Applied Microeconometrics - Assignment 3

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Construct a variable full-time equivalent for both waves, which is the number of full-time employees plus the number of part-time employees divided by two and also add the number of managers. I will simply refer to employees for this outcome variable.

(i) Compute separately for New Jersey and Pennsylvania the average number of employees in both waves, and compute the difference-in-difference estimate

STATE	mean_before	mean_after
PA	23.33117	21.16558
NJ	20.43941	21.02743

```
did$mean_after[2] - did$mean_after[1] - (did$mean_before[2] - did$mean_before[1])
```

[1] 2.753606

Next repeat this, but only considering the restaurants that responded in both waves of the survey.

[1] 2.75

(ii) Estimate this model and next subsequently add characteristics of the restaurants observed in the first wave. But think carefully which characteristics can be included. How does the latter affect the estimate for the coefficient δ ?

```
model1 <- lm(data = dataset,
   formula = changeemployees ~ STATE)
model2 <- lm(data = dataset,</pre>
             formula = changeemployees ~ STATE + SOUTHJ + CENTRALJ + SHORE + PA1)
model3 <- update(model1, . ~ . + NCALLS + WAGE_ST + INCTIME + FIRSTINC +</pre>
                    BONUS + MEALS + OPEN + HRSOPEN + PSODA + PFRY + NREGS + NREGS11)
model4 <- update(model2, . ~ . + NCALLS + WAGE_ST + INCTIME + FIRSTINC +</pre>
                    BONUS + MEALS + OPEN + HRSOPEN + PSODA + PFRY + NREGS + NREGS11)
models <- list(model1, model2, model3, model4)</pre>
# Adjust standard errors
cov1 <- vcovHC(model1, type = "HC1")</pre>
robust_se1
            <- sqrt(diag(cov1))</pre>
cov2 <- vcovHC(model2, type = "HC1")</pre>
             <- sqrt(diag(cov2))</pre>
robust_se2
cov3 <- vcovHC(model3, type = "HC1")</pre>
robust_se3
            <- sqrt(diag(cov3))</pre>
cov4 <- vcovHC(model4, type = "HC1")</pre>
robust_se4
            <- sqrt(diag(cov4))</pre>
stargazer(models, omit.stat = c("ll", "ser", "rsq"), df=F,
          omit = c("SOUTHJ", "CENTRALJ", "SHORE", "PA1"),
          add.lines = list(c("Region Dummies", "No", "Yes", "No", "Yes")),
          header = FALSE, font.size = "footnotesize", title='Estimated model for Q2 with robust standard errors.',
          se=list(robust_se1, robust_se2, robust_se3,
                  robust_se4))
```

We want to isolate the effect of the minimum wage by attributing it to the coefficient belonging to STATE, which means we have to account for all possible sources of variation not due to the minimum wage. This also means we cannot control for PCTAFF, because this is the mechanism we care about: if we conditioned on this variable, that would absorb all variation due to the minimum wage policy changes and would change our interpretation of the STATE coefficient to a partial instead of a total effect and bias it towards zero.

Table 1: Estimated model for Q2 with robust standard errors.

	Dependent variable:							
		changeer	mployees					
	(1)	(2)	(3)	(4)				
STATE	-2.750** (1.338)	-1.785 (2.053)	-1.200 (1.717)	0.736 (2.541)				
NCALLS			-0.117 (0.337)	-0.136 (0.332)				
WAGE_ST			2.353 (1.496)	2.800* (1.551)				
INCTIME			-0.072 (0.051)	-0.076 (0.052)				
FIRSTINC			-1.132 (5.868)	-1.460 (6.034)				
BONUS			0.209 (1.249)	0.230 (1.236)				
MEALS			-0.434 (0.928)	-0.269 (0.952)				
OPEN			-1.217^* (0.658)	-1.274^* (0.695)				
HRSOPEN			-0.576 (0.514)	-0.595 (0.551)				
PSODA			1.060 (8.929)	0.134 (9.179)				
PFRY			-6.126 (6.352)	-6.962 (6.643)				
NREGS			-0.544 (0.514)	-0.432 (0.520)				
NREGS11			0.458 (0.635)	0.419 (0.643)				
Constant	2.283* (1.248)	0.970 (1.921)	16.376 (16.048)	14.339 (17.653)				
Region Dummies Observations Adjusted R ² F Statistic	No 384 0.012 5.675**	Yes 384 0.014 2.056*	No 303 -0.002 0.962	Yes 303 0.005 1.091				

Note:

*p<0.1; **p<0.05; ***p<0.01

(iii) Provide a balancing table, i.e. show the sample mean of characteristics observed in the first survey separately for the restaurants in New Jersey and Pennsylvania. What is your opinion about the balancing table?

We think that the table indicates that the covariates are balanced across the two groups, approaching the situation of an experiment. There are, however, a couple of variables that are significantly different across the two groups: MEALS, PSODA and PFRY. Omitting them could have consequences for the estimated effect of the minimum wage on employment, because those variables (i.e. MEALS, PSODA and FRY) could be correlated or even have a causal effect on employment. For example, if the price is higher in one country, they might serve a higher segment of the market, and higher less personnel in general, and pay above the minimum wage.

	NJ (N=309)		PA	(N=75)		
	Mean	Std. Dev.	Mean	Std. Dev.	Diff. in Means	p
changeemployees	-0.467	8.452	2.283	10.854	2.750	0.043
NCALLS	1.214	1.464	0.747	0.960	-0.467	0.001
WAGEST	4.609	0.343	4.630	0.358	0.021	0.647
INCTIME	17.905	10.625	19.279	13.183	1.373	0.420
FIRSTINC	0.228	0.110	0.210	0.096	-0.018	0.199
BONUS	0.239	0.427	0.293	0.458	0.054	0.357
PCTAFF	49.157	34.789	45.571	36.935	-3.586	0.465
MEALS	1.874	0.570	2.027	0.402	0.153	0.008
OPEN	8.100	2.182	7.807	2.164	-0.294	0.295
HRSOPEN	14.398	2.818	14.513	2.960	0.115	0.761
PSODA	1.063	0.086	0.975	0.069	-0.087	0.000
PFRY	0.941	0.103	0.843	0.089	-0.099	0.000
PENTREE	1.360	0.657	1.232	0.635	-0.127	0.131
NREGS	3.697	1.285	3.373	1.100	-0.324	0.029
NREGS11	2.709	0.915	2.811	0.753	0.102	0.321

None of this is particularly plausible in our opinion, but it makes sense to control for these factors anyway, so that the eventual effect takes into account these possible causal links.

(iv) Check for the different characteristics if there is a common support for restaurants in New Jersey and Pennsylvania. And estimate a propensity score for being a restaurant in New Jersey.

```
# Check for common support
dataset2 <- dataset %>%
   mutate(STATE = if_else(STATE == 0, "PA", "NJ")) %>%
   filter(!is.na(changeemployees)) %>%
   rename_with(.fn = ~ stringr::str_replace(.x, "_", "")) %>%
   select(changeemployees, STATE, NCALLS, WAGEST, INCTIME, FIRSTINC, BONUS, PCTAFF, MEALS,
           OPEN, HRSOPEN, PSODA, PFRY, PENTREE, NREGS, NREGS11)
emptycol = function(x) " "
boxplot1 <- lapply(dataset2 %>%
                     filter(STATE == "NJ") %>%
                  select(-STATE), na.omit) %>% lapply(scale)
boxplot2 <- lapply(dataset2 %>%
                     filter(STATE == "PA") %>%
                  select(-STATE), na.omit) %>% lapply(scale)
modelsummary::datasummary(
 data = dataset2,
  NCALLS + WAGEST + INCTIME + FIRSTINC + BONUS + PCTAFF + MEALS +
          OPEN + HRSOPEN + PSODA + PFRY + PENTREE + NREGS + NREGS11 ~
   STATE * (Mean + SD + Heading("Boxplot")*emptycol + Heading("Histogram")*emptycol),
 title = 'Common support table.') %>%
 column_spec(column = 4, image = spec_boxplot(boxplot1)) %>%
 column_spec(column = 8, image = spec_boxplot(boxplot2)) %>%
 column_spec(column = 5, image = spec_hist(boxplot1)) %>%
 column_spec(column = 9, image = spec_hist(boxplot2)) %>%
 kableExtra::kable_styling(latex_options = "hold_position")
```

Table 2: Common support table.

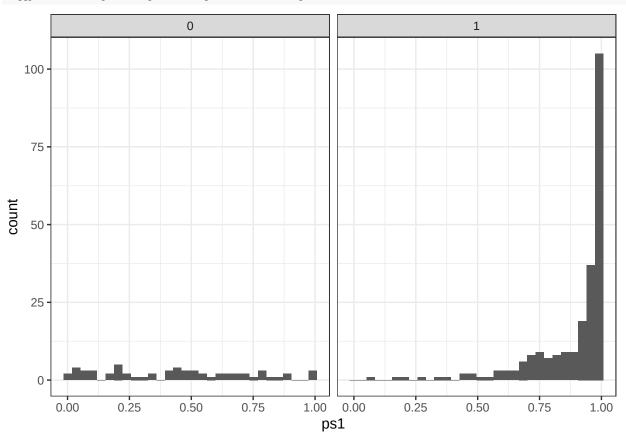
	NJ					PA				
	Mean	SD	Boxplot	Histogram	Mean	SD	Boxplot	Histogram		
NCALLS	1.21	1.46	• • • • • • • • • • • • • • • • • • • •		0.75	0.96	•——••			
WAGEST	4.61	0.34	··	L	4.63	0.36				
INCTIME	17.91	10.63			19.28	13.18				
FIRSTINC	0.23	0.11	⊢ ⊢ •		0.21	0.10	⊢ ⊢ •			
BONUS	0.24	0.43	├	\mathbb{A}	0.29	0.46	H •			
PCTAFF	49.16	34.79			45.57	36.93	—			
MEALS	1.87	0.57	HIH		2.03	0.40	H			
OPEN	8.10	2.18	• • • •		7.81	2.16	• • •			
HRSOPEN	14.40	2.82	• 🗕 +		14.51	2.96	⊢			
PSODA	1.06	0.09	⊢ •		0.98	0.07	⊢ ⊢ •	<u></u>		
PFRY	0.94	0.10	•—•		0.84	0.09	• 🖂			
PENTREE	1.36	0.66	⊢ ∎••		1.23	0.64	\vdash			
NREGS	3.70	1.28	⊢	<u></u>	3.37	1.10	⊢) ••• •			
NREGS11	2.71	0.92	• H H • • •		2.81	0.75	⊢ ⊢ •			

As indicated in the table, there is no common support for any of the variables, as we are dealing with continuous variables, so that the probability of realizing two zero outcomes is practically zero. We estimate two propensity scores, one extensive model, which sacrifices many observations, and one parsimonious model,

which does not.

We show the distribution of propensity scores across the two groups:

```
dataset %>%
   ggplot(aes(x = ps1)) + geom_histogram()+ facet_wrap(~as.factor(STATE)) + theme_bw()
```



(v) Use propensity score matching to estimate the average treatment effect on the treated for the employment before and after the minimum wage increase in New Jersey, so on E_{0i} and E_{1i} separately.

We report the results of (v) and (vi) in table 3. We use the MatchIt package to estimate the propensity-score again and subsequently match using the nearest neighbor algorithm to compute E_{0i} :

```
matched_data1 <- MatchIt::matchit(STATE ~ MEALS + OPEN + HRSOPEN + PSODA + PFRY,</pre>
                 data = dataset %>%
                   select(employees, STATE, MEALS, OPEN, HRSOPEN, PSODA, PFRY) %>%
                   na.omit(),
                 method = "nearest") %>%
  match.data()
e_Oi <- lm(employees ~ STATE + MEALS + OPEN + HRSOPEN + PSODA + PFRY, data = matched_data1)
# Adjust standard errors
cov0i
             <- vcovHC(e_0i, type = "HC1")</pre>
robust_se0i <- sqrt(diag(cov0i))</pre>
And for E_{1i}:
matched_data2 <- MatchIt::matchit(STATE ~ MEALS + OPEN + HRSOPEN + PSODA + PFRY,
                 data = dataset %>%
                   select(employees2, STATE, MEALS, OPEN, HRSOPEN, PSODA, PFRY) %>%
                   na.omit(),
                 method = "nearest") %>%
  match.data()
e_1i <- lm(employees2 ~ STATE + MEALS + OPEN + HRSOPEN + PSODA + PFRY, data = matched_data2)
# Adjust standard errors
cov1i
            <- vcovHC(e_1i, type = "HC1")</pre>
robust_se1i <- sqrt(diag(cov1i))</pre>
```

(vi) Now use propensity score matching to estimate the average treatment effect on the treated on the change in employment in the restaurants, so $E_{1i} - E_{0i}$.

```
matched_data3 <- MatchIt::matchit(STATE ~ MEALS + OPEN + HRSOPEN + PSODA + PFRY,</pre>
                 data = dataset %>%
                  select(changeemployees, STATE, MEALS, OPEN, HRSOPEN, PSODA, PFRY) %>%
                  na.omit(),
                 method = "nearest") %>%
 match.data()
ate <- lm(changeemployees ~ STATE + MEALS + OPEN + HRSOPEN + PSODA + PFRY, data = matched_data3)
# Adjust standard errors
cov_ate
         <- vcovHC(ate, type = "HC1")</pre>
robust_se_ate <- sqrt(diag(cov_ate))</pre>
stargazer(e_0i, e_1i, ate, header = F,
         omit.stat = c("ll", "ser", "rsq"), df = F,
          font.size = "footnotesize",
          label="tab:hoi",
         title='Application of propensity score matching to estimate the average treatment
         effect on the treated on the change in employments in the restaurants. Standard erros are robust',
         se= list(robust_se0i, robust_se1i, robust_se_ate))
```

Table 3: Application of propensity score matching to estimate the average treatment effect on the treated on the change in employments in the restaurants. Standard errors are robust

		Dependent var	riable:
	employees	employees2	changeemployees
	(1)	(2)	(3)
STATE	-1.262	1.410	-2.133
	(3.042)	(2.494)	(3.334)
MEALS	-0.095	1.360	-1.243
	(1.277)	(1.304)	(1.407)
OPEN	0.914	1.223	-0.436
	(0.895)	(0.750)	(0.988)
HRSOPEN	2.804***	2.518***	0.232
	(0.630)	(0.474)	(0.648)
PSODA	-18.029	-16.884	-1.719
	(11.010)	(16.133)	(15.524)
PFRY	1.868	3.983	-2.654
	(8.215)	(7.062)	(9.723)
Constant	-7.829	-14.547	8.992
	(22.649)	(20.303)	(25.688)
Observations	150	150	146
Adjusted R ²	0.331	0.321	0.009
F Statistic	13.276***	12.756***	1.231

Note:

 $^*\mathrm{p}{<}0.1;\ ^{**}\mathrm{p}{<}0.05;\ ^{***}\mathrm{p}{<}0.01$

(vii) Now check the sensitivity of the propensity score matching estimate by also computing the weighting estimators for the average treatment effect on the treated.

For transparency, we show (the first rows of) intermediate datasets we use. We first calculate the nearest neighbour weighting estimate, using the matched_data3 data.frame, which implements nearest-neighbor matching:

Table 4: Nearest neigbour weighting estimate.

changeemployees	STATE	MEALS	OPEN	HRSOPEN	PSODA	PFRY	distance	weights	subclass
-4.00	0	2	11	10.0	0.94	0.73	0.5233362	1	1
-0.50	1	1	9	12.5	1.12	1.02	0.9952999	1	1
3.50	0	3	7	16.0	1.00	0.87	0.4749829	1	2
-3.50	1	2	10	14.0	1.06	0.98	0.9940832	1	2
-16.00	0	2	6	16.0	0.94	0.90	0.2123310	1	3
22.75	1	1	7	15.0	1.17	0.95	0.9898831	1	3

```
treated_subjects <- sum(matched_data3$STATE)

differences <- matched_data3 %>%
    group_split(subclass) %>%
    map_dbl(.f =~ .x$changeemployees[2]- .x$changeemployees[1])

paste('The mean difference in employees is', mean(differences))
```

```
## [1] "The mean difference in employees is -3.1472602739726"
paste('The standard deviation is', sd(differences))
```

[1] "The standard deviation is 14.6013481360335"

Now, we calculate the neighborhood matching weighting estimate, with k=4, implemented again in the MatchIt package:

```
matched data4 <- Matchit::matchit(STATE ~ MEALS + OPEN + HRSOPEN + PSODA + PFRY,
                 data = dataset %>%
                   select(changeemployees, STATE, MEALS, OPEN, HRSOPEN, PSODA, PFRY) %>%
                  na.omit().
                 method = "nearest",
                 replace = TRUE,
                 ratio = 4,
                 min.controls = 3) %>%
 get_matches()
matched_data4 %>%
 arrange(subclass) %>%
 head(6) %>%
 kable(booktabs=T, caption = "Neigborhood
                            weighting estimate, with $k=4$.") %>%
 kableExtra::kable_styling(latex_options = c("hold_position", "scale_down"))
differences <- matched_data4 %>%
 group_split(subclass) %>%
 map_dbl(.f = ~.x$changeemployees[1] - 0.25*(.x$changeemployees[2] +
                                                 .x$changeemployees[3] +
                                                 .x$changeemployees[4] +
                                                 .x$changeemployees[5])
```

Table 5: Neigborhood weighting estimate, with k = 4.

id	subclass	weights	changeemployees	STATE	MEALS	OPEN	HRSOPEN	PSODA	PFRY	distance
74	1	1.00	-12.0	1	2	7.0	16.0	1.06	0.95	0.9155793
34	1	0.25	-3.0	0	2	10.0	12.0	1.01	0.89	0.9235344
22	1	0.25	0.0	0	2	7.0	15.0	1.05	1.01	0.9002948
1	1	0.25	16.5	0	2	6.5	16.5	1.03	1.03	0.8989450
64	1	0.25	18.5	0	2	8.5	13.0	1.05	0.91	0.8924194
75	2	1.00	-6.5	1	2	7.0	14.5	1.06	0.95	0.8359729

```
paste('The mean difference in employees is', mean(differences))
```

```
## [1] "The mean difference in employees is -2.82320819112628" paste('The standard deviation is', sd(differences))
```

[1] "The standard deviation is 10.0384946681803"

Finally, we manually implement the normal kernel density with $\Sigma = I$:

```
library(mvtnorm)
normalkernel <- function(dataset){</pre>
  treated_outcomes <- dataset %>%
   filter(STATE ==1) %>%
    select(changeemployees)
  untreated_outcomes <- dataset %>%
    filter(STATE == 0) %>%
    select(changeemployees)
  treated_obs <- dataset %>%
    filter(STATE == 1) %>%
    select(MEALS, OPEN, HRSOPEN, PSODA, PFRY)
  untreated_obs <- dataset %>%
    filter(STATE == 0) %>%
    select(MEALS, OPEN, HRSOPEN, PSODA, PFRY)
  outcomes <- vector(length = nrow(treated_obs))</pre>
  w <- matrix(nrow = nrow(treated_obs), ncol = nrow(untreated_obs))</pre>
  for (i in 1:nrow(treated_obs)){
    for(j in 1:nrow(untreated_obs)){
      # Create the weight matrix
      w[i,j] <- mvtnorm::dmvnorm(as.numeric(untreated_obs[j,]),
                                  mean = as.numeric(treated_obs[i,]))
   }
    # Normalize the weights
  weightstotal <- sum(w[i,], na.rm = TRUE)</pre>
  w[i,] = w[i,] / weightstotal
  # Compute the estimated outcomes
  outcomes[i] <- treated_outcomes[i,] - sum(w[i,]*untreated_outcomes[,1], na.rm = TRUE)</pre>
  outcomes <- unlist(outcomes)</pre>
```

```
print(paste('The estimated ATT is equal to:', mean(outcomes, na.rm = TRUE)))
print(paste('The std. deviation is equal to:', sd(outcomes, na.rm = TRUE)))
# Compute the average - final number
}
normalkernel(dataset)
```

```
## [1] "The estimated ATT is equal to: -2.4880507067192" ## [1] "The std. deviation is equal to: 8.98329827881984"
```