Take Home Exercise 1

Replicating an Experimental Study (THE 1)

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options(repos = c(CRAN = "https://cloud.r-project.org")) # set CRAN mirror

Exercise 1: Summary of the Paper and Main Findings Provide a brief summary of the paper you are replicating. Describe the main findings, especially those related to Study 1.

Disclaimer! For this exercise I made use of the following AI Generation tools: Deepl.com for translation purposes, the integrated Copilot tool in R Studio for generating code and text and the free versions of OpenAI's GPT-40 Mini, Anthropics Claude 3.5 Sonnet and Mistral's Codestral AI model for generating code and assisting me in solving the tasks.

The authors posit that according to the Homonationalist thesis, support for LGBT+ issues among anti-immigration nativists and right-wing populists is not entirely genuine (organic liberalism thesis) and is instead used strategically (instrumental liberalism thesis) as a tool to justify further exclusion of immigrants, especially Muslims, who are perceived to be more homophobic, than the average western citizen. The rational behind this could be summed up as: "The enemy of my enemy is my friend" or more drastically: "I hate them (Muslims) more than I hate the other group (LGBT+ people)". In order to test this hypothesis, the authors conducted two survey experiments in the UK and Spain. The dependent variable is "Support for LGBT+ inclusive education in school" and was measured by means of survey response on an order randomized scale from 0 to 11, but then for the sake of simplicity were turned into a dichotomous variable for support (0 or 1).

In the UK, for study 1, the experimental research design randomly assigned 1200 participants into a control group and a treatment group. Those in the control group were exposed to text and pictures of nondenominational people with conventional British names protesting against LGBT+ inclusive education in schools. The treatment group was exposed to the same pictures and text, but the names and pictures were changed to appear Muslim. The authors expected that individuals who harbor negative predispositions toward immigration in the treatment group, will display more socially liberal preferences toward LGBT+ education than individuals with comparable anti-immigrant sentiment in the control group. Indeed, the results showed that (with an average treatment effect of 9.7% points or a 22% increase against

the baseline) respondents with an anti-immigrant disposition in the treatment group showed increased support for LGBT+ inclusive education in schools than comparable individuals in the control group. On the other hand, among those with positively predisposed toward immigration, treatment exhibits no significant effect below the 5% significance level. But overall, it can be said, that support for LGBT+ inclusive education is positively correlated with liberal attitudes towards immigrants.

In Study 2, conducted in Spain, chosen because of generally more positive attitude levels towards immigration and LGBT+ issues than in the UK, and a right-wing populist party (VOX), that unlike its counter-part in the UK (UKIP), is not following a homonationalist strategy and is more openly outspoken against LGBT+ issues. They found that regardless of immigration sentiments, treatment resulted in an 10-11% increase in support for LGBT+ inclusive education in schools. However, the effect was stronger among those with anti-immigrant sentiments (21% vs. 14%), when compared against the baseline. Furthermore, the authors also looked at shifts in citizens pride in "Western Culture". Only among individuals with nativist's views could they reasonably demonstrate increased levels of "pride in Western Culture" in the treatment group.

With these findings in mind, the authors argue that the homonationalist thesis is correct and that the increased levels of support for LGBT+ issues among the wider general public in western countries is not entirely genuine, but also in part the result of a strategic move by right-wing populists to deradicalize their image by defining themselves by "who they are not" rather than "who they are". They further speculate that as soon as the political context changes and LGBT+ rights are at stake (but absent of the perceived threat of "immigrants"), the support for LGBT+ issues among right-wing populists will decrease.

Exercise 2: Data Preparation and Exploration Use the data file study1_data.csv to begin the replication process. Identify and describe the experimental variables (i.e., treatment and immigration attitudes) and provide visualizations of their distribution. Then, select up to four covariates (e.g., gender, age, etc.) and plot their distribution too. If necessary, clean or transform variables. Document any changes.

From looking at the csv table for Study1 as well as looking at the study1.R script, one can deduce which variables are likely of interest for the replication. As a first step, I need to find the dependent variable (DV) support for LGBT+ inclusive education, which according to the text had a 0-11 range with a randomized scale order that was transformed to 0 and 1 for easier interpretation. Secondly, I need to find the experimental variables for treatment and immigration attitudes. The treatment variable concerns the exposure to Muslim names and pictures and is most likely operationalized as a binary variable (0,1). The variable for immigration attitudes is likely to be operationalized as an ordinal scale (0-10) or as a binary variable.

A look at the Csv table revealed the following variables as of interest for the replication:

• Treatment variable: outcome treat (categorical with NAs), treatment (ordinal)

- Immigration attitudes: imm_1, imm_2, (both ordinal) imm1mean (continuous, same value), immbelow, imm3 (both ordinal)
- Dependent Variable: support (ordinal), support2 (categorical)

By also looking at the study1.R script in the code chunk below, I can see that the following variables are of interest for the replication, while the rest can likely be ignored:

- Treatment variable: treatment is transformed twofold -> treat (converted to cat. variable for model transformation), treatnum (converted to numeric binary variable for graph)
- Immigration attitudes: immbelow (cat., converted to binary variable for graph), imm_1 (ordinal, 0-10 with 10 standing for maximal positive attitude towards immigrants)
- Dependent Variable: support (ordinary)

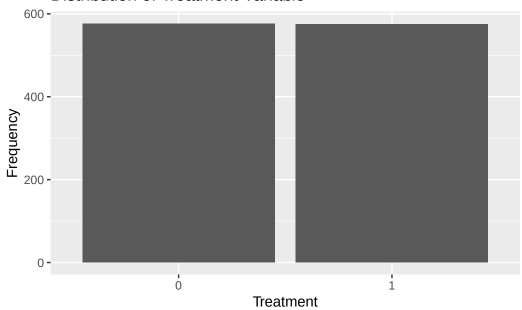
From this I conclude that the following variables are of interest for the replication:

- Treatment variable operationalization: treatment (0,1)
- Immigrant attitudes operationalization: imm_1 (ordinal scale 0-10) ((or immbelow, for binary distribution Pro-/AntiImmigration))
- Dependent Variable operationalization: support (0,1)

Below I will provide visualizations of the distribution of these variables. Let's start with the treatment variable. But first, I need to load the data sets and the necessary packages. I will then transform the treatment variable for better visualization and provide a table and a bar plot of the distribution of the treatment variable.

```
study1 <-
  read.csv("/Users/nicolaswaser/New-project-GitHub-first/R/Take-Home-Ex.-1/Input Data/study1
### Please insert your own data path to run the code correctly ##
library(tidyverse)
-- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
v dplyr 1.1.4 v readr
                                    2.1.5
v forcats 1.0.0 v stringr 1.5.1
v ggplot2 3.5.1 v tibble 3.2.1
v lubridate 1.9.3 v tidyr 1.3.1
           1.0.2
v purrr
-- Conflicts ----- tidyverse conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()
                 masks stats::lag()
i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become
library(ggplot2)
#is.factor(study1$treatment)
study1 <-study1%>%
  mutate(treatment= as.factor(treatment))
#is.factor(study1$treatment)
Variable distribution of "treatment" in the form of a table and a bar plot:
table(study1$treatment)
  0 1
576 575
```

Distribution of Treatment Variable



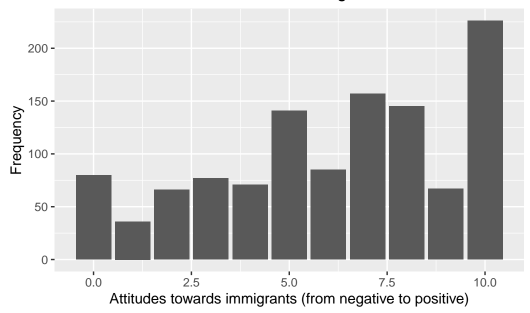
0 = control group, 1 = treatment group Unsurprisingly, the treatment variable is distributed evenly across the two groups, as it should be in a randomized experiment.

Next, I will visualize the distribution of the Immigration attitude variable "imm_1"::

```
table(study1$imm_1)
```

```
0 1 2 3 4 5 6 7 8 9 10
80 36 66 77 71 141 85 157 145 67 226
```

Distribution of attitudes towards immigrants



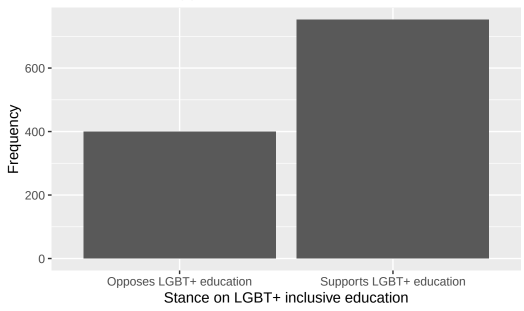
Ordinal scale: 0 = most negative attitude towards immigrants, 1 = most positive attitude towards immigrants The distribution of attitudes towards immigrants is skewed towards the positive side, with more respondents having a positive attitude.

Lastly, I will visualize the distribution of the dependent variable "support for LGBT+ inclusive education":

table(study1\$support2)

Opposes LGBT+ education Supports LGBT+ education 399 752

Distribution of support for LGBT+ inclusive education



The distribution of support for LGBT+ inclusive education is skewed towards the positive side, with most respondents supporting LGBT+ inclusive education.

Selecting covariates - I pick the following covariates for the analysis: - age: Age is often associated with a change in political attitudes over time -> age (numeric) or agecat (categorical) -> I pick age so that I have a continuous variable in the analysis

- gender: Gender is associated with different attitudes towards LGBT+ rights and immigration and could impact that relationship as well -> I pick the binary variable gender (categorical, Man, Woman)
- sexual identity/orientation: I expect that identifying as LGBT+ will have a big influence on the relationship of the experimental variables and the DV. -> I pick the variable queer (categorical, Heterosexual, Not heterosexual)
- race: not being white (in the case of the UK) might also impact the relationship between IV's and the DV. I also wanted to add a non-binary categorical variable -> I pick the variable race.

Age:

table(study1\$agecat)

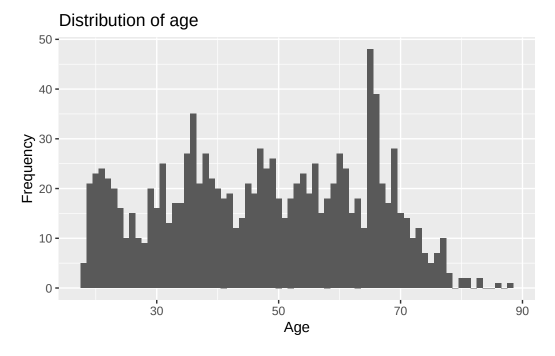
18-24 25-34 35-44 45-54 55-64 65+

```
3 131 152 215 212 194 244
```

Data Cleaning: There are 3 Nas in the data frame that affect all the covariates I selected. I will remove from the data set:

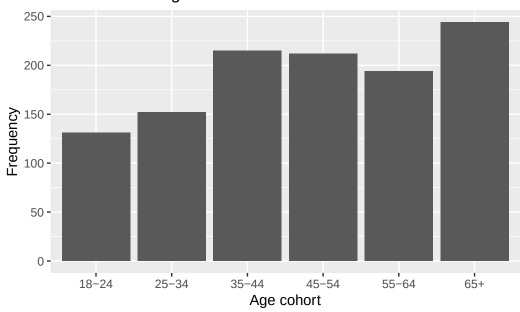
```
nas_by_column <-
  colSums(is.na(
    study1[, c("age", "agecat", "gender", "queer", "race")]))
#print(nas_by_column)
rows_with_nas <-
  which(rowSums(is.na(
    study1[, c("age", "agecat", "gender", "queer")])) > 0)
#print(rows_with_nas)
study1_cleaned <- study1[-rows_with_nas,]
#print(nrow(study1))
#print(nrow(study1_cleaned))</pre>
```

```
ggplot(study1_cleaned, aes(x=age)) +
  geom_histogram(binwidth = 1) + scale_x_continuous() +
  labs(title="Distribution of age", x="Age", y="Frequency")
```



Also looking at age cohorts:

Distribution of age cohorts



The distribution of age is skewed towards older respondents, with most respondents being in their 30s and above.

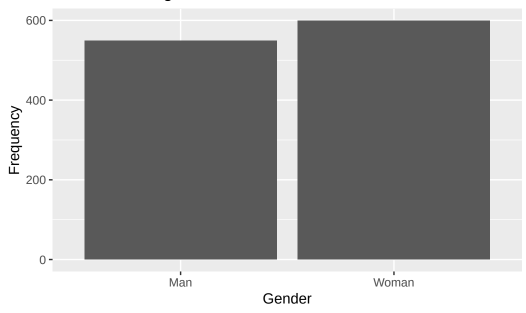
Gender:

```
table(study1_cleaned$gender)
```

Man Woman 549 599

```
ggplot(study1_cleaned, aes(x=gender)) + geom_bar() +
labs(title="Distribution of gender", x="Gender", y="Frequency")
```

Distribution of gender



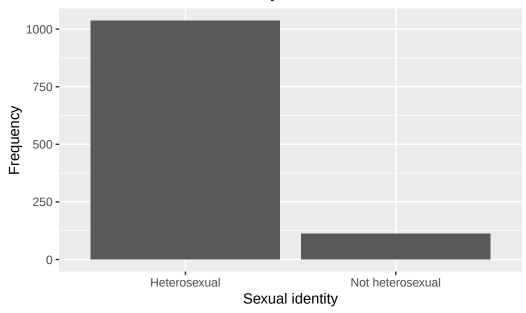
There are slightly more Men than Women in the sample, but overall they are distributed evenly.

Sexual Identity:

```
table(study1_cleaned$queer)
```

Heterosexual Not heterosexual 1036 112

Distribution of sexual identity



The large majority of respondents identify as heterosexual, with only a small minority identifying as not heterosexual.

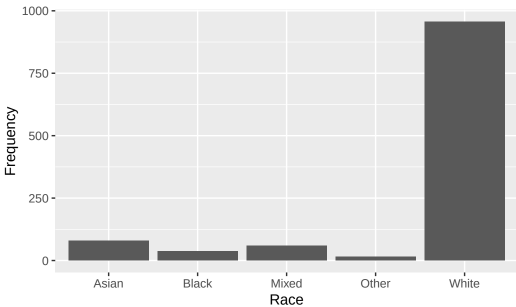
Race:

```
table(study1_cleaned$race)
```

```
Asian Black Mixed Other White 80 38 59 15 956
```

```
ggplot(study1_cleaned, aes(x=race)) + geom_bar() +
labs(title="Distribution of different races", x="Race", y="Frequency")
```





Overwhelming majority of respondents identify as white, with the rest distributing evenly.

Exercise 3: Covariate Balance Check for balance across covariates and report the results in a table and a plot. Explain your findings. Why would you expect randomization to lead to balance across covariates?

```
study1_cleaned_transformed <-study1_cleaned%>%
mutate(treatment= as.factor(treatment))
```

Checking for balance across covariates:

```
install.packages("dplyr")
```

The downloaded binary packages are in /var/folders/3k/4r5z06ts7bl41nkd_6d0dcyh0000gn/T//Rtmp60PFFi/downloaded_packages

```
library(dplyr)
install.packages("ggplot2")
```

The downloaded binary packages are in /var/folders/3k/4r5z06ts7bl41nkd_6d0dcyh0000gn/T//Rtmp60PFFi/downloaded_packages

```
library(ggplot2)
install.packages("tableone")
```

The downloaded binary packages are in /var/folders/3k/4r5z06ts7b141nkd_6d0dcyh0000gn/T//Rtmp60PFFi/downloaded_packages

```
library(tableone)
install.packages("knitr")
```

The downloaded binary packages are in /var/folders/3k/4r5z06ts7bl41nkd_6d0dcyh0000gn/T//Rtmp60PFFi/downloaded_packages

```
library(knitr)
treatment_var <- "treatment"
covariates <- c("gender", "age", "queer", "race")
balance_table <- CreateTableOne(vars = covariates, strata = treatment_var, data = study1_cleaterint(balance_table, smd = TRUE)  # SMD = standardized mean differences</pre>
```

Stratified by treatment 0 1 test SMD p 574 574 gender = Woman (%) 299 (52.1) 300 (52.3) 1.000 0.003 age (mean (SD)) 48.06 (15.95) 46.98 (16.85) 0.265 0.066 queer = Not heterosexual (%) 52 (9.1) 60 (10.5) 0.486 0.047 race (%) 0.904 0.060 Asian 40 (7.0) 40 (7.0) 21 (3.7) Black 17 (3.0) Mixed 29 (5.1) 30 (5.2) Other 6 (1.0) 9 (1.6) 478 (83.3) White 478 (83.3)

Treatment group outcome statistically distinct at: * p < 0.1, ** p < 0.05, *** p < 0.01 The sample size in each treatment group is n = 574 participants per group.

Distribution of covariates across treatment groups: - gender (Woman): In terms of gender distribution the sample is quite balanced with women accounting for about 52% in both treatment groups - age: The mean age is slightly different with 48.06 years for group 0 and 46.98 years for group 1, but this difference is minor. - queer (Not heterosexual): The proportion

of participants identifying as "Not heterosexual" is 9.1% in group 0 and 10.5% in group 1 and therefore quite well balanced. - Distribution of race across treatment groups is consistent: - Asian: 7.0% in each group - Black: 3.7% in group 0 and 3.0% in group 1 - Mixed: 5.1% in group 0 and 5.2% in group 1 - Other: 1.0% in group 0 and 1.6% in group 1 - White: 83.3% in both groups Although there are small variations in the counts of some racial groups, these differences are not statistically significant.

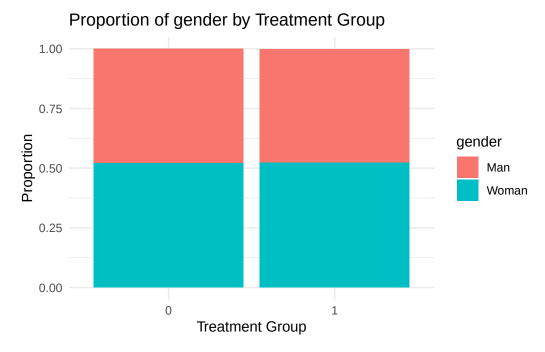
All SMD Values are well below 0.1, which typically indicates that covariates are well-balanced between groups. Also, none of the p-values are significant, meaning there are no statistically significant differences in these covariates between groups, which further confirms balance.

Why Randomization Should Lead to Balance: In randomized experiments, each participant has an equal probability of being assigned to any treatment group. This process should, on average, distribute participants with different covariate values equally across treatment groups. Since the distribution of covariates is similar, any observed effects are more likely attributable to the treatment rather than pre-existing differences. Some minor imbalance can still occur due to random chance, but with sufficiently large sample sizes (as is the case here), these differences should be minimal, as seen in the low SMDs.

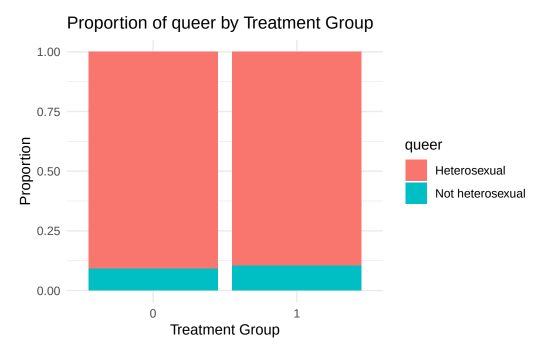
Considering the table, the results confirm that randomization has effectively created comparable groups across gender, age, and queer. This balanced setup make it possible to assess the treatment effect with confidence, knowing that pre-treatment covariate differences are not driving any observed outcome differences.

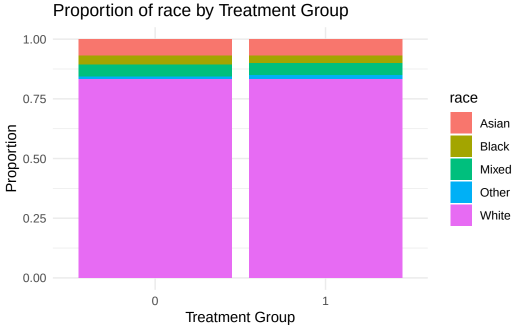
Plots for all Covariates across treatment groups:

```
for (covariate in covariates) { # looping through each covariate
  if (is.numeric(study1 cleaned transformed[[covariate]]) &&
      length(unique(study1_cleaned_transformed[[covariate]])) > 2) {
    p <- ggplot(data = study1_cleaned_transformed, aes(</pre>
      x = .data[[treatment_var]], y = !!sym(covariate))) +
      geom_boxplot() + # boxpolot for continuous covariates
      labs(title = paste("Distribution of", covariate, "by Treatment Group"),
           x = "Treatment Group", y = covariate) +
      theme minimal()
  } else {
    p <- ggplot(data = study1_cleaned_transformed, aes(</pre>
      x = .data[[treatment_var]], fill = factor(!!sym(covariate)))) +
      geom_bar(position = "fill") + # barplot for categorical covariates
      labs(title = paste("Proportion of", covariate, "by Treatment Group"),
           x = "Treatment Group", y = "Proportion", fill = covariate) +
      theme minimal()
  print(p)
```



Distribution of age by Treatment Group Treatment Group Treatment Group





All the plots show nicely that samples are evenly distributed across both treatment groups.

Exercise 4 (additional): Estimate Treatment Effect This exercise is not mandatory, but it serves only to opt for the maximum grade (6). Estimate the average effect of the treatment on the outcome variable support conditional on the pre-treatment immigration attitudes imm_1.

For this, use an interaction term in an OLS regression model. Compare your results to those in the original paper. Then, repeat this analysis with three iteratively smaller random samples of the treatment ($n=200,\ n=100,\ n=10$) and control groups ($n=200,\ n=100,\ n=10$); total $N=400,\ 200,\ and\ 20,\ respectively.$ Explain your findings. Finally, discuss how sample size impacts the results and what this implies about the role of randomization for selection bias.

Estimating the average effect of the treatment on the outcome variable support conditional on the pre-treatment variable imm_1:

```
Study1_OLS <-study1_cleaned_transformed%>%
   mutate(treat= as.factor(treatment)) # preparing treatment variable
OLS_model <- lm(support ~ treat*imm_1, data = Study1_OLS) # interaction term
summary(OLS_model, robust=TRUE)</pre>
Call:
```

Residuals:

```
Min 1Q Median 3Q Max -0.8916 -0.4358 0.1526 0.3037 0.7595
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
              0.240487
                         0.041816
                                    5.751 1.14e-08 ***
              0.146177
                         0.058195
                                    2.512
                                            0.0121 *
treat1
imm_1
              0.065114
                         0.006066 10.734 < 2e-16 ***
treat1:imm_1 -0.019038
                         0.008520 -2.234
                                            0.0256 *
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
```

lm(formula = support ~ treat * imm_1, data = Study1_OLS)

Residual standard error: 0.444 on 1144 degrees of freedom Multiple R-squared: 0.1328, Adjusted R-squared: 0.1305 F-statistic: 58.39 on 3 and 1144 DF, p-value: < 2.2e-16

The model suggests that treat, imm_1, and the interaction between them all significantly influence support. The treatment (treat1) has a positive effect, and higher values of imm_1 also increase the likelihood of support. The negative interaction term suggests that the effect of imm_1 on support is slightly weaker when the treatment is applied. Although all the predictors are significant, the model explains only a modest proportion of the variance in the outcome.

My results are quite similar to those of Model 2 in the original paper! Results from study1 in the original paper:

```
###|-----|----|-----|
###/
                | Model 1 | Model 2 | Model 3 | Model 4 |
###/ (Intercept)
                | 0.642*** | 0.238*** | 0.830*** | 0.447*** |
###|
                | (0.020) | (0.042) | (0.023) | (0.030) |
###| treat1
                | 0.022 | 0.148** | -0.056* | 0.097**
                | (0.028) | (0.058) | (0.033) | (0.042)
###|
###| imm_1
                       | 0.065*** |
###|
                       | (0.006) |
###| treat1 × imm_1 |
                        | -0.019** |
###|
                       | (0.009) |
             | 556
###| Num.Obs.
###/ R2
              | 0.000 | 0.132
###| R2 Adj.
                                | 0.003 | 0.008
###| RMSE
               ###Note: * p < 0.1, ** p < 0.05, *** p < 0.01
```

For Model 2, they used a similar code as me:

```
#models2 [['Model 2']] <-lm (support ~ treat*imm_1, data=df)</pre>
```

Comparison:

```
##/
                  | My Model | Model 2 from paper |
              ----/----/
##| (Intercept) | 0.240*** | 0.238*** |
##/
                  | (0.041) | (0.042) |
                 | 0.146* | 0.148** |
##| treat1
##/
                 | (0.058) | (0.058) |
\#\#/ imm 1
                 | 0.065*** | 0.065*** |
##/
                 | (0.006) | (0.006) |
##| treat1 × imm_1 | -0.019* | -0.019** |
##|
                 | (0.008) | (0.009) |
##| Num.Obs. | 1148
                           / 1151
##| R2
                 | 0.132 | 0.134
##| R2 Adj.
##| RMSE
                 | 0.135 | 0.132
                  1 0.44
                           1 0.44
##Note: * p < 0.1, ** p < 0.05, *** p < 0.01
```

The results are quite similar, with the coefficients for treat1 and imm_1 being almost identical. The p-values are also very close, with the coefficient for treat1 being significant at the 5% level in both models. The coefficient for the interaction term treat1*imm_1 is also significant at the 5% level in both models. Minor differences in significance levels are likely due to small variations in standard errors and the slightly smaller sample in my model. The overall similarity in coefficients and model fit measures indicates that the results are robust and aligned. Both models suggest a moderate positive effect of treatment on support, a strong positive association with imm_1, and a small dampening effect of treatment on the relationship between imm_1 and support.

Repeat the analysis with three iteratively smaller random samples of the treatment (n = 200, n = 100, n = 10) and control groups (n = 200, n = 100, n = 10); total N = 400, 200, and 20, respectively. n = 400:

```
create_balanced_sample <- function(data, n_per_group) {</pre>
  treat_data <- subset(data, treat == 1) # splitting data</pre>
  control_data <- subset(data, treat == 0)</pre>
  set.seed(123) # for reproducibility
  treat_sample <- treat_data[sample(nrow(treat_data), n_per_group), ]</pre>
  control_sample <- control_data[sample(nrow(control_data), n_per_group), ]</pre>
  balanced_sample <- rbind(treat_sample, control_sample)</pre>
  return(balanced_sample) # Combining samples
}
check_balance <- function(sample_data) {</pre>
  print("Treatment distribution:")
  print(table(sample_data$treat))
  print("Means by treatment group:")
  aggregate(cbind(support, imm_1) ~ treat, data = sample_data, mean)
}
sample 400 <- create balanced sample (Study1 OLS, 200) # 200 per t = 400 total
model_400 <- lm(support ~ treat*imm_1, data = sample_400)</pre>
print("Balance check for N=400:")
```

[1] "Balance check for N=400:"

```
check_balance(sample_400)
```

```
[1] "Treatment distribution:"
    0    1
200  200
[1] "Means by treatment group:"
```

```
treat support imm_1
1
     0
         0.615 6.14
2
     1
         0.660 5.98
print("Results for N=400:")
[1] "Results for N=400:"
summary(model_400)
Call:
lm(formula = support ~ treat * imm_1, data = sample_400)
Residuals:
   Min
            1Q Median
                           3Q
                                 Max
-0.8310 -0.4953 0.1821 0.3392 0.7286
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.27142 0.07199 3.770 0.000188 ***
                       0.10272 1.497 0.135229
treat1
             0.15375
             0.05596
                       0.01046 5.348 1.51e-07 ***
imm_1
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.4592 on 396 degrees of freedom
Multiple R-squared: 0.0966,
                             Adjusted R-squared: 0.08975
F-statistic: 14.11 on 3 and 396 DF, p-value: 9.301e-09
n = 200:
sample_200 <- create_balanced_sample(Study1_OLS, 100) # 100 per t = 200 total</pre>
model_200 <- lm(support ~ treat*imm_1, data = sample_200)</pre>
print("Balance check for N=200:")
```

[1] "Balance check for N=200:"

check_balance(sample_200)

```
[1] "Treatment distribution:"
 0
   1
100 100
[1] "Means by treatment group:"
 treat support imm_1
    0
         0.75 6.86
2
    1
         0.69 6.47
print("Results for N=200:")
[1] "Results for N=200:"
summary(model_200)
Call:
lm(formula = support ~ treat * imm_1, data = sample_200)
Residuals:
   Min
          1Q Median
                        3Q
                             Max
-0.9102 -0.5530 0.2429 0.3052 0.6001
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.39992 0.12474 3.206 0.00157 **
treat1
           imm_1
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.4421 on 196 degrees of freedom
Multiple R-squared: 0.04982, Adjusted R-squared: 0.03527
F-statistic: 3.425 on 3 and 196 DF, p-value: 0.01822
```

```
n = 20:
sample_20 <- create_balanced_sample(Study1_OLS, 10) # 10 per t = 20 total</pre>
model_20 <- lm(support ~ treat*imm_1, data = sample_20)</pre>
print("Balance check for N=20:")
[1] "Balance check for N=20:"
check_balance(sample_20)
[1] "Treatment distribution:"
0 1
10 10
[1] "Means by treatment group:"
 treat support imm_1
            0.5
      0
                  5.7
     1
            0.6
                 7.0
print("Results for N=20:")
[1] "Results for N=20:"
summary(model_20)
Call:
lm(formula = support ~ treat * imm_1, data = sample_20)
Residuals:
    Min
             1Q Median
                             3Q
                                    Max
-0.7552 -0.4187 0.1872 0.3612 0.5034
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.23298 0.36743 -0.634 0.5350
treat1
             1.19505
                         0.59216 2.018 0.0607 .
```

2.188

0.0438 *

 imm_1

0.12859

0.05876

Regression table for all sample sizes:

```
install.packages("stargazer")
```

The downloaded binary packages are in /var/folders/3k/4r5z06ts7bl41nkd_6d0dcyh0000gn/T//Rtmp60PFFi/downloaded_packages

```
library(stargazer)
```

Please cite as:

Hlavac, Marek (2022). stargazer: Well-Formatted Regression and Summary Statistics Tables.

 ${\tt R} \ {\tt package} \ {\tt version} \ {\tt 5.2.3.} \ {\tt https://CRAN.R-project.org/package=stargazer}$

```
Regression Results for OG Sample
```

Dependent variable:

support -----

treat1 0.146** (0.058)

```
0.065***
imm_1
                         (0.006)
treat1:imm_1
                        -0.019**
                         (0.009)
Constant
                        0.240***
                         (0.042)
Observations
                          1,148
R2
                          0.133
Adjusted R2
                          0.131
Residual Std. Error 0.444 (df = 1144)
                 58.386*** (df = 3; 1144)
F Statistic
_____
Note:
                 *p<0.1; **p<0.05; ***p<0.01
```

Regression Results for Different Sample Sizes

______ Dependent variable: _____ support (1) (2) (3) 0.232 treat1 0.154 1.195* (0.103)(0.165)(0.592)0.056*** 0.051*** 0.129** imm 1 (0.010)(0.017)(0.059)treat1:imm_1 -0.017 -0.042* -0.180* (0.015)(0.023)(0.086)0.271*** 0.400*** Constant -0.233 (0.072)(0.125)(0.367)

Observations	400	200	20
R2	0.097	0.050	0.262
Adjusted R2	0.090	0.035	0.124
Residual Std. Error	0.459 (df = 396)	0.442 (df = 196)	0.478 (df = 16)
F Statistic	14.114*** (df = 3; 396)	3.425** (df = 3; 196)	1.896 (df = 3; 16)
Note:		*p<0.1;	**p<0.05; ***p<0.01

Treatment Effect (treat1): As sample size decreases, the effect of treatment on support grows larger, but standard errors increase, while statistical significance fluctuates. This suggests that smaller samples are less precise and more prone to random fluctuations in treatment effect estimates. Similar observations can be made for R-squared and adjusted R-squared values and F-Values, which all decrease as sample size decreases.

The findings illustrate that larger sample sizes provide more stable and precise estimates. As sample size decreases, estimates for treatment and the interaction term become increasingly variable, with higher standard errors and changing significance levels. This instability suggests that smaller samples may fail to capture the true treatment effect accurately, leading to overor underestimation of effects. When it comes to random sampling and random assignment in experiments, they typically reduce selection bias. However, in smaller samples, the risk of imbalances in covariates increases even with randomization, because smaller groups are more prone to random variation that may not reflect the population accurately. For example, the inflated treatment effect observed with the smallest sample size (20 observations) likely reflects random sampling noise rather than a genuine increase in the treatment effect. This implies that randomization alone is insufficient to prevent selection bias in small samples. Adequate sample size is necessary to ensure that randomization leads to balanced groups and that results are generalizable.