

# Tutorial for the WGCNA package for R:

## I. Network analysis of liver expression data in female mice

### 1. Data input and cleaning

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### Contents

|          |   |          |
|----------|---|----------|
| <b>1</b> | <b>Data input, cleaning and pre-processing</b>  | <b>1</b> |
| 1.a      | Loading expression data . . . . .   | 1        |
| 1.b      | Checking data for excessive missing values and identification of outlier microarray samples . . . . . | 2        |
| 1.c      | Loading clinical trait data . . . . .   | 3        |

## 1 Data input, cleaning and pre-processing

This is the first step of any network analysis. We show here how to load typical expression data, pre-process them into a format suitable for network analysis, and clean the data by removing obvious outlier samples as well as genes and samples with excessive numbers of missing entries.

### 1.a Loading expression data

The expression data is contained in the file `LiverFemale3600.csv` that comes with this tutorial. After starting an R session, we load the requisite packages and the data, after appropriately setting the working directory:

```
# Display the current working directory
getwd();
# If necessary, change the path below to the directory where the data files are stored.
# "." means current directory. On Windows use a forward slash / instead of the usual \.
workingDir = ".";
setwd(workingDir);
# Load the package
library(WGCNA);
# The following setting is important, do not omit.
options(stringsAsFactors = FALSE);
# Read in the female liver data set
femData = read.csv("LiverFemale3600.csv");
# Take a quick look at what is in the data set:
dim(femData);
names(femData);
```

In addition to expression data, the data files contain extra information about the surveyed probes we do not need. One can inspect larger data frames such as `femData` by invoking R data editor via `fix(femData)`. The expression data set contains 135 samples. Note that each row corresponds to a gene and column to a sample or auxiliary information. We now remove the auxiliary data and transpose the expression data for further analysis.

```
datExpr0 = as.data.frame(t(femData[, -c(1:8)]));
names(datExpr0) = femData$substanceBXH;
rownames(datExpr0) = names(femData)[-c(1:8)];
```

## 1.b Checking data for excessive missing values and identification of outlier microarray samples

We first check for genes and samples with too many missing values:

```
gsg = goodSamplesGenes(datExpr0, verbose = 3);
gsg$allOK
```

If the last statement returns TRUE, all genes have passed the cuts. If not, we remove the offending genes and samples from the data:

```
if (!gsg$allOK)
{
  # Optionally, print the gene and sample names that were removed:
  if (sum(!gsg$goodGenes)>0)
    printFlush(paste("Removing genes:", paste(names(datExpr0)[!gsg$goodGenes], collapse = ", ")));
  if (sum(!gsg$goodSamples)>0)
    printFlush(paste("Removing samples:", paste(rownames(datExpr0)[!gsg$goodSamples], collapse = ", ")));
  # Remove the offending genes and samples from the data:
  datExpr0 = datExpr0[gsg$goodSamples, gsg$goodGenes]
}
```

Next we cluster the samples (in contrast to clustering genes that will come later) to see if there are any obvious outliers. We use the function `flashClust` that provides faster hierarchical clustering than the standard function `hclust`.

```
sampleTree = flashClust(dist(datExpr0), method = "average");
# Plot the sample tree: Open a graphic output window of size 12 by 9 inches
# The user should change the dimensions if the window is too large or too small.
sizeGrWindow(12,9)
#pdf(file = "Plots/sampleClustering.pdf", width = 12, height = 9);
par(cex = 0.6);
par(mar = c(0,4,2,0))
plot(sampleTree, main = "Sample clustering to detect outliers", sub="", xlab="", cex.lab = 1.5,
      cex.axis = 1.5, cex.main = 2)
```

It appears there is one outlier (sample F2\_221, see Fig. 1). One can remove it by hand, or use an automatic approach. Choose a height cut that will remove the offending sample, say 15 (the red line in the plot), and use a branch cut at that height.

```
# Plot a line to show the cut
abline(h = 15, col = "red");
# Determine cluster under the line
clust = cutreeStatic(sampleTree, cutHeight = 15, minSize = 10)
table(clust)
# clust 1 contains the samples we want to keep.
keepSamples = (clust==1)
datExpr = datExpr0[keepSamples, ]
nGenes = ncol(datExpr)
nSamples = nrow(datExpr)
```

The variable `datExpr` now contains the expression data ready for network analysis.

## 1.c Loading clinical trait data

We now read in the trait data and match the samples for which they were measured to the expression samples.

```
traitData = read.csv("ClinicalTraits.csv");
dim(traitData)
names(traitData)

# remove columns that hold information we do not need.
allTraits = traitData[, -c(31, 16)];
allTraits = allTraits[, c(2, 11:36) ];
dim(allTraits)
names(allTraits)

# Form a data frame analogous to expression data that will hold the clinical traits.

femaleSamples = rownames(datExpr);
traitRows = match(femaleSamples, allTraits$Mice);
datTraits = allTraits[traitRows, -1];
rownames(datTraits) = allTraits[traitRows, 1];

collectGarbage();
```

We now have the expression data in the variable `datExpr`, and the corresponding clinical traits in the variable `datTraits`. Before we continue with network construction and module detection, we visualize how the clinical traits relate to the sample dendrogram.

```
# Re-cluster samples
sampleTree2 = flashClust(dist(datExpr), method = "average")
# Convert traits to a color representation: white means low, red means high, grey means missing entry
traitColors = numbers2colors(datTraits, signed = FALSE);
# Plot the sample dendrogram and the colors underneath.
plotDendroAndColors(sampleTree2, traitColors,
                    groupLabels = names(datTraits),
                    main = "Sample dendrogram and trait heatmap")
```

In the plot, shown in Fig. 2, white means a low value, red a high value, and grey a missing entry. The last step is to save the relevant expression and trait data for use in the next steps of the tutorial.

```
save(datExpr, datTraits, file = "FemaleLiver-01-dataInput.RData")
```



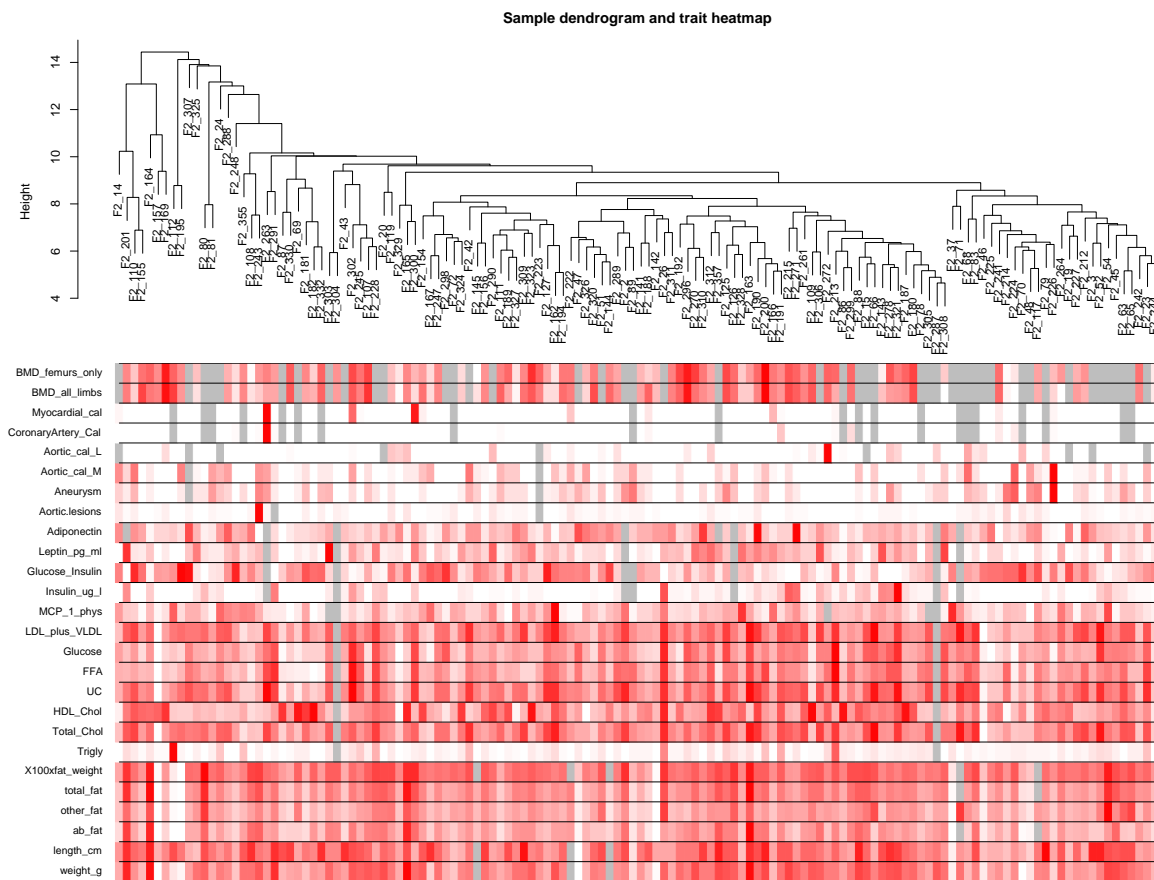


Figure 2: Clustering dendrogram of samples based on their Euclidean distance.