Statistical Methods for High Dimensional Biology

More limma & multiple testing

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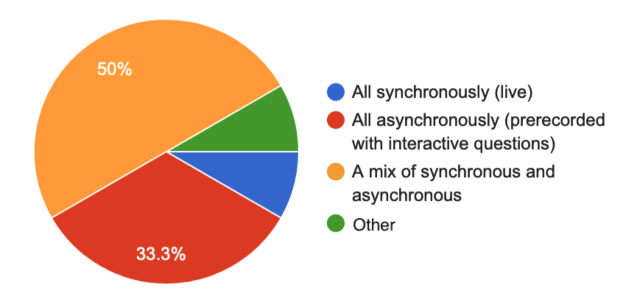
8 February 2021

with slide contributions from Gabriela Cohen Freue, Jenny Bryan, and Sara Mostafavi

Announcements

- Feedback posted on your initial project proposals (see the issue in your team's repository)
- Reminder that the finalized project proposals are due Feb 18
- Instructions on how to submit your deliverables have been updated for clarity on the website
 - in particular pay attention to the recommended way to grab the link to the submitted files from your latest commit

Course format feedback



- Preference for mix of synchronous & asynchronous
- © Positive feedback on interactive quiz questions / polls / chat
- 😉 Several people felt a lack of communication let's utilize the Discussion repo more!

Learning objectives

- How to use limma to perform genome-wide differential expression testing on microarray data
- Understand the key differences between limma and lm
- Explain why multiple testing increases the number of errors we make by chance
- Be able to adjust for multiple comparisons by controlling the False Discovery Rate
 - o e.g. using Benjamini-Hochberg or Storey's q-value

Recall: The hybrid estimator in limma

$$ilde{s}_{g}^{2}=rac{d_{0}s_{0}^{2}+ds_{g}^{2}}{d_{0}+d}$$

• recall that (s_0,d_0) are the *prior* parameters for σ_g^2 (random variable):

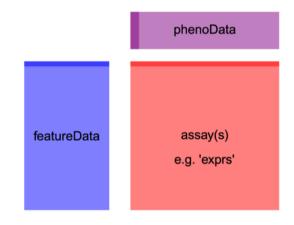
$$rac{1}{\sigma_g^2} \sim rac{1}{d_0 s_0^2} \chi_{d_0}^2$$

- the prior parameters incorporate information from all genes which allows us to shrink/nudge the gene-specific variances toward a common consensus
 - \circ they are estimated from the data the formulas for (s_0,d_0) and their derivations are beyond the scope of the course (but limma takes care of the details for us)
 - \circ note that (s_0,d_0) do not depend on g

limma quickstart

$$\mathbf{Y}_g = \mathbf{X}oldsymbol{lpha}_g + oldsymbol{arepsilon}_g, \ E(oldsymbol{arepsilon}_g) = 0, \; Var(oldsymbol{arepsilon}_g) = \sigma_g^2, \; arepsilon_{ig} \perp arepsilon_{jg},$$

- Within each gene observations are iid / constant variance
- lmFit() function in limma carries out multiple linear regression on each gene
- Usage: lmFit(myDat, desMat)
 - myDat is a data frame, matrix with features in rows and samples in columns (*G* genes by *N* samples), or ExpressionSet object
 - desMat is a design matrix (output of model.matrix(y ~ x); N samples by p parameters)



Bioconductor's ExpressionSet class

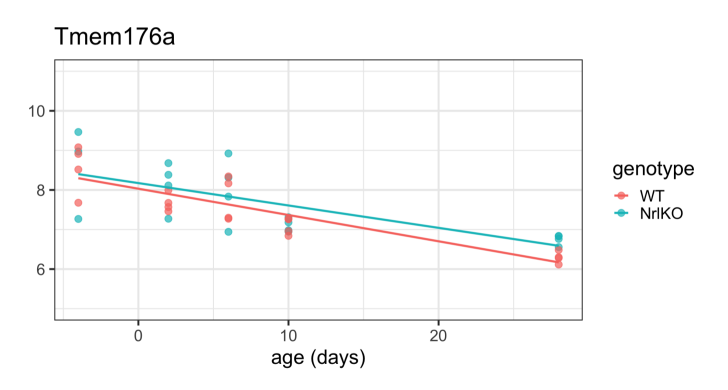
Let's run limma for the interactive model with age

$$y_{ig} = heta + au_{KO} x_{ig,KO} + au_{Age} x_{ig,Age} + au_{KO:Age} x_{ig,KO} x_{ig,Age}$$

- $ullet \ i \ {\it indexes} \ {\it mouse}, \ {\it g} \ {\it indexes} \ {\it genes}$
- ullet $x_{iq,KO}$ is the dummy/indicator variable for the NrlKO group
- ullet $x_{ig,Age}$ is the continuous age variable

Let's run limma for the interactive model with age

Example gene (but we want to fit this model on all genes):



Arranging input for lmFit: Bioconductor way

```
eset
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 45101 features, 39 samples
##
   element names: exprs
## protocolData: none
## phenoData
##
    sampleNames: GSM92610 GSM92611 ... GSM92648 (39 total)
    varLabels: sample_id dev_stage genotype age
##
##
    varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
    pubMedIds: 16505381
##
## Annotation: GPI 1261
```

Arranging input for \lambda mF i t: Separate expression & metadata

```
myDat <- assayData(eset)$exprs</pre>
myDat[1:5,1:5]
##
               GSM92610 GSM92611
                                   GSM92612 GSM92613
                                                      GSM92614
## 1415670 at
              7.108863 7.322392
                                  7.420947 7.351444
                                                     7,240428
                                  9.831072 9.658442
## 1415671 at
               9.714002 9.797742
                                                      9,708906
## 1415672 at 9.429030 9.846977 10.003092 9.914112 10.174548
## 1415673 at
               8.426974 8.404206
                                 8.594600 8.404206
                                                      8,835334
## 1415674 a at 8.498338 8.458287 8.426651 8.372776 8.541722
myMeta <- pData(eset)</pre>
myMeta[1:5,]
##
            sample_id dev_stage genotype age
## GSM92610 GSM92610
                                   NrlKO
                                         28
                                   NrlKO
## GSM92611 GSM92611
                             4W
                                         28
## GSM92612 GSM92612
                                   NrlKO
                                         28
                             4W
                                   NrlKO
## GSM92613 GSM92613
                             4W
                                         28
## GSM92614 GSM92614
                                   NrlKO
                            E16
```

Formulating input for \lambda mFit: Design Matrix

Bioconductor way:

Equivalently, if using the separate way:

desMat									
##		(Intercept)	genotypeNrlKO	age	<pre>genotypeNrlKO:age</pre>	!			
##	GSM92610	1	1	28	28				
##	GSM92611	1	1	28	28				
##	GSM92612	1	1	28	28				
##	GSM92613	1	1	28	28				
##	GSM92614	1	1	-4	-4				
##	GSM92615	1	1	-4	-4				
##	GSM92616	1	1	-4	-4	•			
##	GSM92617	1	1	10	10	1			
##	GSM92618	1	1	10	10	1			
##	GSM92619	1	1	10	10	1			
##	GSM92620	1	1	10	10	1			
##	GSM92621	1	1	2	2	•			
##	GSM92622	1	1	2	2	•			
##	GSM92623	1	1	2	2				
##	GSM92624	1	1	2	2				
##	GSM92625	1	1	6	6	i			
##	GSM92626	1	1	6	6	i			
##	GSM92627	1	1	6	6	i			
##	GSM92628	1	1	6	6	i			
##	GSM92629	1	0	28	0	1			
##	GSM92630	1	0	28	0	1			
##	GSM92631	1	0	28	0	1			
##	GSM92632	1	0	28	Θ	1			
##	GSM92633	1	0	-4	0	1			
##	GSM92634	1	0	-4	0	1			
##	GSM92635	1	0	-4	0	1			
##	GSM92636	1	Θ	-4	Θ	1			
##	GSM92637	1	0	10	0	1			
##	GSM92638	1	0	10	0	1			

Computation is fast

• Equivalently, using the 'separate' way:

- Using lmFit to fit an interactive model on 45K probesets takes a fraction of a second
- The time-intensive parts of an analysis lie in selecting the model and covariates, choosing how to parameterize it, and interpreting the output

Output of lmFit

```
summary(gFit)
##
                    Length Class Mode
## coefficients
                    180404 -none- numeric
## rank
                         1 -none- numeric
## assign
                         4 -none- numeric
## qr
                         5 qr
                                   list
## df.residual
                     45101 -none- numeric
## sigma
                     45101 -none- numeric
## cov.coefficients
                        16 -none- numeric
## stdev.unscaled
                    180404 -none- numeric
## pivot
                         4 -none- numeric
## Amean
                     45101 -none- numeric
## method
                         1 -none- character
## design
                       156 -none- numeric
```

```
nrow(eset)

## Features
## 45101

nrow(eset)*4

## Features
## 180404
```

Output of lmFit

method

design

```
summary(gFit)
##
                    Length Class Mode
## coefficients
                    180404 -none- numeric
## rank
                          1 -none- numeric
## assign
                         4 -none- numeric
## qr
                         5 qr
                                   list
## df.residual
                     45101 -none- numeric
## sigma
                     45101 -none- numeric
## cov.coefficients
                        16 -none- numeric
## stdev.unscaled
                    180404 -none- numeric
## pivot
                         4 -none- numeric
                     45101 -none- numeric
## Amean
```

1 -none- character

156 -none- numeric

```
nrow(eset)

## Features
## 45101

nrow(eset)*4

## Features
## 180404
```

- OK... but where are the shrunken variable estimates?? How do I pull out p-values??
- Actually, we haven't carried out the empirical Bayesian computation yet -- need to run eBayes()!

eBayes()

summary(gFit)

```
##
                    Length Class Mode
## coefficients
                    180404 -none- numeric
## rank
                         1 -none- numeric
## assign
                         4 -none- numeric
## qr
                         5 qr
                                  list
## df.residual
                     45101 -none- numeric
## sigma
                     45101 -none- numeric
## cov.coefficients
                        16 -none- numeric
## stdev.unscaled
                    180404 -none- numeric
## pivot
                         4 -none- numeric
## Amean
                     45101 -none- numeric
## method
                         1 -none- character
## design
                       156 -none- numeric
```

summary(ebFit <- eBayes(gFit))</pre>

```
##
                    Length Class Mode
## coefficients
                    180404 -none- numeric
## rank
                         1 -none- numeric
## assign
                         4 -none- numeric
                                  list
## qr
                         5 ar
## df.residual
                     45101 -none- numeric
## sigma
                     45101 -none- numeric
## cov.coefficients
                        16 -none- numeric
## stdev.unscaled
                    180404 -none- numeric
## pivot
                         4 -none- numeric
## Amean
                     45101 -none- numeric
## method
                         1 -none- character
## design
                       156 -none- numeric
## df.prior
                         1 -none- numeric
## s2.prior
                         1 -none- numeric
## var.prior
                         4 -none- numeric
## proportion
                         1 -none- numeric
## s2.post
                     45101 -none- numeric
## t
                    180404 -none- numeric
## df.total
                     45101 -none- numeric
## p.value
                    180404 -none- numeric
## lods
                    180404 -none- numeric
## F
                     45101 -none- numeric
## F.p.value
                     45101 -none- numeric
```

Components of the empirical Bayes estimators

math	plain english	limma	numerical result	also in lm?
\boldsymbol{s}_g^2	gene-specific residual variance	gFit\$sigma^2	45K numbers	\checkmark
d	residual degrees of freedom $\left(n-p ight)$	gFit\$df.residual	$39 - 4 = 35^*$	\checkmark
\boldsymbol{s}_0^2	mean of inverse χ^2 prior for s_g^2	ebFit\$s2.prior	0.048	
d_0	degrees of freedom for the prior	ebFit\$df.prior	2.61	
$ ilde{oldsymbol{s}}_g^2$	posterior mean of s_g^2 (i.e. moderated residual variance)	ebFit\$s2.post	45K numbers	

^{*} limma can handle more complicated models where this is not the same for each gene, so this is actually a vector of 45K copies of the number 35

topTable() will help us extract relevant output in a convenient format!

topTable(fit, coef=NULL, number=10, genelist=fit\$genes, adjust.method="BH", sort.by="B", resort.by=NULL, p.value=1, lfc=0, confint=FALSE) fit list containing a linear model fit produced by lmFit, lm.series, gls.series or mrlm. For topTable, fit should be an object of class MArrayLM as produced by lmFit and eBayes. coef column number or column name specifying which coefficient or contrast of the linear model is of interest. For topTable, can also be a vector of column subscripts, in which case the gene ranking is by F-statistic for that set of contrasts. number maximum number of genes to list adjust.method method used to adjust the p-values for multiple testing. Options, in increasing conservatism, include "none", "BH", "BY" and "holm". See p. adjust for the complete list of options. A NULL value will result in the default adjustment method, which is "BH". sort.by character string specifying statistic to rank genes by. Possible values for topTable and toptable are "logFC", "AveExpr", "t", "p", "p", "B" or "none". (Permitted synonyms are "M" for "logFC", "A" or "Amean" for "AveExpr", "T" for "t" and "p" for "P".) Possibilities for topTableF are "F" or "none". Possibilities for topTreat are as for topTable except for "B".

^{... (}truncated - see ?topTable for full listing)

Summary of topTable function

- coef is the argument where you specify the coefficient(s) you want to test for equality with zero (default is NULL; must be specified)
- number lets you control size of hit list (default is 10)
- p.value lets you specify a minimum adjusted p-value cutoff (default is 1)
- 1fc lets you specify a minimum observed effect size log2 fold change (default is 0)
- sort.by and resort.by give control over the ordering (default is by "B": log-odds that the gene is differentially expressed)
- adjust.method specifies how/if to adjust p-values for multiple testing (default is BH)

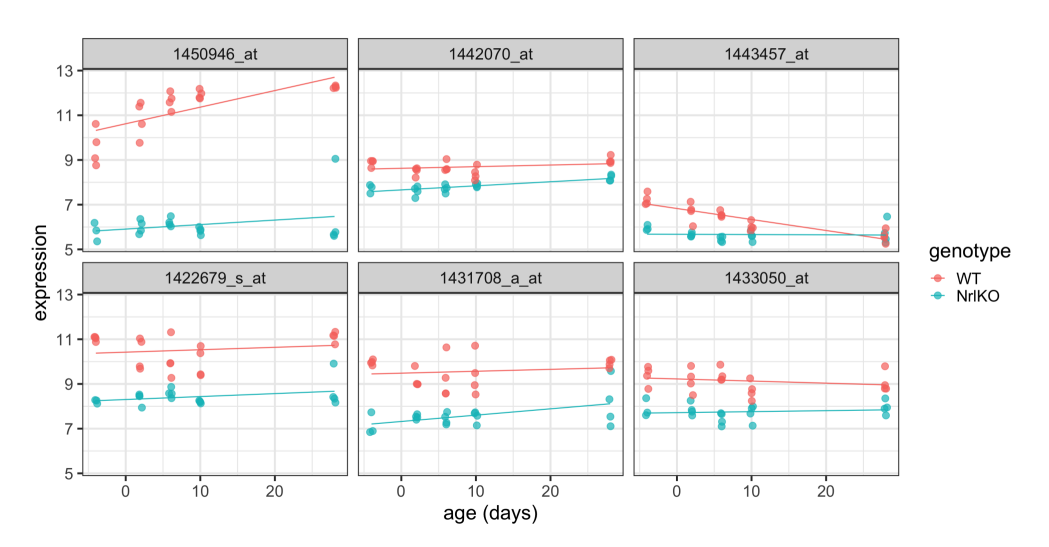
topTable in action: genotypeNrlKO

```
topTable(ebFit, coef = "genotypeNrlKO")
```

```
##
                   logFC AveExpr t
                                                 P.Value
                                                            adi.P.Val
## 1450946_at -4.7134891 8.733103 -15.583683 5.390931e-18 2.431364e-13 28.40938
## 1442070 at -0.9669400 8.268122 -9.787407 6.896220e-12 1.555132e-07 16.46086
## 1443457 at -1.1530057 6.049428
                                   -9.091412 4.947029e-11 6.029428e-07 14.68843
## 1422679 s at -2.1187394 9.495435 -9.002625 6.388954e-11 6.029428e-07 14.45703
## 1431708 a at -2.1610625 8.591450
                                   -8.967950 7.062007e-11 6.029428e-07 14.36634
## 1433050 at -1.5083887 8.468891 -8.923932 8.021234e-11 6.029428e-07 14.25096
## 1426288 at
               -4.2100657 9.323796
                                   -8.772209 1.246491e-10 8.031144e-07 13.85106
## 1418108 at 0.8662091 8.260647 8.537984 2.475294e-10 1.395478e-06 13.22713
## 1450770 at 1.3103576 7.357033 8.128847 8.332369e-10 3.747976e-06 12.11870
## 1457802_at
               -0.8362030 7.778189 -8.117112 8.629984e-10 3.747976e-06 12.08657
```

- topTable(ebFit, coef = 2) is equivalent here, but much less informative!!
- What is the null hypothesis here?
 - $\bullet H_0: \tau_{KO}=0$
 - this finds genes where the knockouts differ from the wild types when age is zero

Plotting the top 6 probes for genotypeNrlKO



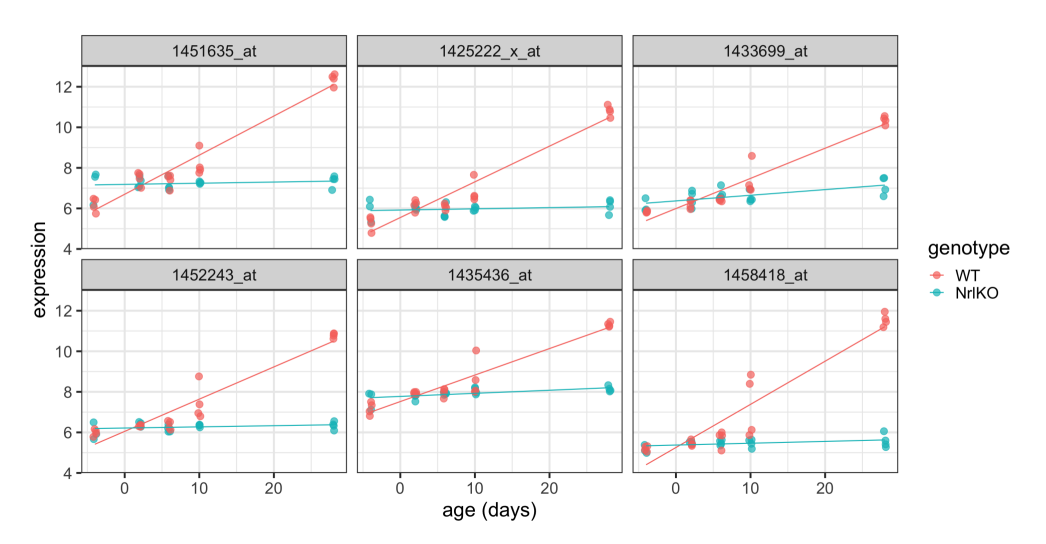
topTable in action: age

```
topTable(ebFit, coef = "age")
```

```
##
                    logFC AveExpr t P.Value
                                                            adi.P.Val
                                                                             В
## 1451635 at
               0.19294061 7.791237 20.78935 3.204717e-22 1.445359e-17 40.60768
## 1425222 x at 0.17604111 6.514475 19.72633 1.962954e-21 4.426561e-17 38.79502
## 1433699 at
               0.14875031 6.939973 17.80467 6.442530e-20 7.957605e-16 35.29412
               0.15889179 6.840975 17.75658 7.057586e-20 7.957605e-16 35.20253
## 1452243 at
## 1435436 at
               0.13067119 8.274676 17.61808 9.187120e-20 8.286966e-16 34.93763
## 1458418 at
               0.21277027 6.270187 16.43686 9.326171e-19 7.006477e-15 32.60718
## 1424977 at
               0.07814988 6.505464 16.36079 1.087456e-18 7.006477e-15 32.45260
               0.16158579 7.418000 16.01901 2.183055e-18 1.230725e-14 31.75106
## 1431174 at
## 1421818 at
               0.11250919 7.982278 15.66666 4.531133e-18 2.270651e-14 31.01562
## 1419069 at
               0.09325234 8.222096 15.54775 5.813459e-18 2.621928e-14 30.76457
```

- topTable(ebFit, coef = 3) is equivalent here, but much less informative!!
- What is the null hypothesis here?
 - $\circ~H_0: au_{Age}=0$
 - this finds genes where age significantly affects gene expression for WT

Plotting the top 6 genes for age



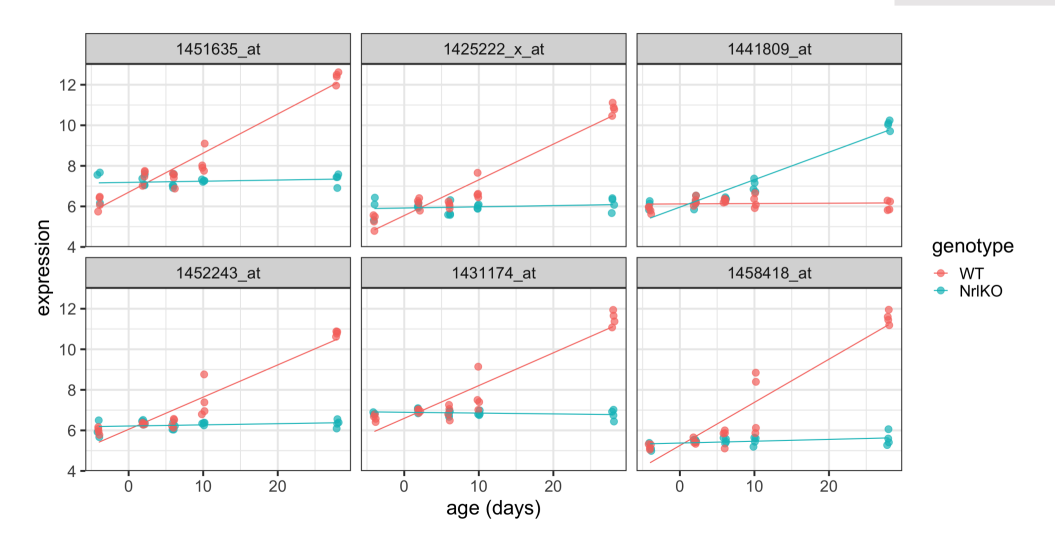
topTable in action: genotypeNrlKO: age

```
topTable(ebFit, coef = "genotypeNrlKO:age")
```

```
##
                    logFC AveExpr t P.Value
                                                             adi.P.Val
## 1451635_at
               -0.1871800 7.791237 -14.00463 1.684302e-16 7.596369e-12 27.52010
## 1425222_x_at -0.1700908 6.514475 -13.23451 9.961393e-16 2.246344e-11 25.74877
## 1441809 at
                0.1336356 6.649440 12.55702 5.037221e-15 7.572790e-11 24.13110
               -0.1531326 6.840975 -11.88284 2.670693e-14 3.011273e-10 22.46414
## 1452243 at
## 1431174 at
               -0.1655894 7.418000 -11.39882 9.161231e-14 8.263614e-10 21.23123
## 1458418 at
               -0.2037314 6.270187 -10.92850 3.122990e-13 2.347500e-09 20.00384
## 1435436 at
               -0.1154500 8.274676 -10.80855 4.289363e-13 2.763637e-09 19.68613
## 1446715 at
               -0.1409774 5.613122 -10.57909 7.912260e-13 4.325169e-09 19.07307
## 1442190 at
                -0.1323038 5.896077 -10.54672 8.630965e-13 4.325169e-09 18.98601
## 1416306 at
                0.1702059 \ 6.560050 \ 10.41285 \ 1.238250e-12 \ 5.584630e-09 \ 18.62456
```

- topTable(ebFit, coef = 4) is equivalent here, but much less informative!!
- What is the null hypothesis here?
 - $\circ H_0: au_{KO:Age} = 0$
 - o this finds genes where the effect of age is significantly different in each genotype

Plotting the top 6 genes for genotypeNrlKO: age



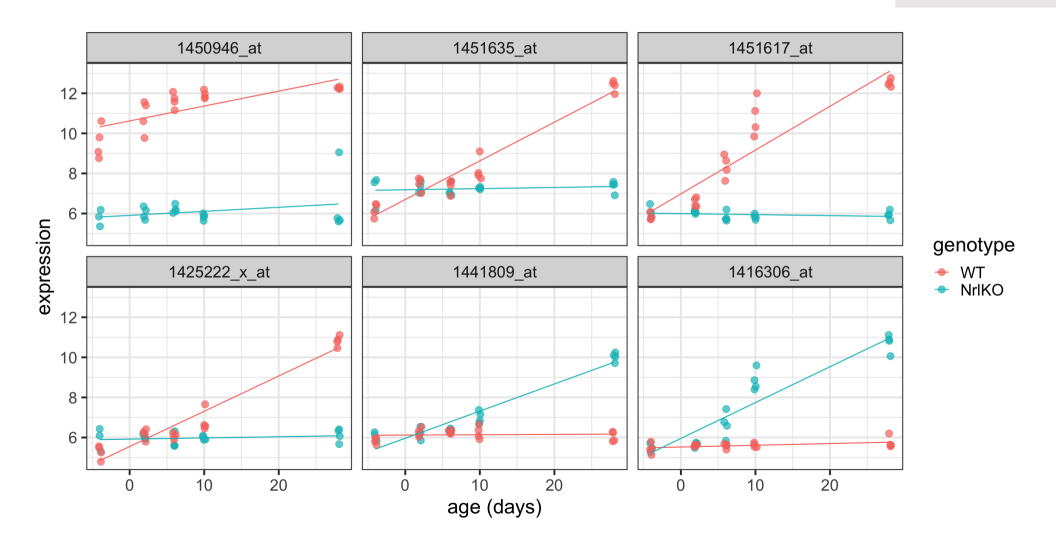
topTable in action: any effect of genotype

```
topTable(ebFit, coef = c("genotypeNrlKO", "genotypeNrlKO:age"))[,-3]
```

```
##
               genotypeNrlKO.age F P.Value
                                                                       adi.P.Val
## 1450946 at
                  -4.7134891
                                   -0.05411298 246.5762 2.403638e-22 1.084065e-17
## 1451635 at
                   0.4868705
                                   -0.18718005 129.8112 1.307185e-17 2.180049e-13
## 1451617 at
                  -0.9885684
                                   -0.22329233 128.9935 1.450111e-17 2.180049e-13
                  0.3698846
## 1425222 x at
                                   -0.17009077 120.0041 4.719613e-17 5.321482e-13
## 1441809 at
                  -0.1576064
                                    0.13363563 117.5571 6.594308e-17 5.948198e-13
## 1416306 at
                                    0.17020589 113.2125 1.212332e-16 9.112895e-13
                  0.4278776
## 1439143 at
                   0.6570269
                                    0.06170447 111.2102 1.616023e-16 1.001158e-12
                   0.9935801
                                    0.04232958 110.5598 1.775851e-16 1.001158e-12
## 1449526 a at
## 1452243 at
                   0.1639373
                                   -0.15313265 106.2784 3.344140e-16 1.675823e-12
## 1451763 at
                                   -0.14010359 102.7010 5.771358e-16 2.442992e-12
                  -0.6819899
```

- topTable(ebFit, coef = c(2,4)) is equivalent here, but much less informative!!
- What is the null hypothesis here?
 - $\circ~H_0: au_{KO}= au_{KO:Age}=0$
 - this finds genes where any (additive/interaction) effect of genotype is significant

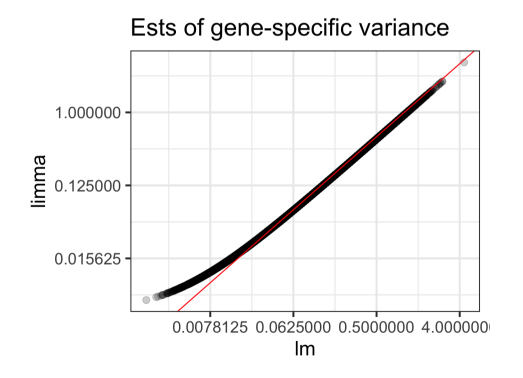
Plotting the top 6 genes for any effect of genotype



Comparison of s_g^2 and \tilde{s}_g^2 (shrinkage!)

For **small** variances, limma _____ the estimates (fill in the blank):

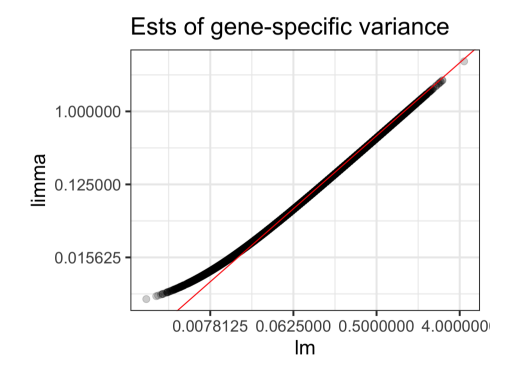
- (a) INCREASES
- (b) DECREASES



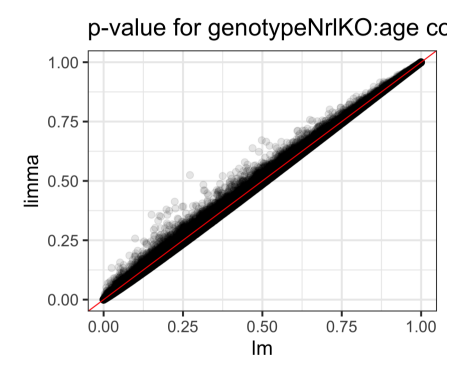
Comparison of s_g^2 and \tilde{s}_g^2 (shrinkage!)

For **large** variances, limma _____ the estimates (fill in the blank):

- (a) INCREASES
- (b) DECREASES



Comparison of interaction coefficient p-values



- 17479 genes where limma p-value is *larger* than lm
- 27622 genes where limma p-value is *smaller* than lm

Multiple testing

Error rates

Actual Situation "Truth" Decision H₀ False H₀ True **Correct Decision Incorrect Decision Do Not Type II Error** 1-α Reject H₀ β **Correct Decision Incorrect Decision** Reject Ho 1-β **Type I Error** α

$$\alpha = P(\text{Type I Error}), \ \beta = P(\text{Type II Error}), \ \text{Power} = 1 - \beta$$

Type I Error rate for m tests

- $P(\text{incorrect decision}|H_0) = \alpha$
 - \circ let lpha=0.05
- $P(\text{correct decision}|H_0) = 1 \alpha = 0.95$

Type I Error rate for m tests

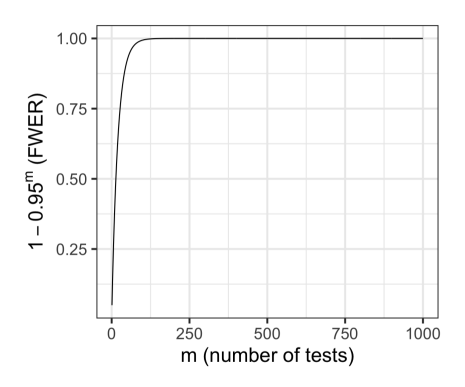
- $P(\text{incorrect decision}|H_0) = \alpha$
 - \circ let $\alpha = 0.05$
- $P(\text{correct decision}|H_0) = 1 \alpha = 0.95$
- $P(\text{correct decision on } m \text{ tests}|H_0) =$

$$(1-\alpha)^m = 0.95^m$$

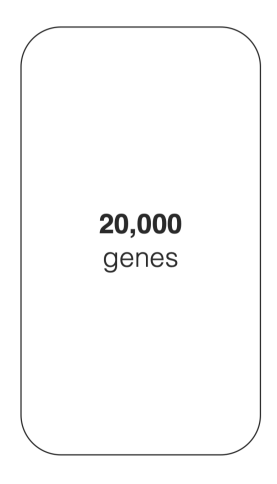
 $P({
m at\ least\ one\ incorrect\ decision\ on\ }m\ {
m tests}|H_0)=$

$$1 - (1 - \alpha)^m =$$

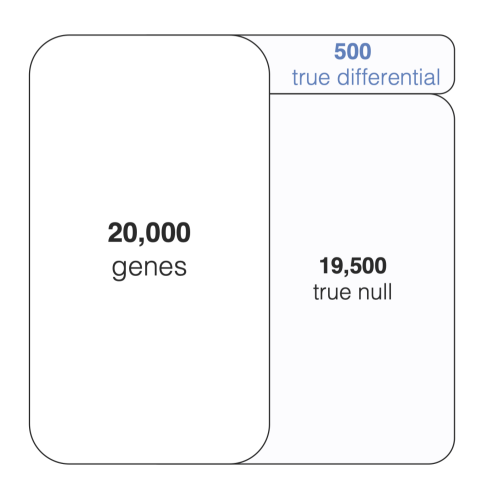
$$1 - 0.95^m = \alpha_{FWER}$$



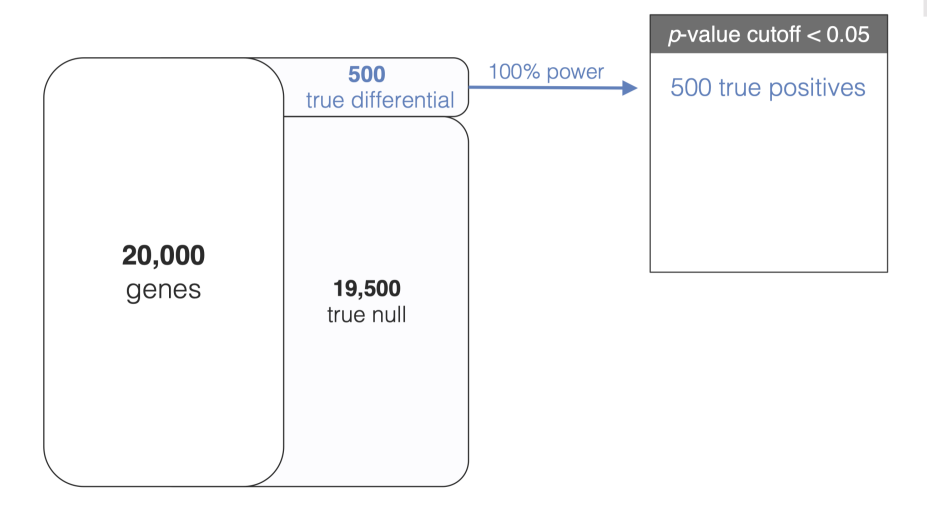
Multiple comparisons and error rates



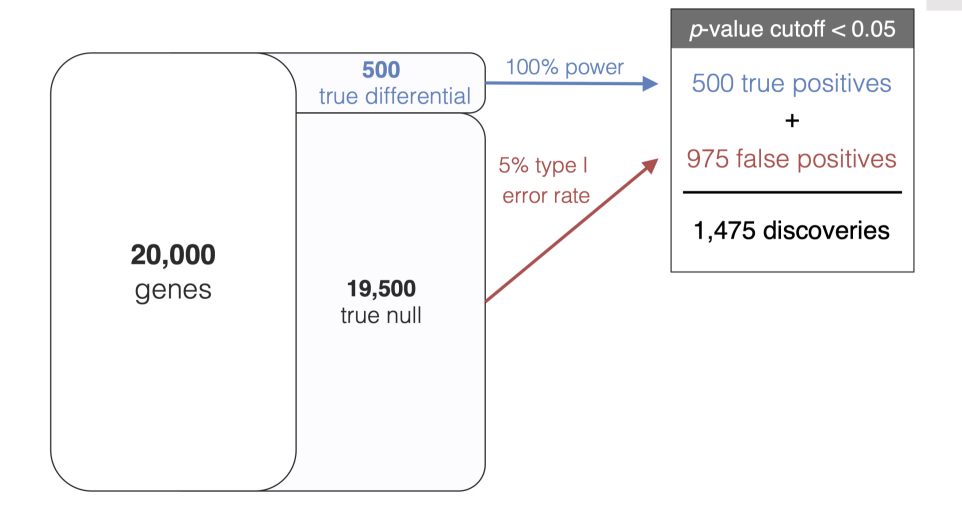
Multiple comparisons and error rates



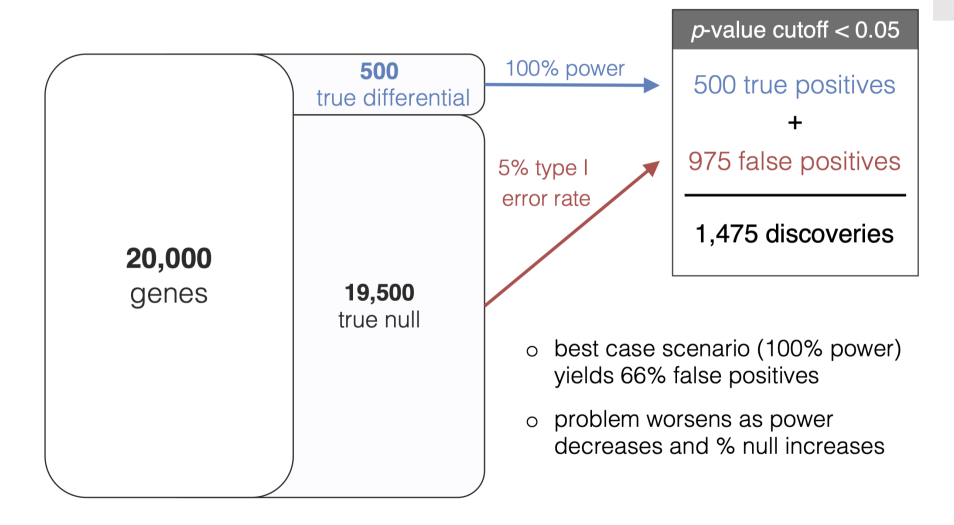
Multiple comparisons and error rates



Multiple comparisons and error rates



Multiple comparisons and error rates



Family-Wise Error Rate (FWER)

- FWER is the probability of making at least one error when testing m tests
- Control the FWER: limit the probability of making at least one incorrect decision
- One example: the **Bonferroni** correction for $\alpha=0.05$:

If
$$P(\text{at least one error on } m \text{ tests}) < \alpha$$

$$\Rightarrow P(ext{at least one error on } m ext{ tests}) < \sum_{i=1}^m P(ext{error on test } i)$$

$$\sum_{i=1}^m P(ext{error on test } i) = mlpha_{Bon}$$

$$lpha_{Bon}=rac{lpha}{m}=rac{0.05}{m}$$

Bonferroni correction: controlling the FWER

Can think of controlling the probability of at least one false positive in two ways:

1. Adjust the p-values; keep same α :

$$p_{Bon,i} = mp_i ext{ (more technically correct: } p_{Bon,i} = min(mp_i,1))$$

Then, threshold $p_{Bon,i}$ at α

2. Adjust the α threshold; keep same p-values:

$$\alpha_{Bon} = rac{lpha}{m}$$

Then, threshold p_i at α_{Bon}

Multiple test correction is an active area of statistical research

- Bonferroni correction is very **conservative** (i.e. controls the FWER even lower than α in many settings)
- Several other options are better
- For example, the Holm procedure: multiplier for p-value correction is not the same for all genes; more powerful

$$egin{aligned} p_{Holm,1} &= mp_1 \ p_{Holm,2} &= (m-1)p_2 \ p_{Holm,3} &= (m-2)p_3 \ &dots \ \end{pmatrix} egin{aligned} &dots \ &dots \ \end{pmatrix} FWER \leq lpha \end{aligned}$$

How practical is the FWER in high-throughput biology?

• Why do we care so much about making one single error??



- One extreme way to ensure no Type I errors: reject no hypotheses! 😃
 - However, then our power is zero... ⁶
- Being overly strict about Type I error leads to greater Type II error (loss of power)

Radical idea: it's OK to make multiple mistakes, as long as you also have some true positives!

Enter: the False Discovery Rate (FDR)

J. R. Statist. Soc. B (1995) 57, No. 1, pp. 289-300

Controlling the False Discovery Rate: a Practical and Powerful Approach to Multiple Testing

By YOAV BENJAMINI† and YOSEF HOCHBERG

Tel Aviv University, Israel

[Received January 1993. Revised March 1994]

SUMMARY

The common approach to the multiplicity problem calls for controlling the familywise error rate (FWER). This approach, though, has faults, and we point out a few. A different approach to problems of multiple significance testing is presented. It calls for controlling the expected proportion of falsely rejected hypotheses—the false discovery rate. This error rate is equivalent to the FWER when all hypotheses are true but is smaller otherwise. Therefore, in problems where the control of the false discovery rate rather than that of the FWER is desired, there is potential for a gain in power. A simple sequential Bonferronitype procedure is proved to control the false discovery rate for independent test statistics, and a simulation study shows that the gain in power is substantial. The use of the new procedure and the appropriateness of the criterion are illustrated with examples.

Benjamini Y, Hochberg Y. "Controlling the false discovery rate: a practical and powerful approach to multiple testing." Journal of the Royal statistical society: series B (Methodological). 1995 Jan;57(1):289-300.

Over 60K citations!!

False Discovery Rate

	Null	Alternative	
	True	True	Total
Not Called Significant	U	τ	m - R
Called Significant	V	S	R
	m_0	<i>m-m</i> ₀	m

V = # Type I errors [false positives]

FDR is designed to control the expected proportion of false positives (V) among all hypotheses where the null has been rejected (R)

False Discovery Rate

	Null	Alternative	
	True	True	Total
Not Called Significant	U	T	m - R
Called Significant	V	S	R
	m_0	<i>m-m</i> ₀	m

V = # Type I errors [false positives]

$$FDR = E\Big[rac{V}{R}\Big]$$

FDR vs FPR vs FWER

• False Discovery Rate (FDR) is the rate that significant features (R) are truly null

$$FDR = E\Big[rac{V}{R}\Big]$$

• False Positive Rate (FPR) is the rate that truly null features (m_0) are called significant

$$FPR = E\Big[rac{V}{m_0}\Big]$$

• Family-Wise Error Rate (FWER) is the probability that the number of truly null features rejected (V) is at least 1

$$\mathrm{FWER}\ = P(V \geq 1)$$

Benjamini-Hochberg FDR (BH procedure)

- Proposed the idea of controlling FDR instead of FWER
- Proposed a procedure for doing so
 - \circ note that we know R, but we don't know V
- Procedure: control FDR at level q
 - 1. order the raw p-values $p_1 \leq p_2 \leq \ldots \leq p_m$
 - 2. find test with highest rank j such that $p_j < rac{jq}{m}$
 - 3. declare all smaller ranks up to j significant

Rank (j)	P-value
1	0.0008
2	0.009
3	0.127
4	0.205
5	0.396
6	0.450
7	0.641
8	0.781
9	0.900
10	0.993

Rank (j)	P-value	(j/m)*q
1	0.0008	0.005
2	0.009	0.010
3	0.127	0.015
4	0.205	0.020
5	0.396	0.025
6	0.450	0.030
7	0.641	0.035
8	0.781	0.040
9	0.900	0.045
10	0.993	0.050

Rank (j)	P-value	(j/m)*q	Reject H_0 ?
1	0.0008	0.005	\checkmark
2	0.009	0.010	\checkmark
3	0.127	0.015	
4	0.205	0.020	
5	0.396	0.025	
6	0.450	0.030	
7	0.641	0.035	
8	0.781	0.040	
9	0.900	0.045	
10	0.993	0.050	

Rank (j)	P-value	(j/m)*q	Reject H_0 ?	$\overline{FWER_{Bon} < 0.05$?
1	0.0008	0.005	\checkmark	\checkmark
2	0.009	0.010	\checkmark	
3	0.127	0.015		
4	0.205	0.020		
5	0.396	0.025		
6	0.450	0.030		
7	0.641	0.035		
8	0.781	0.040		
9	0.900	0.045		
10	0.993	0.050		

Where $lpha_{Bon}=0.05/10=0.005$

BH FDR values given in limma by default

```
topTable(ebFit, coef = "genotypeNrlKO")
##
                    logFC AveExpr
                                                   P.Value
                                                             adi.P.Val
## 1450946_at -4.7134891 8.733103 -15.583683 5.390931e-18 2.431364e-13 28.40938
              -0.9669400 8.268122 -9.787407 6.896220e-12 1.555132e-07 16.46086
## 1442070 at
## 1443457 at
               -1.1530057 6.049428 -9.091412 4.947029e-11 6.029428e-07 14.68843
## 1422679 s at -2.1187394 9.495435
                                    -9.002625 6.388954e-11 6.029428e-07 14.45703
## 1431708 a at -2.1610625 8.591450
                                    -8.967950 7.062007e-11 6.029428e-07 14.36634
## 1433050 at
               -1.5083887 8.468891
                                    -8.923932 8.021234e-11 6.029428e-07 14.25096
## 1426288 at
               -4.2100657 9.323796
                                    -8.772209 1.246491e-10 8.031144e-07 13.85106
## 1418108 at 0.8662091 8.260647
                                    8.537984 2.475294e-10 1.395478e-06 13.22713
## 1450770 at 1.3103576 7.357033
                                    8.128847 8.332369e-10 3.747976e-06 12.11870
## 1457802 at
               -0.8362030 7.778189
                                    -8.117112 8.629984e-10 3.747976e-06 12.08657
```

Or, obtain them yourself for any vector of p-values p with p.adjust(p, method="BH")

Other ways to control FDR

- BH is just one (the first) method to control FDR
- Since the publication of the BH method, other methods have been proposed
- One of the most popular is Storey's q-value

Statistical significance for genomewide studies

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Edited by Philip P. Green, University of Washington School of Medicine, Seattle, WA, and approved May 30, 2003 (received for review January 28, 2003)

With the increase in genomewide experiments and the sequencing of multiple genomes, the analysis of large data sets has become commonplace in biology. It is often the case that thousands of features in a genomewide data set are tested against some null hypothesis, where a number of features are expected to be significant. Here we propose an approach to measuring statistical significance in these genomewide studies based on the concept of the false discovery rate. This approach offers a sensible balance between the number of true and false positives that is automatically calibrated and easily interpreted. In doing so, a measure of statistical significance called the q value is associated with each tested feature. The q value is similar to the well known p value, except it is a measure of significance in terms of the false discovery rate rather than the false positive rate. Our approach avoids a flood of false positive results, while offering a more liberal criterion than what has been used in genome scans for linkage.

false discovery rates | genomics | multiple hypothesis testing | q values

to the method in ref. 5 under certain assumptions. Also, ideas similar to FDRs have appeared in the genetics literature (1, 13).

Similarly to the p value, the q value gives each feature its own individual measure of significance. Whereas the p value is a measure of significance in terms of the false positive rate, the q value is a measure in terms of the FDR. The false positive rate and FDR are often mistakenly equated, but their difference is actually very important. Given a rule for calling features significant, the false positive rate is the rate that truly null features are called significant. The FDR is the rate that significant features are truly null. For example, a false positive rate of 5% means that on average 5% of the truly null features in the study will be called significant. A FDR of 5% means that among all features called significant, 5% of these are truly null on average.

The \dot{q} value provides a measure of each feature's significance, automatically taking into account the fact that thousands are simultaneously being tested. Suppose that features with q values 5% are called significant in some genomewide test of significance. This results in a FDR of 5% among the significant features. A

• qvalue package implementation: provides adjusted p-values

Storey's q-value vs BH (Conceptual)

- Just like BH, is focused on the proportion of discoveries that are false positives
- Conceptual difference between BH and Storey's q-value is:
 - BH **controls** the FDR
 - q-values give an unbiased **estimate** of the FDR (will control the FDR on eaverage)

Storey's q-value vs BH (Mathematical)

- Mathematically, the difference between the two is in how m_0 is estimated
 - \circ Or equivalently, how $\pi_0=rac{m_0}{m}$ is estimated (since m is known)
 - \circ π_0 represents the proportion of tests that are truly null
- q-value:

$$\hat{q}\left(p_{i}
ight)=\min_{i}\left(rac{\hat{\pi}_{0}m}{rank(p_{i})}p_{i},\,1
ight)$$

ullet q-value and BH-adjusted p-values are equivalent when $\pi_0=1$

$$\hat{p}_{BH}(p_i) = \min_i \left(rac{m}{rank(p_i)}p_i,\,1
ight).$$

(BH conservatively assumes that $\pi_0=1$)

BH vs q-value in our example

Rank (j)	P-value	$\hat{p}_{BH}(p_i)$	$\hat{q}\left(p_{i} ight)$
1	0.0008	0.008	0.008
2	0.009	0.045	0.045
3	0.127	0.423	0.423
4	0.205	0.513	0.513
5	0.396	0.792	0.750
6	0.450	0.750	0.750
7	0.641	0.916	0.916
8	0.781	0.976	0.976
9	0.900	1.000	0.993
10	0.993	0.993	0.993

Compounding issues of multiple comparisons

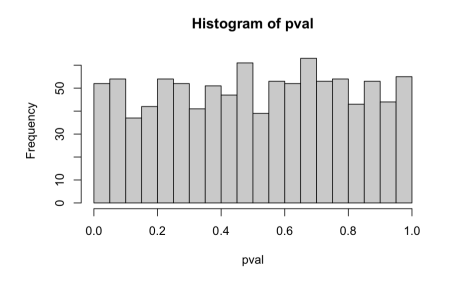
- What if you're not only testing 45K genes, but also multiple tests per gene (e.g. multiple contrasts, such as several two-group comparisons)?
- Classical procedures for adjustment:
 - Tukey multiple comparison procedure
 - Scheffe multiple comparison procedure
 - Bonferroni or Holm FWER correction
- In our setting, we can also apply BH to all p-values globally
 - o limma::decideTests(pvals, method="global") for a matrix of p-values or eBayes output (e.g. rows = genes, columns = contrasts)
 - o p-values are combined, adjusted globally, then separated back out and sorted

Assumptions about p-values

• Implicit assumption for all multiple testing correction methods:

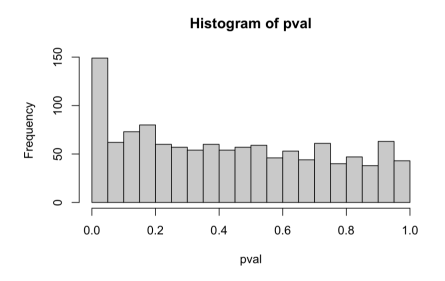
p-value distribution is "well-behaved"

- What does this mean?
 - primarily, that the distribution of p-values under the null is **uniform**



p-value distributions

Spike of small p-values indicates non-null tests:



Great primer on how things can go wrong: http://varianceexplained.org/statistics/interpreting-pvalue-histogram/

What if p-values are *poorly behaved*?

- FDR estimates can be invalid (assumptions are violated)
- Solution: compute p-values "empirically" using resampling/permutation/bootstrap techniques
- **Bootstrap**: take repeated random samples with replacement from your data and compute statistic; repeat many times and use bootstrap statistics as your sampling distribution rather than a t, Normal, F, χ^2 , etc
- **Permutation**: construct a simulated version of your dataset that satisfies the null hypothesis and compute statistic (e.g. shuffle group labels for a two-group comparison); repeat many times and use permutation statistics as your sampling distribution rather than a t, Normal, F, χ^2 , etc
- Downside: often computationally intensive for genomics