

class16

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```
library(tximport)

folders <- dir(pattern="SRR21568*")
samples <- sub("_quant", "", folders)
files <- file.path( folders, "abundance.h5" )
names(files) <- samples

txi.kallisto <- tximport(files, type = "kallisto", txOut = TRUE)
```

1 2 3 4

```
head(txi.kallisto$counts)
```

| | SRR2156848 | SRR2156849 | SRR2156850 | SRR2156851 |
|-----------------|------------|------------|------------|------------|
| ENST00000539570 | 0 | 0 | 0.00000 | 0 |
| ENST00000576455 | 0 | 0 | 2.62037 | 0 |
| ENST00000510508 | 0 | 0 | 0.00000 | 0 |
| ENST00000474471 | 0 | 1 | 1.00000 | 0 |
| ENST00000381700 | 0 | 0 | 0.00000 | 0 |
| ENST00000445946 | 0 | 0 | 0.00000 | 0 |

```
colSums(txi.kallisto$counts)
```

| SRR2156848 | SRR2156849 | SRR2156850 | SRR2156851 |
|------------|------------|------------|------------|
| 2563611 | 2600800 | 2372309 | 2111474 |

```
sum(rowSums(txi.kallisto$counts)>0)
```

[1] 94561

```
to.keep <- rowSums(tx1.kallisto$counts) > 0  
kset.nonzero <- tx1.kallisto$counts[to.keep,]
```

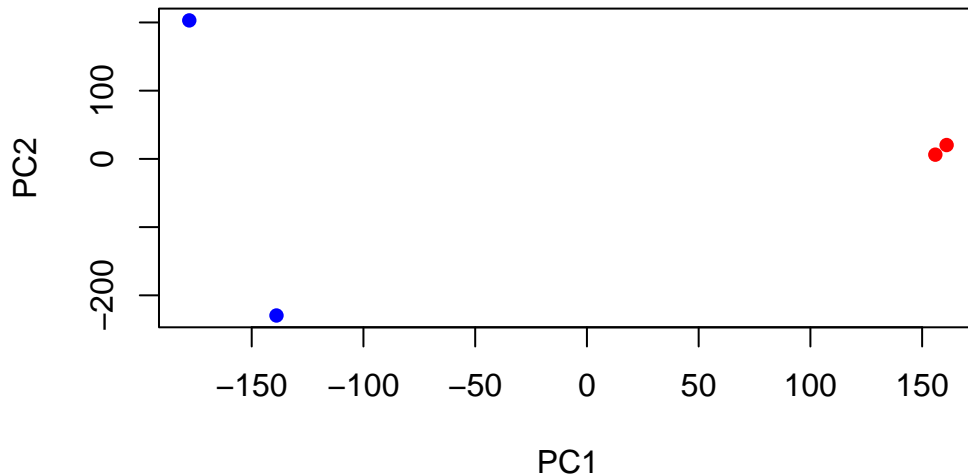
```
keep2 <- apply(kset.nonzero, 1, sd) > 0  
x <- kset.nonzero[keep2, ]
```

```
pca <- prcomp(t(x), scale=TRUE)  
summary(pca)
```

Importance of components:

| | PC1 | PC2 | PC3 | PC4 |
|------------------------|----------|----------|----------|-------|
| Standard deviation | 183.6379 | 177.3605 | 171.3020 | 1e+00 |
| Proportion of Variance | 0.3568 | 0.3328 | 0.3104 | 1e-05 |
| Cumulative Proportion | 0.3568 | 0.6895 | 1.0000 | 1e+00 |

```
plot(pca$x[, 1], pca$x[, 2],  
     col=c("blue", "blue", "red", "red"),  
     xlab="PC1", ylab="PC2", pch=16)
```



```

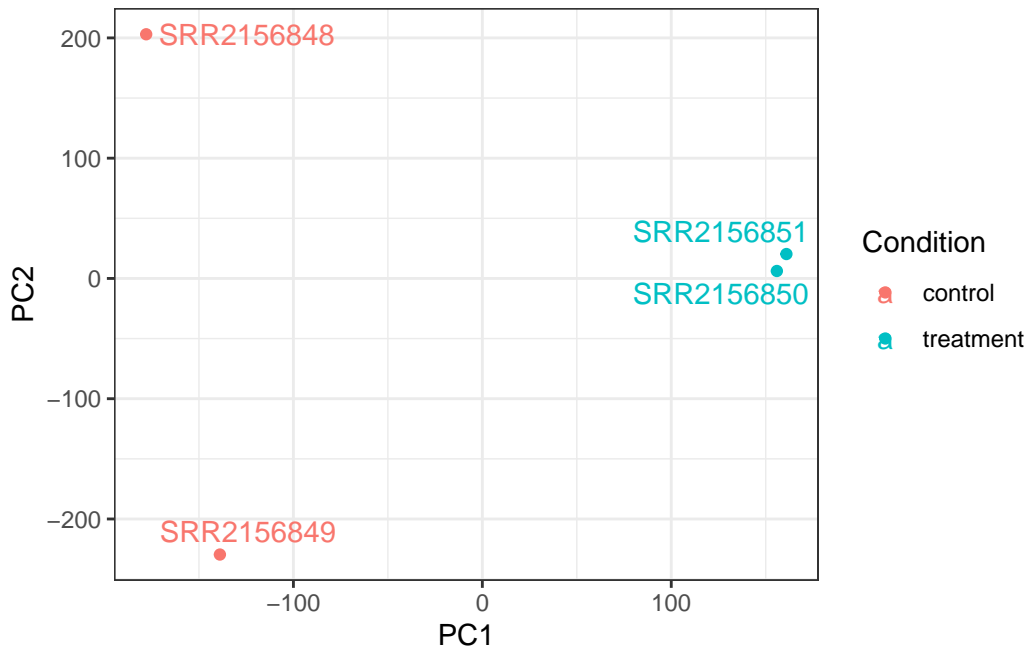
library(ggplot2)
library(ggrepel)

colData <- data.frame(condition = factor(rep(c("control", "treatment"), each = 2)))
rownames(colData) <- colnames(tx1.kallisto$counts)

y <- as.data.frame(pca$x)
y$Condition <- as.factor(colData$condition)

ggplot(y) +
  aes(PC1, PC2, col=Condition) +
  geom_point() +
  geom_text_repel(label=rownames(y)) +
  theme_bw()

```



```
library(DESeq2)
```

Loading required package: S4Vectors

Loading required package: stats4

Loading required package: BiocGenerics

```
Loading required package: generics
```

```
Attaching package: 'generics'
```

```
The following objects are masked from 'package:base':
```

```
as.difftime, as.factor, as.ordered, intersect, is.element, setdiff,  
setequal, union
```

```
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, aperm, append, as.data.frame, basename, cbind,  
colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,  
get, grep, grepl, is.unsorted, lapply, Map, mapply, match, mget,  
order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank,  
rbind, Reduce, rownames, sapply, saveRDS, table, tapply, 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
```

```
Loading required package: GenomicRanges
```

```
Loading required package: Seqinfo
```

```
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
```

```
sampleTable <- data.frame(condition = factor(rep(c("control", "treatment"), each = 2)))
rownames(sampleTable) <- colnames(tx1.kallisto$counts)
dds <- DESeqDataSetFromTximport(tx1.kallisto,
                                 sampleTable,
                                 ~condition)
```

```
using counts and average transcript lengths from tximport
```

```
dds <- DESeq(dds)
```

```
estimating size factors
```

```
using 'avgTxLength' from assays(dds), correcting for library size
```

```
estimating dispersions
```

```
gene-wise dispersion estimates
```

```
mean-dispersion relationship
```

```
-- note: fitType='parametric', but the dispersion trend was not well captured by the
       function: y = a/x + b, and a local regression fit was automatically substituted.
       specify fitType='local' or 'mean' to avoid this message next time.
```

```
final dispersion estimates
```

```
fitting model and testing
```

```

res <- results(dds)
head(res)

log2 fold change (MLE): condition treatment vs control
Wald test p-value: condition treatment vs control
DataFrame with 6 rows and 6 columns
  baseMean log2FoldChange    lfcSE      stat     pvalue
  <numeric>      <numeric> <numeric> <numeric> <numeric>
ENST00000539570  0.000000        NA       NA       NA       NA
ENST00000576455  0.761453  3.155061  4.86052  0.6491203  0.516261
ENST00000510508  0.000000        NA       NA       NA       NA
ENST00000474471  0.484938  0.181923  4.24871  0.0428185  0.965846
ENST00000381700  0.000000        NA       NA       NA       NA
ENST00000445946  0.000000        NA       NA       NA       NA
  padj
  <numeric>
ENST00000539570    NA
ENST00000576455    NA
ENST00000510508    NA
ENST00000474471    NA
ENST00000381700    NA
ENST00000445946    NA

```