## Analysis Report (Part6): Epigenetics Analysis

(Last updated: 11/05/21)

The following R script gets the percentage of overlap between the promoters of differentially expressed genes found in Report-P5-Differential\_Expression\_Analysis.pdf and the epigenetically marked (H3K4me3) promoters in the fetal and adult brain cells, and in liver cell lines which are retrieved from Annotationhub (EpigenomeRoadMap).

## 1. Loading the Necessary Libraries and Data found in Report-P5-Differential\_Expression\_Analysis.pdf

## 2. Get the Promoter Associated Histone Modifications (H3k4me3) Narrow Peaks Data from Annotationhub (EpigenomeRoadMap)

```
# initialize AnnotationHub object
ah = AnnotationHub()
ah = subset(ah, species == "Homo sapiens")
# get the narrow peaks data (promoter associated histone modification H3K4me3) for the fetal brain cell
fetal.brain = AnnotationHub::query(ah, c("EpigenomeRoadMap", "H3K4me3", "E081"))
fetal.brain.gr = fetal.brain[[2]]
# get the narrow peaks data (promoter associated histone modification H3K4me3) for the adult brain cell
adult.brain = AnnotationHub::query(ah, c("EpigenomeRoadMap", "H3K4me3", "E073"))
adult.brain.gr = adult.brain[[2]]
# get the narrow peak data (promoter associated histone modification H3K4me3) for liver cell line
liver.line = AnnotationHub::query(ah, c("EpigenomeRoadMap", "H3K4me3", "Liver"))
liver.line.gr = liver.line[[2]]
```

## 3. Change ENSEMBL ID to ENTREZID that Matches gene\_id in TXDB

```
sorted\_significant\_differ\_res\_deseq2\_shrunk\$gene = sub("\\\\"", as.character(sorted\_significant\_differ\_res\_deseq2\_shrunk\$gene) = sub("\\\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\) = sub("\\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\\) = sub("\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\\) = sub("\\\) = sub("\\\\) = sub("\\\) = sub("\\) = sub("\\\) = sub("\\) = sub("\\) = sub("\\) = sub("\\) = sub("\\) = sub("\\) = sub("\) = sub
AnnotationDbi::keytypes(EnsDb.Hsapiens.v86)
              [1] "ENTREZID"
                                                                                                                                          "EXONID"
##
                                                                                                                                                                                                                                                "GENEBIOTYPE"
             [4] "GENEID"
                                                                                                                                          "GENENAME"
                                                                                                                                                                                                                                                "PROTDOMID"
##
               [7] "PROTEINDOMAINID"
                                                                                                                                          "PROTEINDOMAINSOURCE" "PROTEINID"
## [10] "SEQNAME"
                                                                                                                                          "SEQSTRAND"
                                                                                                                                                                                                                                                "SYMBOL"
## [13] "TXBIOTYPE"
                                                                                                                                          "TXID"
                                                                                                                                                                                                                                                "TXNAME"
## [16] "UNIPROTID"
sorted_significant_differ_res_deseq2_shrunk.map = AnnotationDbi::select(EnsDb.Hsapiens.v86,
                                                                                           keys = as.character(sorted_significant_differ_res_deseq2_shrunk$gene), keytype = "G
```

#### 4. Get the Promoters of DGEs Found in Report-P5-Differential\_Expression\_

```
# get the known genes from Tx database
txdb = TxDb.Hsapiens.UCSC.hg19.knownGene
txdb.genes = genes(txdb)
# get the promoters of differentially expressed genes
diff.expressed.gene.promoters = promoters(txdb.genes[txdb.genes$gene_id %in% sorted_significant_differ_identially.genes$gene_id %in% sorted_signif
```

# 5. Calculate the Percentage of Overlap Between the Promoters of DEGs Found in Report-P5-Differential\_Expression\_Analysis.pdf and the Epigenetically Marked (H3K4me3) Promoters in the 3 Different Kind of Cells

The percentage of overlap between the promoters of differentially expressed genes found in Report-P5-Differential\_Expression\_Analysis.pdf and the epigenetically marked (H3K4me3) promoters in the fetal brain cells is calculated as following:

```
# subsetByOverlaps() extracts the elements in the query (the first argument) that overlap at least one
fetal.brain.overlap.H3K4me3 = subsetByOverlaps(diff.expressed.gene.promoters, fetal.brain.gr)
(fetal.brain.overlap.percentage.H3K4me3 = length(fetal.brain.overlap.H3K4me3) / length(sorted_significant)
```

```
## [1] 34.77612
```

The percentage of overlap between the promoters of differentially expressed genes found in Report-P5-Differential\_Expression\_Analysis.pdf and the epigenetically marked (H3K4me3) promoters in the adult brain cells is calculated as following:

```
# subsetByOverlaps() extracts the elements in the query (the first argument) that overlap at least one
adult.brain.overlap.H3K4me3 = subsetByOverlaps(diff.expressed.gene.promoters, adult.brain.gr)
(adult.brain.overlap.percentage.H3K4me3 = length(adult.brain.overlap.H3K4me3) / length(sorted_significant)
```

## [1] 52.60448

The percentage of overlap between the promoters of differentially expressed genes found in Report-P5-Differential\_Expression\_Analysis.pdf and the epigenetically marked (H3K4me3) promoters in the liver cell line is calculated as following:

```
# subsetByOverlaps() extracts the elements in the query (the first argument) that overlap at least one
liver.line.overlap.H3K4me3 = subsetByOverlaps(diff.expressed.gene.promoters, liver.line.gr)
(liver.line.overlap.percentage.H3K4me3 = length(liver.line.overlap.H3K4me3) / length(sorted_significant)
```

## [1] 47.87313