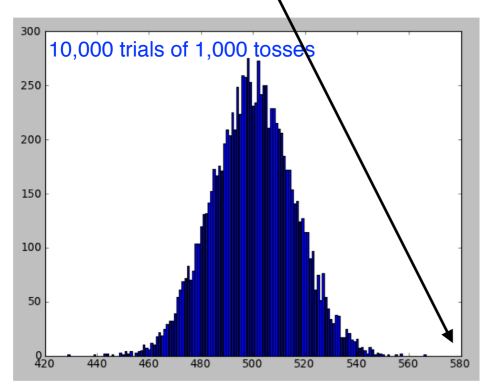
Lectures 5: null hypothesis tests I.

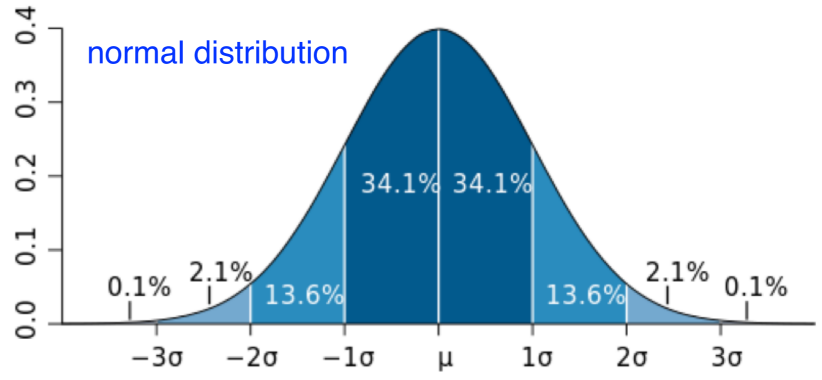
Null hypothesis testing is reductio ad absurdum argument:

- Null hypothesis testing is a reductio ad absurdum argument adapted to statistics: a hypothesis is shown to be valid by demonstrating the improbability of the consequence that results from assuming the counter-claim to be true (fair coin, normal distribution).
- The only hypothesis that needs to be specified in this test and which embodies the counterclaim is referred to as the null hypothesis.
- A result is said to be statistically significant if it allows us to reject the null hypothesis. That is, as per the reductio ad absurdum reasoning, the statistically significant result should be highly improbable if the null hypothesis is assumed to be true.
- The rejection of the null hypothesis implies that the correct hypothesis lies in the logical complement of the null hypothesis. However, unless there is a single alternative to the null hypothesis, the rejection of null hypothesis does not tell us which of the alternatives might be the correct one.

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Null hypothesis testing:

- A statistical hypothesis refers to a probability distribution that is assumed to govern the observed data. If X is a random variable representing the observed data and H is the statistical hypothesis under consideration, then the notion of statistical significance can be quantified by the conditional probability P(X I H), which gives the likelihood of the observation if the hypothesis is assumed to be correct.
- The p-values should not be confused with probability on hypothesis (as is done in Bayesian Hypothesis Testing) such as P(H I X), the probability of the hypothesis given the data, or P(H), the probability of the hypothesis being true, or P(X), the probability of observing the given data.

Null hypothesis testing:

The p-value is defined as the probability, under the assumption of hypothesis H, of obtaining a result equal to or more extreme than what was actually observed. Depending on how it is looked at, the "more extreme than what was actually observed" can mean

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\{ X \ge x \} (right-tail event) or \{ X \le x \} (left-tail event) or the "smaller" of \{ X \le x \} and \{ X \ge x \} (double-tailed event).
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Thus, the p-value is given by

- P($X \ge x \mid H$) for right tail event,
- P($X \le x \mid H$) for left tail event,
- $2*min \{ P(X \le x \mid H), P(X \ge x \mid H) \}$ for double tail event.

The smaller the p-value, the larger the significance because it tells the investigator that the hypothesis under consideration may not adequately explain the observation. The hypothesis H is rejected if any of these probabilities is less than or equal to a small, fixed but arbitrarily pre-defined threshold value α , which is referred to as the level of significance.

Unlike the p-value, the α level is not derived from any observational data and does not depend on the underlying hypothesis; the value of α is instead determined by the consensus of the research community that the investigator is working in.

Null hypothesis testing:

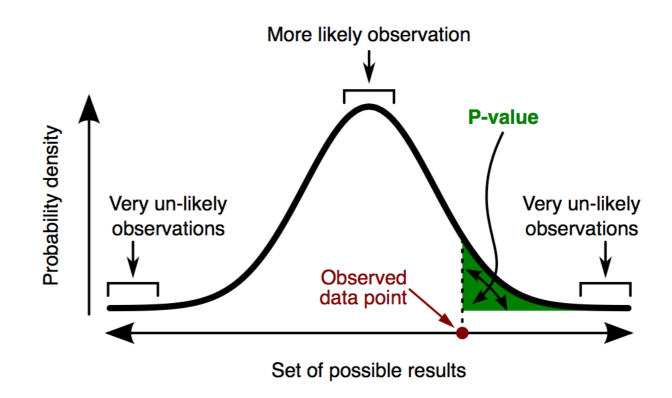
Computing a p-value requires a null hypothesis, a test statistic (together with deciding whether the researcher is performing a one-tailed test or a two-tailed test), and data. Even though computing the test statistic on given data may be easy, computing the sampling distribution under the null hypothesis, and then computing its cumulative distribution function (CDF) is often a difficult computation. Today, this computation is done using statistical software and computational power.

Important:

Pr (observation | hypothesis) ≠ Pr (hypothesis | observation)

The probability of observing a result given that some hypothesis is true is *not equivalent* to the probability that a hypothesis is true given that some result has been observed.

Using the p-value as a "score" is committing an egregious logical error: the transposed conditional fallacy.



A **p-value** (shaded green area) is the probability of an observed (or more extreme) result assuming that the null hypothesis is true.

simplified coin flipping test null hypothesis testing:

As an example of a statistical test, an experiment is performed to determine whether a coin flip is fair (equal chance of landing heads or tails) or unfairly biased (one outcome being more likely than the other).

Suppose that the experimental results show the coin turning up heads 14 times out of 20 total flips. The null hypothesis is that the coin is fair, and the test statistic is the number of heads. If a right-tailed test is considered, the p-value of this result is the chance of a fair coin landing on heads at least 14 times out of 20 flips. That probability can be computed from binomial coefficients as

$$ext{Prob}(14 ext{ heads}) + ext{Prob}(15 ext{ heads}) + \cdots + ext{Prob}(20 ext{ heads}) \\ = rac{1}{2^{20}} \left[inom{20}{14} + inom{20}{15} + \cdots + inom{20}{20}
ight] = rac{60,460}{1,048,576} pprox 0.058 \end{aligned}$$

This probability is the p-value, considering only extreme results that favor heads. This is called a one-tailed test. However, the deviation can be in either direction, favoring either heads or tails. The two-tailed p-value, which considers deviations favoring either heads or tails, may instead be calculated. As the binomial distribution is symmetrical for a fair coin, the two-sided p-value is simply twice the above calculated single-sided p-value: the two-sided p-value is 0.116. In the above example:

null hypothesis (H_0) p(head) = 0.5

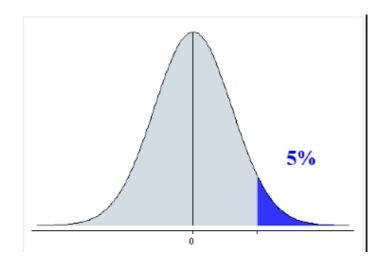
- Test statistic: number of heads
- Level of significance: 0.05
- Observation O: 14 heads out of 20 flips; and
- Two-tailed p-value of observation O given $H_0 = 2*min(Prob(no. of heads \ge 14 heads), Prob(no. of heads \le 14 heads)) = 2*min(0.058, 0.978) = 2*0.058 = 0.116.$

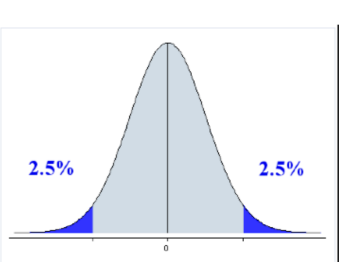
Note that the Prob(no. of heads \leq 14 heads) = 1 - Prob(no. of heads \geq 14 heads) + Prob(no. of head = 14) = 1 - 0.058 + 0.036 = 0.978; however, symmetry of the binomial distribution makes that an unnecessary computation to find the smaller of the two probabilities. Here, the calculated p-value exceeds 0.05, so the observation is consistent with the null hypothesis, as it falls within the range of what would happen 95% of the time were the coin is in fact fair. Hence, the null hypothesis at the 5% level is not rejected. Although the coin did not fall evenly, the deviation from expected outcome is small enough to be consistent with chance.

However, had one more head been obtained, the resulting p-value (two-tailed) would have been 0.0414 (4.14%). The null hypothesis is rejected when a 5% cut-off is used.

Null hypothesis testing summary:

- "null hypothesis"
- "the statistic" (e.g., t-value or χ²)
 - calculable for the null hypothesis
 - intuitively should be "deviation from" in some way
- "the critical region" α
 - biologists use 0.05
 - physicists use 0.0026 (3 σ)
- one-sided or two?
 - somewhat subjective
 - use one-sided only when the other side has an understood and innocuous interpretation
- if the data is in the critical region, the null hypothesis is ruled out at the α significance level
- after seeing the data you
 - may adjust the significance level α
 - may not try a different statistic, because any statistic can rule out at the α level in $1/\alpha$ tries ("data dredging" for a significant result!)
- if you decided in advance to try N tests, then the critical region for α significance is α/N (Bonferroni correction).





Phys. Rev. Lett. discovery threshold: 5 σ (0.000057 percent)

Count nucleotides A,C,G,T on SacCer Chr4:

Take the file **SacSerChr4.txt** (on course web site).

Count the letters **A,C,G,T**.

You should get:

A = 476750

C = 289341

G = 291352

T = 474471



Are these counts consistent with the model

$$p_A = p_C = p_C = p_T = 0.25$$
?

(Of course not! But we'll check.)

Are they consistent with the model

$$p_A = p_T \approx 0.31$$
 $p_C = p_G \approx 0.19$?

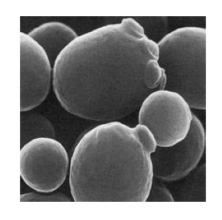
That's a deeper question! You might think yes, because of A-T and C-G base pairing.

but what is the model?

For practice with p- and t-values, let's look at the Sac cer genome. We'll use as a data set all of Chromosome 4.

Yeast and Human are very close relatives in the great scheme of things.

Saccharomyces cerevisiae = baker's yeast



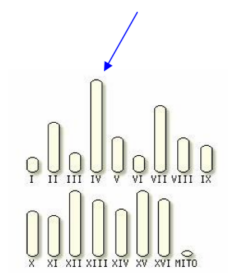


goal is to build probability models for chromosome 4 from four nucleobases

ACGT and subject them to null hypothesis

Chromosome 4: ACACCACACC (1531894 omitted) TAGCTTTTGG

Watson-Crick base pairs: (guanine-cytosine and adenine-thymine)

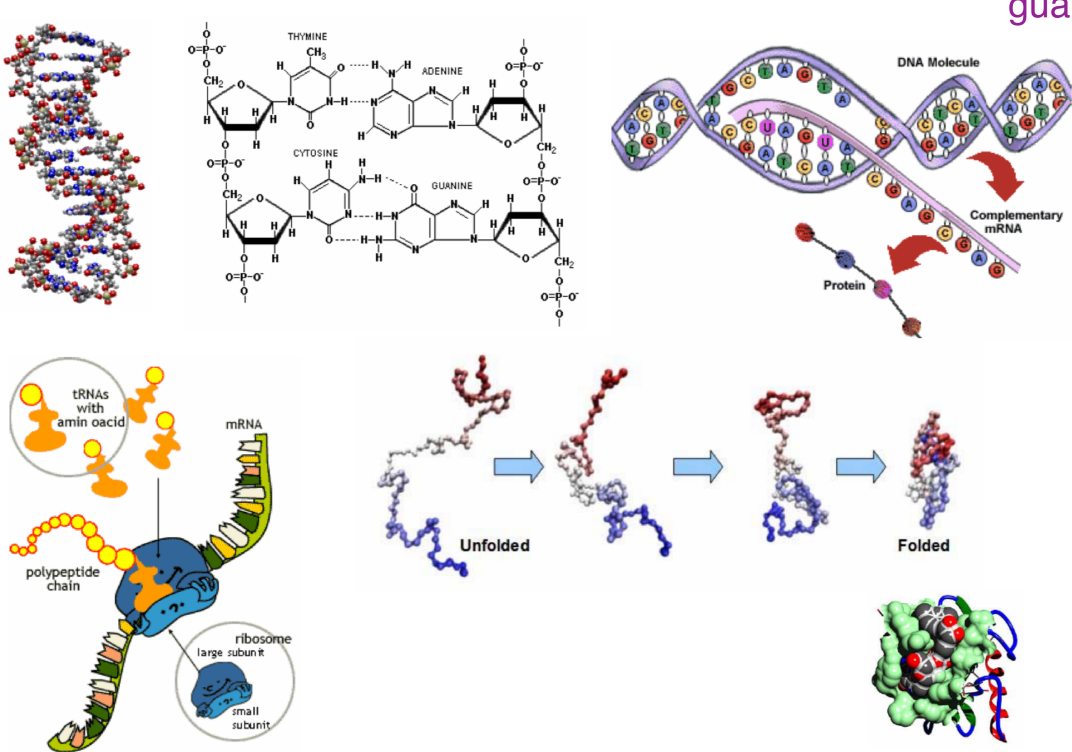


molecular biology on one slide:

https://en.wikipedia.org/wiki/DNA#Base_pairing

https://en.wikipedia.org/wiki/Base_pair

adenine-thymine guanin-cytosin



As always, the starting point is to write down a model. Bayesian: What is the probability of hypothesis. Frequentist: What is the probability of a test statistic for a null hypothesis.

A possible model is multinomial: At each position an i.i.d. choice of A,C,G,T, with respective probabilities adding up to 1.

Almost equivalent (and simpler for now) is 4 separate binomial models: At each position an i.i.d. choice of A vs. not A with some probability p_A . Then do separately for p_C , p_G , p_T .

The counts are all so large that the normal approximation is highly accurate:

$$Bin(n, p) \approx Normal(np, \sqrt{np(1-p)})$$

Why? CLT applies to binomial because it's sum of Bernoulli r.v.'s: N tries of an r.v. with values 1 (prob p) or 0 (prob 1-p).

$$\mu = p \times 1 + (1 - p) \times 0 = p$$

$$\sigma^2 = p \times (1 - \mu)^2 + (1 - p) \times (0 - \mu)^2 = p(1 - p)$$

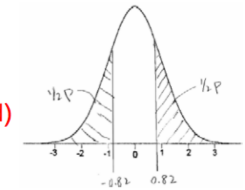
Let's dispose of the silly (all p's = 0.25):

The test statistic: the value of the observed count under the null hypothesis that it is binomially (or equivalent normally) distributed with p=0.25.

$$egin{aligned} \mu &= 0.25\,N \ \sigma &= \sqrt{0.25 imes 0.75\,N} \ t &= rac{n-\mu}{\sigma} \ p &= 2[1-P_{ ext{Normal}}(|t|)] \end{aligned}$$

t-value = number of standard deviations

p-value = tail probability (here, 2-tailed)



	t-value	p-value
Α	174.965	≈ 0
С	-174.715	≈ 0
G	-170.963	≈ 0
Т	170.713	≈ 0

The null hypothesis is (totally, infinitely, beyond any possibility of redemption!) ruled out.

The not-silly model: A and T occur with identical probabilities, as do C and G.

The test statistic: Difference between A and T (or C and G) counts under the null hypothesis that they have the same p, which we will estimate in the obvious way (which is actually an MLE).

$$\hat{p}_{AT} = \frac{1}{2}(n_A + n_T)/N$$

$$\hat{p}_{CG} = \frac{1}{2}(n_C + n_G)/N$$

$$n_A \sim \text{Normal}(N\hat{p}_{AT}, \sqrt{N\hat{p}_{AT}(1 - \hat{p}_{AT})})$$

$$n_T \sim \text{Normal}(N\hat{p}_{AT}, \sqrt{N\hat{p}_{AT}(1 - \hat{p}_{AT})})$$

$$\Rightarrow n_A - n_T \sim \text{Normal}(0, \sqrt{2N\hat{p}_{AT}(1 - \hat{p}_{AT})})$$

the difference of two Normals is itself Normal

the variance of the sum (or difference) is the sum of the variances

In MATLAB the calculation now looks like this:

```
dif = [count(1)-count(3); count(2)-count(4) ]
                                                                     A = 476750
pdiff = [pnuc(1); pnuc(2)]
                                                                     C = 289341
mu = [0; 0];
                                                                     G = 291352
sig = sqrt(2 .* pdiff .* (1 - pdiff) .* len)
                                                                     T = 474471
tval = (dif - mu) ./ sig
pval = 2*(1-normcdf(abs(tval),0,1))
                          2-tailed
 dif =
         -2279
                                                  Why? Because, we're discovering genes!
         -2011
 pdiff =
                                                                                          + strand
      0.3097
      0.1889

    strand

 mu =
       0
                                                 The fluctuating "units" are indeed not single bases.
       0
                                                 Rather, they are genes which, individually, do not
 sig =
    809.3402
                                                 have (or prefer) A=T, C=G. Their placement on
    685.1154
                                                 one strand or the other is random.
 tva1 =
     -2.8159
                      Surprise!
     -2.9353
                      The model is ruled out
 pva1 =
      0.0049
                      with high significance
      0.0033
                      (small p-value)!
```