

## Statistics for Biologists

Lecture 2: Inferential statistics: Pairwise comparisons, power and multiple testing

> George Bell, Ph.D. Senior Bioinformatics Scientist Bioinformatics and Research Computing Whitehead Institute

# Outline

- Lecture 1 review
- Intro to pairwise comparisons
- The t-test and other tests
- Statistical power
- Intro to multiple hypothesis testing
- Methods to correct for multiple hypothesis testing
- Using the R Commander
- Exercises

# From lecture 1

- Why use statistics?
- Descriptive statistics - central tendency + variability
- Visualization of quantitative data

   What are you trying to show?
- Inferential statistics:  $H_0$ ,  $H_a$ ,  $\alpha$ ,  $\beta$
- False positives and false negatives
- Software for statistics
- Exercises

# Exercise 1 question: boxplot



## Exercise 1 questions: R

• Place multiple figures on the same plot:

par(mfrow=c(2,1))
hist(obese, col="red")
hist (lean, col="blue")
par() # reset

 Save a plot to a file: right click or savePlot(filename="myPlot.pdf", device=2, type="pdf")





• Open another window for a new figure



## Intro to pairwise comparisons

- A common research question is,
  - "Is one measurement equal to another measurement?"
- More specifically, Q: "Is the mean of values in set X equal to the mean in set Y?
  - sample answers: no; yes; X > Y; X:Y = 1.5
- To get an answer that includes some measure of confidence, we need to do a statistical test:
  - T-test (parametric: uses actual values)
  - Wilcoxon Rank Sum (Mann-Whitney) test
     (nonparametric: uses ranks; less powerful: greater β)

### Student's t-test

- History: Gossett ("Student") created this as a new technique to analyze Guinness beer using a small sample size.
- Assumptions:
  - measurements are independent
  - input data are normally distributed
  - Samples are from populations with equal variances (but test can be modified ("Welch's test") when this isn't true)

Fortunately the test is robust, but...

- Two-tailed (testing for ≠) or one-tailed (testing for > or <)
- T statistic =  $\frac{\text{mean}_1 \text{mean}_2}{\text{SE}}$  df = n<sub>1</sub> + n<sub>2</sub> 2

# Performing the t-test

Data sets:  $a = \{ 2 4 6 \}$   $b = \{ 7 8 9 \}$ 

- Excel command: =TTEST(array1, array2, tails, type) example: =TTEST(A1:A3, B1:B3, 2, 3)
- R command: t.test(a, b, alternative="two.sided", var.equal=FALSE)

Output:



### Comparing variances

• Use the F test

– Excel: =FTEST(array1, array2)

- R: var.test(a, b) # a=c(2, 4, 6); b=c(7, 8, 9)
- H<sub>0</sub>: True ratio of variances is 1

F test to compare two variances Prob (H<sub>a</sub> is false) data: a and b F = 4, num df = 2, denom df = 2, p-value = 0.4 alternative hypothesis: true ratio of variances is not equal to 1 95 percent confidence interval: 0.1025641 156.0000000 sample estimates: ratio of variances 4 # To calculate variance: =VAR or 'var' 9

# The paired t-test

- Used when one measurement in each group comes from the same "experimental unit"
- example:
  - mouse assay after treatment by injection
  - one leg is treated; other leg is control
- Method (done by paired test):
  - get difference between each measurement pair
  - Are these differences  $\neq 0$ ?
  - (Do a one-sample t-test)
- Use when experimental design warrants it
- R command: t.test(a, b, paired=TRUE)

Set A	Set B		
100	105		
50	55		
20	23		
250	260		
0.94 vs. 0.03			

### T-test: summary

- Tests for a difference between two means
- Assumes data are normally distributed
- Test comes in multiple flavors:
  - -1 vs. 2 tails
  - equality of variances?
- Use the paired test when appropriate
- Excel just gives the take-home message
   Or see Tools >> Data Analysis

### Moderated t-test

- Generic t statistic:  $t = \frac{x_1 x_2}{s}$
- Moderated t statistic
  - Best known in microarray analysis
  - Corrects gene expression standard deviations toward a pooled estimate
  - Modify all standard deviations used in the t-test
  - $s_0 = 90^{\text{th}}$  percentile of all s (limma R package)
  - $s_0 =$ exchangeability constant (SAM)
  - gains power from sharing variation data across genes

 $s + s_0$ 

### Power and sample size for the t-test

- Power reflects the probability of finding a true difference.
- Determinants of power:  $n \ge \frac{2s_p^2}{\delta^2} (t_{\alpha,\upsilon} + t_{\beta(1),\upsilon})^2$
- Calculating power in R using log<sub>2</sub>-transformed data:
  - How many measurements are needed to be 80% sure that a 2-fold difference  $[\log_2(2)=1]$  can be determined when using  $\alpha=0.05$  on data with a sd of 0.45 (standard deviation of log2-transformed measurements)?

power.t.test(n=NULL, delta=1, sd=0.45, sig.level=0.05, power=0.80, type="two.sample", alternative="two.sided")

### Power and sample size for the t-test

R command: power.t.test(n=NULL, delta=1, sd=0.45, sig.level=0.05, power=0.80, type="two.sample", alternative="two.sided")

• Output:

Two-sample t test power calculation

```
n = 4.3824
delta = 1
sd = 0.45
sig.level = 0.05
power = 0.8
alternative = two.sided
```

#### NOTE: n is number in \*each\* group

# Wilcoxon Rank Sum test

- A non-parametric test:
  - An alternative to a t-test
  - Ranks of data (rather than actual values) are used.
- Also known as the Mann-Whitney test
- No assumptions about normally distributed data are required.
- Data sets  $a = \{ 0.2 \ 0.4 \ 0.6 \}$   $b = \{ 70 \ 80 \ 90 \}$ become  $a = \{ 6 \ 5 \ 4 \}$   $b = \{ 3 \ 2 \ 1 \}$
- R command:

wilcox.test(a, b, alternative="two.sided") => p-value = 0.1

- But this test doesn't have much power:
  - same data in t-test yield a p-value of 0.005

### Intro to multiple hypothesis testing

- A p-value reflects the probability of a false positive call in a statistical test.
- When multiple hypotheses are tested on the same data, the rate of false positives greatly increases

Example for  $\alpha = 0.05$ :

Number of genes tested (n)	Expected number of FPs = $\alpha * n$	Probability of at least one FP = $100(1 - (1 - \alpha)^n)$		
1	0.05	5%		
2	0.1	9.75%		
100	5	99.4%		

• So what to do?

# Choices for multiple hypothesis testing

- How many true positives do you expect?
- How costly is a false positive call?
- How costly is a false negative call?
- Major choices:
  - Family-wise error rate (ex: Bonferroni)
  - False discovery rate (ex: Benjamini and Hochberg)

# Corrections with the FWER

- Family-wise error rate = the probability of at least one false positive in the "family" of positive tests.
- "Bonferroni correction"
- A good choice if you predict that there are very few if any true positive tests
- Large correction means that
  - the false positive rate is very low
  - the false negative rate can be very high
- Practical implication: p-value must be really low to make it past a FWER correction.

### Performing a Bonferroni correction

- Corrected p-value = raw p-value \* n
  - n is the number of tests
  - if corrected p-value > 1, set to 1
- Example:
  - a microarray assays 10,000 genes
  - All are tested for differential expression with the t-test
  - If the raw p-value for one gene =  $10^{-5}$ 
    - The corrected p-value:
      - $= 10^{-5} * 10,000 = 0.1$
    - If  $\alpha = 0.05$ , then this gene cannot be described as differentially expressed.

# Corrections with the FDR

- False discovery rate = the frequency of false positives among the positive tests
- Introduced by Benjamini and Hochberg, 1995
- Tolerates a chosen proportion of false positives
- Much less conservative than Bonferroni:
  - false positive rate is higher
  - false negative rate is lower
- A good choice if you predict that there are many truly positive tests
- Practical implications:
  - more tests stay below  $\alpha$  compared to FWER
  - appropriate correction for many tests of differential expression in microarray experiments

# Performing a FDR correction

- Sort list of p-values in increasing order and starting at the bottom row
- Corrected p-value = the minimum between
  - 1: raw p-value \* (n/rank)
  - 2: corrected p-value below
  - n is the number of tests
  - rank is the position in the sorted list
- Example: a microarray assays 5 genes for differential expression

cion /	$\land$	Gene	Rank	Raw p-value	Formula	Corrected p-value	
order of calculat		С	1	0.001	min (0.001 * (5/1), 0.0125)	0.005	
		А	2	0.005	min (0.005 * (5/2), 0.017)	0.0125	
		В	3	0.01	min (0.01 * (5/3), 0.063)	0.017	
		Е	4	0.05	min (0.05 * (5/4), 0.1)	0.063	
		D	5	0.1	0.1 * (5/5)	0.1	

### Performing multiple hypothesis corrections in R

# Read the data (tab-delimited text, with header fields; p-values in second field)
data.raw.pvals = read.delim("pvals\_raw.txt", h=T)

# Correct the p-values using one of 7 methods ('?p.adjust' to see them) #  $[,2] \rightarrow$  use the data in column 2

```
FDR.p.vals = p.adjust(data.raw.pvals[,2], "fdr")
```

```
# Combine original file with corrected p-value output
data.adjp = cbind(data.raw.pvals, FDR.p.vals)
```

	id	Raw.p	bonferroni	BY	holm	hochberg	BH	fdr
1	Α	0.0400	0.1600	0.1389	0.1200	0.1000	0.0667	0.0667
2	В	0.0010	0.0040	0.0083	0.0040	0.0040	0.0040	0.0040
3	С	0.2000	0.8000	0.4167	0.2000	0.2000	0.2000	0.2000
4	D	0.0500	0.2000	0.1389	0.1200	0.1000	0.0667	0.0667

### Summary: multiple hypothesis corrections

- Beware that with raw  $\alpha = 0.05$ , 5% of tests will be positive just by chance
- Whenever performing >1 statistical test together, corrections should be done
- Select method based on desired FP and FN error rates
- Use R, Excel or BaRC web tool

# The R Commander

- Graphical interface designed to facilitate learning R
- Pull down menus and multiple-choice interface for common statistics and graphics
- Created by John Fox from McMaster University
- Installed with R on barra
- To start => begin R => library(Rcmdr)
- For Macs (but installation may be difficult), Windows, and Unix/Linux
- Other R graphical tools: affylmGUI, limmaGUI



### Summary

- Intro to pairwise comparisons
- The t-test and other pairwise tests
- Intro to multiple hypothesis testing
- Methods to correct for multiple hypothesis testing
- Using the R Commander
- Exercises

### References

- Zar JH. *Biostatistical Analysis*. Prentice Hall, 1998. [or any general biostatistics textbook]
- Dalgaard P. *Introductory Statistics with R*. Springer, 2002.
- Venables W.N. and Ripley B.D. Modern Applied Statistics with S. Springer, 2002.
- Tufte E. *The Visual Display of Quantitative Information*. Graphics Press, 1992.
- Lots of web sites
- R documentation

# Exercise 2 - To do

### Using Excel and R:

- Transform data to get a more normal distribution
- Perform different t-test flavors for several different types of data
- Compute power for some t-tests
- Perform a Wilcoxon rank sum test
- Given a series p-values, perform multiple hypothesis testing
  - Bonferroni
  - FDR

# Helpful R commands



```
# Read a tab-delimited data table
energy = read.delim("energy.txt", header=T)
dat = read.delim(file("clipboard"), h=T)
rats = read.delim("http://.../rats.txt", h=T)
```

**Google mailing lists, etc. => r-project t-test** 

### Exercise 2 functions

- Excel
- TTEST
- RANK
- SORT
- LOG
- R
- t.test
- wilcox.test
- p.adjust
- log
- power.t.test