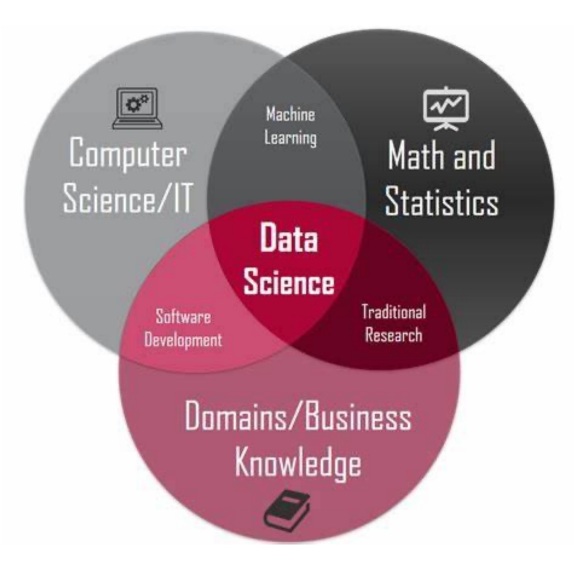
STATISTICS REVIEW

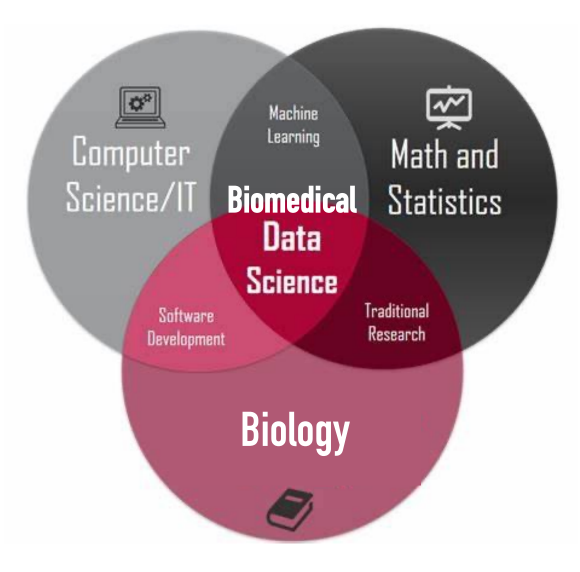
OUTLINE

- Random variable
- Probability distribution
- Central limit theorem
- Hypothesis testing
- P-value
- Multiple testing correction
- Type I and type II errors
- False discovery rate (FDR)

Data Science



Computational Biology



Why statistics is important

- Statistics is the theoretical foundation of machine learning and data science
- Statistics is the bridge between experiments and theories
- Statistics is the bridge between observations (data) and discoveries (science)

What is a model

• A model describes the relationship between quantities

– Quantitative/mathematical models

- Statistical models
 - Relationship between random variables

• "All models are wrong; but some are useful." (George Box)

Random variables

- Randomness
- All quantities obtained from experimental measurements can be considered as random variables
- Discrete vs. continuous random variables
- Probability distribution

Discrete probability distributions

Bernoulli distribution

$$\Pr(X = 1) = p = 1 - \Pr(X = 0) = 1 - q.$$

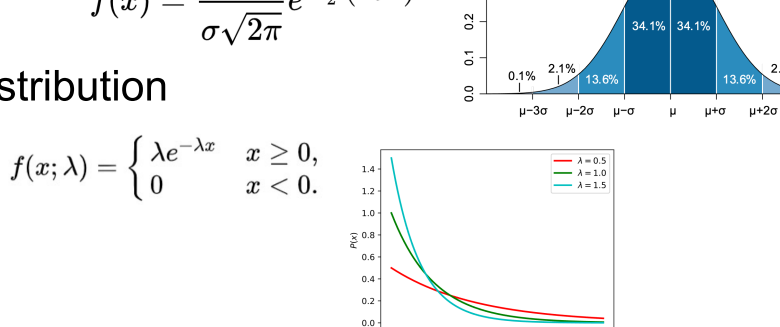
- Binomial distribution $\Pr(X = k) = \binom{n}{k} p^k (1-p)^{n-k}$
- Negative binomial distribution $\Pr(X = k) = \binom{k+r-1}{k} (1-p)^k p^r$
- Poisson distribution $Pr(X=k) = \frac{\lambda^k e^{-\lambda}}{k!}$

Continuous probability distributions

- Uniform distribution
- Normal distribution (Gaussian distribution)

$$f(x)=rac{1}{\sigma\sqrt{2\pi}}e^{-rac{1}{2}\left(rac{x-\mu}{\sigma}
ight)^2}$$





0.4

0.3

f(x)

<u>1</u> *b*− a

0

а

Х

2.1%

0.1%

μ+3σ

Use of probability distributions in computational biology

- Null models for hypothesis testing
- Noise or background models for signal detection and differential enrichment analysis
- Error models in regression/machine learning

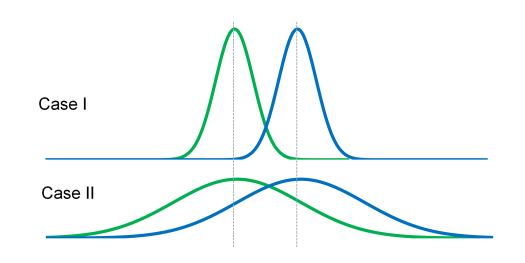
Central limit theorem

 Regardless of the population distribution, the sample mean will follow a standard normal distribution, if the samples are independent and equal size.

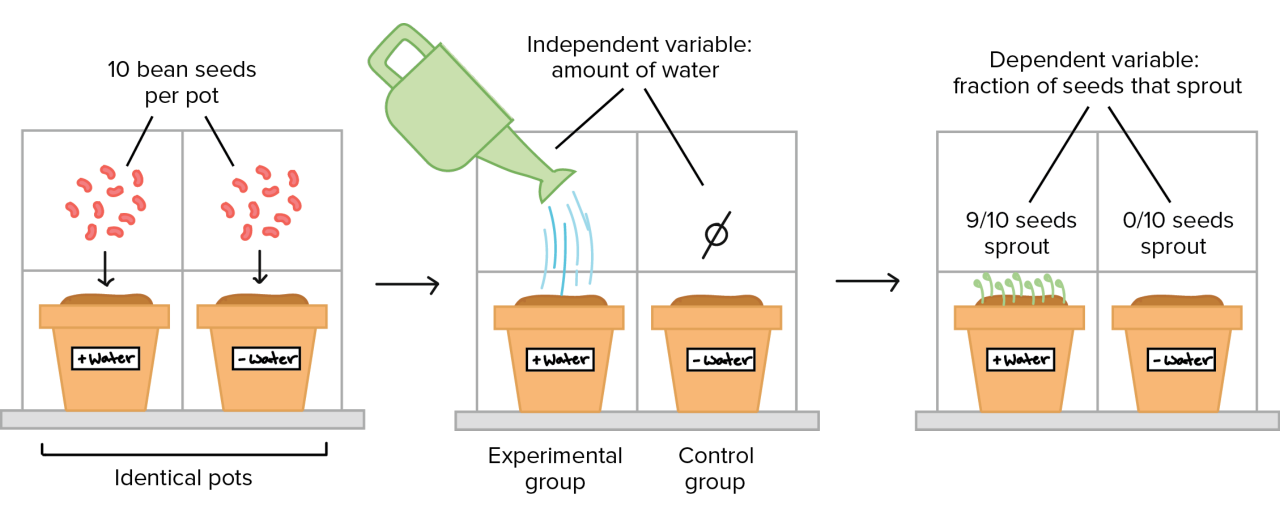
• Theoretical foundation of t-test

Hypothesis testing

- General thinking:
 - Are they different?
 - Is the difference "statistically significant"?
- Statistical thinking:
 - Null hypothesis
 - Alternative hypothesis



Negative control in experiments



Source: <u>https://www.khanacademy.org/science/biology/intro-to-biology/science-of-biology/a/experiments-and-observations</u> Slide fr

Slide from Jessica Li

Null hypothesis in statistical hypothesis testing

AwesomeFinTech

A null hypothesis is a type of conjecture used in statistics that proposes that there is no difference between certain characteristics of a population or data-generating process.

Null Hypothesis : Testing & Examples

ww.awesomefintech.com/terms/null_hypothesis/

The null hypothesis does not depend on a test procedure

Source: <u>https://www.awesomefintech.com/term/null_hypothesis/</u>

Null hypothesis in statistical hypothesis testing

AwesomeFinTech

Hypothesis testing provides a method to reject a null hypothesis within a certain confidence level. (Null hypotheses cannot be proven, though.)

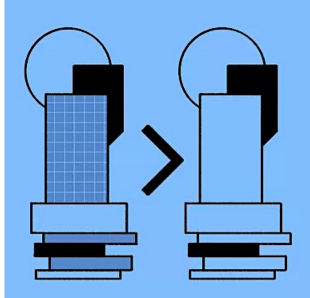
Null Hypothesis : Testing & Examples

ww.awesomefintech.com/terms/null_hypothesis/

Statistics cannot tell us everything b/c data is random

Source: <u>https://www.awesomefintech.com/term/null_hypothesis/</u>

Null hypothesis is often misunderstood



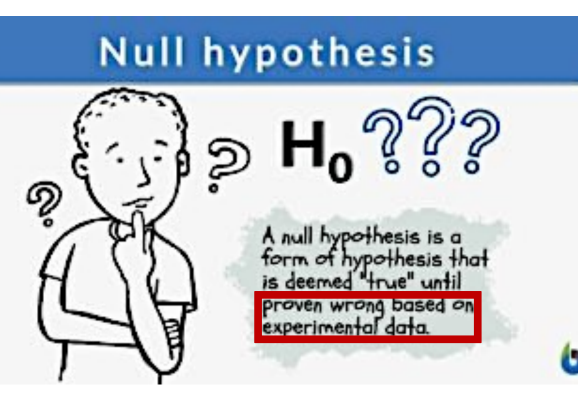
Null Hypthesis

[ˈnəl hī-ˈpä-thə-səs]

A hypothesis that proposes that no statistical significance exists in a set of given observations and is used to assess the credibility of a hypothesis by using sample data.

Investopedia

Source: https://www.investopedia.com/terms/n/null_hypothesis.asp



Source: https://www.biologyonline.com/dictionary/null-hypothesis

Common hypothesis tests

- Student's t-test (parametric)
 - Null (2-sample): The sample means are equal.
- Fisher's exact test
 - Null: The two groups are equally likely for an event/feature.
- Wilcoxon (rank-sum) test
 - Null: The two samples X and Y, P(X>Y) = P(X<Y) = 0.5
- Kolmogorov-Smirnov test (K-S test)
 - Null: The two samples have the same cumulative distribution.
- Hypothesis test statistic, p-value

Which of the following statements about p-values is true?

- A. P-values measure how big the difference is between the datasets compared.
- B. P-value is the probability of observing the data by random chance.
- C. P-value is the least probability of observing the data under the assumption that the null hypothesis is true.

ASA statement on statistical significance and p-values

- 1. P-values can indicate how incompatible the data are with a specified statistical model.
- 2. P-values do not measure the probability that the studied hypothesis is true, or the probability that the data were produced by random chance alone.
- Scientific conclusions and business or policy decisions should not be based only on whether a p-value passes a specific threshold.

ASA statement on statistical significance and p-values

- 4. Proper inference requires full reporting and transparency.
- 5. A p-value, or statistical significance, does not measure the size of an effect or the importance of a result.
- 6. By itself, a p-value does not provide a good measure of evidence regarding a model or hypothesis.

Multiple testing correction

- High-throughput experiments
 - RNA-seq: 500 Differentially Expressed genes from 20K genes
 - ChIP-seq: 10,000 TF binding sites from the genome

Bonferroni correction

- Controls family-wise error rate
- N tests
- Adjusted P-value = N * P
- Very strict



Carlo Emilio Bonferroni (1892-1960)

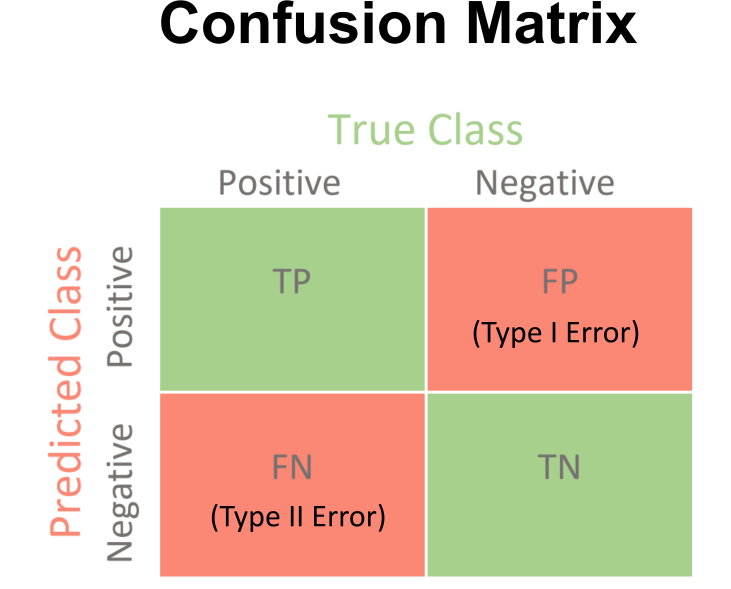
Benjamini-Hochberg (B-H) Correction

- N observations w/ various p-values
- Rank N
- Adjusted P = P * N / R
- Moderate



Yoav Benjamini (1949-)

and Yosef Hochberg



Source: https://towardsdatascience.com/confusion-matrix-for-your-multi-class-machine-learning-model-ff9aa3bf7826

Summary

| | | CONDITION determined by "Gold Standard" | | | |
|----------------------|--|---|---|--|---|
| | TOTAL POPULATION | CONDITION POS | CONDITION NEG | PREVALENCE CONDITION POS TOTAL POPULATION | |
| TEST OUT- COME | TEST POS | True Pos TP | <i>Type I Error</i> False Pos FP | Precision Pos Predictive Value PPV = <u>TP</u> TEST P | False Discovery Rate FDR = <u>FP</u> TEST P |
| | TEST NEG | <i>Type II Error</i> False Neg FN | True Neg TN | False Omission Rate FOR = <u>FN</u> TEST N | Neg Predictive Value NPV = <u>TN</u> TEST N |
| | ACCURACY ACC ACC = <u>TP + TN</u> TOT POP | Sensitivity (SN), Recall Total Pos Rate TPR TPR = <u>TP</u> CONDITION POS | Fall-Out False Pos Rate FPR FPR = <u>FP</u> CONDITION NEG | Pos Likelihood Ratio LR + LR + = <u>TPR</u> FPR | Diagnostic Odds Ratio DOR DOR = <u>LR +</u> LR - |
| | | <i>Miss Rate</i> False Neg Rate FNR FNR = <u>FN</u> CONDITION POS | Specificity (SPC) True Neg Rate TNR TNR = <u>TN</u> CONDITION NEG | Neg Likelihood Ratio LR - LR - = <u>TNR</u> FNR | |

Source: Wikimedia. Author: Lavender888000

SHORT REPORT



Open Access

Exaggerated false positives by popular differential expression methods when analyzing human population samples

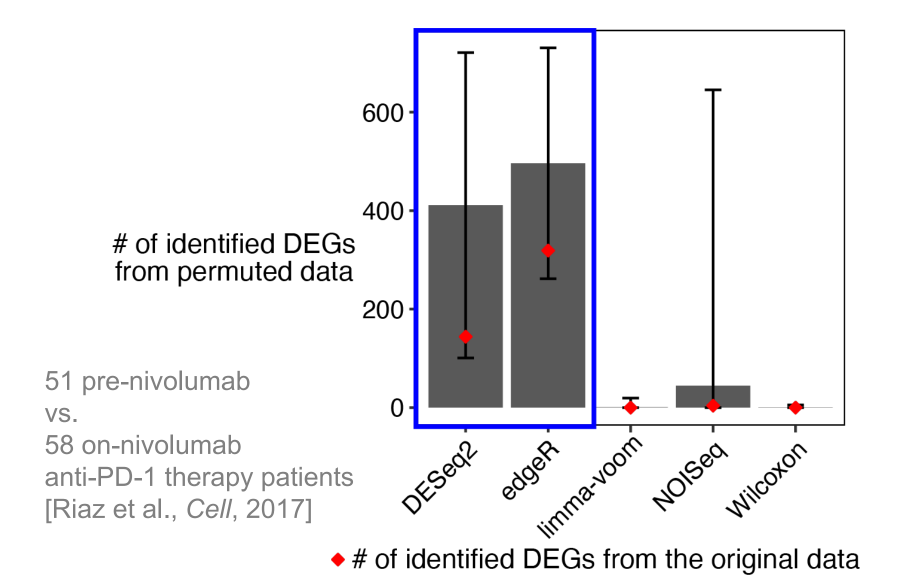
Yumei Li^{1†}, Xinzhou Ge^{2†}, Fanglue Peng³, Wei Li^{1*} and Jingyi Jessica Li^{2,4,5,6,7*}

*Correspondence: wei.li@uci.edu; lijy03@g. ucla.edu [†]Yumei Li and Xinzhou Ge contributed equally to this work. ¹ Division of Computational Biomedicine, Department of Biological Chemistry, School of Medicine, University of California, Irvine, Irvine, CA 92697, USA ² Department of Statistics, University of California, Los

Abstract

When identifying differentially expressed genes between two conditions using human population RNA-seq samples, we found a phenomenon by permutation analysis: two popular bioinformatics methods, DESeq2 and edgeR, have unexpectedly high false discovery rates. Expanding the analysis to limma-voom, NOISeq, dearseq, and Wilcoxon rank-sum test, we found that FDR control is often failed except for the Wilcoxon rank-sum test. Particularly, the actual FDRs of DESeq2 and edgeR sometimes exceed 20% when the target FDR is 5%. Based on these results, for population-level RNA-seq studies with large sample sizes, we recommend the Wilcoxon rank-sum test.

Why there are many DE genes identified from permuted data?



Li et al. Genome Biol 2022

Important assumption in DESeq2 and edgeR

Both DESeq2 and edgeR assume a **negative binomial (NB)** distribution per gene and condition

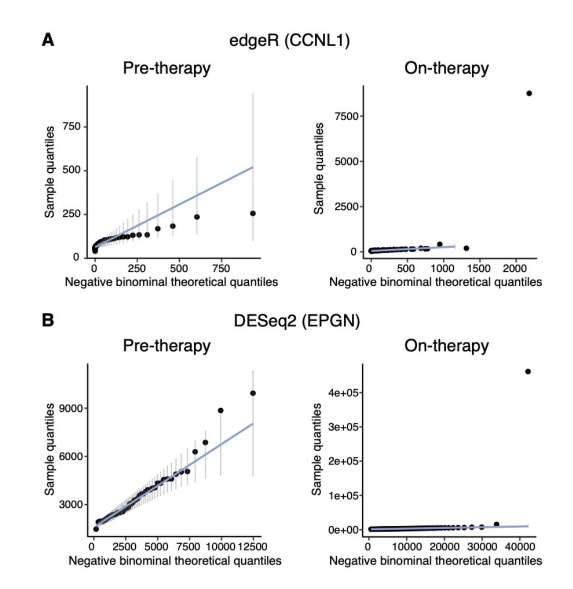
For each gene,

- Condition 1: $X_i \stackrel{\text{ind}}{\sim} \text{NB}(\mu_1 s_i, \sigma_1), i = 1, \dots, n$
- Condition 2: $Y_j \stackrel{\text{ind}}{\sim} \text{NB}(\mu_2 s_j, \sigma_2), \ j = 1, \dots, m$

Null hypothesis $H_0: \mu_1 = \mu_2$

appropriate only if the NB assumption is reasonable

Gene expression can deviate from NB distribution



Li et al. Genome Biol 2022

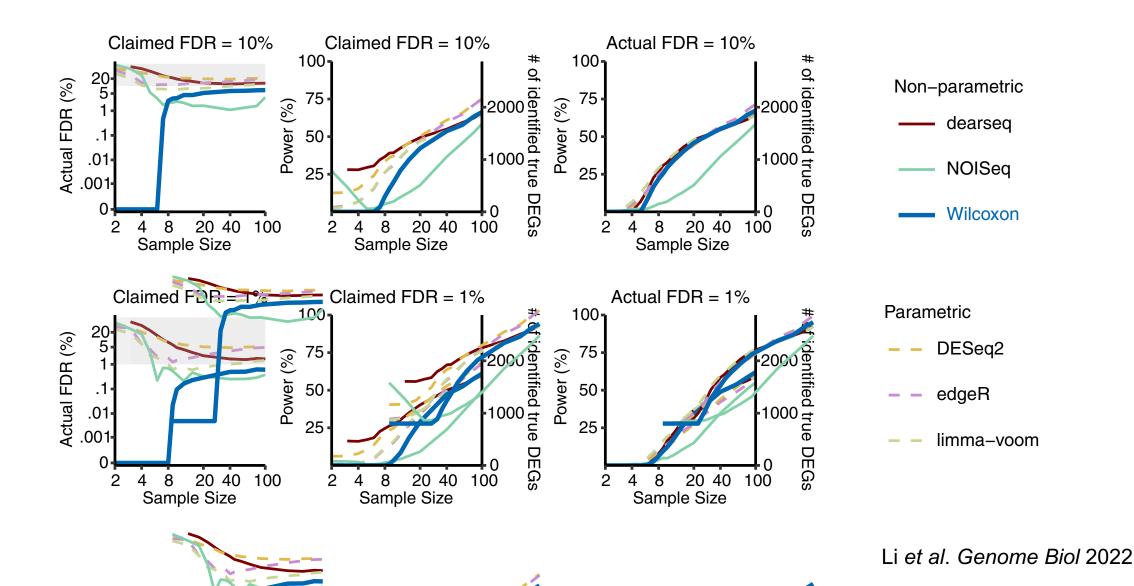
Why does Wilcoxon test work in this scenario?

For each gene, the normalized counts Condition 1: \widetilde{X}_i , i = 1, ..., nCondition 2: \widetilde{Y}_j , j = 1, ..., m

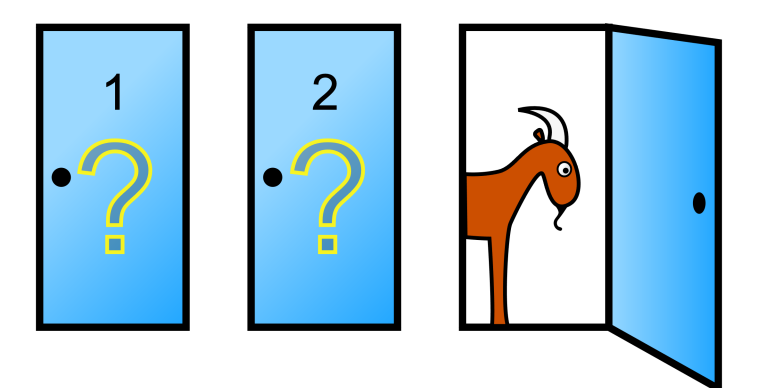
Null hypothesis (approximate, ignoring ties): $H_0: \mathbb{P}(\widetilde{X}_i > \widetilde{Y}_j) = 0.5, \text{ for all } i, j$

which does NOT have the NB assumption

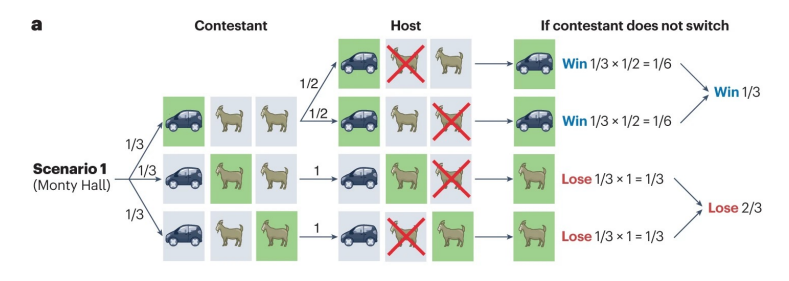
Wilcoxon test is better when sample size > 8

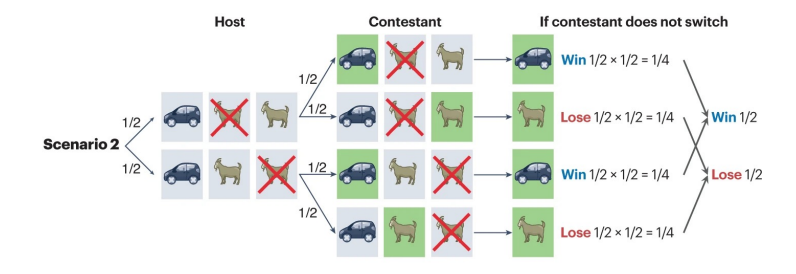


Monty Hall Problem



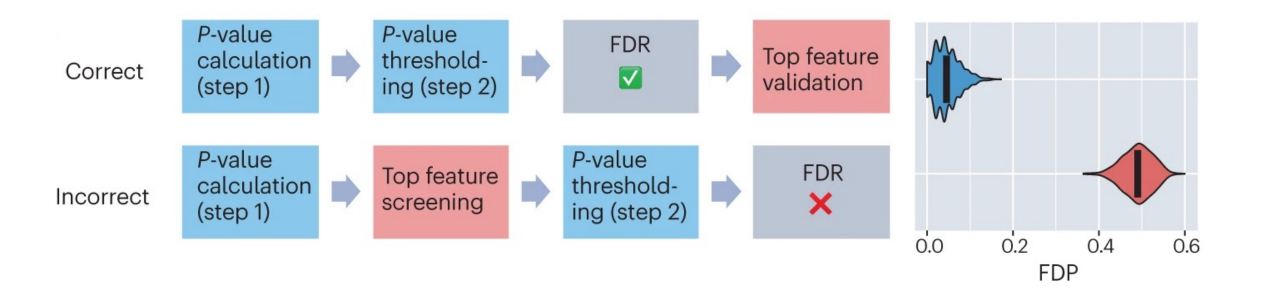
Monty Hall Problem





Li Nat Biotech 2023

The order of action matters





EDITORIAL

Ten Simple Rules for Effective Statistical Practice

Robert E. Kass¹, Brian S. Caffo², Marie Davidian³, Xiao-Li Meng⁴, Bin Yu⁵, Nancy Reid⁶*

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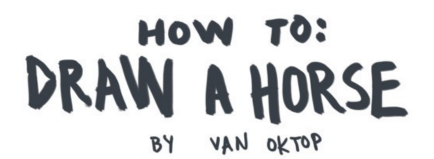
* reid@utstat.utoronto.ca

Ten Simple Rules for Effective Statistical Practice

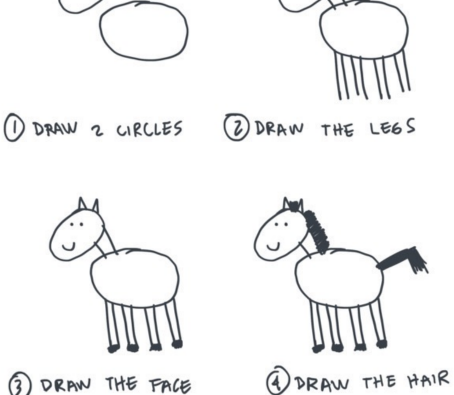
- 1. Statistical methods should enable data to answer scientific questions.
- 2. Signals always come with noise.
- 3. Plan ahead, really ahead.
- 4. Worry about data quality.
- 5. Statistical analysis is more than a set of computations.

Ten Simple Rules for Effective Statistical Practice

- 6. Keep it simple.
- 7. Provide assessments of variability.
- 8. Check your assumptions.
- 9. When possible, replicate!
- 10.Make your analysis reproducible.



Record procedure details!



DRAW THE FACE 3



About Assignment 1 (Section B)

- Record all code and results
- Include sufficient comments and explanations
- Submit in any format (RMD preferred)
- Assignment 1 can be emailed to <u>zang@virginia.edu</u>