Dimension_reduction

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```
library(factoextra)

## Loading required package: ggplot2

## Welcome! Want to learn more? See two factoextra-related books at https://goo.gl/ve3WBa

library(purrr)
library(NMF)

## Loading required package: pkgmaker

## Loading required package: registry

## Loading required package: rngtools

## Loading required package: cluster

## NMF - BioConductor layer [OK] | Shared memory capabilities [NO: bigmemory] | Cores 11/12

## To enable shared memory capabilities, try: install.extras('

## NMF

## ")
```

load data

```
gene <- read.delim("/Users/rrrrrita/Documents/GitHub/PHS597DeepLearning/gene_expression_sample/GEUVADIS
geneexp <- gene[,c(1,5:ncol(gene))]
genotype <- read.delim("/Users/rrrrrita/Documents/GitHub/PHS597DeepLearning/gene_expression_sample/GEUV</pre>
```

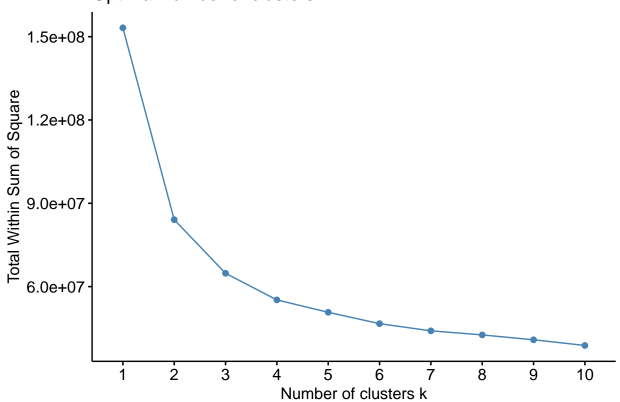
Vector quantization (VQ)

VQ is one type of k means. First I determined the optimal k based on total within-cluster sum of square:

```
wss <- function(k) {
   kmeans(t(geneexp[,-1]), k, nstart = 10 )$tot.withinss
}

fviz_nbclust(t(geneexp[-1]), kmeans, method = "wss")</pre>
```

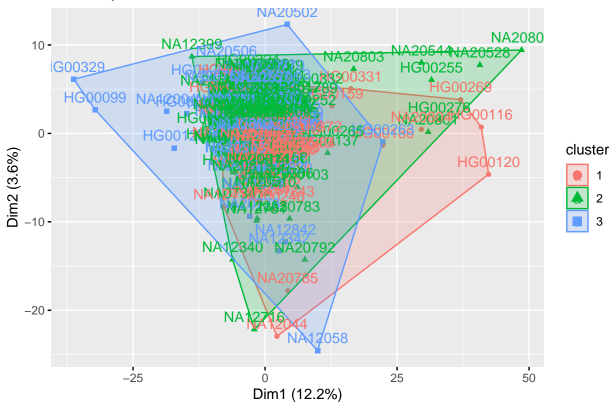
Optimal number of clusters



The elbow point happens at k=2 or 3. Here I chose k=3:

```
kres <- kmeans(t(geneexp[-1]), 3, nstart = 1)
fviz_cluster(kres, data = t(geneexp[-1]))</pre>
```

Cluster plot



The cluster plot tells that it is hard to discern these samples based on gene expression.

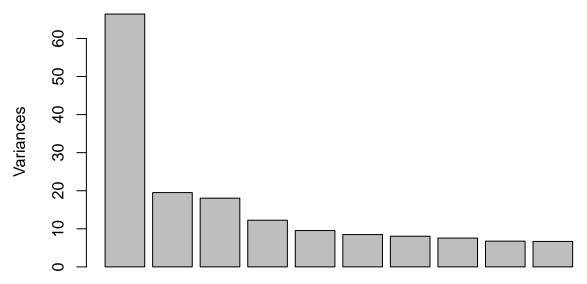
I used the centroid values of each cluster to represent samples within each cluster:

```
y_vq <- kres$centers[kres$cluster,]
rownames(y_vq) <- colnames(geneexp[,-1])</pre>
```

PCA

```
pca <- prcomp(t(geneexp[-1]), center=TRUE, scale = TRUE)
plot(pca, npcs=10)</pre>
```





As we can tell, first 10 PCs have represented most of the total variances, so I used the first 10 pcs to recover the uncompressed gene expression:

(It is also reasonable to choose first 3 PCs)

```
pca <- prcomp(t(geneexp[-1]), center=TRUE, scale = FALSE)
pc <- pca$x
rotation <- pca$rotation

npc <- 10
colm <- colMeans(t(geneexp[,-1]))

y_pca <- pc[,1:npc] %*% t(rotation[,1:npc]) + colm</pre>
```

NMF

First, run several experiments to select the optimal rank:

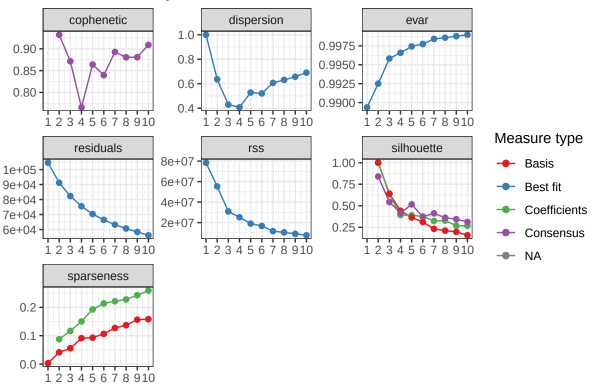
```
geneexp_nmf <- t(geneexp[-1])

# transfer into nonnegative data
geneexp_nmf <- nneg(geneexp_nmf)
estim.r <- nmf(geneexp_nmf, 1:10, nrun=20, seed=123456)
plot(estim.r)
estim.r</pre>
```

```
## Warning: Removed 3 row(s) containing missing values (geom_path).
```

Warning: Removed 5 rows containing missing values (geom_point).

NMF rank survey



Factorization rank For

optimal rank, (Frigyesi 2008) considered the smallest value at which the decrease in the RSS is lower than the decrease of the RSS obtained from random data. From the above RSS plot, the decrease from 3 to 4 is lower than the decrease from 1 to 2 and 2 to 3. Therefore, I chose rank = 4.

```
geneexp_nmf <- t(geneexp[-1])
# transfer into nonnegative data
geneexp_nmf <- nneg(geneexp_nmf)

nmffit <- nmf(geneexp_nmf, rank=4, "brunet", seed=123456) # higher rank -> better results
w <- nmffit@fit@W
h <- nmffit@fit@H
y_nmf <- w %*% h</pre>
```

Results comparison

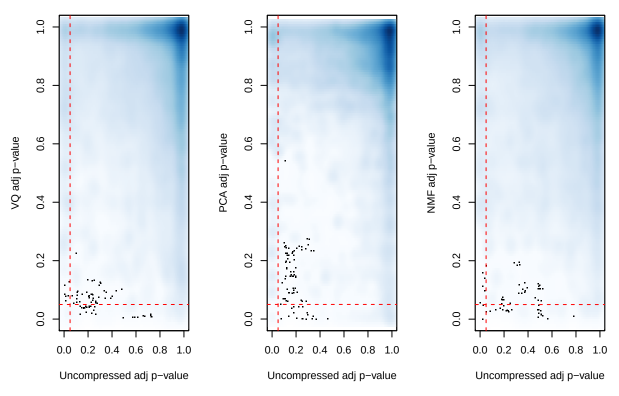
eQTL analysis

To conduct the eQTL analysis, I filtered the data within 500000 downstream and upstream of gene start and end sites. I regressed each gene expression (SNP) on each genotype. The Benjamini-Hochberg method was applied

```
resdf <- c()
eqtl <- function(gexp){

for ( i in 1:nrow(gene)){
   gtype <- subset(genotype, POS >= (gene$start[i] - 500000) & POS <= (gene$end[i] + 500000))</pre>
```

```
gtype <- gtype[,-(1:6)]
    y \leftarrow gexp[,i]
    resdf1 <- c()
    for ( j in 1:nrow(gtype)){
      x <- t(gtype[j,])</pre>
      lmfit \leftarrow lm(y \sim x)
      out <- data.frame(gene=gene$gene_id[i],</pre>
                          genotype = genotype$SNP[j],
                          pval = summary(lmfit)$coefficients[2,4])
      resdf1 <- rbind(resdf1, out)</pre>
    }
    resdf1$pval <- p.adjust(resdf1$pval, method = "BH")</pre>
    resdf <- rbind(resdf, resdf1)</pre>
  return(resdf)
# Read saved results
vq <- readRDS("vq_dopar.rds")</pre>
pca <- readRDS("pca_dopar.rds")</pre>
nmf <- readRDS("nmf_dopar.rds")</pre>
uncomp <- readRDS("uncomp_dopar.rds")</pre>
par(mfrow=c(1,3))
smoothScatter(x=uncomp$pval, y=vq$pval,
               xlab = "Uncompressed adj p-value", ylab = "VQ adj p-value")
abline(v = 0.05, h=0.05, lty = 2, col = "red")
smoothScatter(x=uncomp$pval, y=pca$pval,
               xlab = "Uncompressed adj p-value", ylab = "PCA adj p-value")
abline(v = 0.05, h=0.05, lty = 2, col = "red")
smoothScatter(x=uncomp$pval, y=nmf$pval,
              xlab = "Uncompressed adj p-value", ylab = "NMF adj p-value")
abline(v = 0.05, h=0.05, lty = 2, col = "red")
```



It's not easy to see a linear relationship between adjusted p-values for uncomparessed dataset and compressed dataset with three dimension reduction methods. Then we further evaluate the relationship with confusion matrices:

Confusion matrix

```
binary.pval <- function(x){</pre>
  y \leftarrow ifelse(x<0.05, 1, 0)
  y \leftarrow factor(y, levels = c(0,1))
  return(y)
vq_cm <- caret::confusionMatrix(data = binary.pval(vq$pval),</pre>
                         reference = binary.pval(uncomp$pval))
## Registered S3 methods overwritten by 'proxy':
     method
##
##
     print.registry_field registry
     print.registry_entry registry
pca_cm <- caret::confusionMatrix(data = binary.pval(pca$pval),</pre>
                         reference = binary.pval(uncomp$pval))
nmf_cm <- caret::confusionMatrix(data = binary.pval(nmf$pval),</pre>
                         reference = binary.pval(uncomp$pval))
byClass <- rbind(VQ =round(vq_cm$byClass, 3),</pre>
                  PCA=round(pca_cm$byClass, 3),
```

```
NMF = round(nmf_cm$byClass, 3))
overall <- rbind(VQ =round(vq_cm$overall, 3),</pre>
                 PCA=round(pca_cm$overall, 3),
                 NMF = round(nmf_cm$overall, 3))
byClass
##
       Sensitivity Specificity Pos Pred Value Neg Pred Value Precision Recall
## VQ
             0.998
                          0.006
                                         0.979
                                                         0.074
                                                                    0.979 0.998
             0.998
                          0.005
                                         0.979
                                                         0.044
                                                                    0.979 0.998
## PCA
## NMF
             0.999
                          0.000
                                         0.979
                                                         0.001
                                                                    0.979 0.999
          F1 Prevalence Detection Rate Detection Prevalence Balanced Accuracy
##
## VQ 0.989
                  0.979
                                  0.978
                                                        0.998
                                                                           0.502
## PCA 0.988
                  0.979
                                  0.977
                                                        0.998
                                                                           0.501
## NMF 0.989
                                  0.978
                                                                           0.499
                  0.979
                                                        0.999
overall
##
                 Kappa AccuracyLower AccuracyUpper AccuracyNull AccuracyPValue
       Accuracy
## VQ
          0.978
                 0.008
                                0.978
                                              0.978
                                                            0.979
## PCA
          0.977 0.005
                                0.977
                                              0.977
                                                            0.979
                                                                                1
## NMF
          0.978 -0.002
                                0.978
                                               0.978
                                                            0.979
                                                                                1
       McnemarPValue
##
## VQ
## PCA
                   0
## NMF
                   0
```

All three methods have high sensitivity but low specificity. NMF has the lowest negative predictive value among the three. For other measurements, these three methods are pretty much the same.