

Journal club

Papers to be selected by 18.00 on 13th October; discuss it with Hubert and I.

Start: Oct 12 or Oct 19

Journal Club schedule to be finalized by 19th October

Given the number of students, groups of 3 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 (give a link to the paper, give initials of group members)

09.11.2020	Kathi	hands-on session #1: RNA-seq	FASTQC/Salmon/etc.	Statistical significance for genomewide studies (RA, HH)	Identifying signaling genes in spatial single cell expression data (YM, KD, GJ)
16.11.2020	Hubert	single-cell 1: preprocessing, dim. reduction, clustering		Testing hypotheses about the microbiome using the linear decomposition model (FS,BO,SA)	Generalizing RNA velocity to transient cell states through dynamical modeling (DP, EH)
23.11.2020	Helena	hands-on session #2: cytometry	cytof null comparison	Genome-wide detection of intervals of genetic heterogeneity associated with complex traits (Richard Affolter, Philip Hartout, Martin Emons)	Empirical Bayes Analysis of a Microarray Experiment (Jennifer Probst, Eljas Röllin, Lisa Herzog)
30.11.2020	Mark	single-cell 2: cell type definition, differential state	scRNA exercise 2	A Bayesian mixture model for the analysis of allelic expression in single cells (Sneha-Sundar,SmaragdaDimitrakopoulou,marinapanteli)	scMET: Bayesian modelling of DNA methylation heterogeneity at single-cell resolution (MW, RM, DW)
07.12.2020	Pierre- Luc	hands-on session #3: single-cell RNA- seq	full scRNA-seq pipeline	ScreenBEAM: a novel meta-analysis algorithm for functional genomics screens via Bayesian hierarchical modeling (Leonor Schubert, Jonathan Haab, Flavio Rump)	Detection of differentially abundant cell subpopulations discriminates biological states in scRNA-seq data (TE, RB, AB)
14.12.2020	Mark	loose ends: HMM, EM, robustness	segmentation, peak finding	NEBULA: a fast negative binomial mixed model for differential expression and co-expression analyses of large-scale multi-subject single-cell data (HML, SCD, SW)	х

- 1



Journal Club procedure

- During/after journal clubs: give the presenters some constructive feedback
- New this year: giving feedback (via Google form) is part of your JC grade!
- Note that they will be recorded this year (<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					
O 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/btg323

Gene expression

Advance Access publication June 26, 2010

Over-optimism in bioinformatics: an illustration

Monika Jelizarow¹, Vincent Guillemot^{1,2}, Arthur Tenenhaus², Korbinian Strimmer³ and Anne-Laure Boulesteix^{1,*}

¹Department of Medical Informatics, Biometry and Epidemiology, University of Munich, Marchioninistr. 15, 81377 Munich, Germany, ²SUPELEC 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 ³Department 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

Essential guidelines for computational method benchmarking



Open Access

Lukas M. Weber^{1,2}, Wouter Saelens^{3,4}, Robrecht Cannoodt^{3,4}, Charlotte Soneson^{1,2,8}, Alexander Hapfelmeier⁵, Paul P. Gardner⁶, Anne-Laure Boulesteix⁷, Yvan Saeys^{3,4*} and Mark D. Robinson^{1,2*}



Breakout room discussion

- 10 minutes in breakout room; read (the excerpt from "Terence's Stuff"), then discuss with your group
- 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?
- Organize some notes to these 4 questions in a Google doc (or similar); These will be shared in "plenary"
- 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.

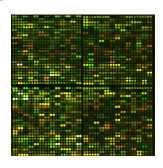
What test might you use? Why is this hard? What issues arise? How much statistical power is there [1]?

```
> 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



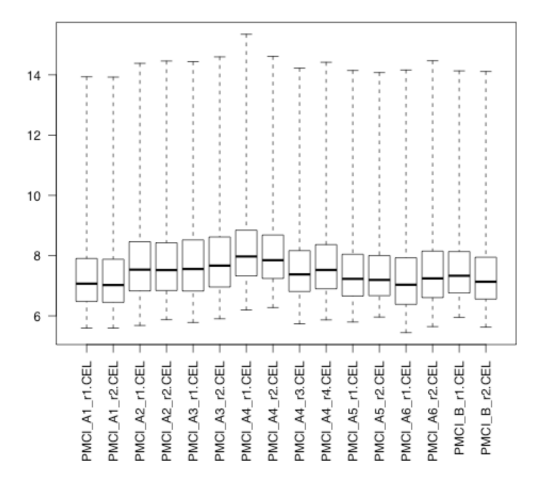


Normalization: one-colour



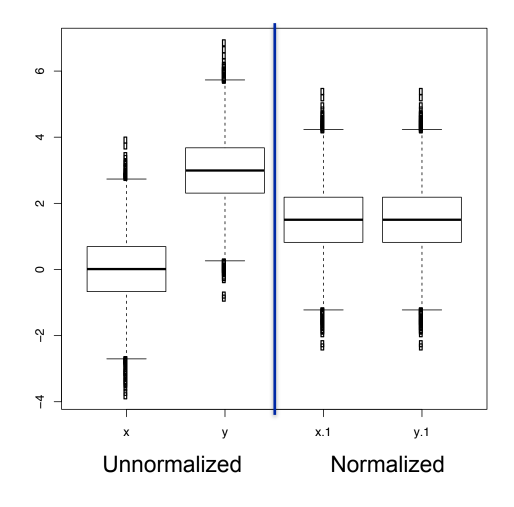


Similarly for single channel data, adjustments need to be made for all samples to be comparable.





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(α)

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

JEREMY D. SILVER

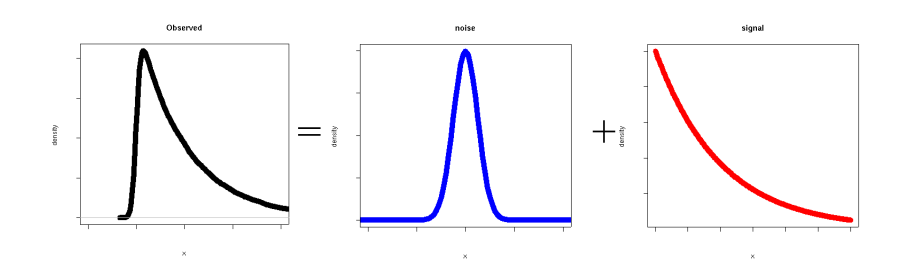
Bioinformatics Division, Walter and Eliza Hall Institute, Parkville 3050, Victoria, Australia and Department of Biostatistics, University of Copenhagen, Øster Farimagsgade 5, Entrance B, PO Box 2099, DK-1014 Copenhagen K, Denmark j.silver@biostat.ku.dk

MATTHEW E. RITCHIE

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GORDON K. SMYTH*

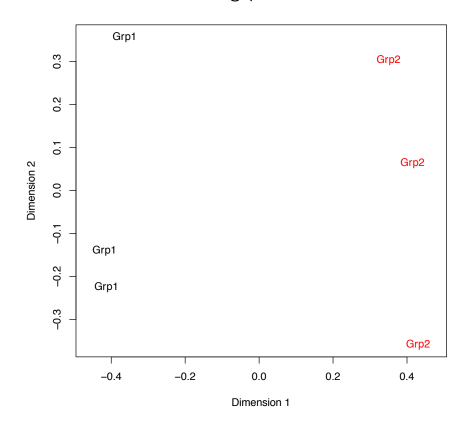
Bioinformatics Division, Walter and Eliza Hall Institute, Parkville 3050, Victoria, Australia smyth@wehi.edu.au





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
```

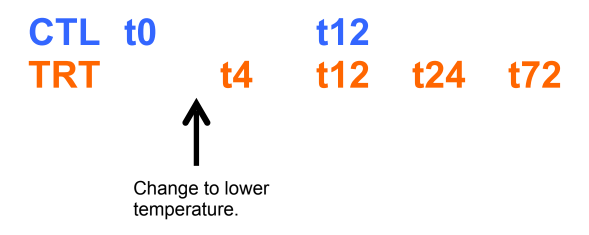
"To consult the statistician after an experiment is finished is often merely to ask him[her] to conduct a post mortem examination. He[She] can perhaps say what the experiment died of." R. A. Fisher

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:

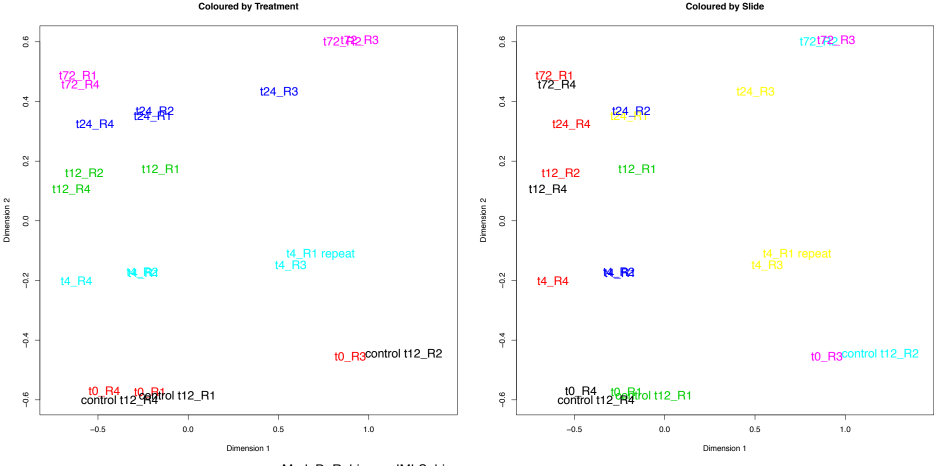


~4 replicates for each condition



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

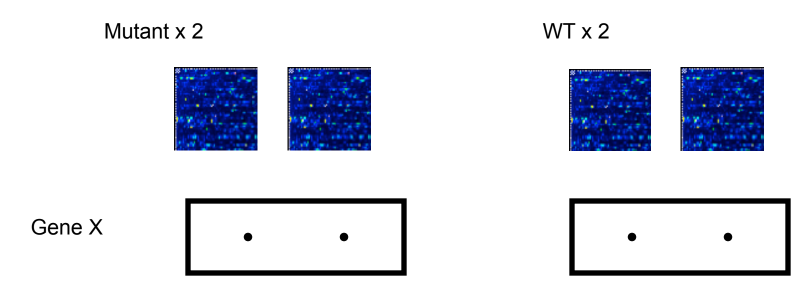


Limma concept: borrowing information across genes

- 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

S₀ pooled

More stable, but ignores gene-specific variability

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



A better compromise

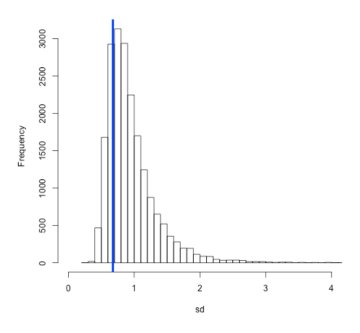
Shrink standard deviations towards common value

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

Moderated t-statistics

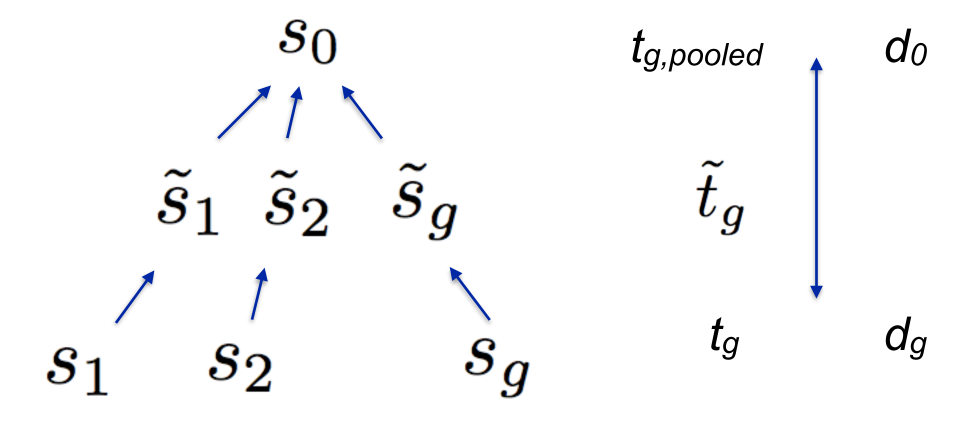
$$ilde{t}_{\!\scriptscriptstyle g} = rac{\overline{y}_{\!\scriptscriptstyle \mathrm{mu}} - \overline{y}_{\!\scriptscriptstyle \mathrm{wt}}}{ ilde{s}_{\!\scriptscriptstyle g} \, u}$$

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, d₀ and s₀) 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 d₀ extra arrays for estimating the variance.

Wright and Simon 2003, Smyth 2004



Aside: Marginal Distributions to calculate

Under usual likelihood model, s_g is independent of the estimated coefficients.

Under the hierarchical model, s_g 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

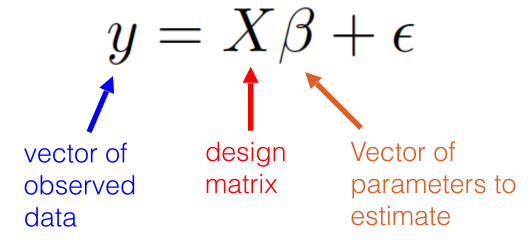
- Each statistical test has an associated false positive rate
- Traditional method in statistics is to control family wise error rate, e.g., by Bonferroni.
- Controlling the false discovery rate (FDR) is more appropriate in microarray studies
- Benjamini and Hochberg method controls expected FDR for 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.)

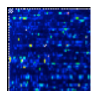
More generally:

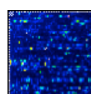


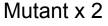


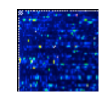
Design → Linear models

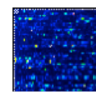
WT x 2











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

$$\beta_2 = \text{mutant} - \text{wt}$$

$$\beta_1$$
 = 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: d₀ + d
- empirical Bayes: how to estimate the hyperparameters (d₀ and s₀)
- Design matrices + contrast matrices in practice



The construction of the classical t-statistic:

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

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

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

Stated another way \rightarrow Exercise (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

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

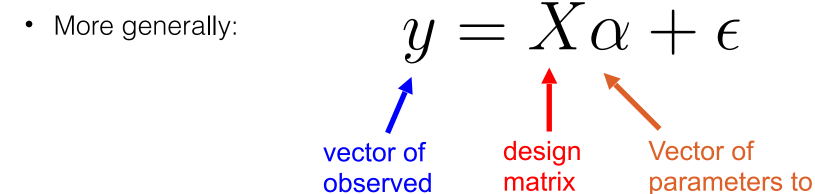
Bonus exercise (= one exercise, i.e., 3 points towards 30 for exercises; submit PDF/scan with Exercise 5)

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 θ); iii) collect terms and see if you can identify the distribution and the parameters of it; iv) What is the mean of this distribution?



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

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



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

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

 σ^{-2} is chi-squared (or gamma)

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



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

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



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

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

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

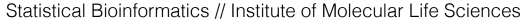
$$p(\tilde{t}, s^2 \mid \beta = 0) = \frac{(d_0 s_0^2)^{d_0/2} d^{d/2} s^{2(d/2 - 1)}}{B(d/2, d_0/2) (d_0 s_0^2 + d s^2)^{d_0/2 + d/2}} \times \frac{(d_0 + d)^{-1/2}}{B(1/2, d_0/2 + d/2)} \left(1 + \frac{\tilde{t}^2}{d_0 + d}\right)^{-(1 + d_0 + d)/2}$$

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

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

and

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





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 α_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_g is independent of the estimated coefficients.

Under the hierarchical model, s_g 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 d₀ and s₀



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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
```