# Hands on practical: Online databases exploration

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# Investigate age-related changes across multiple tissues

Over the past years, an enormous amount of omics data has become publically available. This data is freely available and can be used by researchers across the world. In this practical we will download genome-wide DNA methylation data from the Gene Expression Omnibus (GEO) for two tissues. We will use this data to investigate the relation between DNA methylation and age.

# Load packages

```
library(Biobase)
library(GEOquery)
library(ggplot2)
```

#### **Datasets**

We use two datasets. The first dataset is from Tsaprouni  $et\ al.$  Note that this is not the largest dataset available on GEO, but for for this pratical large enough. The accession number of this dataset is GSE50660. The accession number is a number that refers to a specific dataset on GEO and can be referred to in scientific publications. The second dataset is from Berko  $et\ al.$  with accession number GSE50759.

There are different ways to download data from GEO. Go to https://www.ncbi.nlm.nih.gov/geo/. On the right you can search for datasets. Search for *GSE50660*. This will take you to the page of the dataset. You can see the summary of the data/study, the authors, the paper the data is from, contact details and the number of samples.

# 1. How many samples are there? What is the tissue the samples were taken from?

There are different ways to get the data. One is GEO2R which allows direct parsing of data in R. You can also manually download the data at the bottom of the page.

#### 2. In what ways is the data available? Which one would you use?

Now we download the data from GEO. We can do this for both accession numbers. Note that this will take a few minutes to download and parse.

```
gset.blood <- getGEO("GSE50660", GSEMatrix =TRUE, getGPL=FALSE)
gset.buccal <- getGEO("GSE50759", GSEMatrix =TRUE, getGPL=FALSE)</pre>
```

This is gives back a list with one slot. This contains a ExpressionSet. This is a way to store multiple datatypes/phenotypes in one object.

```
length(gset.blood)
class(gset.blood[[1]])
eSet.blood <- gset.blood[[1]]
eSet.buccal <- gset.buccal[[1]]</pre>
```

We want to extract the DNA methylation levels of this dataset. That can be achieved with exprs(). The phenotypes can be extracted with pheno, which returns an AnnotatedDataFrame from which the phenotypes can be extracted using @data. See code below.

```
exprs.blood <- exprs(eSet.blood)
exprs.buccal <- exprs(eSet.buccal)

pheno.blood <- phenoData(eSet.blood)@data
pheno.buccal <- phenoData(eSet.buccal)@data</pre>
```

3. The DNA methylation data contains quite some loci. How many (code below)? Are the number of loci measured equal? If not, why do you think not?

```
dim(exprs.blood)
dim(exprs.buccal)
```

For the sake of time, only a subset of the data is investigated for a relation with age.

Note that in the chunk above, for buccal the values are still in M-values, which are different from beta-values. M-values are on a continuous scale, while beta values are easier to interpret, but less optimal in statistical analyses. For clarity we use beta values only here.

4. What are the dimensions of the new datasets?

```
dim(exprs.buccal)
dim(exprs.blood)
```

# Run the analysis

We are now ready for the actual analysis. For this we use function to loop over the CpGs. Otherwise we would have to many tests manually while now we can run them at once. See the function below.

```
get.aDMP <- function(CpG, data.in, samplesheet, phenotype)
{
    #Just to be sure make the phenotype numeric
    age <- as.numeric(as.character(samplesheet[,phenotype]))
    fit <- lm(data.in[CpG,]~age) # Run the LM
    fit.anova <- anova(fit) #Run the anova

coef <- fit$coefficients["age"] #Extract coef
    p.val <- fit.anova["age",] #Extract pvalue

out <- data.frame(CpG, coef, p.val) #Create a dataframe to output
    return(out)
}</pre>
```

5. Run the model for CpG cg16867657. What do you observe?

6. Plot this one CpG (see code below). What do you observe? Is this locus really associated with age? Compare the slope of the line, what do you observe?

7. Run the model for all CpGs. Look at the top results. What is the top locus for blood? Is it the same for buccal?

```
#Combine data to dataframe
results.blood <- as.data.frame(do.call(rbind, res.blood))
results.buccal <- as.data.frame(do.call(rbind, res.buccal))

#Sort on P-value
results.blood <- results.blood[order(results.blood$Pr..F., decreasing=F),]
results.buccal <- results.buccal[order(results.buccal$Pr..F., decreasing=F),]

#Look at the top
head(results.blood)
head(results.buccal)</pre>
```

8. How many CpGs are significant in each of the tissues? Adjust for the number of tests performed!

```
alpha <- 0.05/nrow(results.blood)
results.blood.sign <- results.blood[results.blood$Pr..F. <= alpha,]
results.buccal.sign <- results.buccal[results.buccal$Pr..F. <= alpha,]
nrow(results.blood.sign)
nrow(results.buccal.sign)</pre>
```

9. Plot the coefficients against each other. What do observe in this subset?

10. How many are shared between blood and buccal?

```
table(results.buccal.sign$CpG %in% results.blood.sign$CpG)
```

11. What do you notice in the figure of Q9 in terms of effect size? Look at the axis range. One could also add an effect size threshold to have biological relevant differences. Set the threshold to 2%/10 years. How many are significant and how many overlap?

```
results.blood.sign.eff <- results.blood.sign[results.blood.sign$coef >= 0.002,]
results.buccal.sign.eff <- results.buccal.sign[results.buccal.sign$coef >= 0.002,]
nrow(results.blood.sign.eff)
nrow(results.buccal.sign.eff)
```

table(results.blood.sign.eff\$CpG %in% results.buccal.sign.eff\$CpG)

- 12. One of the loci is cg16867657 which we looked at before. Go to UCSC (https://genome.ucsc.edu), Genomes Buid hg19. Look-up cg16867657. Zoom out, what protein coding gene(s) are close to this CpG? Is the CpG in a CpG island?
- 13. Go to GeneCards (https://www.genecards.org), search for the gene found in 11.. What is the function of the gene? Does that make sense?