

# WGCNA Analysis

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Call libraries

```
library(WGCNA)
```

```
## Loading required package: dynamicTreeCut

## Loading required package: fastcluster

##
## Attaching package: 'fastcluster'

## The following object is masked from 'package:stats':
##
##      hclust

## Warning: replacing previous import 'utils::findMatches' by
## 'S4Vectors::findMatches' when loading 'AnnotationDbi'

##

##
## Attaching package: 'WGCNA'

## The following object is masked from 'package:stats':
##
##      cor
```

```
library(DESeq2)
```

```
## Loading required package: S4Vectors

## Loading required package: stats4

## Loading required package: BiocGenerics

##
## Attaching package: 'BiocGenerics'
```

```

## The following objects are masked from 'package:stats':
##
##     IQR, mad, sd, var, xtabs

## The following objects are masked from 'package:base':
##
##     anyDuplicated, append, as.data.frame, basename, cbind, colnames,
##     dirname, do.call, duplicated, eval, evalq, Filter, Find, get, grep,
##     grepl, intersect, is.unsorted, lapply, Map, mapply, match, mget,
##     order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank,
##     rbind, Reduce, rownames, sapply, setdiff, sort, table, tapply,
##     union, unique, unsplit, which.max, which.min

##
## Attaching package: 'S4Vectors'

## The following object is masked from 'package:utils':
##
##     findMatches

## The following objects are masked from 'package:base':
##
##     expand.grid, I, unname

## Loading required package: IRanges

##
## Attaching package: 'IRanges'

## The following object is masked from 'package:grDevices':
##
##     windows

## Loading required package: GenomicRanges

## Loading required package: GenomeInfoDb

## Loading required package: SummarizedExperiment

## Loading required package: MatrixGenerics

## Loading required package: matrixStats

##
## Attaching package: 'MatrixGenerics'

## The following objects are masked from 'package:matrixStats':
##
##     colAlls, colAnyNAs, colAnys, colAvgsPerRowSet, colCollapse,
##     colCounts, colCummaxs, colCummins, colCumprods, colCumsums,

```

```
##      colDiffs, colIQRDiffs, colIQRs, colLogSumExps, colMadDiffs,
##      colMads, colMaxs, colMeans2, colMedians, colMins, colOrderStats,
##      colProds, colQuantiles, colRanges, colRanks, colSdDiffs, colSds,
##      colSums2, colTabulates, colVarDiffs, colVars, colWeightedMads,
##      colWeightedMeans, colWeightedMedians, colWeightedSds,
##      colWeightedVars, rowAlls, rowAnyNAs, rowAnys, rowAvgsPerColSet,
##      rowCollapse, rowCounts, rowCummaxs, rowCummins, rowCumprods,
##      rowCumsums, rowDiffs, rowIQRDiffs, rowIQRs, rowLogSumExps,
##      rowMadDiffs, rowMads, rowMaxs, rowMeans2, rowMedians, rowMins,
##      rowOrderStats, rowProds, rowQuantiles, rowRanges, rowRanks,
##      rowSdDiffs, rowSds, rowSums2, rowTabulates, rowVarDiffs, rowVars,
##      rowWeightedMads, rowWeightedMeans, rowWeightedMedians,
##      rowWeightedSds, rowWeightedVars
```

```
## Loading required package: Biobase
```

```
## Welcome to Bioconductor
```

```
##
##      Vignettes contain introductory material; view with
##      'browseVignettes()'. To cite Bioconductor, see
##      'citation("Biobase")', and for packages 'citation("pkgname")'.
```

```
##
## Attaching package: 'Biobase'
```

```
## The following object is masked from 'package:MatrixGenerics':
##
##      rowMedians
```

```
## The following objects are masked from 'package:matrixStats':
##
##      anyMissing, rowMedians
```

```
library(tidyverse)
```

```
## -- Attaching packages ----- tidyverse 1.3.1 --
```

```
## v ggplot2 3.4.2      v purrr  1.0.1
## v tibble  3.2.1      v dplyr  1.1.1
## v tidyr   1.3.0      v stringr 1.5.0
## v readr   2.1.2      v forcats 1.0.0
```

```
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::collapse() masks IRanges::collapse()
## x dplyr::combine()  masks Biobase::combine(), BiocGenerics::combine()
## x dplyr::count()    masks matrixStats::count()
## x dplyr::desc()     masks IRanges::desc()
## x tidyr::expand()   masks S4Vectors::expand()
## x dplyr::filter()   masks stats::filter()
## x dplyr::first()    masks S4Vectors::first()
## x dplyr::lag()      masks stats::lag()
```

```
## x ggplot2::Position() masks BiocGenerics::Position(), base::Position()
## x purrr::reduce()      masks GenomicRanges::reduce(), IRanges::reduce()
## x dplyr::rename()      masks S4Vectors::rename()
## x dplyr::slice()       masks IRanges::slice()
```

```
library(CorLevelPlot)
library(gridExtra)
```

```
##
## Attaching package: 'gridExtra'

## The following object is masked from 'package:dplyr':
##
##      combine

## The following object is masked from 'package:Biobase':
##
##      combine

## The following object is masked from 'package:BiocGenerics':
##
##      combine
```

```
library(magrittr)
```

```
##
## Attaching package: 'magrittr'

## The following object is masked from 'package:purrr':
##
##      set_names

## The following object is masked from 'package:tidyr':
##
##      extract
```

```
library(ggplot2)
allowWGCNAThreads()
```

```
## Allowing multi-threading with up to 8 threads.
```

```
setwd('E:/WGCNA')
```

## 1. Read data

```
data<- read.csv('Counts.csv', row.names = 1)
Sample_info<- read.csv('Sample_info.csv', row.names = 1)
```

## 2. Quality control

```
gg<- goodSamplesGenes(t(data))
```

Step1: Detect outlier genes with WGCNA package

```
## Flagging genes and samples with too many missing values...
## ..step 1
## ..step 2
```

```
summary(gg)
```

```
##           Length Class  Mode
## goodGenes   28089  -none- logical
## goodSamples    52  -none- logical
## allOK         1  -none- logical
```

```
gg$allOK
```

```
## [1] FALSE
```

#Since it indicates tat all genes are not good genes let's quantify the outlier genes #We'll do it for samples as well

```
table(gg$goodGenes)
```

```
##
## FALSE  TRUE
##  3576 24513
```

```
table(gg$goodSamples)
```

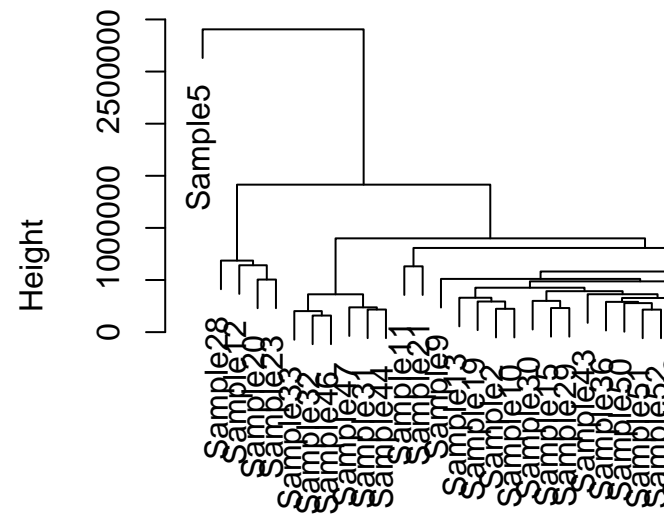
```
##
## TRUE
##   52
```

#Around 3500 outlier genes but no outlier samples. Let's remove those genes

```
data<- data[gg$goodGenes=='TRUE',]
```

```
htree <- hclust(dist(t(data)), method = "average")
plot(htree)
```

## Cluster D



**Step2: Identify outlier samples with hierachical clustering**

#### Step3: Identify outlier samples with PCA

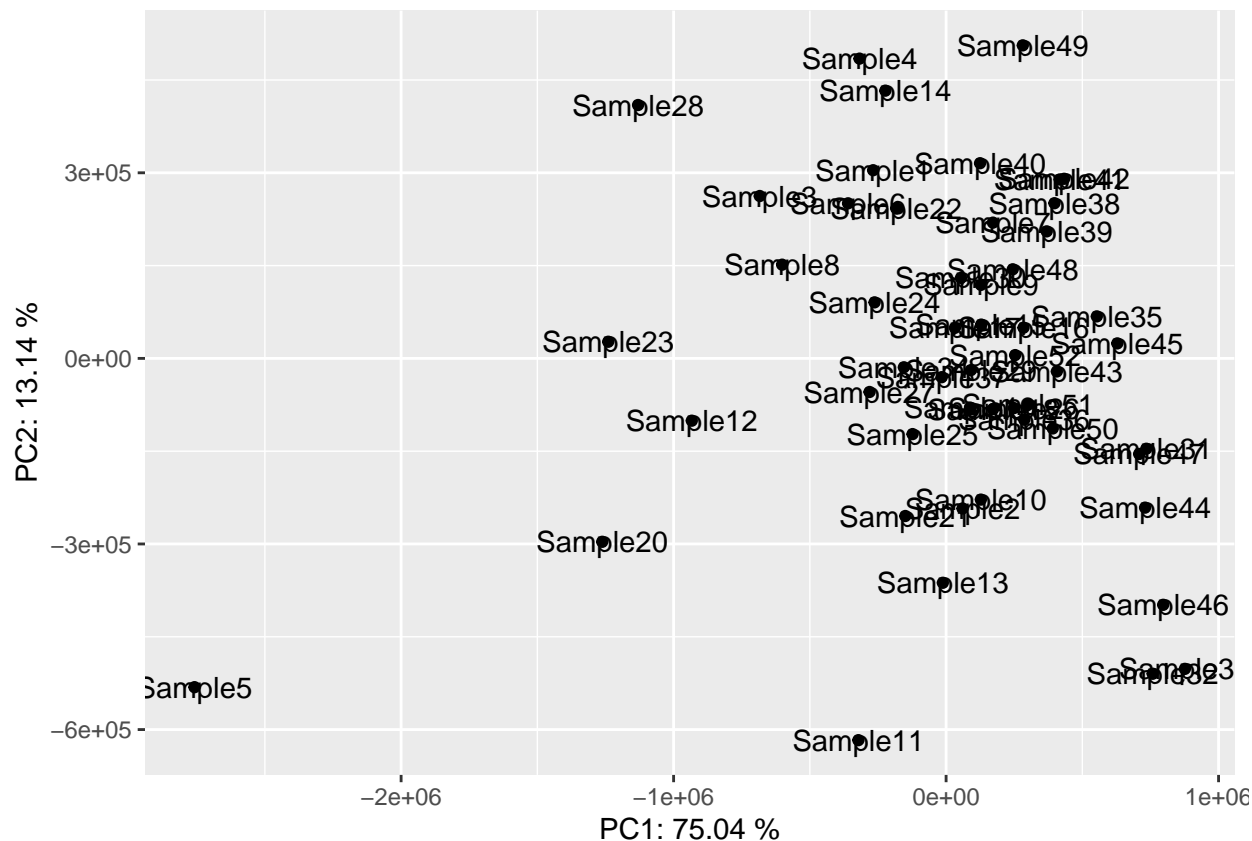
```
pca <- prcomp(t(data))
pca.dat <- pca$x

pca.var <- pca$sdev^2
pca.var.percent <- round(pca.var/sum(pca.var)*100, digits = 2)

pca.dat <- as.data.frame(pca.dat)

ggplot(pca.dat, aes(PC1, PC2)) +
  geom_point() +
  geom_text(label = rownames(pca.dat)) +
  labs(x = paste0('PC1: ', pca.var.percent[1], ' %'),
       y = paste0('PC2: ', pca.var.percent[2], ' %'))
```

dist(t  
hclust (\*,



```
#N.B. We need to remove batch effect if there's any at this stage before we proceed further

#In both these methods we found Sample5 is distantly related
#So we'll exclude them
discard_samples<- c('Sample5')
data.final<- data[,!colnames(data) %in% discard_samples]
Sample_info_final<- Sample_info[!row.names(Sample_info) %in% discard_samples,]
```

#### 4. Perform VST normalization with DESeq2 since it's a count matrix

#N.B. FPKM or RPKM matrices need to be log transformed #Before performing normalization make sure if the rownames in matrix and colnames in #Sample info are identical

```
all(rownames(Sample_info_final)%in%colnames(data.final)) #If present
```

```
## [1] TRUE
```

```
all(rownames(Sample_info_final)==colnames(data.final)) #If present in same order
```

```
## [1] TRUE
```

```
dds <- DESeqDataSetFromMatrix(countData = data.final,
                              colData = Sample_info_final,
                              design = ~ 1) #Specifying no design
nrow(dds)
```

Create deseq2 object

```
## [1] 24513
```

**Remove samples that have <15 counts in more than 75% of the samples** #Recommended by WGCNA #However, the downstream analysis with higher # of genes will depend largely on #the available ram of PC. So, we'll consider gene numbers based on our PC capacity.

```
keep <- dds[rowSums(counts(dds) >= 15)>=39,] #75% of 52 samples is 52*0.75=39
nrow(keep)
```

```
## [1] 12226
```

```
dds.vst<- vst(keep)
dds.vst<- dds.vst[1:5000,] #We reduced the # of genes to 8000 because of low RAM
nrow(dds.vst)
```

Perform VST transformation

```
## [1] 5000
```

Get normalized counts

```
norm.counts <- assay(dds.vst) %>%
  t()
```

5. Constructing network

Set softthreshold power

```
power <- c(c(1:10), seq(from = 12, to = 50, by = 2))
```

Set the network topology analysis function



```
sft <- pickSoftThreshold(norm.counts,
                          powerVector = power,
                          networkType = "signed",
                          verbose = 5)
```

```
## pickSoftThreshold: will use block size 5000.
## pickSoftThreshold: calculating connectivity for given powers...
## ..working on genes 1 through 5000 of 5000
## Power SFT.R.sq slope truncated.R.sq mean.k. median.k. max.k.
## 1 1 0.1310 32.20 0.974 2.50e+03 2.50e+03 2580.00
## 2 2 0.0456 -5.56 0.951 1.34e+03 1.33e+03 1470.00
## 3 3 0.2070 -5.36 0.962 7.52e+02 7.47e+02 931.00
## 4 4 0.3250 -4.12 0.966 4.43e+02 4.36e+02 627.00
## 5 5 0.3900 -3.25 0.960 2.71e+02 2.64e+02 443.00
## 6 6 0.4500 -2.67 0.962 1.72e+02 1.64e+02 324.00
## 7 7 0.5190 -2.36 0.966 1.12e+02 1.05e+02 244.00
## 8 8 0.5990 -2.33 0.975 7.56e+01 6.88e+01 191.00
## 9 9 0.6550 -2.37 0.977 5.21e+01 4.59e+01 153.00
## 10 10 0.7020 -2.35 0.980 3.67e+01 3.12e+01 124.00
## 11 12 0.7560 -2.39 0.983 1.92e+01 1.51e+01 85.80
## 12 14 0.7910 -2.47 0.982 1.08e+01 7.73e+00 61.50
## 13 16 0.8010 -2.47 0.965 6.33e+00 4.13e+00 45.40
## 14 18 0.8090 -2.55 0.977 3.89e+00 2.27e+00 34.20
## 15 20 0.8280 -2.47 0.982 2.48e+00 1.28e+00 26.30
## 16 22 0.8380 -2.42 0.975 1.63e+00 7.48e-01 20.50
## 17 24 0.8280 -2.41 0.948 1.10e+00 4.44e-01 16.20
## 18 26 0.8410 -2.35 0.949 7.63e-01 2.70e-01 13.00
## 19 28 0.8500 -2.29 0.947 5.39e-01 1.66e-01 10.50
## 20 30 0.3950 -2.99 0.321 3.87e-01 1.04e-01 8.52
## 21 32 0.3950 -2.91 0.321 2.83e-01 6.61e-02 6.99
## 22 34 0.4190 -2.90 0.426 2.10e-01 4.29e-02 5.78
## 23 36 0.4150 -2.82 0.407 1.58e-01 2.79e-02 4.81
## 24 38 0.9330 -1.95 0.994 1.20e-01 1.83e-02 4.03
## 25 40 0.9280 -1.90 0.962 9.27e-02 1.23e-02 3.39
## 26 42 0.9340 -1.88 0.956 7.23e-02 8.30e-03 3.11
## 27 44 0.9460 -1.86 0.957 5.69e-02 5.56e-03 2.92
## 28 46 0.9470 -1.85 0.950 4.52e-02 3.73e-03 2.74
## 29 48 0.9400 -1.82 0.932 3.63e-02 2.54e-03 2.58
## 30 50 0.4160 -2.32 0.276 2.94e-02 1.75e-03 2.43
```

#N.B. unsigned -> nodes with positive & negative correlation are treated equally #N.B. signed -> nodes with negative correlation are considered *unconnected*, treated as zero

```
sft.data <- sft$fitIndices
```

## visualization to pick power

```
Plot1 <- ggplot(sft.data, aes(Power, SFT.R.sq, label = Power)) +
  geom_point() +
```

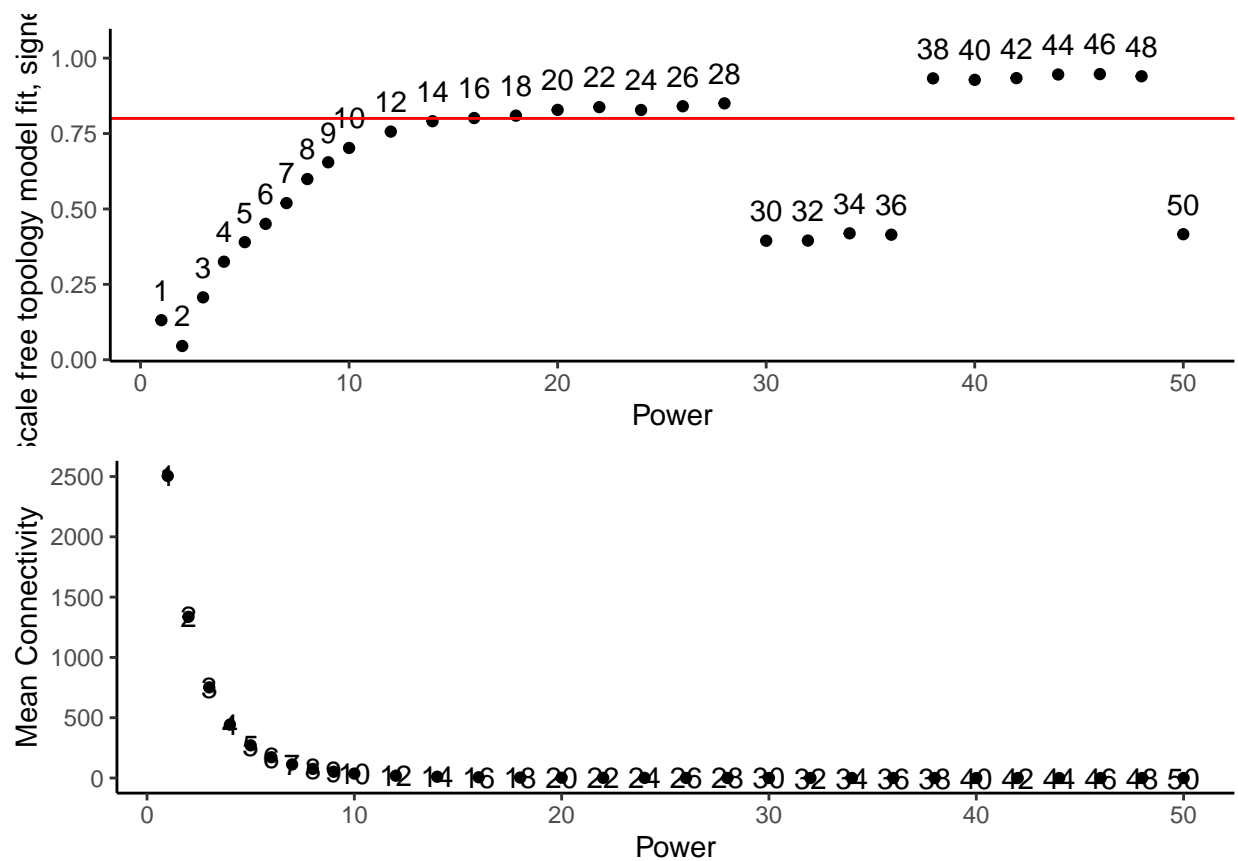
```

geom_text(nudge_y = 0.1) +
geom_hline(yintercept = 0.8, color = 'red') +
labs(x = 'Power', y = 'Scale free topology model fit, signed R^2') +
theme_classic()

Plot2 <- ggplot(sft.data, aes(Power, mean.k., label = Power)) +
  geom_point() +
  geom_text(nudge_y = 0.1) +
  labs(x = 'Power', y = 'Mean Connectivity') +
  theme_classic()

grid.arrange(Plot1, Plot2, nrow = 2)

```



convert matrix to numeric and assign power

```

norm.counts[] <- sapply(norm.counts, as.numeric)
soft_power <- 14
temp_cor <- cor
cor <- WGCNA::cor

```

## Estimate w.r.t blocksize memory depends on available ram of PC

```
bwnet <- blockwiseModules(norm.counts,
                           maxBlockSize = 5000,
                           TOMType = "signed",
                           power = soft_power,
                           mergeCutHeight = 0.25,
                           numericLabels = FALSE,
                           randomSeed = 1234,
                           verbose = 3)

## Calculating module eigengenes block-wise from all genes
##   Flagging genes and samples with too many missing values...
##   ..step 1
##   ..Working on block 1 .
##   TOM calculation: adjacency..
##   ..will not use multithreading.
##   Fraction of slow calculations: 0.000000
##   ..connectivity..
##   ..matrix multiplication (system BLAS)..
##   ..normalization..
##   ..done.
##   ....clustering..
##   ....detecting modules..
##   ....calculating module eigengenes..
##   ....checking kME in modules..
##   ..reassigning 1 genes from module 3 to modules with higher KME.
##   ..merging modules that are too close..
##   mergeCloseModules: Merging modules whose distance is less than 0.25
##   Calculating new MEs...

cor <- temp_cor
```

### 6. Identify Module Eigengenes

```
module_eigengenes <- bwnet$MEs
head(module_eigengenes)

##           MEturquoise    MEbrown    MEgreen    MEblack    MEred
## Sample1 -0.10375879  0.21685398  0.020542538 -0.09222291 -0.100192258
## Sample2 -0.01255866  0.12839304  0.032163029 -0.07031597 -0.221029610
## Sample3  0.04519388  0.03717920  0.066635498 -0.05908388  0.017316170
## Sample4  0.06835863 -0.08730094 -0.003943952 -0.06046904  0.074580135
## Sample6  0.15559583 -0.04275675  0.087524806 -0.07252025  0.041854527
## Sample7  0.05497992  0.13879174 -0.144092037 -0.04763668 -0.004967255
##           MEblue      MEyellow      MEgrey
## Sample1 -0.06997022 -0.167852659 -0.0823958709
## Sample2 -0.12854895 -0.149656308  0.0066448007
## Sample3 -0.02317673 -0.073541884 -0.0006775664
```

```
## Sample4 -0.06173939 -0.047079887 0.0677114636
## Sample6 -0.14128791 -0.004358101 0.1408660417
## Sample7 -0.13322219 -0.090824750 0.0951961513
```

Check the number of genes for each module

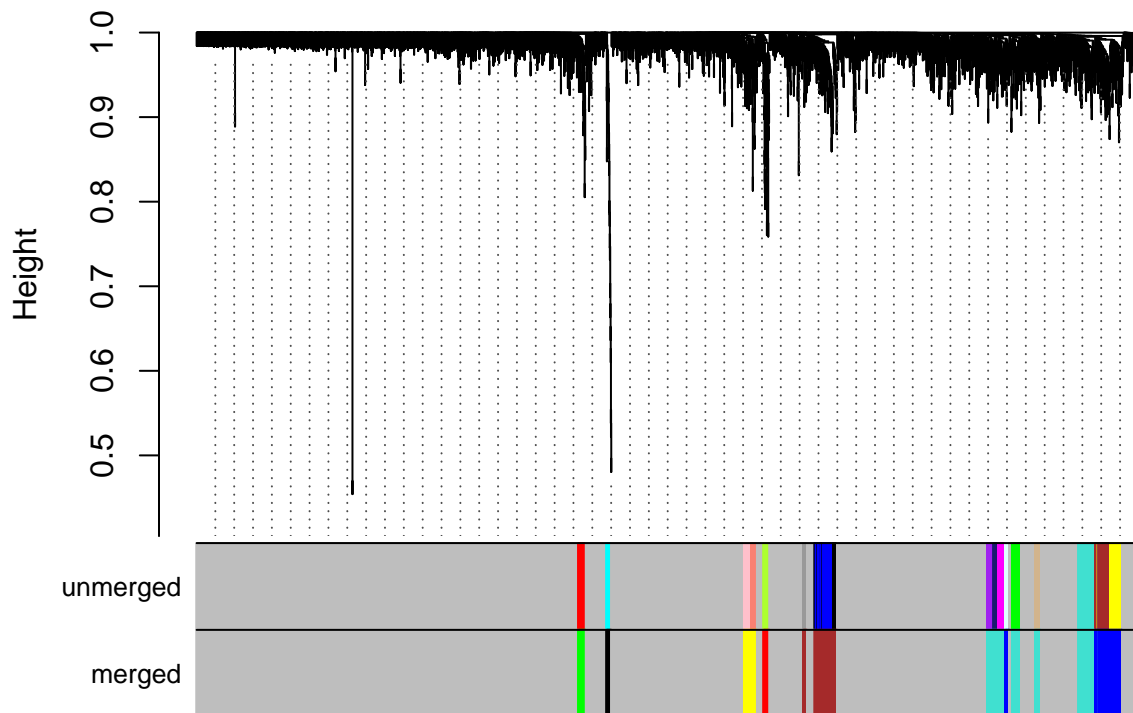
```
table(bwnet$colors)
```

```
##
##      black      blue      brown      green      grey      red turquoise      yellow
##         26        164        141         39       4262         32        267         69
```

Plot the dendrogram and the module colors before and after merging underneath

```
plotDendroAndColors(bwnet$dendrograms[[1]], cbind(bwnet$unmergedColors, bwnet$colors),
  c("unmerged", "merged"),
  dendroLabels = FALSE,
  addGuide = TRUE,
  hang= 0.03,
  guideHang = 0.05)
```

## Cluster Dendrogram



grey module = all genes that doesn't fall into other modules were assigned to the grey module

#### 7. Extract genes and their associated color modules and save them

```
mergedColors<- labels2colors(bwnet$colors)
module_df <- data.frame(
  gene_id <- names(bwnet$colors),
  colors <- labels2colors(bwnet$colors)
)

module_df[1:5,]
```

```
##   gene_id...names.bwnet.colors. colors...labels2colors.bwnet.colors.
## 1          ENSG00000000419                      green
## 2          ENSG00000000457                      green
## 3          ENSG00000000460                      green
## 4          ENSG00000000938                      green
## 5          ENSG00000001036                      green
```

```
write_delim(module_df,
             file = "gene_modules.txt",
             delim = "\t")
```

#### 8. Merging and clustering modules from eigengenes

##### Calculate eigengenes

```
MEList <- moduleEigengenes(norm.counts, colors = mergedColors)
MEs <- MEList$eigengenes
```

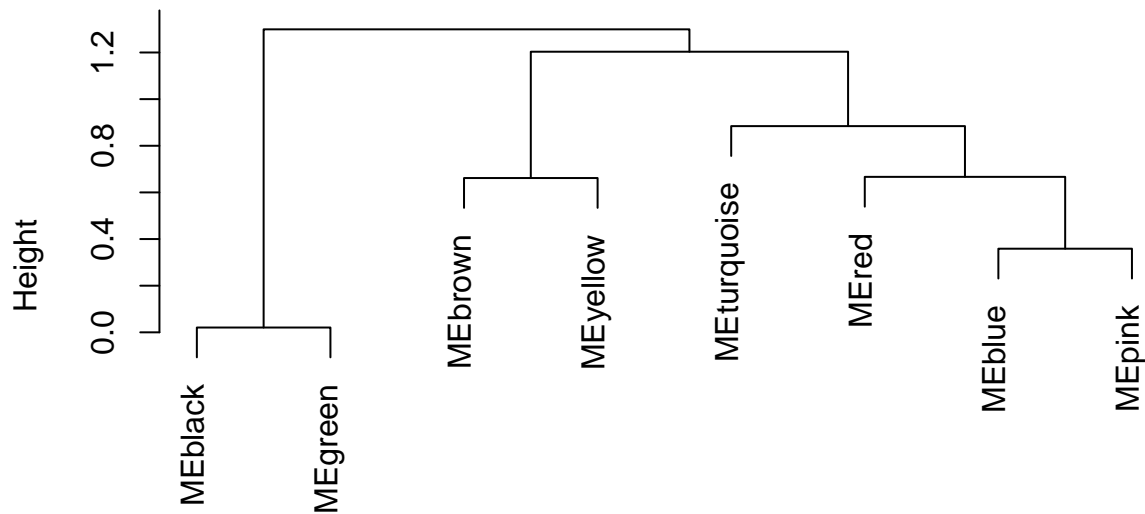
##### Calculate dissimilarity of module eigengenes

```
MEDiss <- 1-cor(MEs)
METree <- hclust(as.dist(MEDiss), method = "average")
```

##### Plot the result

```
plot(METree, main = "Clustering of Module eigengenes",
     xlab = "", sub = "")
```

## Clustering of Module eigengenes



9. Show the correlation between modules and each sample

## Get Module Eigengenes per cluster

```
MEs0 <- moduleEigengenes(norm.counts, mergedColors)$eigengenes
```

#Reorder modules so similar modules are next to each other

```
MEs0 <- orderMEs(MEs0)  
module_order <- names(MEs0) %>% gsub("ME", "", .)
```

## Add the treatment names

```
MEs0$treatment <- row.names(MEs0)
```

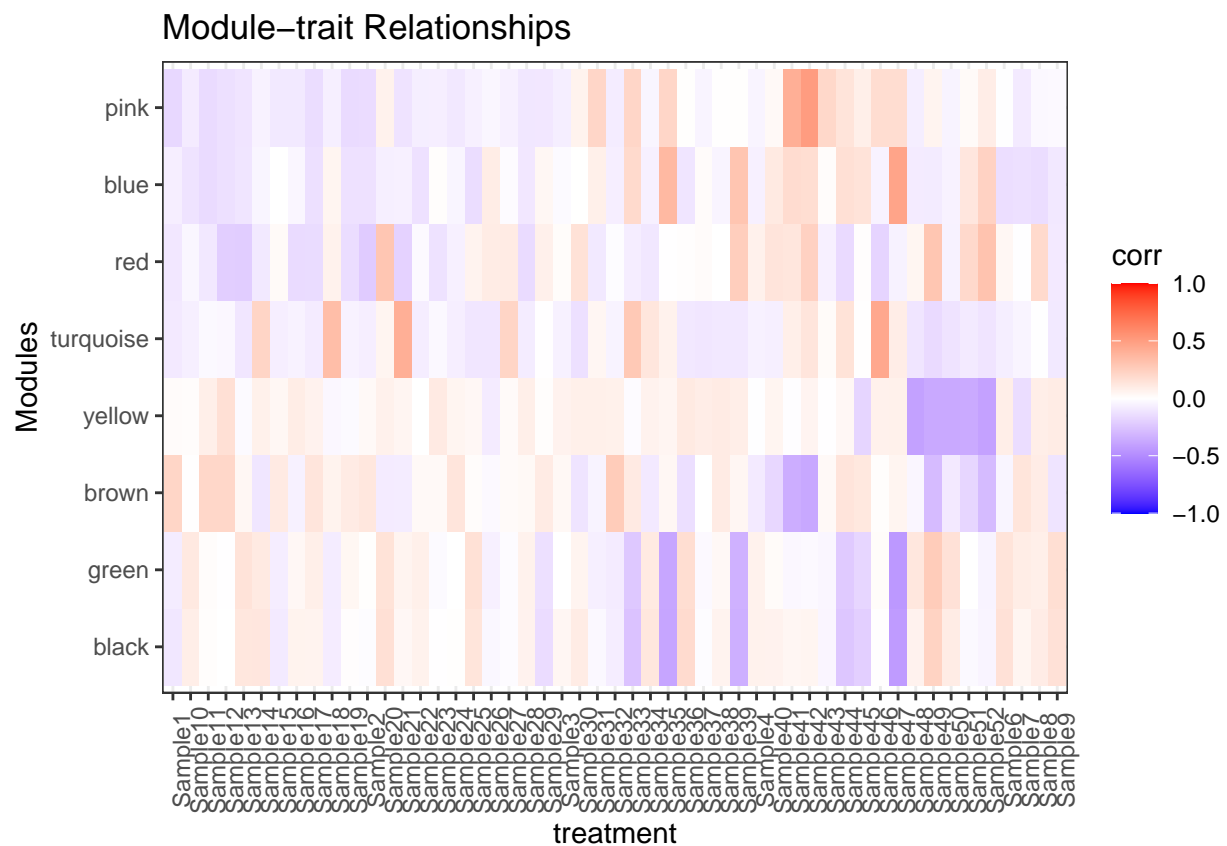
## Tidy & plot data

```

mME = MEs0 %>%
  pivot_longer(-treatment) %>%
  mutate(
    name = gsub("ME", "", name),
    name = factor(name, levels = module_order)
  )

mME %>% ggplot(., aes(x=treatment, y=name, fill=value)) +
  geom_tile() +
  theme_bw() +
  scale_fill_gradient2(
    low = "blue",
    high = "red",
    mid = "white",
    midpoint = 0,
    limit = c(-1,1)) +
  theme(axis.text.x = element_text(angle=90)) +
  labs(title = "Module-trait Relationships", y = "Modules", fill="corr")

```



#### 10. Showing gene expression level in specific modules

```

modules_of_interest = c("green", "turquoise", "red")

```

## Pull out list of genes in that module

```
submod = module_df %>%  
  subset(colors %in% modules_of_interest)  
  
row.names(module_df) = module_df$gene_id
```

## Get normalized expression for those genes

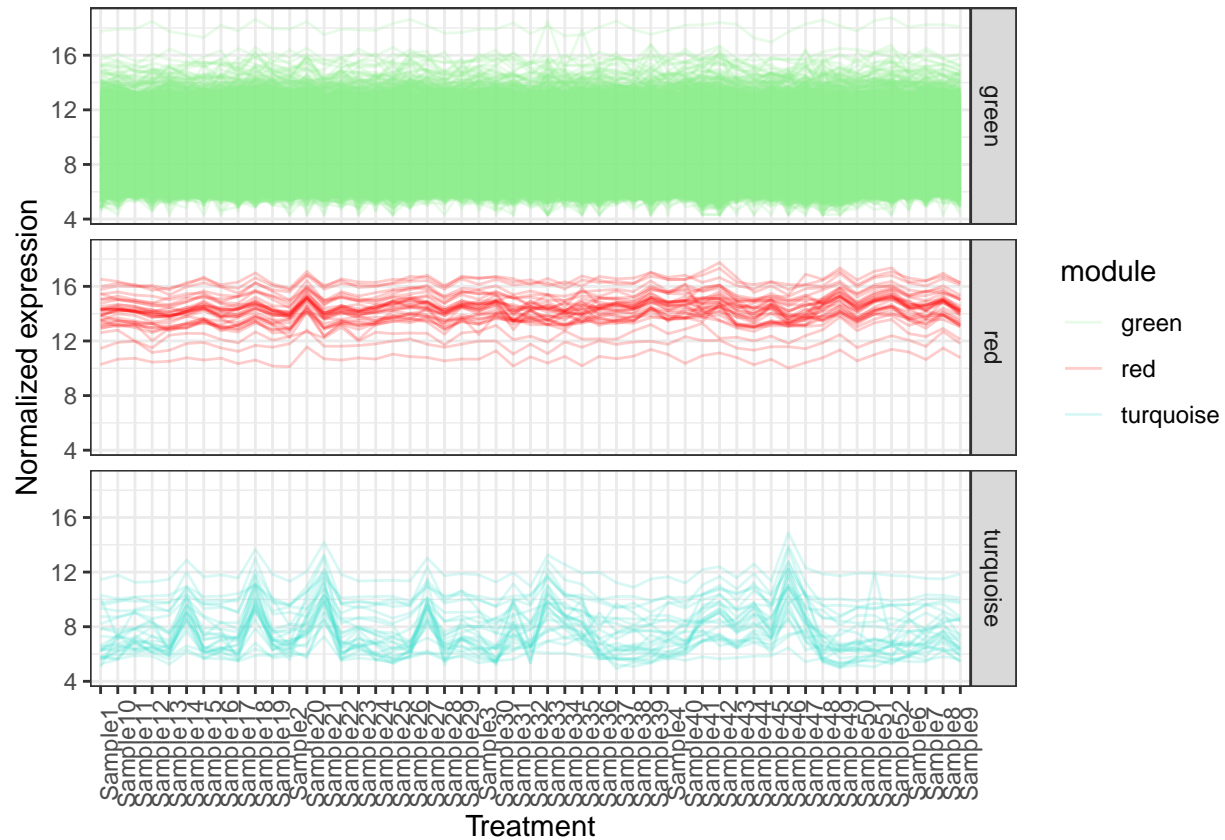
```
norm.counts[1:5,1:10]
```

```
##      ENSG00000000419 ENSG00000000457 ENSG00000000460 ENSG00000000938  
## Sample1      7.817808      9.010102      8.278295      13.14962  
## Sample2      7.860089      9.828239      8.520372      13.56801  
## Sample3      8.152240      8.428289      8.494442      11.96361  
## Sample4      8.430191      9.276900      8.532905      12.65407  
## Sample6      8.316706      8.996154      8.293994      12.79312  
##      ENSG000000001036 ENSG000000001084 ENSG000000001167 ENSG000000001460  
## Sample1      7.925833      9.816597      9.853574      7.460652  
## Sample2      8.382598      9.414906      9.628366      7.404398  
## Sample3      8.657241      9.982769      9.601348      6.924517  
## Sample4      8.300237      10.903511      10.299930      7.517321  
## Sample6      8.890079      10.308196      9.615493      7.567784  
##      ENSG000000001461 ENSG000000001497  
## Sample1      9.763363      10.061901  
## Sample2      10.019682      9.610658  
## Sample3      9.858863      10.066467  
## Sample4      9.777715      10.017410  
## Sample6      9.932600      10.083488
```

```
norm.exprs<- data.frame(t(norm.counts))  
norm.exprs$gene_id<- row.names(norm.exprs)  
norm.exprs<- norm.exprs %>% select(gene_id,everything())  
subexpr <- norm.exprs[submod$gene_id,]  
  
submod_df <- data.frame(subexpr) %>%  
  mutate(  
    gene_id = row.names(.)  
  ) %>%  
  pivot_longer(-gene_id) %>%  
  mutate(  
    module = module_df[gene_id,]$colors  
  )  
  
submod_df %>% ggplot(., aes(x=name, y=value, group=gene_id, colour=module)) +  
  geom_line(alpha=0.2) +  
  theme_bw() +  
  theme(  
    axis.text.x = element_text(angle = 90)
```



```
) +
  facet_grid(rows = vars(module)) +
  labs(x = "Treatment",
       y = "Normalized expression") +
  scale_color_manual(values = c("lightgreen", "red", "turquoise"))
```



### 11. Finding the association between module and traits # Prepare and manipulate data for association analysis # Converting categorical variable into binary group variables

```
type <- Sample_info_final %>%
  mutate(type = ifelse(grepl('Lung cancer', Group), 1, 0)) %>%
  select(4)
```

```
Sample_info_final$Status <- factor(Sample_info_final$Status,
  levels = c("None", "Newly diagnosed",
             "Post Chemotherapy", "On Doxorubicin"))
```

```
Status<- binarizeCategoricalColumns(Sample_info_final$Status,
  includePairwise = FALSE,
  includeLevelVsAll = TRUE,
  minCount = 1)
```

```
Status.final<- cbind(type, Status)
```

## Define numbers of genes and samples and draw correlation

```
nSamples <- nrow(norm.counts)
nGenes <- ncol(norm.counts)
status.module.corr <- cor(module_eigengenes, Status.final, use = 'p')
status.module.corr.pvals <- corPvalueStudent(status.module.corr , nSamples)
nrow(module_eigengenes)
```

```
## [1] 51
```

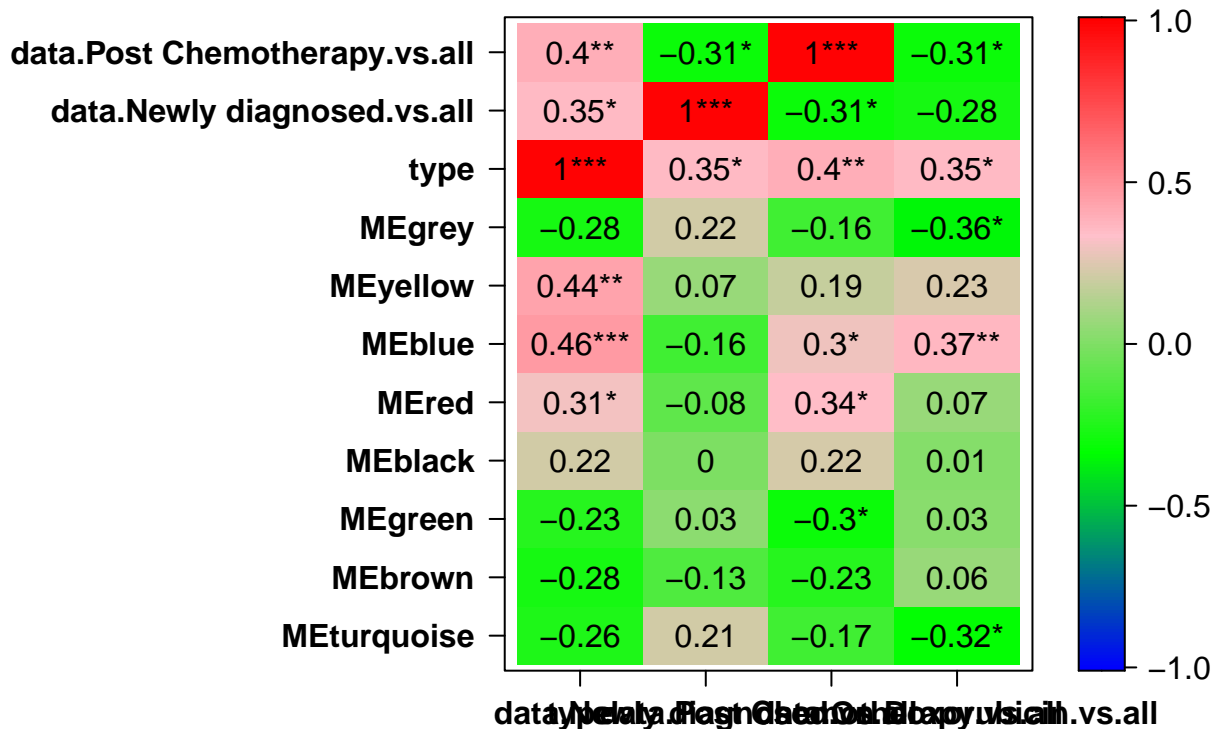
```
heatmap.data <- merge(module_eigengenes, Status.final, by = 'row.names')
head(heatmap.data)
```

### Visualize module-trait association as a heatmap

```
## Row.names MEturquoise MEbrown MEgreen MEblack MERed
## 1 Sample1 -0.1037587873 0.21685398 0.02054254 -0.09222291 -0.10019226
## 2 Sample10 0.0895341492 0.00293838 0.01367637 -0.06928702 -0.03836467
## 3 Sample11 0.0099912350 0.20925609 0.08402187 -0.02660053 -0.09621342
## 4 Sample12 0.0006960986 0.20967381 0.16100147 -0.03369510 -0.20749830
## 5 Sample13 0.1272433032 0.04066019 -0.02093506 -0.10491016 -0.21409801
## 6 Sample14 0.1288343520 -0.10802289 0.07607466 0.22150429 -0.09486251
## MEblue MEyellow MEgrey type data.Newly diagnosed.vs.all
## 1 -0.06997022 -0.16785266 -0.082395871 0 0
## 2 -0.11979305 -0.08319146 0.116674013 0 0
## 3 -0.15188447 -0.15462433 0.014037255 0 0
## 4 -0.13145467 -0.13190268 -0.003316252 0 0
## 5 -0.10896400 -0.11249850 0.148243795 0 0
## 6 -0.04248159 -0.05396714 0.110840769 0 0
## data.Post Chemotherapy.vs.all data.On Doxorubicin.vs.all
## 1 0 0
## 2 0 0
## 3 0 0
## 4 0 0
## 5 0 0
## 6 0 0
```

```
heatmap.data <- heatmap.data %>%
  column_to_rownames(var = 'Row.names')

CorLevelPlot(heatmap.data,
  x = names(heatmap.data)[9:12],
  y = names(heatmap.data)[1:11],
  col = c("blue1", "green", "pink", "red"))
```



```
module.gene.mapping <- as.data.frame(bwnet$colors)
module.gene.mapping %>%
  filter(`bwnet$colors` == 'blue') %>%
  rownames()
```

```
## [1] "ENSG00000005483" "ENSG00000006607" "ENSG000000010072" "ENSG000000010165"
## [5] "ENSG000000029363" "ENSG000000035141" "ENSG000000043093" "ENSG000000044574"
## [9] "ENSG000000048649" "ENSG000000051382" "ENSG000000051620" "ENSG000000053254"
## [13] "ENSG000000053770" "ENSG000000055070" "ENSG000000056586" "ENSG000000059769"
## [17] "ENSG000000067064" "ENSG000000067082" "ENSG000000069345" "ENSG000000069956"
## [21] "ENSG000000070831" "ENSG000000072415" "ENSG000000075568" "ENSG000000076053"
## [25] "ENSG000000076248" "ENSG000000077152" "ENSG000000079332" "ENSG000000080371"
## [29] "ENSG000000080824" "ENSG000000081320" "ENSG000000084463" "ENSG000000087074"
## [33] "ENSG000000088682" "ENSG000000089597" "ENSG000000090432" "ENSG000000090863"
## [37] "ENSG000000091527" "ENSG000000092094" "ENSG000000092871" "ENSG000000095002"
## [41] "ENSG000000095370" "ENSG000000095787" "ENSG000000099246" "ENSG000000099797"
## [45] "ENSG000000100221" "ENSG000000100354" "ENSG000000100393" "ENSG000000100395"
## [49] "ENSG000000100401" "ENSG000000100532" "ENSG000000100603" "ENSG000000100614"
## [53] "ENSG000000100982" "ENSG000000101166" "ENSG000000101337" "ENSG000000101367"
## [57] "ENSG000000103061" "ENSG000000103415" "ENSG000000103429" "ENSG000000104164"
## [61] "ENSG000000104312" "ENSG000000104889" "ENSG000000105835" "ENSG000000105856"
## [65] "ENSG000000106245" "ENSG000000106263" "ENSG000000106615" "ENSG000000106682"
## [69] "ENSG000000107560" "ENSG000000107897" "ENSG000000107937" "ENSG000000108061"
## [73] "ENSG000000108106" "ENSG000000108829" "ENSG000000108960" "ENSG000000109332"
## [77] "ENSG000000110696" "ENSG000000111802" "ENSG000000112081" "ENSG000000112118"
## [81] "ENSG000000112149" "ENSG000000112245" "ENSG000000113163" "ENSG000000113575"
```

```
## [85] "ENSG00000113734" "ENSG00000113811" "ENSG00000114125" "ENSG00000114796"
## [89] "ENSG00000114988" "ENSG00000115165" "ENSG00000115520" "ENSG00000115840"
## [93] "ENSG00000116455" "ENSG00000116731" "ENSG00000116747" "ENSG00000116752"
## [97] "ENSG00000116830" "ENSG00000116906" "ENSG00000117036" "ENSG00000117505"
## [101] "ENSG00000117614" "ENSG00000118894" "ENSG00000119392" "ENSG00000119559"
## [105] "ENSG00000119725" "ENSG00000119801" "ENSG00000119899" "ENSG00000119929"
## [109] "ENSG00000119950" "ENSG00000119953" "ENSG00000120690" "ENSG00000120705"
## [113] "ENSG00000120709" "ENSG00000120727" "ENSG00000121578" "ENSG00000121864"
## [117] "ENSG00000122068" "ENSG00000122257" "ENSG00000122882" "ENSG00000123091"
## [121] "ENSG00000123505" "ENSG00000123728" "ENSG00000124209" "ENSG00000124380"
## [125] "ENSG00000125835" "ENSG00000126524" "ENSG00000126945" "ENSG00000128245"
## [129] "ENSG00000128590" "ENSG00000128881" "ENSG00000128989" "ENSG00000129235"
## [133] "ENSG00000129315" "ENSG00000129691" "ENSG00000130311" "ENSG00000130522"
## [137] "ENSG00000130803" "ENSG00000130844" "ENSG00000131263" "ENSG00000131323"
## [141] "ENSG00000131381" "ENSG00000132507" "ENSG00000132603" "ENSG00000132661"
## [145] "ENSG00000132823" "ENSG00000133026" "ENSG00000133606" "ENSG00000133773"
## [149] "ENSG00000134375" "ENSG00000134644" "ENSG00000134686" "ENSG00000134970"
## [153] "ENSG00000135334" "ENSG00000136527" "ENSG00000136819" "ENSG00000137075"
## [157] "ENSG00000137502" "ENSG00000137575" "ENSG00000137876" "ENSG00000137947"
## [161] "ENSG00000138032" "ENSG00000138069" "ENSG00000138166" "ENSG00000138433"
```

## 12. Identifying significant genes against different features of data

Calculating the module membership and the associated p-values

The module membership/intramodular connectivity is calculated as the correlation of the eigengene and the gene expression profile.

This quantifies the similarity of all genes on the array to every module.

```
module.membership.measure <- cor(module_eigengenes, norm.counts, use = 'p')
module.membership.measure.pvals <- corPvalueStudent(module.membership.measure, nSamples)
module.membership.measure.pvals[1:5,1:10]
```

```
##          ENSG00000000419 ENSG00000000457 ENSG00000000460 ENSG00000000938
## MEturquoise      0.08179877      3.935824e-06      7.873861e-01      0.16687163
## MEbrown          0.01031164      3.801793e-01      1.015938e-01      0.02125898
## MEgreen          0.27024680      2.434798e-02      4.255289e-05      0.07970018
## MEblack          0.95418281      7.291502e-01      3.064485e-01      0.46636890
## MERed            0.06741831      1.364833e-01      6.076142e-02      0.03317124
##          ENSG000000001036 ENSG000000001084 ENSG000000001167 ENSG000000001460
## MEturquoise      1.066509e-05      1.211570e-02      0.1373691154      0.03960014
## MEbrown          1.158562e-02      7.932893e-01      0.0255369951      0.02068358
## MEgreen          3.976450e-02      2.249371e-07      0.0006352417      0.77101215
## MEblack          3.100530e-01      6.265870e-02      0.2118802323      0.83164735
## MERed            7.442618e-01      8.543609e-02      0.4044932254      0.89784399
```

##	ENSG000000001461	ENSG000000001497
## MEturquoise	2.468861e-05	0.0001716473
## MEbrown	8.252186e-01	0.7602053664
## MEgreen	4.901483e-01	0.1947561939
## MEblack	5.127292e-04	0.7311565906
## MERed	6.122777e-01	0.8786558002

```
gene.signf.corr <- cor(norm.counts, Status.final$'data.Post Chemotherapy.vs.all', use = 'p')
gene.signf.corr.pvals <- corPvalueStudent(gene.signf.corr, nSamples)
```

Calculating the gene significance and associated p-values against a particular trait

Sub-setting based on significant association: X1:corr, X2:pval

```
Sig_chemo<-data.frame(cbind(gene.signf.corr, gene.signf.corr.pvals))
Sig_chemo_final<- filter(Sig_chemo, Sig_chemo$X1> 0.3 | Sig_chemo$X1< -0.3
                        & Sig_chemo$X2<0.05)

write.csv(Sig_chemo_final, 'sig_genes_chemo.csv')
```