



Part II: Statistical Inference

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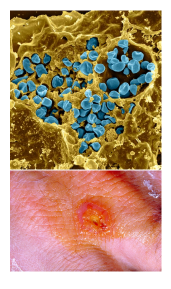
Proteomics Data Analysis Shortcourse

Statistical Inference

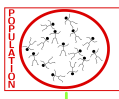
- Francisella tularensis Example
- Hypothesis testing
- Multiple testing
- Moderated statistics
- Experimental design



Francisella tularensis experiment



- Pathogen: causes tularemia
- Metabolic adaptation key for intracellular life cycle of pathogenic microorganisms.
- Upon entry into host cells quick phasomal escape and active multiplication in cytosolic compartment.
- Francisella is auxotroph for several amino acids, including arginine.
- Inactivation of arginine transporter delayed bacterial phagosomal escape and intracellular multiplication.
- Experiment to assess difference in proteome using 3 WT vs 3 ArgP KO mutants



Effect of treatment in population

EXP. DESIGN





Effect of treatment in sample

DATA EXPLORATION



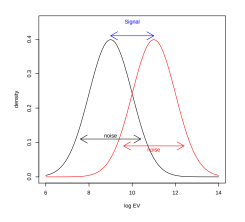
Summarized data structure

- WT vs KO
- 3 vs 3 repeats
- 882 proteins

Protein	WT_1	WT_2	WT ₃	KO ₁	KO ₂	KO ₃
gi 118496616	29.83	29.77	29.91	29.70	29.86	29.80
gi 118496617	31.28	31.23	31.51	31.30	31.51	31.76
gi 118496635	32.39	32.27	32.24	32.25	32.14	32.22
gi 118496636	30.74	30.54	30.64	30.65	30.49	30.60
gi 118496637	29.56	29.35	29.56	29.30	29.24	29.14
gi 118498323	31.38	30.52	30.62	31.04	27.38	NA
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Hypothesis testing: a single protein

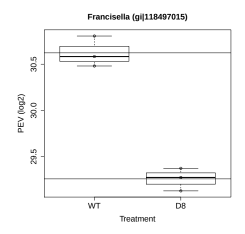


$$\Delta = \bar{z}_{p1} - \bar{z}_{p2}$$
 $T_g = \frac{\Delta}{\sec_{\Delta}}$
 $T_g = \frac{\widehat{\text{signal}}}{\widehat{\text{Noise}}}$

If we can assume equal variance in both treatment groups:

$$\operatorname{se}_{\Delta} = \operatorname{SD}\sqrt{\frac{1}{n_1} + \frac{1}{n_2}}$$

Hypothesis testing: a single protein



$$t = \frac{\log_2 \widehat{\mathsf{FC}}}{\mathsf{se}_{\log_2 \widehat{\mathsf{FC}}}} = \frac{-1.4}{0.118} = -11.9$$

Is t = -11.9 indicating that there is an effect?

How likely is it to observe t=-11.8 when there is no effect of the argP KO on the protein expression?

Null hypothesis and alternative hypothesis

- In general we start from alternative hypothese H_A : we want to show an effect of the KO on a protein
 - On average the protein abundance in WT is different from that in KO



Null hypothesis and alternative hypothesis

- In general we start from alternative hypothese H_A : we want to show an effect of the KO on a protein
 - On average the protein abundance in WT is different from that in KO
- But, we will assess it by falsifying the opposite: null hypothesis H₀
 - On average the protein abundance in WT is equal to that in KO



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Two Sample t-test

data: z by treat

t = -11.449, df = 4, p-value = 0.0003322
alternative hypothesis: true difference in means is not equal to 0

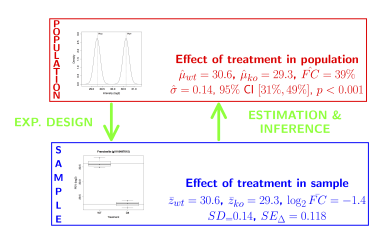
95 percent confidence interval:
-1.031371 -1.691774
sample estimates:
mean in group D8 mean in group WT

29.26094
30.62251
```

- How likely is it to observe an equal or more extreme effect than the one observed in the sample when the null hypothesis is true?
- When we make assumptions about the distribution of our test statistic we can quantify this probability: p-value. The p-value will only be calculated correctly if the underlying assumptions hold!
- When we repeat the experiment, the probability to observe a fold change more extreme than a 2.6 fold ($\log_2 FC = -1.36$) down or up regulation by random change (if H_0 is true) is 3 out of 10.000.
- If the p-value is below a significance threshold α we reject the null hypothesis. We control the probability on a false positive result at the α -level (type I error)



Hypothesis testing: a single protein





Multiple hypothesis testing



Problem of multiple hypothesis testing

- Consider testing DA for all m = 882 proteins simultaneously
- What if we assess each individual test at level α ?
- \rightarrow Probability to have a false positive among all $\it m$ simultatenous test $>>> \ \alpha = 0.05$

Suppose that 600 proteins are non-DA, then we could expect to discover on average $600 \times 0.05 = 30$ false positive proteins. Hence, we are bound to call false positive proteins each time we run the experiment.

FDR: False discovery rate

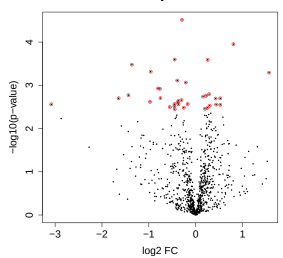
- FDR: Expected proportion of false positives on the total number of positives you return.
- An FDR of 1% means that on average we expect 1% false positive proteins in the list of proteins that are called significant.
- Defined by Benjamini and Hochberg in 1995

$$\begin{split} \mathsf{FDR}(|t_{\mathsf{thres}}|) &= \mathsf{E}\left[\frac{\mathit{FP}}{\mathit{FP} + \mathit{TP}}\right] = \frac{\pi_0 \mathit{Pr}(|\mathit{T}| \geq t_{\mathsf{thres}}|\mathit{H}_0)}{\mathit{Pr}(|\mathit{T}| \geq t_{\mathsf{thres}})} \\ &\quad \mathsf{FDR}_{\mathsf{BH}}(|t_{\mathsf{thres}}|) = \frac{1 \times \mathit{p}_{\mathsf{thres}}}{\#|t_i| \geq t_{\mathsf{thres}}} \end{split}$$

• FDR adjusted p-values can be calculated (e.g. Perseus, R, ...)



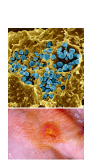
Ordinary t-test



Moderated Statistics



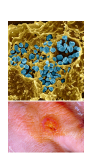
Problems with ordinary t-test



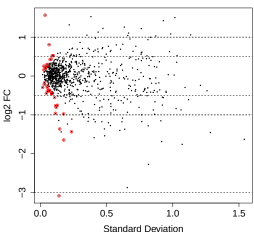
Ordinary t-test -log10(p-value) 0 -1 log2 FC



Problems with ordinary t-test



Original t-test



A moderated *t*-test

A general class of moderated test statistics is given by

$$T_g^{mod} = rac{ar{Y}_{g1} - ar{Y}_{g2}}{c\left(ilde{S}_g
ight)},$$

where \tilde{S}_g is a moderated standard deviation estimate.

- empirical Bayes theory provides formal framework for borrowing strength across genes,
- Implemented in popular bioconductor package limma

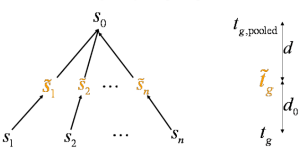
$$\tilde{S}_{g} = \sqrt{\frac{d_{g}S_{g}^{2} + d_{0}S_{0}^{2}}{d_{g} + d_{0}}},$$

- S_0^2 : common variance (over all proteins)
- Moderated t-statistic is t-distributed with $d_0 + d_g$ degrees of freedom.
- → Note that the degrees of freedom increase by borrowing strength across genes!



Shrinkage of the variance and moderated t-statistics

Shrinkage of Standard Deviations

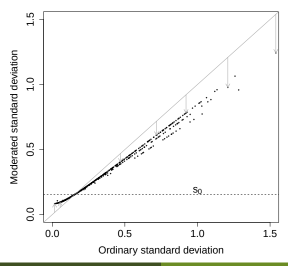


The data decides whether $ilde{t}_{m{g}}$

should be closer to $t_{g,pooled}$ or to t_g

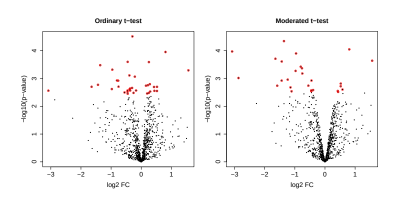


Shrinkage of the variance with limma



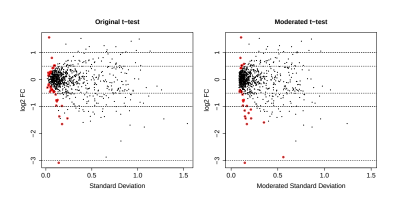


Problems with ordinary t-test solved by moderated EB t-test

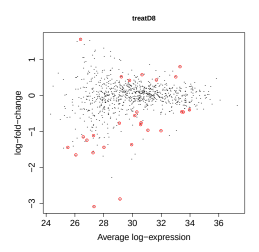




Problems with ordinary t-test solved by moderated EB t-test



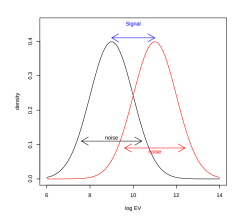




Experimental Design



Power?



$$\Delta = \bar{z}_{p1} - \bar{z}_{p2}$$
 $T_g = \frac{\Delta}{\stackrel{\text{signal}}{\widehat{\text{Noise}}}}$

If we can assume equal variance in both treatment groups:

$$\operatorname{se}_{\Delta} = \operatorname{SD}\sqrt{\frac{1}{n_1} + \frac{1}{n_2}}$$

→ Design: if number of bio-repeats increases we have a higher power!



- Study on tamoxifen treated Estrogen Receptor (ER) positive breast cancer patients
- Proteomes for tumors of patients with good and poor outcome upon recurrence.
- Assess difference in power between 3vs3, 6vs6 and 9vs9 patients.



Experimental Design: Blocking



Sources of variability

$$\sigma^2 = \sigma_{bio}^2 + \sigma_{lab}^2 + \sigma_{extraction}^2 + \sigma_{run}^2 + \dots$$

- Biological: fluctuations in protein level between mice, fluctations in protein level between cells, ...
- Technical: cage effect, lab effect, week effect, plasma extraction, MS-run, ...



Blocking Example: mouse T-cells

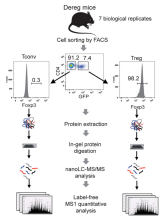
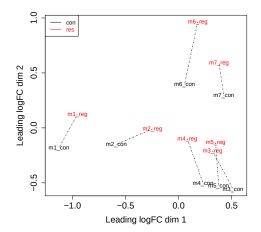


Fig. 1. Label-free quantitative analysis of conventional and regulatory T cell proteomes. General analytical workflow based on cell sorting by flow cytometry using the DEREG mouse model and parallel proteomic analysis of Toonv and Treg cell populations by nanoLC-MS/MS and label-free relative quantification.



Blocking Example: mouse T-cells





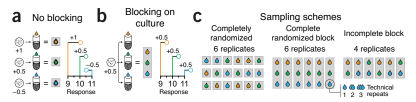
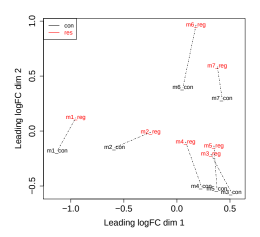


Figure 2 | Blocking improves sensitivity by isolating variation in samples that is independent from treatment effects. (a) Measurements from treatment aliquots derived from different cell cultures are differentially offset (e.g., 1, 0.5, -0.5) because of differences in cultures. (b) When aliquots are derived from the same culture, measurements are uniformly offset (e.g., 0.5). (c) Incorporating blocking in data collection schemes. Repeats within blocks are considered technical replicates. In an incomplete block design, a block cannot accommodate all treatments.

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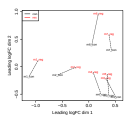
Blocking

$$\sigma^2 = \sigma_{\rm within\ mouse}^2 + \sigma_{\rm between\ mouse}^2$$



Blocking

$$\sigma^2 = \sigma_{\rm within\ mouse}^2 + \sigma_{\rm between\ mouse}^2$$



- → All treatments of interest are present within block!
- → We can estimate the effect of the treatment within block!
- → We can isolate the between block variability from the analysis
- → linear model:

$$y \sim \mathsf{type} + \mathsf{mouse}$$

→ Not possible with Perseus!



Power gain of blocking

- Completely randomized design (CRD): 8 mice, 4 conventional T-cells, 4 regulatory T-cells.
- Randomized complete block desigh (RBC): 4 mice, for each mouse conventional and regulatory T-cells.

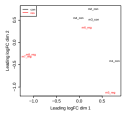


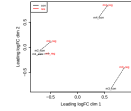
Power gain of blocking CRD

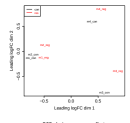


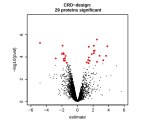
RCB $y \sim \text{type} + \text{mouse}$

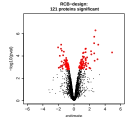
RCB $y \sim \text{type}$

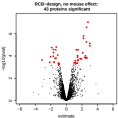






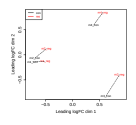








Anova table: P24452, Capg, Macrophage-capping protein



```
### RCB design ###

Df Sum Sq Mean Sq F value Pr(>F)

type 1 15.2282 15.2282 3720.035 9.71e-06 ***
mouse 3 0.2179 0.0726 17.747 0.02058 *

Residuals 3 0.0123 0.0041
```

```
### RCB design: no mouse effect ###

Df Sum Sq Mean Sq F value Pr(>F)

type 1 15.2282 15.2282 396.87 1.038e-06 ***

Residuals 6 0.2302 0.0384

### CRD design ###
```

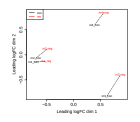
```
### Ord Ussign ###

Df Sum Sq Mean Sq F value Pr(>F)

type 1 11.6350 11.6350 122.86 3.211e-05 ***
Residuals 6 0.5682 0.0947
```

Anova table: P24452, Capg, Macrophage-capping protein

2.75937



```
### RCB design ###
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 22.21485
                       0.05058 439.190 2.60e-08 ***
            2.75937
                       0.04524 60.992 9.71e-06 ***
typereg
            0.30560
                       0.06398 4.776
                                         0.0174 *
mouse2
monse3
           -0.15193
                       0.06398 -2.375
                                         0.0981
            0.07331
                       0.06398
                                 1.146
                                         0.3350
monse4
Residual standard error: 0.06398 on 3 degrees of freedom
### RCB design: no mouse effect ###
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 22.27160
                       0.09794 227.40 4.88e-13 ***
```

0.13851 Residual standard error: 0.1959 on 6 degrees of freedom

19.92 1.04e-06 ***

```
### CRD design ###
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 23.3012
                        0.1557 149.65 6.00e-12 ***
                        0.2251
                               11.08 3.21e-05 ***
typereg
              2.4956
```

Residual standard error: 0.3077 on 6 degrees of freedom



typereg

Comparison residual variance

