidin domain receptor tyrosine kinase 1
## 5 1053_at   RFC2      5982      replication factor C subunit 2
## 6 1053_at   RFC2      5982      replication factor C subunit 2
```

To summarize what has just been done: A bioconductor database was used to find the corresponding gene for every probe. It's important to note that for yet unknown reasons some probes were not recognized. The proper database was used, as can be seen by the database name.

GSM

The GSM columns indicate the different samples used in the paper this data is derived from. The values under these columns represent sequencing concentration, which is the result of a normalisation algorithm called MAS5.0. This algorithm is also developed by Affymetrix. These values are also not log transformed. A dedicated column was made for that.

The first two samples, GSM701542 and GSM701543 are iPSC sequences derived from Sporadic Parkinson's disease patients. The latter two, GSM701544 and GSM701545 are iPSC sequences from familial Alzheimer's disease (FAD) patients. The referenced paper aimed to compare the two conditions and their gene expressions.

No mutation & Mutation

The first column, no mutation, signifies the average of the first two non mutated parkinson's samples. The second column, mutation, shows the average of the two FAD mutant type samples.

Log2 Fold Change

These are subtracted 2log fold change values, showing which of the two averages are up regulated and down regulated. In case of a positive number, the mutation type samples are up regulated and the non-mutant types are down regulated. The reverse is true in case of a negative number.

Fold change

The ratio between the mutation and no mutation values. Mutation being divided by no mutation in this case.

Data variance

The original paper aimed to compare two groups and their expressions. By looking at the calculated Log2FC data, it'll be possible to see how much the two groups differ. Let's first look at the most significant differently expressing genes.

```
main <- main[order(main$log.2.fold.change),] # Reordering the DF by log2FC
head(main[,c(10,6,7,8)], 10) #Showing the 10 most significant down regulated genes
```

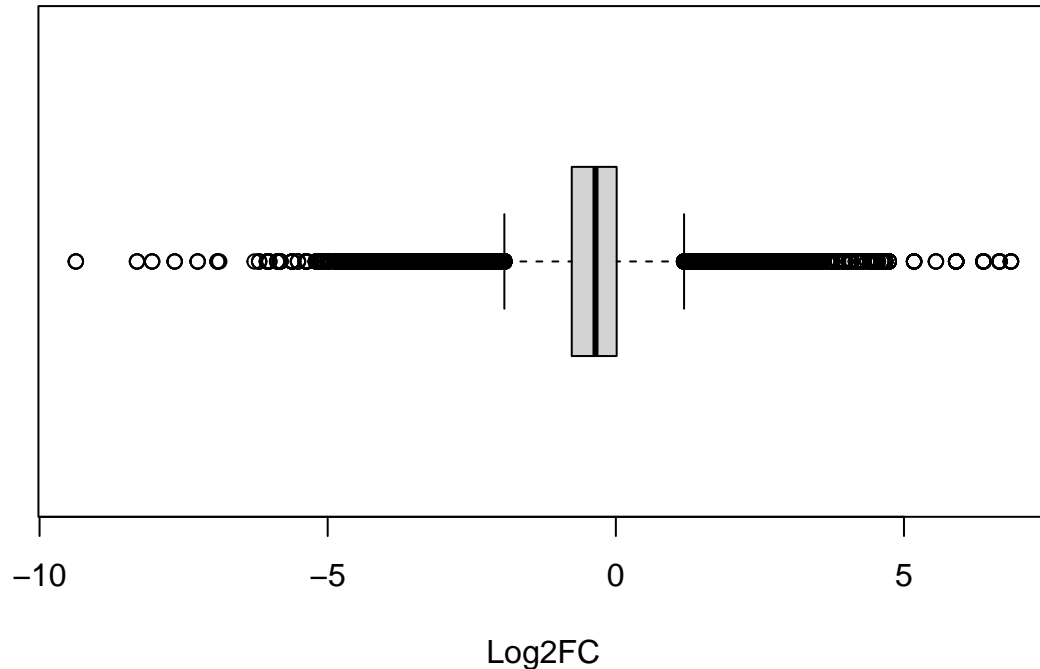
##	SYMBOL	no.mutation	mutation	log.2.fold.change
## 34889	RPS4Y1	3377.05241	5.1017696	-9.370551
## 34890	RPS4Y1	3377.05241	5.1017696	-9.370551
## 34891	RPS4Y1	3377.05241	5.1017696	-9.370551
## 54303	DDX3Y	1358.47492	4.2837637	-8.308893
## 54304	DDX3Y	1358.47492	4.2837637	-8.308893
## 54305	DDX3Y	1358.47492	4.2837637	-8.308893
## 50657	EIF1AY	217.13481	0.8207200	-8.047485
## 50658	EIF1AY	217.13481	0.8207200	-8.047485
## 50659	EIF1AY	217.13481	0.8207200	-8.047485
## 89983	ZNF257	45.97356	0.2280066	-7.655585

```
tail(main[,c(10,6,7,8)], 10) #Showing the 10 most significant up regulated genes
```

##	SYMBOL	no.mutation	mutation	log.2.fold.change
## 94100	CASP1	1.0538908	106.37357	6.65727
## 94101	CASP1	1.0538908	106.37357	6.65727
## 94102	CASP1	1.0538908	106.37357	6.65727
## 94103	CASP1	1.0538908	106.37357	6.65727
## 56957	BMP5	0.4355889	50.47331	6.85641
## 56958	BMP5	0.4355889	50.47331	6.85641
## 56959	BMP5	0.4355889	50.47331	6.85641
## 56960	BMP5	0.4355889	50.47331	6.85641
## 56961	BMP5	0.4355889	50.47331	6.85641
## 56962	BMP5	0.4355889	50.47331	6.85641

By taking a glance at the tables above, it seems that down regulation consists of more extreme values than up regulation. While this may be telling of how expression is affected by the mutation in general, it's not enough on its own to draw any conclusions yet. Let's further explore the log2fc values by creating a boxplot.

```
boxplot(main$log.2.fold.change, horizontal = TRUE, xlab = "Log2FC", title = "Boxplot of Log2FC")
```



As can be seen from the boxplot, there appears to be great variance in the log2FC values. This may be further examined by looking at a summary.

```
summary(main$log.2.fold.change)
```

```
##      Min.   1st Qu.   Median     Mean   3rd Qu.     Max.
## -9.37055 -0.76497 -0.35343 -0.38615  0.01466  6.85641
```

Judging by the boxplot and summary table, the genes seem to be mostly down regulated.

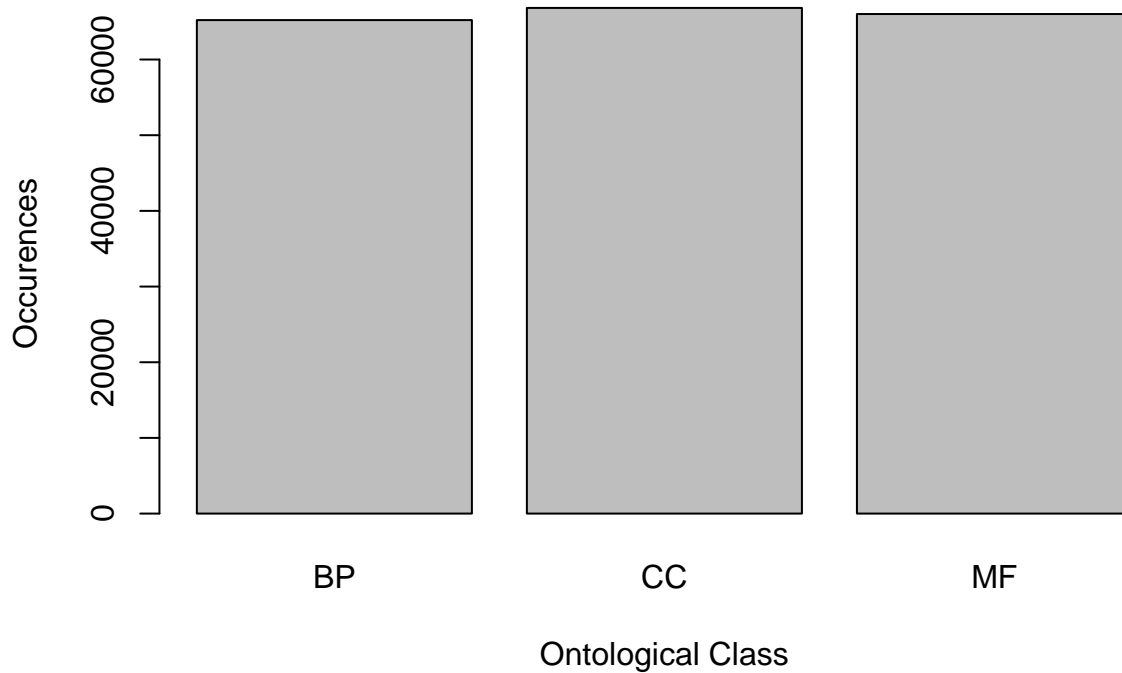
Relations

Since the data provided is that of solely DEG's, there will not be any variables that affect each other, besides the obvious calculation of fold changes and means. That being said: There could be a connection between up/down regulation of certain genetic regions. Taking RPS4Y1, a gene encoding for a ribosomal sub unit, as an example: Could there be a trend in the mutation affecting down/up regulation of specifically ribosomal genes? Questions like these are not suited for EDA, but the addition of gene symbols, KEGG

pathway, ontology and entrezID's columns in this document will help in answering such questions, which will eventually mean answering the research question.

Plotting the different KEGG IDs and ontological vlsddrd can however offer insight into which pathways and classes are most prevalent.

```
barplot(table(main$ONTOLOGY), xlab = "Ontological Class", ylab = "Occurences")
```



The differing classes appear to be close in the amount of occurrences. It may be possible to find a relationship when looking at the Log2FC values for every different class.

```
BP <- subset(main, ONTOLOGY == "BP")
CC <- subset(main, ONTOLOGY == "CC")
MF <- subset(main, ONTOLOGY == "MF")
```

```
summary(BP$log.2.fold.change)
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## -9.3706 -0.7440 -0.3463 -0.3775  0.0141  6.8564
```

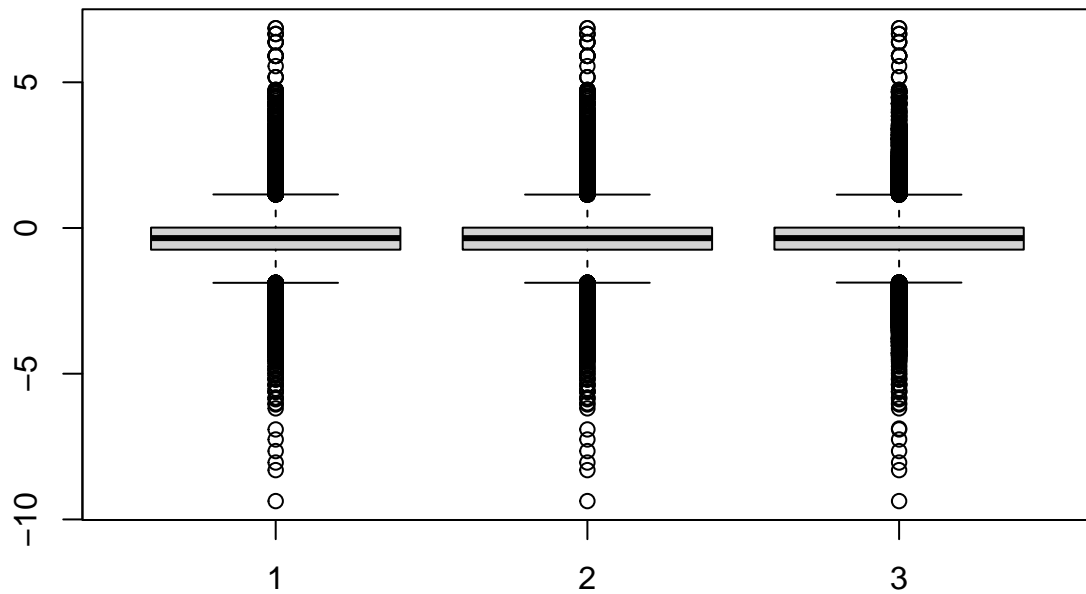
```
summary(CC$log.2.fold.change)
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## -9.37055 -0.74445 -0.34809 -0.37869  0.01193  6.85641
```

```
summary(MF$log.2.fold.change)
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## -9.37055 -0.74151 -0.34626 -0.37680  0.01279  6.85641
```

```
boxplot(BP$log.2.fold.change, CC$log.2.fold.change, MF$log.2.fold.change)
```



Both the summaries and boxplots are very similar for every classification. At first glance this indicates that the classes are not a detriment in the fold change values of genes.