p-hacking
Test statistics and hypothesis tests
Compare two classifiers
A/B testing
Summary

Lecture 8: Hypothesis testing part II Statistical Methods for Data Science

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Today

- p-hacking
- Test statistics and hypothesis tests
 - z-test
 - One-sample t-test
 - Two-sample t-test
 - Paired t-test
 - Binomial test
 - McNemar's test
- Compare two classifiers
- 4 A/B testing
- Summary

Learning outcome

- Be able to explain the concept of p-hacking
- Be able to explain the following hypothesis tests
 - One-sample and two-sample z-test
 - One-sample and two-sample t-test
 - Paired t-test
 - Binomial test (exact, approximate)
 - McNemar's test (exact, approximate)

For each of these tests, be able to describe the typical set up for the experiment, the general purpose of the test, data produced by the experiment, random variables, parameter of interest, null hypothesis, alternative hypothesis, test statistic, null distribution, the computation of *p*-value

 Be able to compare two classifiers using the paired t-test and McNemar's test for different scenarios



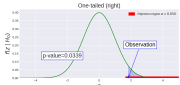


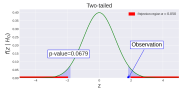
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Recall: one-tailed vs two-tailed tests

- p-value indicates how "surprising" the observation is
- "Surprising" observation usually means potential novelty
- In one of the examples from the previous lecture, we have shown that we reject
 the null hypothesis for the one-tailed test and we fail to reject the null
 hypothesis for the two-tailed test given the same significance level





- In this example, if we use the two-tailed test, we will not claim that we have observed potential novelty with the experiment, whereas if we use the one-tailed test, we claim that we do observe potential novelty
- The conclusion we draw depends on which test we conduct





Variation of the p-value

- *p*-value is computed from data
- Data is random p-value is random
- With the same experiment set up, if we switch to a different sample,
 p-value will be different



p-hacking

- Many factors can result in a different p-value
- p-hacking refers to situations where researchers are trying multiple things until they get the desired result
- This action can be a conscious decision, a subconscious decision or even an unconscious action
- p-hacking can be tricky to identify
- Suggestions to avoid p-hacking, e.g. one should always report effect sizes and confidence intervals
- Reference:
 - https://www.nature.com/news/ scientific-method-statistical-errors-1.14700
 - Why Most Published Research Findings Are False?





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Disclaimer

 Recall that in this course, we only consider H₀ with an equal sign in them, i.e. the null distribution is fully specified; the description of H₀ is based on this assumption



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- For symmetric null distributions, e.g. standard Gaussian distribution, student's t distribution, binomial distribution with p=0.5, etc, we only illustrate examples with the two-tailed alternative hypothesis H_A in this lecture without loss of generality; the one-tailed version can be easily derived



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- For the exact binomial test with $p \neq 0.5$, the null distribution is not symmetric; in this case, the computation of the two-tailed p-value is not uniquely defined; in this lecture, we will not go into details for these cases; we will only look at the one-tailed tests for asymmetric binomial null distributions



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- For each hypothesis test, the purpose of the Python code snippet is to provide a better understanding of the calculation; in practice, there are alternative libraries and built-in functions for these tests that might result in a more compact implementation
- We will come back to this lecture next week for some examples (to give you some time getting familiar with the concepts before the exercises)





Disclaimer (cont.)





Disclaimer (cont.)

- Typical set up for the experiment
 - Test subjects, e.g. the number of samples, the number of groups, etc
 - Description of the experiment and the result
 - Description of the data type produced in the result



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- Typical set up for the experiment
 - Test subjects, e.g. the number of samples, the number of groups, etc
 - Description of the experiment and the result
 - Description of the data type produced in the result
- Purpose: the general purpose of the test





Disclaimer (cont.)

- Typical set up for the experiment
 - Test subjects, e.g. the number of samples, the number of groups, etc
 - Description of the experiment and the result
 - Description of the data type produced in the result
- Purpose: the general purpose of the test
- Data: symbolic description of the data produced by the experiment
- Random variable and assumption corresponding to the data



Disclaimer (cont.)

- Typical set up for the experiment
 - Test subjects, e.g. the number of samples, the number of groups, etc
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- Parameter of interest and the estimates



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- Typical set up for the experiment
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- Data: symbolic description of the data produced by the experiment
- Random variable and assumption corresponding to the data
- Parameter of interest and the estimates
- Hypotheses H_0 and H_A

Disclaimer (cont.)

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- Test statistic
- Null distribution
 - PDF/PMF: description of the PDF/PMF
 - Python: code snippet of the PDF/PMF





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- Typical set up for the experiment
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- Parameter of interest and the estimates
- Hypotheses H_0 and H_A
- Test statistic
- Null distribution
 - PDF/PMF: description of the PDF/PMF
 - Python: code snippet of the PDF/PMF
- p-value
 - Definition: an expression of p-value in terms of a probability
 - Python: code snippet to illustrate the computation of the p-value (see page 9)





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Compare two classifiers
A/B testing

z-test
One-sample t-test
Two-sample t-test
Paired t-test
Binomial test
McNemar's test

z-test





One-sample z-test

- Typical set up for the experiment:
 - One sample of independent test subjects, e.g. a sample of patients, a sample of customers, etc
 - Run the same experiment on each subject and collect the outcomes, e.g. give a new drug to a sample of patients and measure the effect on each individual patient; test a new web design on a sample of customers and record the time they spend on the web page, etc.
 The result contains one i.i.d. sample with continuous numerical values
- Purpose: to test if the mean of the result differs from a predefined constant
- Data: x_1, \dots, x_N , e.g. blood pressure after taking a new drug
- Random variable and assumption: X_1, \dots, X_N
 - Xi i.i.d.
 - X_i Gaussian or large N (CLT)
 - X_i standard deviation σ known
- Parameter of interest: μ
- Parameter estimate: \bar{x} , $\bar{X} \sim \mathcal{N}(\mu, \sigma^2/N)$
- Hypotheses H_0 and H_A : given c a constant

 H_0 :

 $H_A: \mu \neq c$

 $\mu = c$





One-sample z-test (cont.)

Test statistic:

$$z_0 = \frac{\bar{x} - c}{\sigma / \sqrt{N}}$$

Null distribution: standard normal distribution

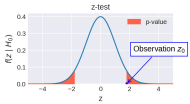
• PDF:
$$f(z \mid H_0) = \frac{1}{\sqrt{2\pi}}e^{-\frac{z^2}{2}}$$

• Python: stats.norm.pdf(z, 0, 1)

p-value

• Definition:
$$p = 2 \min (P(Z \le z_0 \mid H_0), P(Z \ge z_0 \mid H_0))$$

• Python: $2 * min(stats.norm.cdf(z_0, 0, 1), 1-stats.norm.cdf(z_0, 0, 1))$







z-test

Two-sample z-test

- Typical set up for the experiment:
 - Two samples of independent test subjects, where the two samples \mathcal{X} and \mathcal{Y} letters with a calligraphic font are typically used to denote sets are independent from one another, e.g. two samples of independent patients, two samples of independent customers, etc.
 - Run two sets of experiments A and B on the test subjects from the two samples \mathcal{X} and \mathcal{Y} , respectively, and collect the outcomes, e.g. give different drugs to the two samples of patients and measure the effect on each individual patient; test two web designs on two samples of customers and record the time they spend on the web page, etc
 - The result contains two i.i.d. samples with continuous numerical values
- Purpose: to test if two alternative options have different effects by testing if the mean of the result from one sample differs from the mean of the other sample
- Data: x_1, \dots, x_{N_V} and y_1, \dots, y_{N_V} , e.g. blood pressure measured after taking two different drugs
- Random variable and assumption: $X_1, \dots, X_{N_x}, Y_1, \dots, Y_{N_y}$
 - X_i and Y_i independent
 - Xi i.i.d.: Yi i.i.d.
 - X_i Gaussian or large N_X; Y_i Gaussian or large N_Y
 - X_i and Y_i have known standard deviation σ_X and σ_Y , respectively
- Parameter of interest: ux. uy
- Parameter estimate: x̄, ȳ
- Hypotheses H_0 and H_A : given c a constant

 $H_0: \mu_X - \mu_Y = c$ $H_A: \mu_X - \mu_Y \neq c$





z-test Binomial test

Two-sample z-test (cont.)

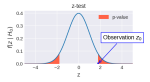
Test statistic:

$$z_0 = \frac{\bar{x} - \bar{y} - c}{\sqrt{\frac{\sigma_X^2}{N_X} + \frac{\sigma_Y^2}{N_Y}}}$$

Hint: $\bar{X} - \bar{Y} \sim \mathcal{N} \left(\mu_X - \mu_Y, \sigma_X^2 / N_X + \sigma_Y^2 / N_Y \right)$

- Null distribution: standard normal distribution
 - PDF: $f(z \mid H_0) = \frac{1}{\sqrt{2\pi}} e^{-\frac{z^2}{2}}$
- Pvthon: stats.norm.pdf(z, 0, 1) p-value
- Definition: $p = 2 \min (P(Z \le z_0 \mid H_0), P(Z \ge z_0 \mid H_0))$

 - Python: $2 * min(stats.norm.cdf(z_0, 0, 1), 1-stats.norm.cdf(z_0, 0, 1))$







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One-sample t-test





One-sample t-test

- Typical set up for the experiment (same as the one-sample z-test):
 - One sample of independent test subjects, e.g. a sample of patients, a sample of customers, etc
 - Run the same experiment on each subject and collect the outcomes, e.g. give a new drug to a sample of patients and measure the effect on each individual patient; test a new web design on a sample of customers and record the time they spend on the web page, etc
 - The result contains one i.i.d. sample with continuous numerical values
- Purpose: to test if the mean of the result differs from a predefined constant
- Data: x_1, \dots, x_N , e.g. blood pressure after taking a new drug
- Random variable and assumption: X_1, \dots, X_N
 - X_i i.i.d.
 - X_i Gaussian or large N
 - X_i standard deviation σ unknown
- Parameter of interest: μ
- Parameter estimate: x̄
- Hypotheses H_0 and H_A : given c a constant

 $H_0: \qquad \mu = c$

 $H_A: \mu \neq c$





One-sample t-test (cont.)

Test statistic:

$$t_0 = \frac{\bar{x} - c}{s / \sqrt{N}}$$

where $s = \sqrt{\frac{1}{N-1} \sum_{i=1}^{N} (x_i - \bar{x})^2}$ is the sample standard deviation

- Null distribution:
 - Student-t distribution with degree of freedom df = N 1
 - Python: stats.t.pdf(t, df = N 1)
- p-value:
 - Definition: $p = 2 \min (P(T \le t_0 \mid H_0), P(T \ge t_0 \mid H_0))$
 - Python: $2 * \min(\text{stats.t.cdf}(t_0, df = N 1), 1 \text{stats.t.cdf}(t_0, df = N 1))$







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Two-sample t-test





Two-sample t-test

- Typical set up for the experiment (same as the two-sample z-test):
 - ullet Two samples of independent test subjects, where the two samples ${\mathcal X}$ and ${\mathcal Y}$ are independent from one another, e.g. two samples of independent patients, two samples of independent customers, etc
 - Run two sets of experiments A and B on the test subjects from the two samples X and Y, respectively, and
 collect the outcomes, e.g. give different drugs to the two samples of patients and measure the effect on each
 individual patient; test two web designs on two samples of customers and record the time they spend on the web
 page, etc
 - The result contains two i.i.d. samples with continuous numerical values
- Purpose: to test if two alternative options have different effects by testing if the mean of the result from one sample differs from the mean of the other sample
- Data: x_1, \dots, x_{N_X} and y_1, \dots, y_{N_Y} , e.g. blood pressure measured after taking two different drugs
- Random variable and assumption: $X_1, \dots, X_{N_X}, Y_1, \dots, Y_{N_Y}$
 - X_i and Y_i independent
 - X_i i.i.d.; Y_i i.i.d.
 - X_i Gaussian or large N_X; Y_i Gaussian or large N_Y
 - X_i and Y_j have unknown standard deviation σ_X and σ_Y , respectively
- Parameter of interest: μ_X , μ_Y
- Parameter estimate: x̄, v̄
- Hypotheses H_0 and H_A : given c a constant

 $H_0: \quad \mu_X - \mu_Y = c$ $H_A: \quad \mu_X - \mu_Y \neq c$





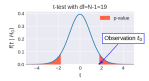
Two-sample t-test (cont.)

Test statistic:

$$t_0 = \frac{\bar{x} - \bar{y} - c}{\sqrt{\frac{s_X^2}{N_X} + \frac{s_Y^2}{N_Y}}}$$

with degree of freedom $\frac{\textit{df}}{\frac{s_X^2/N_X + s_Y^2/N_Y)^2}{(\frac{s_X^2}{N_Y})^2/(N_X - 1) + (\frac{s_Y^2}{N_Y})^2/(N_Y - 1)}}$

- Null distribution:
 - Student-t distribution with degree of freedom df
 - Python: stats.t.pdf(t, df = df)
- p-value:
 - Definition: $p = 2 \min (P(T \le t_0 \mid H_0), P(T \ge t_0 \mid H_0))$
 - Python: $2 * min(stats.t.cdf(t_0, df=df), 1-stats.t.cdf(t_0, df=df))$







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Paired t-test





Paired t-test

- Typical set up for the experiment:
 - One sample of independent test subjects, e.g. one sample of independent patients
 - Run two sets of experiments A and B on all subjects from the sample and collect the outcomes,
 e.g. measure the blood pressure of the patients before giving them a new drug (experiment
 A); measure the blood pressure of the patients after giving them the new drug (experiment B)
 - The result contains two samples with continuous numerical values
- Purpose: to test if two alternative options have different effects by testing if the mean of the difference between two results differs from a predefined constant
- Data: x₁,..., x_N, y₁,..., y_N
 Random variable and assumption: X₁,..., X_N, Y₁,..., Y_N
 - $X_i Y_i$ iid
 - $X_i Y_i \sim \mathcal{N}\left(\mu_{X-Y}, \sigma_{X-Y}^2\right)$ with unknown standard deviation
- Parameter of interest: μ_{X-Y}
- Parameter estimate: $m_{X-Y} = \frac{1}{N} \sum_{i=1}^{N} (x_i y_i)$
- Hypotheses H_0 and H_A : given c a constant

$$H_0: \mu_{X-Y} = c$$

$$H_A: \mu_{X-Y} \neq c$$





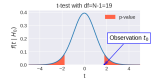
Paired t-test

Test statistic:

$$t_0 = \frac{m_{X-Y} - c}{s_{X-Y}/\sqrt{N}}$$

where
$$s_{X-Y} = \sqrt{\frac{1}{N-1} \sum_{i=1}^{N} (x_i - y_i - m_{X-Y})^2}$$

- Null distribution:
 - ullet Student-t distribution with degree of freedom N-1
 - Python: stats.t.pdf(t, df = N 1)
- p-value:
 - Definition: $p = 2 \min (P(T \le t_0 \mid H_0), P(T \ge t_0 \mid H_0))$
 - Python: $2 * min (stats.t.cdf(t_0, df = N 1), 1-stats.t.cdf(t_0, df = N 1))$





Exercise 1

 A company claims that a new drug E they have developed can increase the average sleeping hours of people with insomnia. Design three different hypothesis tests to test this statement.



z-test
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Exercise 2

One of the tests you have designed is a two-sample test. After the
experiments, you realized the test subjects being selected in the
second group are parents or siblings of the first group. Would that be
a problem? Can you still use the result somehow?



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Binomial test





Binomial distribution

- Discrete distribution
- Applies to discrete numerical data the number of success from n independent Bernoulli trials with probability of success p
- PMF:
 - Equation

$$f_X(k \mid n, p) = P(X = k) = \binom{n}{k} p^k (1 - p)^{n-k}, \ k = 0, \dots, n, \ p \in [0, 1]$$

where $\binom{n}{k} = \frac{n!}{k!(n-k)!}$ is the binomial coefficient (n choose k)

- Shape
 - When p = 0.5, the PMF is symmetric
 - When $p \neq 0.5$, the PMF is asymmetric



Parameters: p and n: n is typically known





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(exact) Binomial test

- Typical set up for the experiment:
 - One sample of independent test subjects, e.g. one sample of independent patients
 - Run the same experiment on all subjects from the sample and collect the outcomes,
 e.g. give a new drug to a sample of patients and measure how many patients are cured
 - The result contains one sample with nominal categorical values with two categories, which is then summarized into one discrete numerical value - the number of "success"
- Purpose: to test if the proportion of "success" differs from a predefined constant
- Data: N independent Bernoulli trials with k_0 "success", e.g. the number of cured patients within the sample of size N
- Random variable and assumption: K ~ Binomial(N, p) with known N and unknown success rate p
- Parameter of interest: p
- Parameter estimate: $\hat{p} = \frac{k_0}{N}$
- Null hypothesis: given π a constant,

$$H_0: p = \pi$$





z-test
One-sample t-tes
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McNemar's test

(exact) Binomial test (cont.)

- Test statistic: k_0
- Null distribution:

$$P(X=k) = \binom{N}{k} \pi^{k} (1-\pi)^{N-k}$$

- ullet Binomial distribution with parameters N and π
- Python: stats.binom.pmf(k, N, π)
- As discussed in the disclaimer (cf. page 9), we only introduce the following scenarios:
 - One-tailed (left) binomial test with any $\pi \in (0,1)$
 - ullet One-tailed (right) binomial test with any $\pi \in (0,1)$
 - ullet Two-tailed binomial test with $\pi=$ 0.5, where the null distribution is symmetric





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(exact) One-tailed (left) binomial test

• Hypotheses H_0 and H_A :

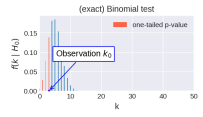
$$H_0: p = \pi$$

 $H_A: p < \pi$

p-value:

• Definition: $p = P(K \le k_0 \mid H_0)$

• Python: stats.binom.cdf(k_0 , n=N, p= π)



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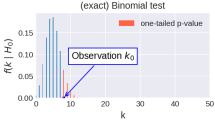
(exact) One-tailed (right) binomial test

• Hypotheses H_0 and H_A :

$$H_0: p = \pi$$

 $H_A: p > \pi$

- p-value:
 - Definition: $p = P(K \ge k_0 \mid H_0)$
 - Python: 1- stats.binom.cdf($k_0, n=N, p=\pi$) + stats.binom.pmf($k_0, n=N, p=\pi$)







(exact) Two-tailed binomial test

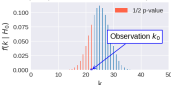
• Hypotheses H_0 and H_A :

$$H_0: p = 0.5$$

 $H_A: p \neq 0.5$

- p-value:
 - Definition: $p = 2 \min (P(K \le k_0 \mid H_0), P(K \ge k_0 \mid H_0))$
 - Python:
 - $c = \text{stats.binom.cdf}(k_0, n = N, p = 0.5)$
 - $2 * min(c, 1 c + stats.binom.pmf(k_0, n = N, p = 0.5))$

(exact) two-tailed Binomial test with p = 0.5



(large N) Binomial test

Same set up as page 29, but with large N

Test statistic:

$$z_0 = \frac{k_0 - N\pi}{\sqrt{N\pi(1-\pi)}}$$

Null distribution: standard normal distribution

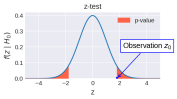
• PDF:
$$f(z \mid H_0) = \frac{1}{\sqrt{2\pi}} e^{-\frac{z^2}{2}}$$

Python: stats.norm.pdf(z, 0, 1)

p-value:

• Definition: $p = 2 \min (P(Z \le z_0 \mid H_0), P(Z \ge z_0 \mid H_0))$

• Python: $2 * min(stats.norm.cdf(z_0, 0, 1), 1-stats.norm.cdf(z_0, 0, 1))$







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Paired t-test
Binomial test
McNemar's test

McNemar's test

- Typical set up for the experiment:
 - One sample of independent test subjects, e.g. one sample of independent patients
 - Within the sample, there are two groups; each subject belongs to one and only one group, e.g. within
 the sample of patients, we have one group with high blood pressure and another group with normal
 blood pressure
 - Run two sets of experiments A and B on all test subjects from the sample and collect the outcomes, e.g. measure the blood pressure (hight or normal) of the patients before giving them a new drug (experiment A); measure the blood pressure (hight or normal) of the patients after giving them the new drug (experiment B)
 - The result contains one sample with nomimal categorical values with two categories measured from each test subject, e.g. high blood pressure and normal blood pressure
- Purpose: to test if an action have different effects on two different groups
- Data: *N* independent Bernoulli trials with outcomes x_1, \dots, x_N and y_1, \dots, y_N for the two experiments, respectively; $x_i, y_i \in \{0, 1\}$

	$x_i = 0$	$x_i = 1$	
$y_j = 0$	n ₀₀	n ₁₀	$n_{00} + n_{10}$
$y_j = 1$	n ₀₁	n ₁₁	$n_{01} + n_{11}$
	$n_{00} + n_{01}$	$n_{10} + n_{11}$	N

where n_{mn} is the count of $x_i = m$ and $y_j = n$

• Random variable and assumption: i.i.d. $X_i \sim Bernoulli(p_X)$ and i.i.d. $Y_i \sim Bernoulli(p_Y)$

McNemar's test (cont.)

Example

- A company is trying to determine the effectiveness of a drug on lowering blood pressure
- The company tested the drug on a sample of 229 independent patients
- There are two groups within this sample: a high blood pressure group (105 patients) and a normal blood pressure group (124 patients); each patient belongs to one and only one of these two groups
- The blood pressure of each patient is measured before (to determine the group) and after (to determine the effect) taking the drug
- The data is summarized as follows:

	After (high blood pressure)	After (normal blood pressure)	
Before (high blood pressure)	90	15	105
Before (normal blood pressure)	22	102	124
	112	117	229





z-test
One-sample t-te
Two-sample t-te
Paired t-test
Binomial test
McNemar's test

(small discordance $n_{01} + n_{10}$) McNemar's test (cont.)

Parameter of interest: discordance

$$p = \min(P(X_i = 0, Y_i = 1 \mid X_i \neq Y_i), P(X_i = 1, Y_i = 0 \mid X_i \neq Y_i))$$

- Parameter estimate: $\hat{p} = \frac{\min(n_{01}, n_{10})}{n_{01} + n_{10}}$
- Hypotheses H_0 and H_A :

$$H_0: p = 0.5$$

 $H_A: p \neq 0.5$

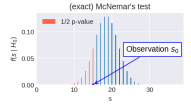
- H_0 : the drug does not have any effect on blood pressure control
- H₁: the drug has effect on blood pressure control





(small discordance $n_{01} + n_{10}$) McNemar's test (cont.)

- Test statistic: $s_0 = \min(n_{01}, n_{10})$
- Null distribution:
 - Binomial distribution with parameters $(n_{01} + n_{10}, 0.5)$
 - Python: stats.binom.pmf(s, $n_{01} + n_{10}$, 0.5)
- p-value:
 - Definition: $p = 2P(S \le s_0 \mid H_0)$
 - Python: $2 * stats.binom.cdf(s_0, n_{01} + n_{10}, 0.5)$







z-test
One-sample t-te
Two-sample t-te
Paired t-test
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(large discordance $n_{01}+n_{10}$) McNemar's test

Same set up as page 36, but with large $n_{01} + n_{10}$, e.g. $n_{01} + n_{10} > 25$

- Parameter of interest: discordance (note: it is different from the previous definition (cf. page 38)) $p_{01} = P(X = 0, Y = 1)$ and $p_{10} = P(X = 1, Y = 0)$
- Parameter estimate: $\hat{p}_{01} = \frac{n_{01}}{N}$ and $\hat{p}_{10} = \frac{n_{10}}{N}$
- Hypotheses H_0 and H_A :

$$H_0: p_{01} = p_{10}$$

 $H_A: p_{01} \neq p_{10}$

Test statistic:

$$s_0 = \frac{\left(|n_{01} - n_{10}| - 1\right)^2}{n_{01} + n_{10}}$$

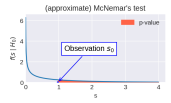
Note: "-1" is called the continuity correction (https://en.wikipedia.org/wiki/Continuity_correction)





(large discordance $n_{01} + n_{10}$) McNemar's test (cont.)

- Null distribution:
 - Large $n_{01} + n_{10}$: s_0 follows a chi-squared distribution with df = 1
 - Python: stats.chi2.pdf(s, df = 1)
- p-value:
 - Definition: $P(S \ge s_0 \mid H_0)$
 - Python: 1-stats.chi2.cdf(s_0 , df = 1)



z-test
One-sample t-tes
Two-sample t-tes
Paired t-test
Binomial test
McNemar's test

Exercise 3

• Run both the exact McNemar's test and the approximate McNemar's test on the data set provided on page 37.



Today

- p-hacking
- 2 Test statistics and hypothesis tests
- 3 Compare two classifiers
- 4 A/B testing
- Summary

K-fold cross validation

- Classifiers: A and B
- Data: evaluation metric; continuous numerical data, e.g. accuracies p_1^A, \cdots, p_K^A and p_1^B, \cdots, p_K^B on the K validation sets

	fold 1	fold 2	 fold K
classifier A	p_1^A	p_2^A	 p_K^A
classifier B	p_1^B	p_2^B	 p_K^B
$p_i^A - p_i^B$	$p_{1}^{A} - p_{1}^{B}$	$p_{2}^{A}-p_{2}^{B}$	 $p_K^A - p_K^B$

- Random variable and assumption: $P_1^A, \dots, P_N^A, P_1^B, \dots, P_N^B$

 - $P_i^A P_i^B$ i.i.d. $P_i^A P_i^B \sim \mathcal{N}\left(\mu_{A-B}, \sigma_{A-B}^2\right)$ with unknown standard deviation
- Parameter of interest: μ_{A-B}
- Parameter estimate: $m_{A-B} = \frac{1}{\kappa} \sum_{i=1}^{K} (p_i^A p_i^B)$
- Hypotheses H₀ and H_A:

$$H_0: \mu_{A-B} = 0$$

$$H_A: \mu_{A-B} \neq 0$$

- Test statistic: $t = \frac{m_{A-B}}{s_{A-B}/\sqrt{K}}$, where $s_{A-B} = \sqrt{\frac{1}{K-1}\sum_{i=1}^{K}\left(p_i^A p_i^B m_{A-B}\right)^2}$
- Hypothesis test: paired t-test





Training-validation split and leave-one-out cross validation

- Classifiers: A and B
- Data: classifiers A and B tested on the validation data; the outcome x_i^A and x_i^B can be either correct (0) or incorrect (1); nominal categorical data with two categories correct or incorrect

	classifier A correct	classifier A incorrect
classifier B correct	n_{00} =count(A correct, B correct)	n_{10} =count(A incorrect, B correct)
classifier B incorrect	n_{01} =count(A correct, B incorrect)	n_{11} =count(A incorrect, B incorrect)

- Random variable and assumption: $X_i^A \sim Bernoulli(p_A), X_i^B \sim Bernoulli(p_B)$
- Test statistic:
 - Small discordance (e.g. $n_{01} + n_{10} < 25$): $s_0 = \min(n_{01}, n_{10})$
 - Large discordance: $s_0 = \frac{(|n_{01} n_{10}| 1)^2}{n_{01} + n_{02}}$
- Hypothesis test: McNemar's test
- Note: why McNemar's test? One consideration while developing hypothesis test is to have a reasonably low type I error





Exercise 4

- You have a labeled data set $\mathcal{D} = \{(x_1, y_1), \cdots, (x_N, y_N)\}$
- You developed two classifiers using the 10-fold validation
- Construct a table of the resulting F1 scores for these two classifiers
- Design a hypothesis test to compare these two classifiers



Exercise 5

- You have a labeled data set $\mathcal{D} = \{(x_1, y_1), \cdots, (x_N, y_N)\}$
- You developed two classifiers using the training-validation split
- Construct a table of the results for these two classifiers
- Design a hypothesis test to compare these two classifiers



Today

- p-hacking
- 2 Test statistics and hypothesis tests
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- Summary

What is A/B testing?

- A/B testing refers to the methodology for testing two variants of the same type of product
- It is widely used for improving marketing strategies
- Example:
 - Web design: a company would like to increase the number of their subscribers by testing a new design for the "subscribe" button on their web page
 - Product pricing: a company would like to introduce a new product to the market; they compare two potential prices to see which one maximizes their sales while maintaining a reasonable margin
- We will revisit this topic next week as part of the exercise



Today

- p-hacking
- 2 Test statistics and hypothesis tests
- Compare two classifiers
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- Summary

Summary

So far:

- Data types and data containers
- Descriptive data analysis: descriptive statistics, visualization
- Probability distributions, events, random variables, PMF, PDF, parameters
- CDF, Q-Q plot, how to compare two distributions (data vs theoretical, data vs data)
- Modeling
- Parameter estimation: maximum likelihood estimation (MLE) and maximum a posteriori estimation (MAP)
- Classification, multinomial naive Bayes classifier, Gaussian naive Bayes classifier
- Central limit theorem, interval estimation
- Hypothesis tests, comparison of two classifiers





Summary

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Next:

• Unsupervised learning, clustering, k-means, Gaussian Mixture Models





Summary

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- Hypothesis tests, comparison of two classifiers

Next:

Unsupervised learning, clustering, k-means, Gaussian Mixture Models

Before next lecture:

- Gaussian distribution
- The Bayes' rule







I heard my name again!