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Section B

Two Sample z-test: Getting a p-value

Hypothesis Test to Compare Two Proportions

- Two sample z-test
- Are the proportions of infants contracting HIV within 18 months-ofbirth equivalent at the population level for those whose mothers are treated with AZT versus untreated (placebo)?
 - $H_0: p_1 = p_2$
 - $H_A: p_1 \neq p_2$
- In other words, is the expected difference in proportions zero?
 - H_0 : $p_1 p_2 = 0$
 - H_A : $p_1 p_2 \neq 0$

Hypothesis Test to Compare Two Independent Groups

- Recall, general "recipe" for hypothesis testing . . .
 - 1. Start by assuming H_o true
 - 2. Measure distance of sample result from μ_0 (here again its 0)
 - 3. Compare test statistic (distance) to appropriate distribution to get p-value

$$z = \frac{(observed dif f) - (null dif f)}{SE of observed dif f erence}$$

$$z = \frac{\hat{p}_1 - \hat{p}_2}{S\hat{E}(\hat{p}_1 - \hat{p}_2)} = \frac{\hat{p}_1 - \hat{p}_2}{\sqrt{\frac{\hat{p}_1 \times (1 - \hat{p}_1)}{n_1} + \frac{\hat{p}_2 \times (1 - \hat{p}_2)}{n_2}}}$$

Infant HIV/ AZT Study

In the infant HIV/AZT study, recall:

$$\hat{p}_1 - \hat{p}_2 = -0.15$$

$$S\hat{E}(\hat{p}_1 - \hat{p}_2) = 0.036$$

So in this study:

$$z = \frac{-0.15}{0.36} \approx -4.2$$

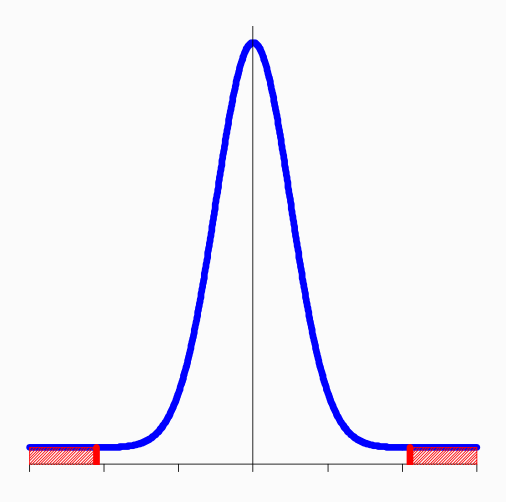
 So this study result was 4.2 standard errors below the null mean of 0 (i.e., 4.2 standard errors from the difference in the proportion of HIV+ infants between the AZT and placebo groups expected if null was true)

How Are p-values Calculated?

- Is a result 4.2 standard errors below 0 unusual?
 - It depends on what kind of distribution we are dealing with
- The p-value is the probability of getting a test statistic as (or more extreme than) what you observed (-4.2) by chance
- The p-value comes from the sampling distribution of the difference in two sample proportions
- What is the sampling distribution of the difference in sample means?
 - If both groups are large then this distribution is approximately normal
 - This sampling distribution will be centered at true difference,
 - Under null hypothesis, this true difference is 0

Diet/Weight Loss Sample

 To compute a p-value, we would need to compute the probability of being 4.2 or more standard errors away from 0 on a standard normal curve



AZT Study

- If we were to look this up on a normal table, we would find a very low p-value (p < .001)
- This method is also essentially equivalent to the chi-square $(\chi 2)$ method
 - Gives about the same answer (p-value)
 - This is how Stata approaches it
 - We will discuss chi-square method in more detail shortly: for now, just "take on faith" that it is equivalent so we can show you how to get the p-value, 95% CI (etc.) using Stata

To Do in Stata: Display Data in a 2x2 Table

- Stata "thinks" of data in a 2x2 (contingency) table
- Two rows and two columns

	Drug Group			
		AZT	Placebo	_
HIV Transmission	Yes	13	40	53
	No	167	143	310
		180	183	363

To Do in Stata: Display Data in a 2x2 Table

- We can get Stata to give us a 95% CI for the difference in proportions, and a p-value by using the csi command
- Syntax csi a b c d
 - Based a 2x2 table using our sample results as such

		Exposure		
		Yes	No	
Outcome	Yes	a	b	
	No	С	d	

csi 13 40 167 143

	Drug Group			
		AZT	Placebo	-
HIV Transmission	Yes	13	40	53
	No	167	143	310
		180	183	363

Results from *csi* command

. csi 13 40 167 143

	Exposed	Unexposed	Total	
Cases Noncases		40 143	53 310	
Total	180	183	363	
Risk	.0722222	.2185792	1 .1460055	
	Point	estimate	[95% Conf.	Interval]
Risk difference Risk ratio Prev. frac. ex. Prev. frac. pop	146357 .3304167 .6695833 .3320248		2171766 .1829884 .4033765	.5966235
1	- 	chi2(1) =	15.59 Pr>chi	2 = 0.0001

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Risk	.0722222	.2185792	.1460055	
	Point	estimate	[95% Conf.	Interval]
Risk difference		46357	2171766	
Risk ratio Prev. frac. ex. Prev. frac. pop	.6695833		'	.5966235 .8170116
_	C	:hi2(1) =	15.59 Pr>chi	2 = 0.0001

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-	(chi2(1) =	15.59 Pr>chi2	2 = 0.0001

Statistical method

"We conducted a randomized, double-blind, placebo-controlled trial of the efficacy and safety of zidovudine (AZT) in reducing the risk of maternal-infant HIV transmission"

Statistical method

- The proportion of infants diagnosed as HIV positive within 18 months of birth was compared between the AZT and placebo groups using a two-sample z-test of proportions
- 95% confidence intervals were computed for the 18-month infection proportion in each group and for the difference in proportions between both groups

Results

- The proportion of infants who tested positive for HIV within 18 months of birth was seven percent (95% CI 4 -12%) in the AZT group and twenty-two percent in the placebo group (95% CI 16 28%)
- This difference is statistically significant (p < .001)

Results

The study results estimate the decrease in the proportion of HIV positive infants born to HIV positive mothers associated with AZT to be as low as 8% and as high as 22%