

UCLA STAT 35
Applied Computational and Interactive Probability

Instructor: Ivo Dinov,
 Asst. Prof. In Statistics and Neurology

Teaching Assistant: Chris Barr

University of California, Los Angeles, Winter 2006

<http://www.stat.ucla.edu/~dinov/>

Slide 1 UCLA, Ivo Dinov

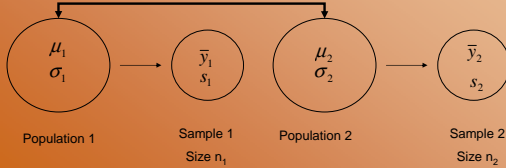
Comparison of Two Independent Samples

Slide 2 UCLA, Ivo Dinov

Comparison of Two Independent Samples

- Many times in the sciences it is useful to compare two groups
 - Male vs. Female
 - Drug vs. Placebo
 - NC vs. Disease

Q: Different?



Slide 3 UCLA, Ivo Dinov

Comparison of Two Independent Samples

- Two Approaches for Comparison
 - Confidence Intervals
 - we already know something about CI's
 - Hypothesis Testing
 - this will be new
- What seems like a reasonable way to compare two groups?
- What parameter are we trying to estimate?

Slide 4 UCLA, Ivo Dinov

Comparison of Two Independent Samples

- RECALL: The sampling distribution of \bar{y} was centered at μ , and had a standard deviation of $\frac{\sigma}{\sqrt{n}}$

- We'll start by describing the sampling distribution of $\bar{y}_1 - \bar{y}_2$

- Mean: $\mu_1 - \mu_2$

- Standard deviation of $\sqrt{\frac{\sigma_1^2}{n_1} + \frac{\sigma_2^2}{n_2}}$

- What seems like appropriate estimates for these quantities?

Slide 5 UCLA, Ivo Dinov

Standard Error of $\bar{y}_1 - \bar{y}_2$

- We know $\bar{y}_1 - \bar{y}_2$ estimates $\mu_1 - \mu_2$
- What we need to describe next is the precision of our estimate, $SE_{(\bar{y}_1 - \bar{y}_2)}$

$$SE_{(\bar{y}_1 - \bar{y}_2)} = \sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}} = \sqrt{SE_1^2 + SE_2^2}$$

Slide 6 UCLA, Ivo Dinov

Standard Error of $\bar{y}_1 - \bar{y}_2$

Example: A study is conducted to quantify the benefits of a new cholesterol lowering medication. Two groups of subjects are compared, those who took the medication twice a day for 3 years, and those who took a placebo. Assume subjects were randomly assigned to either group and that both groups data are normally distributed. Results from the study are shown below:

	Medication	Placebo
\bar{y}	209.8	224.3
n	10	10
s	44.3	46.2
SE	14.0	14.6

Slide 7 UCLA, Jon Dinnar

Standard Error of $\bar{y}_1 - \bar{y}_2$

Example: Cholesterol medicine (cont')

(e.g., <http://ftp.nist.gov/pub/dataplot/other/reference/CHOLEST2.DAT>)

Calculate an estimate of the true mean difference between treatment groups and this estimate's precision.

■ First, denote medication as group 1 and placebo as group 2

$$(\bar{y}_1 - \bar{y}_2) = 209.8 - 224.3 = -14.5$$

	Medication	Placebo
\bar{y}	209.8	224.3
n	10	10
s	44.3	46.2
SE	14.0	14.6

$$SE_{(\bar{y}_1 - \bar{y}_2)} = \sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}} = \sqrt{\frac{44.3^2}{10} + \frac{46.2^2}{10}} = 20.24$$

Slide 8 UCLA, Jon Dinnar

Pooled vs. Unpooled

- $\sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}}$ is known as an **unpooled** version of the

standard error

- there is also a "pooled" SE

- First we describe a "pooled" variance, which can be thought of as a weighted average of s_1^2 and s_2^2

$$s_{pooled}^2 = \frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n_1 + n_2 - 2}$$

Slide 9 UCLA, Jon Dinnar

Pooled vs. Unpooled

- Then we use the pooled variance to calculate the pooled version of the standard error

$$SE_{pooled} = \sqrt{s_{pooled}^2 \left(\frac{1}{n_1} + \frac{1}{n_2} \right)}$$

- NOTE: If $(n_1 = n_2)$ and $(s_1 = s_2)$ the pooled and unpooled will give the same answer for $SE_{(\bar{y}_1 - \bar{y}_2)}$

- It is when $n_1 \neq n_2$ that we need to decide whether to use pooled or unpooled:

- if $s_1 = s_2$ then use pooled (unpooled will give similar answer)
- if $s_1 \neq s_2$ then use unpooled (pooled will NOT give similar answer)

Slide 10 UCLA, Jon Dinnar

Pooled vs. Unpooled

- RESULT: Because both methods are similar when $s_1 = s_2$ and $n_1 = n_2$, and the pooled version is not valid when

- Why all the torture? This will come up again in chapter 11.

- Because the **df** increases a great deal when we do pool the variance.

Slide 11 UCLA, Jon Dinnar

CI for $\mu_1 - \mu_2$

- RECALL: We described a CI earlier as: the estimate \pm (an appropriate multiplier) \times (SE)

- A $100(1 - \alpha)\%$ confidence interval for $\mu_1 - \mu_2$

$$(p.227) \quad (\bar{y}_1 - \bar{y}_2) \pm t(df)_{\alpha/2} (SE_{\bar{y}_1 - \bar{y}_2})$$

$$\text{where } df = \frac{(SE_1^2 + SE_2^2)^2}{SE_1^4/(n_1 - 1) + SE_2^4/(n_2 - 1)}$$

Slide 12 UCLA, Jon Dinnar

CI for $\mu_1 - \mu_2$

Example: Cholesterol medication (cont')

Calculate a 95% confidence interval for $\mu_1 - \mu_2$

We know $\bar{y}_1 - \bar{y}_2$ and $SE_{(\bar{y}_1 - \bar{y}_2)}$ from the previous slides.
Now we need to find the t multiplier

$$df = \frac{(14^2 + 14.6^2)^2}{14^4 / (10-1) + 14.6^4 / (10-1)} = \frac{167411.9056}{9317.021} = 17.97 \approx 17$$

Round down to be conservative

NOTE: Calculating that df is not really that fun, a quick rule of thumb for checking your work is:

$$n_1 + n_2 - 2$$

Slide 13

UCLA, Jon Dineen

CI for $\mu_1 - \mu_2$

$$\begin{aligned} & (\bar{y}_1 - \bar{y}_2) \pm t(df)_{\alpha/2} (SE_{\bar{y}_1 - \bar{y}_2}) \\ & = -14.5 \pm t(17)_{0.025} (20.24) \\ & = -14.5 \pm 2.110 (20.24) \\ & = (-57.21, 28.21) \end{aligned}$$

CONCLUSION: We are highly confident at the 0.05 level, that the true mean difference in cholesterol between the medication and placebo groups is between -57.02 and 28.02 mg/dL.

Note the change in the conclusion of the parameter that we are estimating. Still looking for the 5 basic parts of a CI conclusion (see slide 38 of lecture set 5).

Slide 14

UCLA, Jon Dineen

CI for $\mu_1 - \mu_2$

- What's so great about this type of confidence interval?
- In the previous example our CI contained zero
 - This interval isn't telling us much because:
 - the true mean difference could be more than zero (in which case the mean of group 1 is larger than the mean of group 2)
 - or the true mean difference could be less than zero (in which case the mean of group 1 is smaller than the mean of group 2)
 - or the true mean difference could even be zero!
 - The ZERO RULE!
 - Suppose the CI came out to be (5.2, 28.1), would this indicate a true mean difference?

Slide 15

UCLA, Jon Dineen

Hypothesis Testing: The independent t test

- The idea of a hypothesis test is to formulate a hypothesis that nothing is going on and then to see if collected data is consistent with this hypothesis (or if the data shows something different)
 - Like innocent until proven guilty
- There are four main parts to a hypothesis test:
 - hypotheses
 - test statistic
 - p-value
 - conclusion

Slide 16

UCLA, Jon Dineen

Hypothesis Testing: #1 The Hypotheses

- There are two hypotheses:
 - Null hypothesis (aka the "status quo" hypothesis)
 - denoted by H_0
 - Alternative hypothesis (aka the research hypothesis)
 - denoted by H_a

Slide 17

UCLA, Jon Dineen

Hypothesis Testing: #1 The Hypotheses

- If we are comparing two group means nothing going on would imply no difference
 - the means are "the same"

$$(\mu_1 - \mu_2) = 0$$
- For the independent t-test the hypotheses are:
 - $H_0: (\mu_1 - \mu_2) = 0$
(no statistical difference in the population means)
 - $H_a: (\mu_1 - \mu_2) \neq 0$
(a statistical difference in the population means)

Slide 18

UCLA, Jon Dineen

Hypothesis Testing: #1 The Hypotheses

Example: Cholesterol medication (cont')

Suppose we want to carry out a hypothesis test to see if the data show that there is enough evidence to support a difference in treatment means.

Find the appropriate null and alternative hypotheses.

$$H_0: (\mu_1 - \mu_2) = 0$$

(no statistical difference the true means of the medication and placebo groups)

$$H_a: (\mu_1 - \mu_2) \neq 0$$

(a statistical difference in the true means of the medication and placebo groups, medication has an effect on cholesterol)

Slide 19

UCLA, Jon Dineen

Hypothesis Testing: #2 Test Statistic

● A test statistic is calculated from the sample data

■ it measures the "disagreement" between the data and the null hypothesis

□ if there is a lot of "disagreement" then we would think that the data provide evidence that the null hypothesis is false

□ if there is little to no "disagreement" then we would think that the data do not provide evidence that the null hypothesis is false

$$t_s = \frac{(\bar{y}_1 - \bar{y}_2) - 0}{SE_{\bar{y}_1 - \bar{y}_2}}$$

subtract 0 because the null says the difference is zero

Slide 20

UCLA, Jon Dineen

Hypothesis Testing: #2 Test Statistic

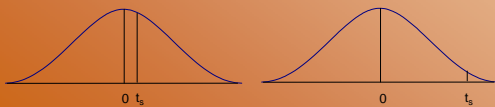
● On a t distribution t_s could fall anywhere

■ If the test statistic is close to 0, this shows that the data are compatible with H_0 (no difference)

□ the deviation can be attributed to chance

■ If the test statistic is far from 0 (in the tails, upper or lower), this shows that the data are incompatible to H_0 (there is a difference)

□ deviation does not appear to be attributed to chance (ie. If H_0 is true then it is unlikely that t_s would fall so far from 0)



Slide 21

UCLA, Jon Dineen

Hypothesis Testing: #2 Test Statistic

Example: Cholesterol medication (cont')

Calculate the test statistic

$$t_s = \frac{(\bar{y}_1 - \bar{y}_2) - 0}{SE_{\bar{y}_1 - \bar{y}_2}} = \frac{(209.8 - 224.3) - 0}{20.24} = -0.716$$

■ Great, what does this mean?

□ \bar{y}_1 and \bar{y}_2 differ by about 0.72 SE's

□ this is because t_s is the measure of difference between the sample means expressed in terms of the SE of the difference

Slide 22

UCLA, Jon Dineen

Hypothesis Testing: #2 Test Statistic

● How do we use this information to decide if the data support H_0 ?

■ Perfect agreement between the means would indicate that $t_s = 0$, but logically we expect the means do differ by at least a little bit.

□ The question is how much difference is statistically significant?

■ If H_0 is true, it is unlikely that t_s would fall in either of the far tails

■ If H_0 is false it is unlikely that t_s would fall near 0

Slide 23

UCLA, Jon Dineen

Hypothesis Testing: #3 P-value

● How far is far?

● For a two tailed test (i.e. $H_a: (\mu_1 - \mu_2) \neq 0$) The p-value of the test is the area under the Student's T distribution in the double tails beyond $-t_s$ and t_s .



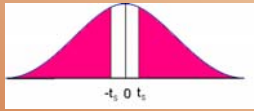
■ Definition (p. 238): The p-value for a hypothesis test is the probability, computed under the condition that the null hypothesis is true, of the test statistic being at least as extreme or more extreme as the value of the test statistic that was actually obtained [from the data].

Slide 24

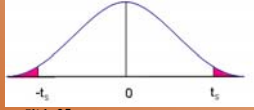
UCLA, Jon Dineen

Hypothesis Testing: #3 P-value

- What this means is that we can think of the p-value as a measure of compatibility between the data and H_0
 - a large p-value (close to 1) indicates that t_s is near the center (data support H_0)



- a small p-value (close to 0) indicates that t_s is in the tail (data do not support H_0)



Slide 25 UCLA, Jon Dineen

Hypothesis Testing: #3 P-value

- Where do we draw the line?
 - how small is small for a p-value?
- The threshold value on the p-value scale is called the significance level, and is denoted by α
 - The significance level is chosen by whomever is making the decision (BEFORE THE DATA ARE COLLECTED!)
 - Common values for α include 0.1, 0.05 and 0.01
- Rules for making a decision:
 - If $p \leq \alpha$ then reject H_0 (statistical significance)
 - If $p > \alpha$ then fail to reject H_0 (no statistical significance)

Slide 26 UCLA, Jon Dineen

Hypothesis Testing: #3 P-value

Example: Cholesterol medication (cont')

Find the p-value that corresponds to the results of the cholesterol lowering medication experiment
 We know from the previous slides that $t = -0.716$ (which is close to 0)
 This means that the p-value is the area under the curve beyond ± 0.716 with 18 df.

Slide 27 UCLA, Jon Dineen

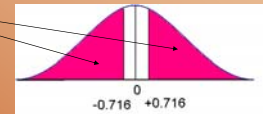
Hypothesis Testing: #3 P-value

Example: Cholesterol medication (cont')

Using SOCR we can find the area under the curve beyond ± 0.716 with 18 df to be:

$$p > 2(0.2) = 0.4$$

NOTE, when H_0 is \neq , the p-value is the area beyond the test statistic in BOTH tails.



Slide 28 UCLA, Jon Dineen

Hypothesis Testing: #4 Conclusion

Example: Cholesterol medication (cont')

Suppose the researchers had set $\alpha = 0.05$
 Our decision would be to fail to reject H_0 because $p > 0.05$ which is > 0.05
 (#4) CONCLUSION: Based on this data there is no statistically significant difference between true mean cholesterol of the medication and placebo groups ($p > 0.4$).
 □ In other words the cholesterol lowering medication does not seem to have a significant effect on cholesterol.
 ■ Keep in mind, we are saying that we couldn't provide sufficient evidence to show that there is a significant difference between the two *population* means.

Slide 29 UCLA, Jon Dineen

Hypothesis Testing Summary

- Important parts of Hypothesis test conclusions:
 1. Decision (significance or no significance)
 2. Parameter of interest
 3. Variable of interest
 4. Population under study
 5. (optional but preferred) P-value

Slide 30 UCLA, Jon Dineen

Was Cavendish's experiment biased?

A number of famous early experiments of measuring physical constants have later been shown to be biased.

Mean density of the earth

True value = 5.517

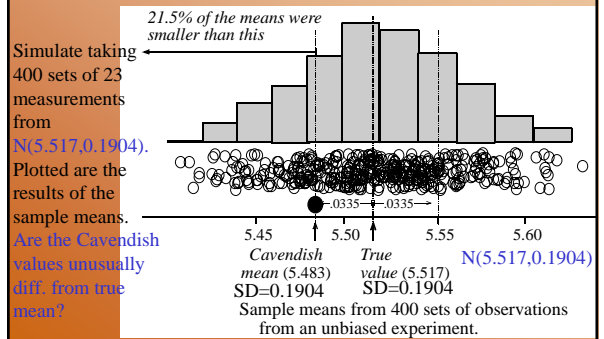
Cavendish's data: (from previous Example 7.2.2)

5.36, 5.29, 5.58, 5.65, 5.57, 5.53, 5.62, 5.29, 5.44, 5.34, 5.79, 5.10,
5.27, 5.39, 5.42, 5.47, 5.63, 5.34, 5.46, 5.30, 5.75, 5.68, 5.85

$n = 23$, sample mean = 5.483, sample SD = 0.1904

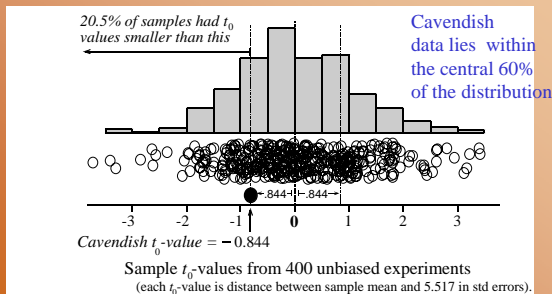
Slide 31 UCLA, Jon Dinger

Was Cavendish's experiment biased?

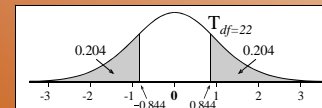
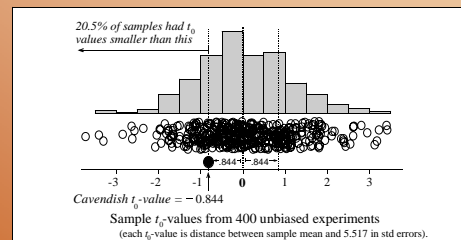


Slide 32 UCLA, Jon Dinger

Cavendish: measuring distances in std errors



Slide 33 UCLA, Jon Dinger



Slide 34 UCLA, Jon Dinger

Measuring the distance between the true-value and the estimate in terms of the SE

- Intuitive criterion: Estimate is credible if it's not **far away** from its hypothesized true-value!
- But how far is **far-away**?
- Compute the distance in standard-terms:

$$T = \frac{\text{Estimator} - \text{TrueParameterValue}}{\text{SE}}$$
- Reason is that the distribution of T is known in some cases (Student's t , or $N(0,1)$). The estimator (obs.-value) is **typical/atypical** if it is close to the **center/tail** of the distribution.

Slide 35 UCLA, Jon Dinger

Comparing CI's and significance tests

- These are **different methods** for coping with the **uncertainty** about the true value of a parameter caused by the sampling variation in estimates.
- **Confidence interval:** A **fixed level of confidence** is chosen. We determine a **range of possible values** for the parameter that are consistent with the data (at the chosen confidence level).
- **Significance test:** **Only one possible value** for the parameter, called the **hypothesized value**, is tested. We determine the **strength of the evidence** (confidence) provided by the data against the proposition that the hypothesized value is the true value.

Slide 36 UCLA, Jon Dinger

Review

- What **intuitive criterion** did we use to determine whether the hypothesized parameter value ($p=0.2$ in the ESP Example 9.1.1, and $\mu = 5.517$ in Example 9.1.2) was **credible in the light of the data**? (Determine if the data-driven parameter estimate is consistent with the pattern of variation we'd expect get if hypothesis was true. If hypothesized value is correct, our estimate should not be far from its hypothesized true value.)
- Why was it that $\mu = 5.517$ was **credible** in Example 9.1.2, whereas $p=0.2$ was **not credible** in Example 9.1.1? (The first estimate is consistent, and the second one is not, with the pattern of variation of the hypothesized true process.)

Slide 37

UCLA, Jon Dineen

Review

- What do t_0 -values tell us? (Our estimate is typical/atypical, consistent or inconsistent with our hypothesis.)
- What is the essential difference between the information provided by a confidence interval (CI) and by a significance test (ST)? (Both are uncertainty quantifiers. CI's use a fixed level of confidence to determine possible range of values. ST's one possible value is fixed and level of confidence is determined.)

Slide 38

UCLA, Jon Dineen

Hypotheses

Guiding principles

We cannot rule in a hypothesized value for a parameter, we can only determine whether there is evidence to rule out a hypothesized value.

The null hypothesis tested is typically a skeptical reaction to a research hypothesis

Slide 39

UCLA, Jon Dineen

Comments

- Why can't we (**rule-in**) prove that a hypothesized value of a parameter is exactly true? (Because when constructing estimates based on data, there's always sampling and may be non-sampling errors, which are normal, and will effect the resulting estimate. Even if we do 60,000 ESP tests, as we saw earlier, repeatedly we are likely to get estimates like 0.2 and 0.200001, and 0.1999999, etc. – non of which may be exactly the theoretically correct, 0.2.)
- Why use the rule-out principle? (Since, we can't use the rule-in method, we try to find compelling evidence against the observed/data-constructed estimate – to reject it.)
- Why is the null hypothesis & significance testing typically used? (H_0 : skeptical reaction to a research hypothesis; ST is used to check if differences or effects seen in the data can be explained simply in terms of sampling variation!)

Slide 40

UCLA, Jon Dineen

Comments

- How can researchers try to demonstrate that effects or differences seen in their data are real? (Reject the hypothesis that there are no effects)
- How does the alternative hypothesis typically relate to a belief, hunch, or research hypothesis that initiates a study? ($H_1=H_a$: specifies the type of departure from the null hypothesis, H_0 (skeptical reaction), which we are expecting (research hypothesis itself).
- In the Cavendish's mean Earth density data, null hypothesis was $H_0 : \mu = 5.517$. We suspected bias, but not bias in any specific direction, hence $H_a : \mu \neq 5.517$.

Slide 41

UCLA, Jon Dineen

Comments

- In the ESP Pratt & Woodruff data, (skeptical reaction) null hypothesis was $H_0 : \mu = 0.2$ (**pure-guessing**). We suspected bias, toward success rate being higher than that, hence the (research hypothesis) $H_a : \mu > 0.2$.
- Other commonly encountered situations are:
 - $H_0 : \mu_1 - \mu_2 = 0 \rightarrow H_a : \mu_1 - \mu_2 > 0$
 - $H_0 : \mu_{rest} - \mu_{activation} = 0 \rightarrow H_a : \mu_{rest} - \mu_{activation} \neq 0$

Slide 42

UCLA, Jon Dineen

The t-test

Using $\hat{\theta}$ to test $H_0: \theta = \theta_0$ versus some alternative H_1 .

STEP 1 Calculate the *test statistic*,

$$t_0 = \frac{\hat{\theta} - \theta_0}{sd(\hat{\theta})} = \frac{\text{estimate} - \text{hypothesized value}}{\text{standard error}}$$

[This tells us how many standard errors the estimate is above the hypothesized value (t_0 positive) or below the hypothesized value (t_0 negative).]

STEP 2 Calculate the *P-value* using the following table.

STEP 3 Interpret the *P-value* in the context of the data.

Slide 43 UCLA, Jon Dinnis

The t-test

Alternative hypothesis	Evidence against $H_0: \theta > \theta_0$ provided by	P-value
$H_1: \theta > \theta_0$	$\hat{\theta}$ too much bigger than θ_0 (i.e., $\hat{\theta} - \theta_0$ too large)	$P = \text{pr}(T \geq t_0)$
$H_1: \theta < \theta_0$	$\hat{\theta}$ too much smaller than θ_0 (i.e., $\hat{\theta} - \theta_0$ too negative)	$P = \text{pr}(T \leq t_0)$
$H_1: \theta \neq \theta_0$	$\hat{\theta}$ too far from θ_0 (i.e., $ \hat{\theta} - \theta_0 $ too large)	$P = 2 \text{pr}(T \geq t_0)$

where $T \sim \text{Student}(df)$

Slide 44 UCLA, Jon Dinnis


Interpretation of the p-value

TABLE 9.3.2 Interpreting the Size of a P-Value

Approximate size of P-Value	Translation
> 0.12 (12%)	No evidence against H_0
0.10 (10%)	Weak evidence against H_0
0.05 (5%)	Some evidence against H_0
0.01 (1%)	Strong evidence against H_0
0.001 (0.1%)	Very Strong evidence against H_0

Slide 45 UCLA, Jon Dinnis

Is a second child gender influenced by the gender of the first child, in families with >1 kid?



First Child	Second Child		Total
	Male	Female	
Male	3,202	2,776	5,978
Female	2,620	2,792	5,412
Total	5,822	5,568	11,390

- Research hypothesis needs to be formulated first before collecting/looking/interpreting the data that will be used to address it. Mothers whose 1st child is a girl are more likely to have a girl, as a second child, compared to mothers with boys as 1st children.
- Data: 20 yrs of birth records of 1 Hospital in Auckland, NZ.

Slide 46 UCLA, Jon Dinnis

Analysis of the birth-gender data – data summary

Group	Second Child	
	Number of births	Number of girls
1 (Previous child was girl)	5412	2792 (approx. 51.6%)
2 (Previous child was boy)	5978	2776 (approx. 46.4%)

- Let p_1 =true proportion of girls in mothers with girl as first child, p_2 =true proportion of girls in mothers with boy as first child. Parameter of interest is $p_1 - p_2$.
- $H_0: p_1 - p_2 = 0$ (skeptical reaction). $H_a: p_1 - p_2 > 0$ (research hypothesis)

Slide 47 UCLA, Jon Dinnis

Hypothesis testing as decision making

Decision made	Actual situation	
	H_0 is true	H_0 is false
Accept H_0 as true	OK	Type II error
Reject H_0 as false	Type I error	OK

- Sample sizes: $n_1=5412$, $n_2=5978$, Sample proportions (estimates) $\hat{p}_1 = 2792/5412 \approx 0.5159$, $\hat{p}_2 = 2776/5978 \approx 0.4644$.
- $H_0: p_1 - p_2 = 0$ (skeptical reaction). $H_a: p_1 - p_2 > 0$ (research hypothesis)

Slide 48 UCLA, Jon Dinnis

Analysis of the birth-gender data

- Samples are large enough to use **Normal-approx.**. Since the two proportions come from totally diff. mothers they are **independent** → use formula 8.5.5.a

$$t_0 = \frac{\text{Estimate} - \text{Hypothesized Value}}{SE} = 5.49986 =$$

$$\frac{\hat{p}_1 - \hat{p}_2 - 0}{SE(\hat{p}_1 - \hat{p}_2)} = \frac{\hat{p}_1 - \hat{p}_2}{\sqrt{\frac{\hat{p}_1(1-\hat{p}_1)}{n_1} + \frac{\hat{p}_2(1-\hat{p}_2)}{n_2}}} =$$

$$P\text{-value} = \Pr(T \geq t_0) = 1.9 \times 10^{-8}$$

Slide 49 UCLA, Jon Dineen

Analysis of the birth-gender data

- We have strong evidence to reject the H_0 , and hence conclude mothers with first child a girl a **more likely** to have a girl as a second child.

- How much more likely? **A 95% CI:**

CI $(p_1 - p_2) = [0.033; 0.070]$. And computed by:

$$\text{estimate} \pm z \times SE = \hat{p}_1 - \hat{p}_2 \pm 1.96 \times SE(\hat{p}_1 - \hat{p}_2) =$$

$$\hat{p}_1 - \hat{p}_2 \pm 1.96 \times \sqrt{\frac{\hat{p}_1(1-\hat{p}_1)}{n_1} + \frac{\hat{p}_2(1-\hat{p}_2)}{n_2}} =$$

$$0.0515 \pm 1.96 \times 0.0093677 = [3\%; 7\%]$$

Slide 50 UCLA, Jon Dineen