

#### Journal club

Papers to be selected by 18.00 on 18th October; please discuss it with Hubert and I before submitting PR.

Start: Oct 25

Journal Club schedule to be finalized by 25th October

Given the number of students, groups of 2 are recommended.

Use the #journal-clubs channel (e.g., to find a group member). I will put some suggestions there.

Sign up by pull request to the 'material' repo. "First come first served"

18.10.2021	Mark	limma + friends	linear model simulation + design matrices		
25.10.2021	Hubert	RNA-seq quantification	RSEM	х	х
01.11.2021	Mark	edgeR+friends 1	basic edgeR/voom	OUTRIDER: A Statistical Method for Detecting Aberrantly Expressed Genes in RNA Sequencing Data (BT, KN)	Powerful and robust non-parametric association testing for microbiome data via a zero-inflated quantile approach (ZINQ) (RM, DS)
08.11.2021	Mark	edgeR+friends 2	advanced edgeR/voom	ZeitZeiger: supervised learning for high- dimensional data from an oscillatory system (TB, OF)	SnapHiC:a computational pipeline to identify chromatin loops from single-cell Hi- C data (JS NH)
15.11.2021	ТВА	hands-on session #1: RNA-seq	FASTQC/Salmon/etc.	Differential abundance testing on single-cell data using k-nearest neighbor graphs (VW,	TedSim: temporal dynamics simulation of single cell RNA- sequencing data and cell



### **Journal Club procedure**

- During/after journal clubs: give the presenters some constructive feedback
- Giving feedback (via Google form) is part of your JC grade! Feedback forms must be submitted within 1 week of presentation; comments will be sent to presenters (anonymously)
- Note that they will be part of the recordings (<u>tube.switch.ch</u> videos only shared with registered students)

# Feedback form: 14.10. Redefining CpG islands using hidden Markov models

Presenters:				
* Required				
How would you rate the presenters' coverage of the topic? *				
O Poor				
O Fair				
Good				
O Very Good				
O Excellent				
How would you rate the presenters' knowledge of the topic? *				
O Poor				
O Fair				
Good				
O Very Good				
Excellent				

## From the feed: "Over-optimism" + Terry's IMS Bulletin

We will see a lot of methods in this course - how do we evaluate what works well in

practice?

BIOINFORMATICS

ORIGINAL PAPER

Vol. 26 no. 16 2010, pages 1990–1998 doi:10.1093/bioinformatics/btq323

Gene expression

Advance Access publication June 26, 2010

#### Over-optimism in bioinformatics: an illustration

Monika Jelizarow<sup>1</sup>, Vincent Guillemot<sup>1,2</sup>, Arthur Tenenhaus<sup>2</sup>, Korbinian Strimmer<sup>3</sup> and Anne-Laure Boulesteix<sup>1,\*</sup>

<sup>1</sup>Department of Medical Informatics, Biometry and Epidemiology, University of Munich, Marchioninistr. 15, 81377 Munich, Germany, <sup>2</sup>SUPELEC Sciences des Systèmes (E3S)-Department of Signal Processing and Electronics Systems - 3, rue Joliot Curie, Plateau de Moulon, 91192 Gif-sur-Yvette Cedex, France and <sup>3</sup>Department of Medical Informatics, Statistics and Epidemiology, University of Leipzig, Härtelstr. 16-18, 04107 Leipzig, Germany Associate Editor: John Quackenbush

"if the improvement of a quantitative criterion such as the error rate is the main contribution of a paper, the superiority of new algorithms should always be demonstrated on independent validation data."



#### REVIEW Open Access

## Essential guidelines for computational method benchmarking



Lukas M. Weber<sup>1,2</sup>, Wouter Saelens<sup>3,4</sup>, Robrecht Cannoodt<sup>3,4</sup>, Charlotte Soneson<sup>1,2,8</sup>, Alexander Hapfelmeier<sup>5</sup>, Paul P. Gardner<sup>6</sup>, Anne-Laure Boulesteix<sup>7</sup>, Yvan Saeys<sup>3,4\*</sup> and Mark D. Robinson<sup>1,2\*</sup>



#### In class discussion

- 5 minutes: read (the excerpt from "Terence's Stuff").
- 10 minutes: Discuss with your neighbour/row and answer the following 4 questions:
  - 1. How do we tell what works in practice?
  - 2. What problems arise using simulated (synthetic) data?
  - 3. What problems arise using real data?
  - 4. What are positive/negative controls?
- n.b. include this (method comparison) context in your Journal Club talks

## limma fundamentals

Mark D. Robinson 5



### The simplistic view: Differential expression, small sample inference

- Table of data (e.g., microarray gene expression data with replicates of each of condition A, condition B)
  - rows = features (e.g., genes), columns = experimental units (samples)
- Most common problem in statistical bioinformatics: want to infer whether there is a change in the response
   a statistical test for each row of the table.

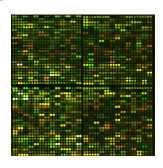
What test might you use? Why is this hard? What issues arise? How much statistical power is there [1]?

```
> head(y)
         group0
                    group0
                            group0
                                           group1
                                                      groupl
                                                                  groupl
genel -0.1874854 0.2584037 -0.05550717 -0.4617966 -0.3563024 -0.03271432
gene2 -3.5418798 -2.4540999
                            0.11750996 - 4.3270442 - 5.3462622 - 5.54049106
gene3 -0.1226303 0.9354707 -1.10537767 -0.1037990 0.5221678 -1.72360854
gene4 -2.3394536 -0.3495697 -3.47742610 -3.2287093 6.1376670 -2.23871974
gene5 -3.7978820 1.4545702 -7.14796503 -4.0500796 4.7235714 10.00033769
gene6
      1.4627078 - 0.3096070 - 0.26230124 - 0.7903434
                                                   0.8398769 - 0.96822312
```

#### **Institute of Molecular Life Sciences**

#### Microarray expression measures array

Two-colour



$$y_{ga} = log_2(R/G)$$
probe or gene

Affymetrix



Illumina



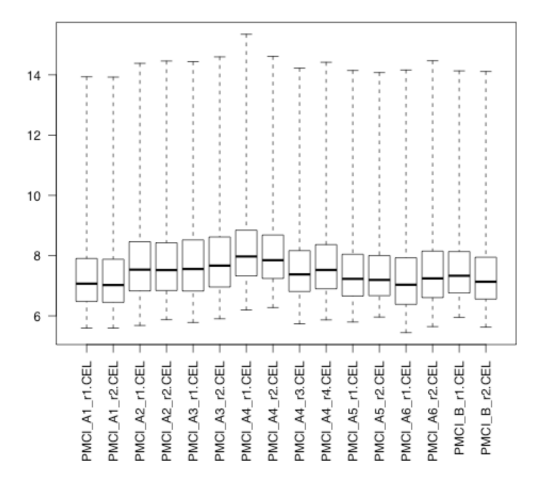


#### Normalization: one-colour



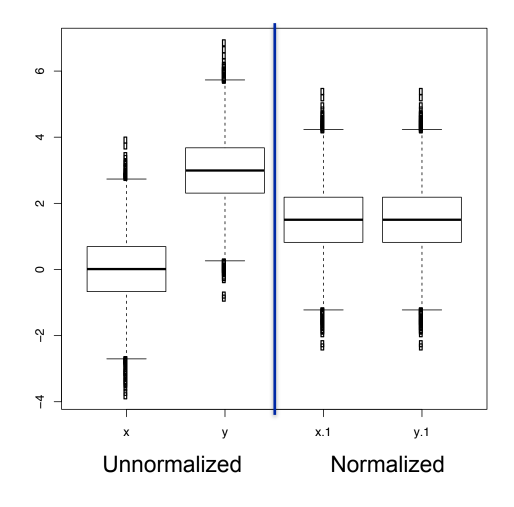


Similarly for single channel data, adjustments need to be made for all samples to be comparable.





#### Quantile normalization



```
x <- rnorm(10000, mean=0, sd=1)
y <- rnorm(10000, mean=3)
z <- cbind(x,y)</pre>
# create "reference" distribution
s <- apply(z,2,sort)</pre>
sm <- rowMeans(s)</pre>
# impose ref. distribution by ranks
r \leftarrow apply(z,2,rank)
n <- apply(r,2,function(u) sm[u])</pre>
boxplot( data.frame(x=x,y=y,n) )
#> library(limma)
#> zn <- normalizeQuantiles(z)</pre>
#> all(zn==n)
#[1] TRUE
```



#### Preprocessing: additive + multiplicative error model

Observe intensity for one probe on one array

$$I = B + S$$
additive additive errors

This idea underlies variance stabilizing transformations vsn (two colour data) and vst (for Illumina data)



## normexp convolution model

Intensity = Background + Signal

 $N(\mu,\sigma^2)$ 

Exponential( $\alpha$ )

#### Microarray background correction: maximum likelihood estimation for the normal-exponential convolution

#### JEREMY D. SILVER

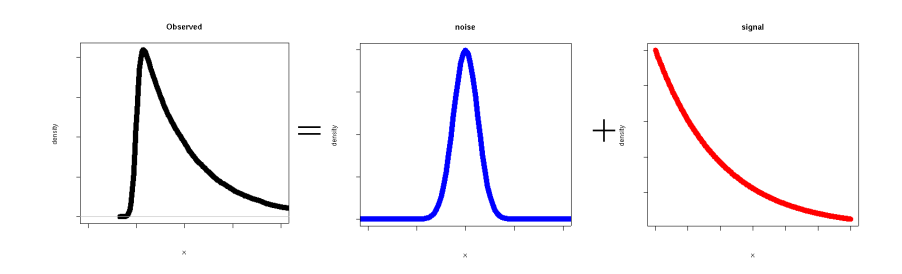
Bioinformatics Division, Walter and Eliza Hall Institute, Parkville 3050, Victoria, Australia and Department of Biostatistics, University of Copenhagen, Øster Farimagsgade 5, Entrance B, PO Box 2099, DK-1014 Copenhagen K, Denmark j.silver@biostat.ku.dk

#### MATTHEW E. RITCHIE

Department of Oncology, University of Cambridge, Cambridge CB2 0RE, UK

#### GORDON K. SMYTH\*

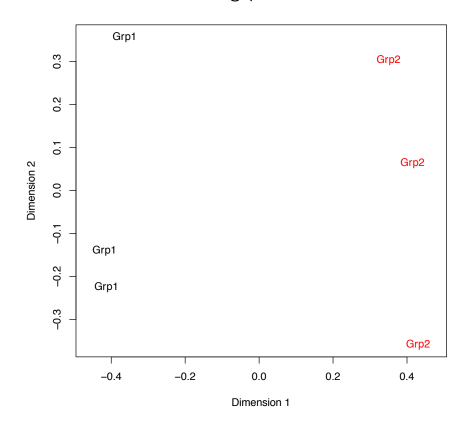
Bioinformatics Division, Walter and Eliza Hall Institute, Parkville 3050, Victoria, Australia smyth@wehi.edu.au





#### Quality assessments / spot checks

#### Multidimensional scaling plot



```
sd <- 0.3*sqrt(4/rchisq(1000,df=4))
x \leftarrow matrix(rnorm(1000*6, sd=sd), 1000, 6)
x[1:50,4:6] \leftarrow x[1:50,4:6] + 2
mds <- plotMDS(x)</pre>
> round(mds$distance.matrix,3)
           [,2] [,3] [,4] [,5] [,6]
[1,] 0.000 0.000 0.000 0.000 0.00
[2,] 0.835 0.000 0.000 0.000 0.00
[3,] 0.850 0.793 0.000 0.000 0.00
[4,] 1.089 1.068 1.058 0.000 0.00
[5,] 1.050 1.058 1.072 0.863 0.00
[6,] 0.991 1.047 1.046 0.865 0.85
                                       0
```

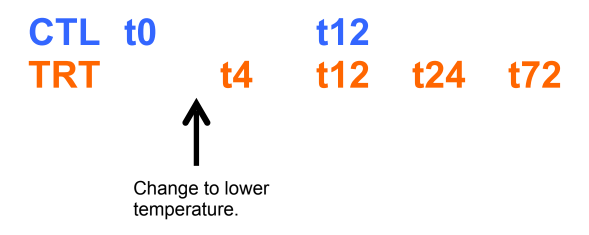
"To consult the statistician after an experiment is finished is often merely to ask him[her] to conduct a post mortem examination. He[She] can perhaps say what the experiment died of." R. A. Fisher

## Motivation for exploratory data analysis: Case Study

(from Stefano, a former M.Sc. student in my Institute)

He is studying gene expression in fruitfly and is interested in transcriptional responses following "heat shock".

#### Basic schematic of experiment:

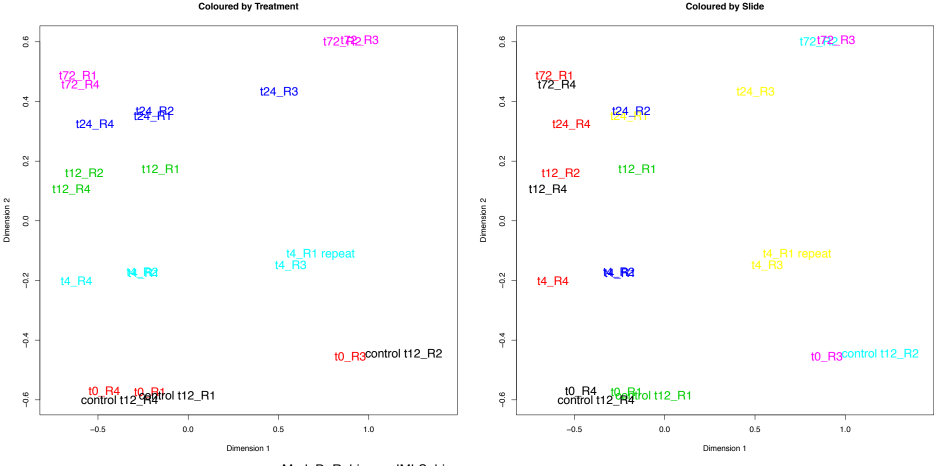


~4 replicates for each condition



library(limma)
plotMDS(d) # 'd' is a matrix

## Take a close look at where the replicates are to each other relative to the X-and Y-axes



22 samples x ~20,000 genes

reduced to 22 samples x 2 dimensions

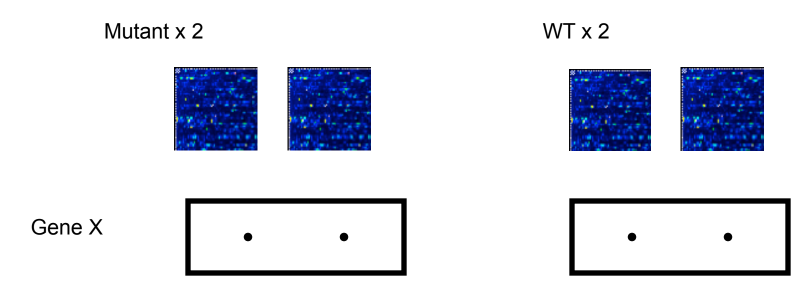


## Limma concept: borrowing information across genes

- Small data sets: few samples, generally under-powered for 1 gene
- Curse of dimensionality: many tests, need to adjust for multiple testing (= loss of power)
- Benefit of parallelism: same model is fit for every gene. Can borrow information from one gene to another
  - Hard: assume parameters are constant across genes
  - Soft: smooth genewise parameters towards a common value in a graduated way, e.g., Bayes, empirical Bayes, Stein shrinkage ...



## A very common experiment (1-colour)



Which genes are differentially expressed?

$$n_1 = n_2 = 2$$
 Affymetrix arrays  
~30,000 probe-sets



#### Ordinary t-tests (1-colour)

$$t_{\!\scriptscriptstyle g} = rac{\overline{y}_{
m mu} - \overline{y}_{
m wt}}{s_{\!\scriptscriptstyle g}\,c}$$

give very high false discovery rates

$$c = \sqrt{\frac{1}{n_1} + \frac{1}{n_2}} \qquad \qquad \text{Residual df = 2}$$



#### t-tests with common variance

$$t_{g, \mathrm{pooled}} = rac{\overline{y}_{\mathrm{mu}} - \overline{y}_{\mathrm{wt}}}{s_{\mathrm{0}} \, c}$$

with residual standard deviation across genes

S<sub>0</sub> pooled

More stable, but ignores gene-specific variability

$$c = \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}$$



### A better compromise

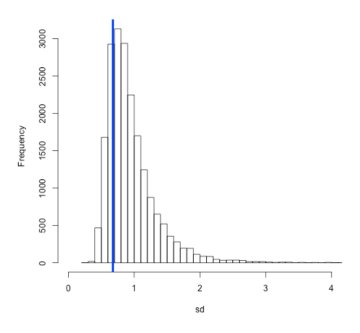
Shrink standard deviations towards common value

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

Moderated t-statistics

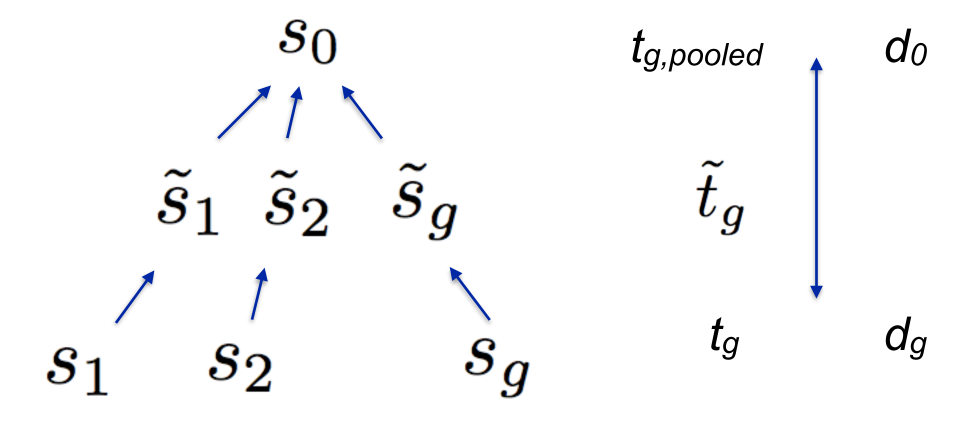
$$ilde{t}_{\!\scriptscriptstyle g} = rac{\overline{y}_{\!\scriptscriptstyle \mathrm{mu}} - \overline{y}_{\!\scriptscriptstyle \mathrm{wt}}}{ ilde{s}_{\!\scriptscriptstyle g} \, u}$$

d = degrees of freedom





## **Shrinkage** of standard deviations



The **data decides** whether  $ilde{t}_g$  should be closer to  $t_{g,pooled}$  or  $t_g$ 



### Why does it work?

- We learn what is the typical variability level by looking at all genes, but allow some flexibility from this for individual genes
- Adaptive data (through hyperparameter estimates, d<sub>0</sub> and s<sub>0</sub>) suggests how much to "squeeze"/"moderate" toward common value



#### Hierarchical model for variances

Data	$s_g^2 \sim \sigma_g^2 rac{\chi_{d_g}^2}{d_g}$
Prior	$rac{1}{\sigma_g^2} \sim s_0^2 rac{\chi_{d_0}^2}{d_0}$
Posterior	$E\left(\frac{1}{\sigma_g^2} \mid s_g^2\right) = \frac{d_0 + d_g}{s_0^2 d_0 + s_g^2 d_g}$



#### **Posterior Statistics**

Posterior variance estimators

$$\tilde{s}_{g}^{2} = \frac{s_{0}^{2}d_{0} + s_{g}^{2}d_{g}}{d_{0} + d_{g}}$$

Moderated t-statistics

$$ilde{t}_{\!\scriptscriptstyle gj} = rac{\hat{eta}_{\!\scriptscriptstyle gj}}{ ilde{s}_{\!\scriptscriptstyle g} \sqrt{c_{\!\scriptscriptstyle gj}}}$$

Baldi & Long 2001, Wright & Simon 2003, Smyth 2004



#### Exact distribution for moderated t

An unexpected piece of mathematics shows that, under the null hypothesis,

$$ilde{t}_g \sim t_{d_0+d_g}$$

#### The degrees of freedom add!

The Bayes prior in effect adds d<sub>0</sub> extra arrays for estimating the variance.

Wright and Simon 2003, Smyth 2004



### Aside: Marginal Distributions to calculate

Under usual likelihood model, s<sub>g</sub> is independent of the estimated coefficients.

Under the hierarchical model, s<sub>g</sub> is independent of the moderated t-statistics instead

$$s_g^2 \sim s_0^2 F_{d,d_0}$$



## Multiple testing and adjusted p-values

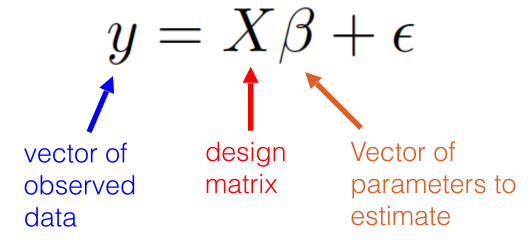
- Each statistical test has an associated false positive rate
- Traditional method in statistics is to control family wise error rate, e.g., by Bonferroni.
- Controlling the false discovery rate (FDR) is more appropriate in microarray studies
- Benjamini and Hochberg method controls expected FDR for independent or weakly dependent test statistics. Simulation studies support use for genomic data.
- All methods can be implemented in terms of adjusted p-values.



#### **Linear Models**

- In general, need to specify:
  - Dependent variable
  - Explanatory variables (experimental design, covariates, etc.)

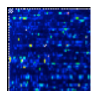
More generally:

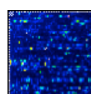


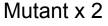


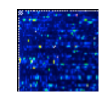
#### Design → Linear models

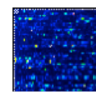
WT x 2











$$\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ 1 & 0 \\ 1 & 1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix} \qquad \beta_1 = \text{wt log-expression}$$
 
$$\beta_2 = \text{mutant} - \text{wt}$$

$$\beta_1$$
 = wt log-expression

$$\beta_2$$
 = mutant – wt

$$\mathsf{E}[\mathsf{y}_1] = \mathsf{E}[\mathsf{y}_2] = \beta$$

$$E[y_1]=E[y_2]=\beta_1$$
  $E[y_3]=E[y_4]=\beta_1+\beta_2$ 



## Layers to add ..

- Where does the moderated variance come from?
- Why the degrees of freedom add: d<sub>0</sub> + d
- empirical Bayes: how to estimate the hyperparameters (d<sub>0</sub> and s<sub>0</sub>)
- Design matrices + contrast matrices in practice



#### The construction of the classical t-statistic:

$$Z = \left(\overline{X}_n - \mu\right) \frac{\sqrt{n}}{\sigma}$$

$$V = (n-1) \frac{S_n^2}{\sigma^2}$$

$$T \equiv \frac{Z}{\sqrt{V/\nu}} = \left(\overline{X}_n - \mu\right) \frac{\sqrt{n}}{S_n},$$

#### Bonus Exercise Part a (optional): what are a, b above?

If T is distributed as  $(a/b)^{1/2}Z/U$  where  $Z \sim N(0,1)$  and  $U \sim \chi_{\nu}$ , then T has density function

$$p(t) = \frac{a^{\nu/2}b^{1/2}}{B(1/2, \nu/2)(a+bt^2)^{1/2+\nu/2}}$$

## **Optional exercise: Derive the posterior**

Data

Prior

 $s_g^2 \sim \sigma_g^2 rac{\chi_{d_g}^2}{d_g}$ 

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

$$p(\theta|x) = \frac{f(x|\theta)p(\theta)}{\int f(x|\theta)p(\theta)d\theta}$$

**Posterior** 

$$E\left(\frac{1}{\sigma_{g}^{2}} \mid s_{g}^{2}\right) = \frac{d_{0} + d_{g}}{s_{0}^{2}d_{0} + s_{g}^{2}d_{g}}$$

#### **Bonus exercise Part b**

Sketch: i) Let  $x=s^2$ ,  $\theta=\sigma^{-2}$ ; ii) Using the functional form of chi-squared distribution, calculate only the numerator (since denominator does not contain  $\theta$ ); iii) collect terms and see if you can identify the distribution and the parameters of it; iv) What is the mean of this distribution?



$$p(\hat{\beta}, s^2 \mid \beta = 0) = \int p(\hat{\beta} \mid \sigma^{-2}, \beta = 0) p(s^2 \mid \sigma^{-2}) p(\sigma^{-2}) d(\sigma^{-2})$$

The integrand is

$$\begin{split} &\frac{1}{(2\pi v\sigma^2)^{1/2}}\exp\left(-\frac{\hat{\beta}^2}{2v\sigma^2}\right) \\ &\times \left[\frac{d}{2\sigma^2}\right]^{d/2}\frac{s^{2(d/2-1)}}{\Gamma(d/2)}\exp\left(-\frac{ds^2}{2\sigma^2}\right) \\ &\times \left[\frac{d_0s_0^2}{2}\right]^{d_0/2}\frac{\sigma^{-2(d_0/2-1)}}{\Gamma(d_0/2)}\exp\left(-\sigma^{-2}\frac{d_0s_0^2}{2}\right) \\ &= \frac{(d_0s_0^2/2)^{d_0/2}(d/2)^{d/2}s^{2(d/2-1)}}{(2\pi v)^{1/2}\Gamma(d_0/2)\Gamma(d/2)} \\ &= \sigma^{-2(1/2+d_0/2+d/2-1)}\exp\left\{-\sigma^{-2}\left(\frac{\hat{\beta}^2}{2v} + \frac{ds^2}{2} + \frac{d_0s_0^2}{2}\right)\right\} \end{split}$$



$$p(\hat{\beta}, s^2 \mid \beta = 0) = \int p(\hat{\beta} \mid \sigma^{-2}, \beta = 0) p(s^2 \mid \sigma^{-2}) p(\sigma^{-2}) d(\sigma^{-2})$$

$$= \frac{(d_0 s_0^2/2)^{d_0/2} (d/2)^{d/2} s^{2(d/2-1)}}{(2\pi v)^{1/2} \Gamma(d_0/2) \Gamma(d/2)}$$

$$\sigma^{-2(1/2+d_0/2+d/2-1)} \exp\left\{-\sigma^{-2} \left(\frac{\hat{\beta}^2}{2v} + \frac{ds^2}{2} + \frac{d_0 s_0^2}{2}\right)\right\}$$

1

 $\sigma^{-2}$  is chi-squared (or gamma)

$$f(x; k) = \begin{cases} \frac{x^{(k/2)-1}e^{-x/2}}{2^{k/2}\Gamma(\frac{k}{2})}, & x \ge 0; \\ 0, & \text{otherwise.} \end{cases}$$



$$p(\hat{\beta}, s^2 | \beta = 0) = \int p(\hat{\beta} | \sigma^{-2}, \beta = 0) p(s^2 | \sigma^{-2}) p(\sigma^{-2}) d(\sigma^{-2})$$

$$\begin{split} p(\hat{\beta}, s^2 \mid \beta &= 0) \\ &= \frac{(1/2v)^{1/2} (d_0 s_0^2/2)^{d_0/2} (d/2)^{d/2} s^{2(d/2-1)}}{D(1/2, d_0/2, d/2)} \left( \frac{\hat{\beta}^2/v + d_0 s_0^2 + ds^2}{2} \right)^{-(1+d_0+d)/2} \end{split}$$



$$p(\hat{\beta}, s^2 \mid \beta = 0)$$

$$= \frac{(1/2v)^{1/2} (d_0 s_0^2 / 2)^{d_0 / 2} (d/2)^{d/2} s^{2(d/2-1)}}{D(1/2, d_0 / 2, d/2)} \left(\frac{\hat{\beta}^2 / v + d_0 s_0^2 + ds^2}{2}\right)^{-(1+d_0+d)/2}$$

The null joint distribution of  $\tilde{t}$  and  $s^2$  is

$$p(\tilde{t}, s^2 | \beta = 0) = \tilde{s}v^{1/2}p(\hat{\beta}, s^2 | \beta = 0)$$

http://en.wikipedia.org/wiki/Random\_variable#Distribution\_functions\_of\_random\_variables

$$f_Y(y) = f_X(g^{-1}(y)) \left| \frac{dg^{-1}(y)}{dy} \right|$$



If T is distributed as  $(a/b)^{1/2}Z/U$  where  $Z \sim N(0,1)$  and  $U \sim \chi_{\nu}$ , then T has density function  $p(t) = \frac{a^{\nu/2}b^{1/2}}{B(1/2,\nu/2)(a+bt^2)^{1/2+\nu/2}}$ 

$$p(\tilde{t}, s^2 \mid \beta = 0) = \frac{(d_0 s_0^2)^{d_0/2} d^{d/2} s^{2(d/2-1)}}{B(d/2, d_0/2) (d_0 s_0^2 + d s^2)^{d_0/2 + d/2}} \times \frac{(d_0 + d)^{-1/2}}{B(1/2, d_0/2 + d/2)} \left(1 + \frac{\tilde{t}^2}{d_0 + d}\right)^{-(1+d_0+d)/2}$$

This shows that  $\tilde{t}$  and  $s^2$  are independent with

$$s^2 \sim s_0^2 F_{d,d_0}$$

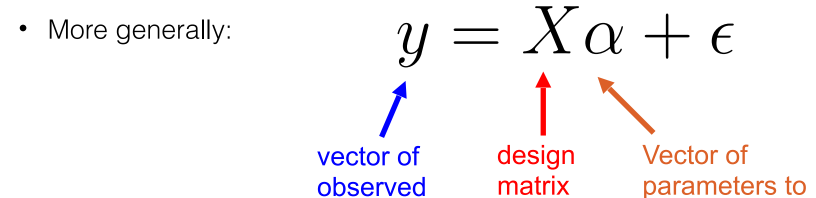
and

$$\tilde{t} \mid \beta = 0 \sim t_{d_0 + d}.$$



#### **Linear Models**

- In general, need to specify:
  - Dependent variable
  - Explanatory variables (experimental design, covariates, etc.)



data

Obtain a linear model for each gene g

$$E(\underline{y}_g) = X\underline{\alpha}_g$$
$$\operatorname{var}(\underline{y}_g) = W_g^{-1}\sigma_g^2$$

estimate



### Contrasts -- contrasts.fit()

A contrast is any linear combination of the coefficients a which we want to test equal to zero.

Define contrasts

$$\beta_g = C^T \alpha_g$$

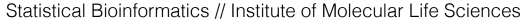
were C is the contrast matrix.

Want to test

$$H_0: \beta_{gj} = 0$$
 
$$H_a: \beta_{gj} \neq 0$$

VS

$$H_a:\beta_{ai}\neq 0$$





#### **Analysis of Variance** → **Linear model**

WT x 2





Cond A x 2





Cond B x 2





$$\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \\ y_6 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 1 & 1 & 0 \\ 1 & 1 & 0 \\ 1 & 0 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} + \begin{bmatrix} \epsilon_1 \\ \epsilon_2 \\ \epsilon_3 \\ \epsilon_4 \\ \epsilon_5 \\ \epsilon_6 \end{bmatrix} \qquad \begin{array}{l} \alpha_1 = \text{ wt log-expression} \\ \alpha_2 = \text{Cond A - wt} \\ \alpha_3 = \text{Cond B - wt} \\ \end{array}$$

$$\alpha_1$$
 = wt log-expression

$$a_2 = Cond A - wt$$

$$a_3 = Cond B - w$$

$$E[y_1]=E[y_2]=\alpha_1$$

$$E[y_3] = E[y_4] = \alpha_1 + \alpha_2$$
  $E[y_5] = E[y_6] = \alpha_1 + \alpha_3$ 

$$E[y_5] = E[y_6] = \alpha_1 + \alpha_3$$





#### **Analysis of Variance** → **Linear model, alternative parameterization**

WT x 2





Cond A x 2





Cond B x 2





$$\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \\ y_6 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} + \begin{bmatrix} \epsilon_1 \\ \epsilon_2 \\ \epsilon_3 \\ \epsilon_4 \\ \epsilon_5 \\ \epsilon_6 \end{bmatrix}$$

$$\alpha_1 = \text{wt log-expression}$$

$$\alpha_2 = \text{Cond A log-expression}$$

$$\alpha_3 = \text{Cond B log-expression}$$

$$\alpha_1$$
 = wt log-expression

$$a_2$$
 = Cond A log-expression

$$a_3$$
 = Cond B log-expression

$$E[y_1] = E[y_2] = a_1$$

$$E[y_1]=E[y_2]=\alpha_1$$
  $E[y_3]=E[y_4]=\alpha_2$   $E[y_5]=E[y_6]=\alpha_3$ 

$$E[y_5] = E[y_6] = a_3$$



#### An example use of design and contrast matrices

design matrix

$$\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \\ y_6 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} + \begin{bmatrix} \epsilon_1 \\ \epsilon_2 \\ \epsilon_3 \\ \epsilon_4 \\ \epsilon_5 \\ \epsilon_6 \end{bmatrix}$$

$$E[y_1] = E[y_2] = \alpha_1$$

$$E[y_3] = E[y_4] = \alpha_2$$

$$E[y_5] = E[y_6] = \alpha_3$$

$$E[y_1] = E[y_2] = \alpha_1$$
  
 $E[y_3] = E[y_4] = \alpha_2$   
 $E[y_5] = E[y_6] = \alpha_3$ 

$$\beta = C\alpha = \begin{bmatrix} -1 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} \alpha_2 - \alpha_1 \\ \alpha_3 - \alpha_2 \end{bmatrix}$$



## Contrasts -- contrasts.fit()

A contrast is any linear combination of the coefficients  $\alpha_i$  that we want to test equal to zero.

Define contrasts

$$\beta_g = C^T \alpha_g$$

were C is the contrast matrix.

Want to test

$$H_0: \beta_{gj} = 0$$
 
$$H_a: \beta_{gj} \neq 0$$

VS

$$H_a:\beta_{ai}\neq 0$$



#### **Limma / Analysis of Variance**

$$F = \frac{\text{variance between treatments}}{\text{variance within treatments}}$$

$$F = \frac{MS_{\text{Treatments}}}{MS_{\text{Error}}} = \frac{SS_{\text{Treatments}}/(I-1)}{SS_{\text{Error}}/(n_T - I)}$$

The moderated t-statistics also lead naturally to moderated F-statistics which can be used to test hypotheses about any set of contrasts simultaneously. Appropriate quadratic forms of moderated t-statistics follow F-distributions just as do quadratic forms of ordinary t-statistics. Suppose that we wish to test all contrasts for a given gene equal to zero, i.e.,  $H_0: \beta_g = 0$ . The correlation matrix of  $\hat{\beta}_g$  is  $R_g = U_g^{-1}C^TV_gCU_g^{-1}$ where  $U_g$  is the diagonal matrix with unscaled standard deviations  $(v_{gj})^{1/2}$  on the diagonal. Let r be the column rank of C. Let  $Q_g$  be such that  $Q_g^TR_gQ_g = I_r$  and let  $\mathbf{q}_g = Q_g^T\mathbf{t}_g$ . Then

$$F_g = \mathbf{q}_g^T \mathbf{q}_g / r = \mathbf{t}_g^T Q_g Q_g^T \mathbf{t}_g / r \sim F_{r,d_0 + d_g}$$



#### **Aside: Marginal Distributions to calculate**

Fun fact: Under usual likelihood model, s<sub>g</sub> is independent of the estimated coefficients.

Under the hierarchical model, s<sub>g</sub> is independent of the moderated t-statistics instead

$$s_g^2 \sim s_0^2 F_{d,d_0}$$

Thus, the set of  $s_g$  can be used to estimated  $d_0$  and  $s_0$ 

Section 6.2 limma paper: other tricks, such as Fisher's z distribution to estimate d<sub>0</sub> and s<sub>0</sub>



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#### Relate to limma objects

$$\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \\ y_6 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} + \begin{bmatrix} \epsilon_1 \\ \epsilon_2 \\ \epsilon_3 \\ \epsilon_4 \\ \epsilon_5 \\ \epsilon_6 \end{bmatrix}$$

$$E[y_1]=E[y_2]=\alpha_1$$
  
 $E[y_3]=E[y_4]=\alpha_2$   
 $E[y_5]=E[y_6]=\alpha_3$ 

$$\beta = C\alpha = \begin{bmatrix} -1 & 1 & 0 \\ 0 & -1 & 1 \end{bmatrix} \begin{bmatrix} \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} \alpha_2 - \alpha_1 \\ \alpha_3 - \alpha_2 \end{bmatrix}$$

$$\begin{bmatrix} 1, ] & -0.07 & 2.03 & -0.16 \\ [2, ] & -4.73 & -5.75 & 2.67 \\ [3, ] & -16.04 & 8.85 & -13.74 \end{bmatrix}$$

```
> design
  alpha1 alpha2 alpha3
> cont.matrix <- makeContrasts(beta1="alpha2-alpha1",</pre>
                beta2="alpha3-alpha2",levels=design)
> cont.matrix
        Contrasts
Levels
        beta1 beta2
 alpha1
 alpha2
                 -1
 alpha3
                  1
fit <- lmFit(y,design)</pre>
fit.c <- contrasts.fit(fit, cont.matrix)</pre>
fit.c <- eBayes(fit.c)</pre>
> head(round(y,2),3)
           [,2] [,3] [,4]
                                       [,6]
[1,] -1.62 1.49 2.50 1.57 -0.71
                                       0.38
[2,] -4.50 -4.95 -3.66 -7.83 -1.59
                                       6.94
[3,] -10.17 -21.90 14.03 3.66 -12.21 -15.26
> head(round(fit$coef,2),3)
     alpha1 alpha2 alpha3
> head(round(fit.c$coef,2),3)
     Contrasts
       beta1 beta2
  [2,] -1.02
              8.42
  [3,] 24.89 -22.59
```



#### **Bonus exercise**

- Alongside Exercise 5 (if you choose to do it, submit together with Exercise 5)
- Optional!
- Counts up to 3 points towards your Exercise mark