

Preparation



• Create the DESeq2 object

```
library(DESeq2)
mr$Group <- factor(mr$Group)
d <- DESeqDataSetFromMatrix(countData=cf,colData=mr,design=~Group)
d</pre>
```

```
## class: DESeqDataSet
## dim: 10573 6
## metadata(1): version
## assays(1): counts
## rownames(10573): ENSMUSG00000098104 ENSMUSG000000033845 ...
## ENSMUSG00000063897 ENSMUSG00000095742
## rowData names(0):
## colnames(6): DSSd00_1 DSSd00_2 ... DSSd07_2 DSSd07_3
## colData names(7): SampleName SampleID ... Group Replicate
```

- Categorical variables must be factors
- Building GLM models: ~var , ~covar+var

Size factors



• Normalisation factors are computed

```
d <- DESeq2::estimateSizeFactors(d,type="ratio")
sizeFactors(d)

## DSSd00_1 DSSd00_2 DSSd00_3 DSSd07_1 DSSd07_2 DSSd07_3
## 1.0136617 0.9570561 0.9965245 1.0354178 1.0780855 1.0017753</pre>
```

Dispersion



- Dispersion is a measure of variability in gene expression for a given mean
- Dispersion is unreliable for low mean counts





Dispersion



- Genes with similar mean values must have similar dispersion
- Estimate likely (ML) dispersion for each gene based on counts
- Fit a curve through the gene-wise estimates
- Shrink dispersion towards the curve

d <- DESeq2::estimateDispersions(d)</pre>







Log2 fold changes changes are computed after GLM fitting FC = counts group B / counts group A

```
dg <- nbinomWaldTest(d)
resultsNames(dg)

## [1] "Intercept" "Group_day07_vs_day00"</pre>
```

- Use results() to customise/return results
 - Set coefficients using contrast or name
 - Filtering results by fold change using lfcThreshold
 - cooksCutoff removes outliers
 - independentFiltering removes low count genes
 - pAdjustMethod sets method for multiple testing correction
 - o alpha set the significance threshold



```
res <- results(dg,name="Group_day07_vs_day00",alpha=0.05)
summary(res)

##

## out of 10573 with nonzero total read count

## adjusted p-value < 0.05

## LFC > 0 (up) : 193, 1.8%

## LFC < 0 (down) : 238, 2.3%

## outliers [1] : 1, 0.0095%

## low counts [2] : 4920, 47%

## (mean count < 21)

## [1] see 'cooksCutoff' argument of ?results

## [2] see 'independentFiltering' argument of ?results
```

Alternative way to specify contrast

```
results(dg,contrast=c("Group","day07","day00"),alpha=0.05)
```



head(res)

```
## log2 fold change (MLE): Group day07 vs day00
## Wald test p-value: Group day07 vs day00
## DataFrame with 6 rows and 6 columns
                       baseMean log2FoldChange
                                                   1.fcSF
##
                                                              stat
                                                                      pvalue
##
                      <numeric>
                                     <numeric> <numeric> <numeric> <numeric>
  FNSMUSG00000098104
                       18.8505
                                     0.205656 0.401543
                                                          0.512164 0.6085362
   ENSMUSG00000033845
                       23.3333
                                     0.653565 0.379627 1.721596 0.0851426
                       37, 1016
   FNSMUSG00000025903
                                     0.672348 0.298923 2.249232 0.0244977
  FNSMUSG000000033793
                       33.3673
                                     0.144833 0.305139
                                                          0.474646 0.6350394
                       22.3875
  FNSMUSG00000025907
                                     0.821006 0.376414
                                                          2.181125 0.0291742
## ENSMUSG00000051285
                       21.1485
                                     0.452451 0.378725 1.194669 0.2322163
##
                          padi
                      <numeric>
##
   ENSMUSG00000098104
                             NA
   ENSMUSG00000033845
                       0.377432
  ENSMUSG00000025903
                       0.177491
  ENSMUSG00000033793
                       0.886264
## ENSMUSG00000025907
                       0.201741
## ENSMUSG00000051285
                             NA
```



- Use IfcShrink() to correct fold changes for genes with high dispersion or low counts
- Does not change number of DE genes



