The Toy Example - A Demo for 500

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0.1 Setup: Loading Packages

```
library(pander); library(tableone); library(broom)
library(Epi); library(survival); library(arm)
library(Hmisc); library(Matching); library(lme4)
library(twang); library(survey); library(rbounds)
library(cobalt); library(tidyverse)
```

1 The Toy Data at Load-In

This document is about showing a (relatively) simple way to do things. In no way would I claim that the approaches provided here are optimal. This is just a first demonstration.

1.1 The Data Set

The Data Set is 100% fictional, and is available as toy.csv on the course website. It contains data on 200 subjects (70 treated - subjects 131-200 and 130 controls - subjects 1-130) on treatment status, six covariates, and three outcomes, with no missing observations anywhere. We assume that a logical argument suggests that the square of covA, as well as the interactions of covB with covC and with covD should be related to treatment assignment, and thus should be included in our propensity model.

Our objective is to estimate the average causal effect of treatment (as compared to control) on each of the three outcomes, without propensity adjustment, and then with propensity matching, subclassification, weighting and regression adjustment using the propensity score.

```
toy <- read_csv("toy.csv")
```

```
Parsed with column specification:
cols(
   subject = col_integer(),
   treated = col_integer(),
   covA = col_double(),
   covC = col_double(),
   covD = col_double(),
   covD = col_double(),
   covE = col_integer(),
```

```
covF = col_character(),
  out1.cost = col_integer(),
  out2.event = col_character(),
  out3.time = col_integer()
toy$out2.event <- factor(toy$out2.event)</pre>
toy$covF <- factor(toy$covF)</pre>
toy
# A tibble: 200 x 11
                                         covD
   subject treated
                      covA covB covC
                                              covE covF
                                                             out1~ out2~ out3~
     <int>
                    <dbl> <int> <dbl> <int> <fctr> <int> <fctr> <int> <fctr> <int> <fctr> <int>
         1
                  0 0.120
                                1 7.60 6.70
                                                   8 3-High
                                                                46 Yes
                                                                             99
1
         2
                  0 0.0500
                                0 11.2 11.7
                                                   7 1-Low
                                                                44 Yes
                                                                            112
                                                                47 No
 3
         3
                  0 0.680
                                0 7.70 9.30
                                                  12 2-Mid~
                                                                             89
 4
         4
                  0 0.460
                                  9.50 9.40
                                                  12 2-Mid~
                                                                42 No
                                                                            121
 5
         5
                  0 0.340
                                1 9.90 10.7
                                                   9 2-Mid~
                                                                30 No
                                                                            120
 6
         6
                  0 0.150
                                0 10.4 12.2
                                                   6 3-High
                                                                28 No
                                                                             91
7
         7
                  0 0.180
                                0 13.7
                                                   9 1-Low
                                                                            101
                                         8.50
                                                                60 Yes
8
         8
                  0 0.280
                                1 11.0
                                         9.20
                                                   6 2-Mid~
                                                                62 No
                                                                            101
9
         9
                                                  10 2-Mid~
                  0 0.160
                                0 9.80 10.2
                                                                48 No
                                                                            103
10
        10
                  0 0.120
                                1 10.8 12.1
                                                  11 1-Low
                                                                54 Yes
                                                                             89
# ... with 190 more rows
```

1.2 The Codebook

Variable	Type	Notes
subject	Subject ID	1-130 = controls, 131-200 = treated
treated	2-level categorical $(0/1)$	0 = control, 1 = treated
covA	Quantitative (2 decimal places)	reasonable values range from 0 to 1
covB	2-level categorical $(0/1)$	0 = no, 1 = yes
covC	Quantitative (1 decimal place)	range $3-20$
covD	Quantitative (1 decimal place)	range $3-20$
covE	Integer	range $3-20$
covF	3-level ordinal factor	1 = Low, 2 = Middle, 3 = High
out1.cost	Quantitative outcome	typical values 20-80
out2.event	Binary outcome (did event occur?)	Yes/No (note: event is bad)
out3.time	Time to event outcome	Time before event is observed or subject
		exits study (censored), range is 75-156
		weeks

With regard to the out3.time variable, subjects with out2.event = No were censored, so that out2.event = Yes indicates an observed event.

1.3 Numerical Summaries

```
summary(toy)

subject treated covA covB
Min. : 1.00 Min. :0.00 Min. :0.0400 Min. :0.00
```

```
1st Qu.: 50.75
                  1st Qu.:0.00
                                  1st Qu.:0.2675
                                                    1st Qu.:0.00
Median :100.50
                  Median:0.00
                                  Median :0.5100
                                                    Median:0.00
Mean
       :100.50
                  Mean
                         :0.35
                                  Mean
                                         :0.4995
                                                    Mean
                                                            :0.37
3rd Qu.:150.25
                                  3rd Qu.:0.7300
                  3rd Qu.:1.00
                                                    3rd Qu.:1.00
       :200.00
Max.
                  Max.
                         :1.00
                                         :0.9900
                                                            :1.00
                       covD
                                         covE
                                                            covF
     covC
Min.
       : 4.200
                  Min.
                         : 5.500
                                    Min.
                                           : 2.00
                                                     1-Low
                                                              :67
1st Qu.: 8.600
                  1st Qu.: 8.500
                                    1st Qu.: 8.00
                                                     2-Middle:84
Median: 9.850
                  Median : 9.900
                                    Median :10.00
                                                     3-High :49
Mean
      : 9.714
                  Mean
                         : 9.863
                                    Mean
                                           :10.21
3rd Qu.:10.725
                  3rd Qu.:11.100
                                    3rd Qu.:12.00
                                            :19.00
Max.
       :16.200
                  Max.
                          :14.400
                                    Max.
                               out3.time
  out1.cost
                 out2.event
Min.
       :24.00
                 No :103
                             Min.
                                    : 79.0
1st Qu.:39.00
                 Yes: 97
                             1st Qu.:101.0
Median :49.00
                             Median :109.0
Mean
       :50.42
                             Mean
                                    :109.2
3rd Qu.:58.00
                             3rd Qu.:118.0
       :80.00
Max.
                             Max.
                                    :151.0
```

1.4 Table 1

S	Stratifi	led by to	reated			
	0		1		p	test
n	130		70			
covA (mean (sd))	0.45	(0.29)	0.59	(0.22)	<0.001	
covB = 1 (%)	40	(30.8)	34	(48.6)	0.020	
covC (mean (sd))	9.76	(2.02)	9.63	(2.02)	0.649	
covD (mean (sd))	9.74	(1.77)	10.10	(1.88)	0.183	
covE (mean (sd))	10.50	(3.54)	9.67	(2.74)	0.090	
covF (%)					0.266	
1-Low	48	(36.9)	19	(27.1)		
2-Middle	54	(41.5)	30	(42.9)		
3-High	28	(21.5)	21	(30.0)		
out1.cost (mean (sd))	44.94	(9.69)	60.61	(16.56)	<0.001	
out2.event = Yes (%)	58	(44.6)	39	(55.7)	0.177	
out3.time (mean (sd))	106.95	(11.94)	113.44	(10.75)	<0.001	

2 Data Management and Cleanup

2.1 Range Checks for Quantitative (continuous) Variables

Checking and cleaning the quantitative variables is pretty straightforward - the main thing I'll do at this stage is check the ranges of values shown to ensure that they match up with what I'm expecting. Here, all of the quantitative variables have values that fall within the "permissible" range described by my codebook, so

we'll assume that for the moment, we're OK on subject (just a meaningless code, really), covA, covC, covD, covE, out1.cost and out3.time, and we see no missingness.

2.2 Restating Categorical Information in Helpful Ways

The cleanup of the toy data focuses, as it usually does, on variables that contain **categories** of information, rather than simple counts or measures, represented in quantitative variables.

2.2.1 Re-expressing Binary Variables as Numbers and Factors

We have three binary variables (treated, covB and out2.event). A major issue in developing these variables is to ensure that the direction of resulting odds ratios and risk differences are consistent and that cross-tabulations are in standard epidemiological format.

It will be useful to define binary variables in two ways:

- as a numeric indicator variable taking on the values 0 (meaning "not having the characteristic being studied") or 1 (meaning "having the characteristic being studied")
- as a text factor with the levels of our key exposure and outcomes arranged so that "having the characteristic" precedes "not having the characteristic" in R when you create a table, but the covariates should still be No/Yes.

So what do we currently have? From the output below, it looks like treated and covB are numeric, 0/1 variables, while out2.event is a factor with levels "No" and then "Yes"

```
toy %>% select(treated, covB, out2.event) %>% summary()
```

```
treated
                     covB
                               out2.event
                               No :103
Min.
       :0.00
               Min.
                       :0.00
1st Qu.:0.00
               1st Qu.:0.00
                               Yes: 97
Median:0.00
               Median:0.00
Mean
       :0.35
               Mean
                       :0.37
3rd Qu.:1.00
               3rd Qu.:1.00
Max.
       :1.00
                       •1 00
               Max.
```

So, we'll create factors for treated and covB:

```
toy$treated_f <- factor(toy$treated, levels = c(1,0), labels = c("Treated", "Control"))
toy$covB_f <- factor(toy$covB, levels = c(0,1), labels = c("No B", "Has B"))</pre>
```

For out2.event, on the other hand, we don't have either quite the way we might want it. As you see in the summary output, we have two codes for out2.event - either No or Yes, in that order. But we want Yes to precede No (and I'd like a more meaningful name). So I redefine the factor variable, as follows.

```
toy$out2_f <- factor(toy$out2.event, levels = c("Yes", "No"), labels = c("Event", "No Event"))</pre>
```

To obtain a numerical (0 or 1) version of out2.event we can use R's as.numeric function - the problem is that this produces values of 1 (for No) and 2 (for Yes), rather than 0 and 1. So, I simply subtract 1 from the result, and we get what we need.

```
toy$out2 <- as.numeric(toy$out2.event) - 1</pre>
```

2.2.2 Testing Your Code - Sanity Checks

Before I move on, I'll do a series of sanity checks to make sure that our new variables are defined as we want them, by producing a series of small tables comparing the new variables to those originally included in the data set.

```
table(toy$treated_f, toy$treated)
            0
                1
  Treated
            0
               70
  Control 130
table(toy$covB_f, toy$covB)
          0
              1
  No B 126
              0
  Has B
          0 74
table(toy$out2_f, toy$out2.event)
            No Yes
             0 97
  Event.
  No Event 103
table(toy$out2, toy$out2.event)
     No Yes
  0 103
          0
      0 97
table(toy$out2, toy$out2_f)
    Event No Event
        0
               103
  0
```

Everything looks OK:

0

97

1

- treated_f correctly captures the information in treated, with the label Treated above the label Control in the rows of the table, facilitating standard epidemiological format.
- covB_f also correctly captures the covB information, placing "Has B" last.
- out2 f correctly captures and re-orders the labels from the original out2.event
- out2 shows the data correctly (as compared to the original out2.event) with 0-1 coding.

2.3 Dealing with Variables including More than Two Categories

When we have a multi-categorical (more than two categories) variable, like covF, we will want to have

- both a text version of the variable with sensibly ordered levels, as a factor in R, as well as
- a series of numeric indicator variables (taking the values 0 or 1) for the individual levels.

```
summary(toy$covF)
```

```
1-Low 2-Middle 3-High
67 84 49
```

From the summary output, we can see that we're all set for the text version of covF, as what we have currently is a factor with three levels, labeled 1-Low, 2-Middle and 3-High. This list of variables should work out well

for us, as it preserves the ordering in a table and permits us to see the names, too. If we'd used just Low, Middle and High, then when R sorted a table into alphabetical order, we'd have High, then Low, then Middle - not ideal.

2.3.1 A Brief Digression

Suppose, for the moment, that a different categorical variable had been included in our data set - this one, which we'll call cat4, has four levels, called (in the imported data: 1, 2, 3 and 4) - I'd turn this into a factor using this command:

```
cat4.f <- factor(cat4, levels=c(1,2,3,4), labels=c("Group 1", "Group 2", "Group 3", "Group 4"))
or, perhaps, instead, something like this:
cat4.f <- factor(cat4, levels=c(1,2,3,4), labels=c("1-Lowest", "2-Low", "3-High", "4-Highest"))</pre>
```

2.3.2 Preparing Indicator Vraiables for covF

So, all we need to do for covF is prepare indicator variables. We can either do this for all levels, or select one as the baseline, and do the rest. Here, I'll show them all.

```
## Re-expressing the Multi-Categorical Variable
toy$covF.Low <- as.numeric(toy$covF=="1-Low")
toy$covF.Middle <- as.numeric(toy$covF=="2-Middle")
toy$covF.High <- as.numeric(toy$covF=="3-High")</pre>
```

And now, some more sanity checks for the covF information:

table(toy\$covF, toy\$covF.Low)

```
0 1

1-Low 0 67

2-Middle 84 0

3-High 49 0

table(toy$covF, toy$covF.Middle)
```

```
0 1

1-Low 67 0

2-Middle 0 84

3-High 49 0

table(toy$covF, toy$covF.High)
```

```
0 1
1-Low 67 0
2-Middle 84 0
3-High 0 49
```

2.4 Creating the Transformation and Product Terms

Remember that we have reason to believe that the square of covA as well as the interaction of covB with covC and also covB with covD will have an impact on treatment assignment. It will be useful to have these

transformations in our data set for modeling and summarizing. I will use covB in its numeric (0,1) form (rather than as a factor - covB.f) when creating product terms, as shown below.

```
toy$Asqr <- toy$covA^2
toy$BC <- toy$covB*toy$covC
toy$BD <- toy$covB*toy$covD</pre>
```

3 Data Set After Cleaning

3.1 Glimpse

```
glimpse(toy)
Observations: 200
Variables: 21
$ subject
             <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15,...
$ treated
             <dbl> 0.12, 0.05, 0.68, 0.46, 0.34, 0.15, 0.18, 0.28, 0....
$ covA
             <int> 1, 0, 0, 0, 1, 0, 0, 1, 0, 1, 0, 1, 0, 0, 1, 1, 0, ...
$ covB
             <dbl> 7.6, 11.2, 7.7, 9.5, 9.9, 10.4, 13.7, 11.0, 9.8, 1...
$ covC
             <dbl> 6.7, 11.7, 9.3, 9.4, 10.7, 12.2, 8.5, 9.2, 10.2, 1...
$ covD
$ covE
             <int> 8, 7, 12, 12, 9, 6, 9, 6, 10, 11, 4, 12, 16, 17, 1...
$ covF
             <fctr> 3-High, 1-Low, 2-Middle, 2-Middle, 2-Middle, 3-Hi...
             <int> 46, 44, 47, 42, 30, 28, 60, 62, 48, 54, 34, 38, 50...
$ out1.cost
$ out2.event
             <fctr> Yes, Yes, No, No, No, Yes, No, No, Yes, Yes, ...
$ out3.time
             <int> 99, 112, 89, 121, 120, 91, 101, 101, 103, 89, 115,...
             <fctr> Control, Control, Control, Control, Control, Cont...
$ treated_f
$ covB_f
             <fctr> Has B, No B, No B, No B, Has B, No B, No B, Has B...
$ out2_f
             <fctr> Event, Event, No Event, No Event, No Event, No Ev...
             <dbl> 1, 1, 0, 0, 0, 0, 1, 0, 0, 1, 1, 1, 1, 1, 1, 1, 1, ...
$ out2
$ covF.Low
             <dbl> 0, 1, 0, 0, 0, 0, 1, 0, 0, 1, 0, 1, 1, 0, 0, 0, 0, ...
$ covF.Middle <dbl> 0, 0, 1, 1, 1, 0, 0, 1, 1, 0, 1, 0, 0, 1, 0, 1, 1,...
$ covF.High
             <dbl> 1, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0,...
             <dbl> 0.0144, 0.0025, 0.4624, 0.2116, 0.1156, 0.0225, 0....
$ Asqr
$ BC
             <dbl> 7.6, 0.0, 0.0, 0.0, 9.9, 0.0, 0.0, 11.0, 0.0, 10.8...
$ BD
             <dbl> 6.7, 0.0, 0.0, 0.0, 10.7, 0.0, 0.0, 9.2, 0.0, 12.1...
```

3.2 Summary

summary(toy)

```
subject
                     treated
                                                         covB
                                       covA
Min.
                                         :0.0400
                                                           :0.00
      : 1.00
                  Min.
                         :0.00
                                  Min.
                                                    Min.
1st Qu.: 50.75
                  1st Qu.:0.00
                                  1st Qu.:0.2675
                                                    1st Qu.:0.00
                  Median:0.00
Median :100.50
                                  Median :0.5100
                                                    Median:0.00
      :100.50
                         :0.35
Mean
                  Mean
                                  Mean
                                         :0.4995
                                                    Mean
                                                           :0.37
3rd Qu.:150.25
                  3rd Qu.:1.00
                                  3rd Qu.:0.7300
                                                    3rd Qu.:1.00
Max.
       :200.00
                  Max.
                         :1.00
                                  Max.
                                         :0.9900
                                                    Max.
                                                           :1.00
                       covD
                                         covE
                                                           covF
     covC
Min.
       : 4.200
                  Min.
                         : 5.500
                                    Min.
                                           : 2.00
                                                     1-Low
                                                             :67
1st Qu.: 8.600
                  1st Qu.: 8.500
                                    1st Qu.: 8.00
                                                     2-Middle:84
```

```
Median: 9.850
                  Median : 9.900
                                    Median :10.00
                                                     3-High :49
                         : 9.863
      : 9.714
Mean
                  Mean
                                    Mean
                                           :10.21
                  3rd Qu.:11.100
3rd Qu.:10.725
                                    3rd Qu.:12.00
                                           :19.00
       :16.200
                  Max.
                         :14.400
Max.
                                    Max.
  out1.cost
                 out2.event
                               out3.time
                                                treated f
                                                              covB f
                 No :103
                                                            No B:126
       :24.00
                                    : 79.0
                                             Treated: 70
Min.
                            Min.
1st Qu.:39.00
                 Yes: 97
                            1st Qu.:101.0
                                             Control:130
                                                            Has B: 74
Median :49.00
                            Median :109.0
Mean
       :50.42
                            Mean
                                    :109.2
3rd Qu.:58.00
                            3rd Qu.:118.0
Max.
       :80.00
                            Max.
                                    :151.0
     out2_f
                                                   covF.Middle
                     out2
                                    covF.Low
                       :0.000
        : 97
                                        :0.000
                                                         :0.00
Event
                Min.
                                 Min.
                                                  Min.
                                                  1st Qu.:0.00
No Event: 103
                1st Qu.:0.000
                                 1st Qu.:0.000
                Median : 0.000
                                 Median : 0.000
                                                  Median:0.00
                Mean
                       :0.485
                                 Mean
                                        :0.335
                                                  Mean
                                                         :0.42
                3rd Qu.:1.000
                                 3rd Qu.:1.000
                                                  3rd Qu.:1.00
                Max.
                       :1.000
                                        :1.000
                                                         :1.00
                                 Max.
                                                  Max.
  covF.High
                                          BC
                                                            BD
                      Asgr
Min.
      :0.000
                 Min.
                        :0.00160
                                    Min.
                                           : 0.000
                                                      Min.
                                                             : 0.000
1st Qu.:0.000
                 1st Qu.:0.07157
                                    1st Qu.: 0.000
                                                      1st Qu.: 0.000
Median :0.000
                 Median :0.26020
                                    Median : 0.000
                                                      Median : 0.000
                 Mean
Mean
       :0.245
                        :0.32389
                                    Mean
                                           : 3.462
                                                             : 3.635
                                                      Mean
3rd Qu.:0.000
                 3rd Qu.:0.53290
                                    3rd Qu.: 8.450
                                                      3rd Qu.: 9.300
                                                             :13.200
Max.
       :1.000
                 Max.
                        :0.98010
                                    Max.
                                           :13.700
                                                      Max.
```

3.3 Table 1

Note that the factors I created for the out2 outcome are not well ordered for a Table 1, but are well ordered for other tables we'll fit later. So, in this case, I'll use the numeric version of the out2 outcome, but the new factor representations of covB and treated.

```
Stratified by treated f
                       Treated
                                       Control
                                                               test
                                                       р
n
                            70
                                           130
covA (mean (sd))
                         0.59 (0.22)
                                          0.45 (0.29)
                                                        <0.001
covB_f = Has B (\%)
                            34 (48.6)
                                            40 (30.8)
                                                         0.020
covC (mean (sd))
                         9.63 (2.02)
                                         9.76 (2.02)
                                                         0.649
covD (mean (sd))
                        10.10 (1.88)
                                         9.74 (1.77)
                                                         0.183
covE (mean (sd))
                         9.67 (2.74)
                                         10.50 (3.54)
                                                         0.090
covF (%)
                                                         0.266
   1-Low
                            19 (27.1)
                                            48 (36.9)
   2-Middle
                            30 (42.9)
                                            54 (41.5)
   3-High
                            21 (30.0)
                                            28 (21.5)
                        60.61 (16.56)
                                        44.94 (9.69)
                                                        < 0.001
out1.cost (mean (sd))
                                            58 (44.6)
                            39 (55.7)
out2 = 1 (\%)
                                                         0.177
out3.time (mean (sd)) 113.44 (10.75) 106.95 (11.94) <0.001
```

4 The 13 Tasks We'll Tackle in this Example

- 1. Ignoring the covariate information, what is the unadjusted point estimate (and 95% confidence interval) for the effect of the treatment on each of the three outcomes (out1.cost, out2.event, and out3.time)?
- 2. Assume that theory suggests that the square of covA, as well as the interactions of covB with covC and covB with covD should be related to treatment assignment. Fit a propensity score model to the data, using the six covariates (A-F) and the three transformations (A², and the B-C and B-D interactions.) Plot the resulting propensity scores, by treatment group, in an attractive and useful way.
- 3. Use Rubin's Rules to assess the overlap of the propensity scores and the individual covariates prior to the use of any propensity score adjustments.
- 4. Use 1:1 greedy matching to match all 70 treated subjects to control subjects without replacement on the basis of the linear propensity for treatment. Evaluate the degree of covariate imbalance before and after propensity matching for each of the six covariates, and present the pre- and post-match standardized differences and variance ratios for the covariates, as well as the square term and interactions, as well as both the raw and linear propensity score in appropriate plots. Now, build a new data frame containing the propensity-matched sample, and use it to first check Rubin's Rules after matching.
- 5. Now, use the matched sample data set to evaluate the treatment's average causal effect on each of the three outcomes. In each case, specify a point estimate (and associated 95% confidence interval) for the effect of being treated (as compared to being a control subject) on the outcome. Compare your results to the automatic versions reported by the Matching package when you include the outcome in the matching process.
- 6. Now, instead of matching, instead subclassify the subjects into quintiles by the raw propensity score. Display the balance in terms of standardized differences by quintile for the covariates, their transformations, and the propensity score in an appropriate table or plot(s). Are you satisfied?
- 7. Regardless of your answer to the previous question, use the propensity score quintile subclassification approach to find a point estimate (and 95% confidence interval) for the effect of the treatment on each outcome.
- 8. Now using a reasonable propensity score weighting strategy, assess the balance of each covariate, the transformations and the linear propensity score prior to and after propensity weighting. Is the balance after weighting satisfactory?
- 9. Using propensity score weighting to evaluate the treatment's effect, developing a point estimate and 95% CI for the average causal effect of treatment on each outcome.
- 10. Finally, use direct adjustment for the linear propensity score on the entire sample to evaluate the treatment's effect, developing a point estimate and 95% CI for each outcome.
- 11. Now, try a double robust approach. Weight, then adjust for linear propensity score.
- 12. Compare your conclusions about the average causal effect obtained in the following six ways to each other. What happens and why? Which of these methods seems most appropriate given the available information?
 - without propensity adjustment,
 - after propensity matching,
 - ullet after propensity score subclassification,
 - after propensity score weighting,
 - after adjusting for the propensity score directly, and
 - after weighting then adjusting for the PS, to each other.
- 13. Perform a sensitivity analysis for your matched samples analysis and the first outcome (out1.cost) if it turns out to show a statistically significant treatment effect.

5 Task 1. Ignoring covariates, estimate the effect of treatment vs. control on...

5.1 Outcome 1 (a continuous outcome)

toy %>%

Our first outcome describes a quantitative measure, cost, and we're asking what the effect of treatment as compared to control is on that outcome. Starting with brief numerical summaries:

```
group_by(treated_f) %>%
    summarize(n = length(out1.cost), mean = mean(out1.cost),
              sd = sd(out1.cost), min = min(out1.cost),
              Q1 = quantile(out1.cost, 0.25), median = median(out1.cost),
              Q3 = quantile(out1.cost, 0.75), max = max(out1.cost))
# A tibble: 2 x 9
  treated f
                n mean
                            sd
                                min
                                        Q1 median
                                                      QЗ
  <fctr>
            <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
1 Treated
               70 60.6 16.6
                                24.0 50.2
                                              62.0 75.8 80.0
              130 44.9 9.69 24.0 38.0
                                              47.0 51.0 78.0
2 Control
It looks like the Treated group has higher costs than the Control group. To model this, we could use a linear
regression model to obtain a point estimate and 95% confidence interval. Here, I prefer to use the numeric
version of the treated variable, with 0 = "control" and 1 = "treated".
unadj.out1 <- lm(out1.cost ~ treated, data=toy)</pre>
summary(unadj.out1); confint(unadj.out1, level = 0.95) ## provides treated effect and CI estimates
Call:
lm(formula = out1.cost ~ treated, data = toy)
Residuals:
                              3Q
    Min
             1Q Median
                                     Max
                          9.062 33.062
-36.614 -7.945
                  2.062
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
                           1.098 40.932 < 2e-16 ***
(Intercept)
              44.938
                           1.856 8.447 6.36e-15 ***
treated
              15.676
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 12.52 on 198 degrees of freedom
Multiple R-squared: 0.2649,
                               Adjusted R-squared: 0.2612
F-statistic: 71.35 on 1 and 198 DF, p-value: 6.364e-15
              2.5 %
                      97.5 %
(Intercept) 42.7734 47.10352
treated
            12.0162 19.33544
We can store these results in a data frame, with the tidy function from the broom package.
temp <- tidy(unadj.out1, conf.int = TRUE, conf.level = 0.95)</pre>
temp
```

term estimate std.error statistic p.value conf.low conf.high

```
1 (Intercept) 44.93846 1.097891 40.931619 1.434004e-98 42.7734 47.10352
2 treated 15.67582 1.855775 8.447051 6.363978e-15 12.0162 19.33544
```

Our unadjusted treatment effect estimate is an increase of 15.68 in cost, with 95% confidence interval (12.02, 19.34).

I should mention that the broom package also has a useful function called glance which lets you get some detailed summaries of the model.

```
glance(unadj.out1)
```

5.2 Outcome 2 (a binary outcome)

5.2.1 Using a 2x2 table in standard epidemiological format

Thanks to our preliminary cleanup, it's relatively easy to obtain a table in standard epidemiological format comparing treated to control subjects in terms of out2:

```
table(toy$treated_f, toy$out2_f)
```

```
Event No Event Treated 39 31 Control 58 72
```

Note that the exposure is in the rows, with "Having the Exposure" or "Treated" at the top, and the outcome is in the columns, with "Yes" or "Outcome Occurred" or "Event Occurred" on the left, so that the top left cell count describes people that had both the exposure and the outcome. That's *standard epidemiological format*, just what we need for the twoby2 function in the Epi package.

```
twoby2(table(toy$treated_f, toy$out2_f))
```

2 by 2 table analysis:

Outcome : Event

Comparing : Treated vs. Control

```
Event No Event P(Event) 95% conf. interval
Treated 39 31 0.5571 0.4398 0.6685
Control 58 72 0.4462 0.3631 0.5324
```

95% conf. interval
Relative Risk: 1.2488 0.9406 1.6579
Sample Odds Ratio: 1.5617 0.8702 2.8028
Conditional MLE Odds Ratio: 1.5582 0.8354 2.9261
Probability difference: 0.1110 -0.0335 0.2489

Exact P-value: 0.1413 Asymptotic P-value: 0.1352

Eventually, we will be interested in at least two measures - the odds ratio and the risk (probability) difference estimates, and their respective confidence intervals.

- For a difference in risk, our unadjusted treatment effect estimate is an increase of 11.1 percentage points as compared to control, with 95% CI of (-3.4, +24.9) percentage points.
- For an *odds ratio*, our unadjusted treatment effect estimate is an odds ratio of 1.56 (95% CI = 0.87, 2.80) for the event occurring with treatment as compared to control.
- For neither measure is the observed unadjusted effect statistically significant at a 95% confidence level.

5.2.2 Using a logistic regression model

For the odds ratio estimate, we can use a simple logistic regression model to estimate the unadjusted treatment effect, resulting in essentially the same answer. We'll use the numerical (0/1) format to represent binary information, as follows.

```
unadj.out2 <- glm(out2 ~ treated, data=toy, family=binomial())</pre>
summary(unadj.out2)
Call:
glm(formula = out2 ~ treated, family = binomial(), data = toy)
Deviance Residuals:
  Min
            1Q Median
                            3Q
                                   Max
-1.276 -1.087 -1.087
                         1.270
                                  1.270
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.2162
                         0.1764
                                 -1.226
                                            0.220
treated
              0.4458
                         0.2984
                                  1.494
                                            0.135
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 277.08 on 199
                                   degrees of freedom
Residual deviance: 274.83 on 198 degrees of freedom
AIC: 278.83
Number of Fisher Scoring iterations: 3
exp(coef(unadj.out2)) # produces odds ratio estimate
(Intercept)
                treated
  0.8055556
              1.5617353
exp(confint(unadj.out2)) # produces 95% CI for odds ratio
Waiting for profiling to be done...
                2.5 %
                        97.5 %
(Intercept) 0.5683258 1.136896
            0.8721205 2.816455
```

And, again, we can use the tidy function in the broom package to build a tibble of the key parts of the output. Note that by including the exponentiate = TRUE command, our results in the treated row describe the odds ratio, rather than the log odds.

```
tidy(unadj.out2, conf.int = TRUE, exponentiate = TRUE)
```

```
term estimate std.error statistic p.value conf.low conf.high
1 (Intercept) 0.8055556 0.1764365 -1.225501 0.2203865 0.5683258 1.136896
2 treated 1.5617353 0.2983755 1.494082 0.1351541 0.8721205 2.816455
```

- Our odds ratio estimate remains about 1.56, with 95% confidence interval ranging from 0.9 to 2.8, again showing no evidence of a statistically significant impact of the treatment on the occurrence of the event described by out2.
- For practical purposes, the odds ratio and 95% confidence interval obtained here matches the methodology for the twoby2 function.
- The approach implemented in the twoby2 function produces slightly less conservative (i.e. narrower) confidence intervals for the effect estimate than does the approach used in the logistic regression model.

Glancing at the summaries of the model may be helpful in some settings, as well.

```
glance(unadj.out2)
```

```
null.deviance df.null logLik AIC BIC deviance df.residual
1 277.0788 199 -137.416 278.832 285.4286 274.832 198
```

5.3 Outcome 3 (a time-to-event outcome with right censoring)

Our out3.time variable is a variable indicating the time before the event described in out2 occurred. This happened to 97 of the 200 subjects in the data set. For the other 103 subjects who left the study before their event occurred, we have the time before censoring. We can see the results of this censoring in the survival object describing each treatment group.

Here, for instance, is the survival object for the *treated* subjects - the third subject listed here is censored - had the event at some point after 124 weeks (124+) but we don't know precisely when after 124 weeks.

```
Surv(toy$out3.time, toy$out2.event == "Yes")[toy$treated == 1]
                                   109
                                        104
                         103+ 112
                                              98+ 129
                                                       114
[15] 137+ 103
              110 106+ 108 136+
                                    96
                                        125+ 118+
                                                   99+ 101
                                                            109
                                                                 128
                                                                      111+
[29] 120+ 118
              115
                   129+ 118+ 106
                                   106+ 118+ 121
                                                  120
                                                       107
                                                            119+ 105
[43] 106+ 132+ 126+ 130+ 115+ 108
                                    99
                                        122 110
                                                  102
                                                       114+ 120+ 100
                                                                      108
[57] 120 118+ 117+ 123+ 109 122 126+ 126+ 118+ 125+ 103
                                                             94
                                                                 106
```

• To see the controls, we could use Surv(toy\$out3.time, toy\$out2.event=="Yes")[toy\$treated==0]

To deal with the right censoring, we'll use the **survival** package to fit a simple unadjusted Cox proportional hazards model to assess the relative hazard of having the event at a particular time point among treated subjects as compared to controls.

```
unadj.out3 <- coxph(Surv(out3.time, out2.event=="Yes") ~ treated, data=toy)
summary(unadj.out3) ## exp(coef) section indicates relative risk estimate and 95% CI</pre>
```

```
Call:
coxph(formula = Surv(out3.time, out2.event == "Yes") ~ treated,
    data = toy)

n= 200, number of events= 97

    coef exp(coef) se(coef) z Pr(>|z|)
treated -0.1535    0.8577    0.2086 -0.736    0.462
```

```
exp(coef) exp(-coef) lower .95 upper .95
           0.8577
                       1.166
                                0.5698
                                           1.291
treated
Concordance= 0.532 (se = 0.028)
Rsquare= 0.003
                 (max possible= 0.988 )
Likelihood ratio test= 0.55
                             on 1 df,
                                        p=0.46
Wald test
                     = 0.54
                             on 1 df,
                                        p=0.462
Score (logrank) test = 0.54 on 1 df,
                                        p=0.4615
```

The relative hazard rate is shown in the exp(coef) section of the output. Our unadjusted treatment model suggests that the hazard of the outcome is smaller (but not significantly smaller) in the treated group than in the control group. Our estimate is that this relative hazard rate for occurrence of the event associated with treatment as compared to control is 0.86 with a 95% confidence interval of (0.57, 1.29).

Yes, you can tidy this model, as well, using the broom package.

```
tidy(unadj.out3, exponentiate = TRUE)
```

term estimate std.error statistic p.value conf.low conf.high 1 treated 0.8577207 0.2086392 -0.7356088 0.4619688 0.5698386 1.29104

Glancing at the summaries of the model may be helpful, too.

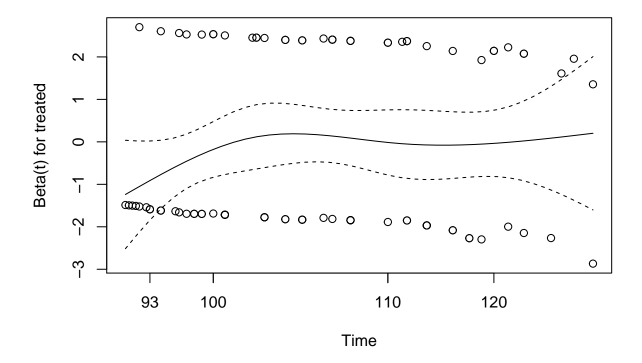
```
glance(unadj.out3)
```

```
n nevent statistic.log p.value.log statistic.sc p.value.sc
1 200
          97
                 0.5460054
                             0.4599545
                                           0.5421382 0.4615479
  statistic.wald p.value.wald
                                r.squared r.squared.max concordance
1
            0.54
                    0.4619688 0.002726304
                                               0.9884724
                                                           0.5318705
                           logLik
  std.error.concordance
                                       AIC
                                                BTC
             0.02796153 -446.0285 894.057 896.6317
1
```

It's wise, whenever fitting a Cox proportional hazards model, to assess the proportional hazards assumption. One way to do this is to run a simple test in R - from which we can obtain a plot, if we like. The idea is for the plot to show no clear patterns over time, and look pretty much like a horizontal line, while we would like the test to be non-significant - if that's the case, our proportional hazards assumption is likely OK.

```
cox.zph(unadj.out3)
```

```
rho chisq p
treated 0.118 1.34 0.247
plot(cox.zph(unadj.out3), var="treated")
```



If the proportional hazards assumption is clearly violated (here it isn't), call a statistician.

5.4 Unadjusted Estimates of Treatment Effect on Outcomes

So, our unadjusted average treatment effect estimates (in each case comparing treated subjects to control subjects) are thus:

Outcome 3				
(Relative Hazard	Outcome 2	Outcome 2 (Risk	Outcome 1 (Cost	Est. Treatment Effect
Rate)	(Odds Ratio)	diff.)	diff.)	(95% CI)
0.86	1.56	+0.11	15.7	No covariate adjustment
(0.57, 1.29)	(0.87, 2.82)	(-0.03, +0.25)	(12.0, 19.3)	(unadjusted)

6 Task 2. Fit the propensity score model, then plot the PS-treatment relationship

I'll use a logistic regression model

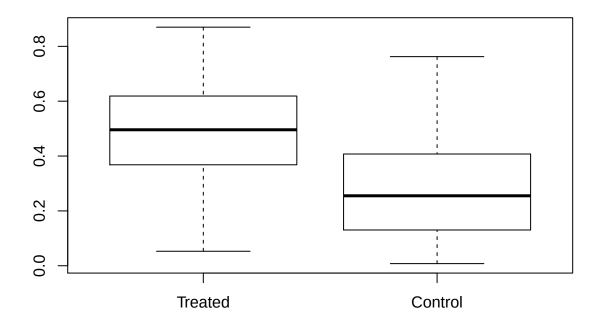
```
glm(formula = treated ~ covA + covB + covC + covD + covE + covF +
    Asqr + BC + BD, family = binomial(), data = toy)
             coef.est coef.se
(Intercept) -5.24
                       1.94
covA
             10.23
                       3.11
covB
             1.07
                       2.47
             0.01
                       0.11
covC
             0.24
                       0.12
covD
covE
             -0.14
                       0.05
                       0.40
covF2-Middle 0.19
covF3-High
             0.59
                       0.45
                       2.83
             -6.91
Asqr
             -0.03
                       0.17
BC
BD
              0.01
                       0.20
  n = 200, k = 11
  residual deviance = 216.9, null deviance = 259.0 (difference = 42.1)
```

Having fit the model, my first step will be to save the raw and linear propensity score values to the main toy example tibble.

```
toy$ps <- psmodel$fitted
toy$linps <- psmodel$linear.predictors</pre>
```

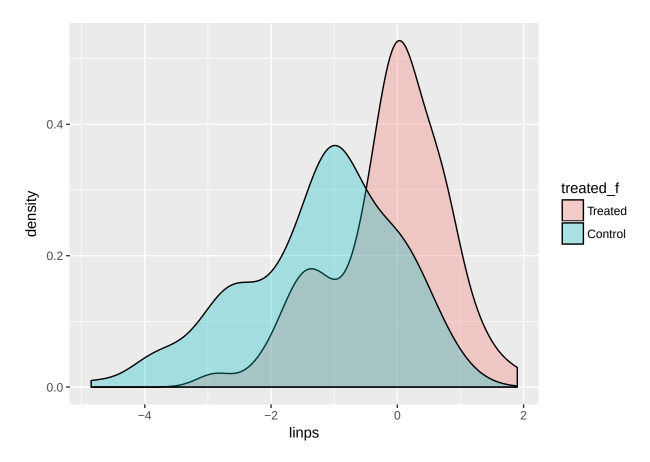
6.1 Comparing the Distribution of Propensity Score Across the Two Treatment Groups

Now, I can use these saved values to assess the propensity model.



I'd rather get a fancier plot to compare the distributions of the propensity score across the two treatment groups, perhaps using a smoothed density estimate, as shown below. Here, I'll show the distributions of the linear propensity score, the log odds of treatment.

```
ggplot(toy, aes(x = linps, fill = treated_f)) +
   geom_density(alpha = 0.3)
```



We see a fair amount of overlap across the two treatment groups. I'll use Rubin's Rules in the next section to help assess the amount of overlap at this point, before any adjustments for the propensity score.

7 Task 3. Rubin's Rules to Check Overlap Before Propensity Adjustment

In his 2001 article¹ about using propensity scores to design studies, as applied to studies of the causal effects of the conduct of the tobacco industry on medical expenditures, Donald Rubin proposed three "rules" for assessing the overlap / balance of covariates appropriately before and after propensity adjustment. Before an outcome is evaluated using a regression analysis (perhaps supplemented by a propensity score adjustment through matching, weighting, subclassification or even direct adjustment), there are three checks that should be performed.

When we do a propensity score analysis, it will be helpful to perform these checks as soon as the propensity model has been estimated, even before any adjustments take place, to see how well the distributions of covariates overlap. After using the propensity score, we hope to see these checks meet the standards below. In what follows, I will describe each standard, and demonstrate its evaluation using the propensity score model we just fit, and looking at the original toy data set, without applying the propensity score in any way to do adjustments.

¹Rubin DB 2001 Using Propensity Scores to Help Design Observational Studies: Application to the Tobacco Litigation. Health Services & Outcomes Research Methodology 2: 169-188. Available on our web site.

7.1 Rubin's Rule 1

Rubin's Rule 1 states that the absolute value of the standardized difference of the linear propensity score, comparing the treated group to the control group, should be close to 0, ideally below 10%, and in any case less than 50%. If so, we may move on to Rule 2.

To evaluate this rule in the toy example, we'll run the following code to place the right value into a variable called rubin1.unadj (for Rubin's Rule 1, unadjusted).

[1] 88.11531

What this does is calculate the (absolute value of the) standardized difference of the linear propensity score comparing treated subjects to control subjects.

- We want this value to be close to 0, and certainly less than 50 in order to push forward to outcomes analysis without further adjustment for the propensity score.
- Clearly, here, with a value of 88%, we can't justify simply running an unadjusted regression model, be it a linear, logistic or Cox model we've got observed selection bias, and need to actually apply the propensity score somehow in order to account for this.
- So, we'll need to match, subclassify, weight or directly adjust for propensity here.

Since we've failed Rubin's 1st Rule, in some sense, we're done checking the rules, because we clearly need to further adjust for observed selection bias - there's no need to prove that further through checking Rubin's 2nd and 3rd rules. But we'll do it here to show what's involved.

7.2 Rubin's Rule 2

Rubin's Rule 2 states that the ratio of the variance of the linear propensity score in the treated group to the variance of the linear propensity score in the control group should be close to 1, ideally between 4/5 and 5/4, but certainly not very close to or exceeding 1/2 and 2. If so, we may move on to Rule 3.

To evaluate this rule in the toy example, we'll run the following code to place the right value into a variable called rubin2.unadj (for Rubin's Rule 2, unadjusted).

```
rubin2.unadj <-with(toy, var(linps[treated==1])/var(linps[treated==0]))
rubin2.unadj</pre>
```

[1] 0.5835438

This is the ratio of variances of the linear propensity score comparing treated subjects to control subjects. We want this value to be close to 1, and certainly between 0.5 and 2. In this case, we pass Rule 2, if just barely.

7.3 Rubin's Rule 3

For Rubin's Rule 3, we begin by calculating regression residuals for each covariate of interest (usually, each of those included in the propensity model) regressed on a single predictor - the linear propensity score. We then look to see if the ratio of the variance of the residuals of this model for the treatment group divided by the variance of the residuals of this model for the control group is close to 1. Again, ideally this will fall between 4/5 and 5/4 for each covariate, but certainly between 1/2 and 2. If so, then the use of regression models seems well justified.

To evaluate Rubin's 3rd Rule, we'll create a little function to help us do the calculations.

```
## General function rubin3 to help calculate Rubin's Rule 3
rubin3 <- function(data, covlist, linps) {
   covlist2 <- as.matrix(covlist)
   res <- NA
   for(i in 1:ncol(covlist2)) {
      cov <- as.numeric(covlist2[,i])
      num <- var(resid(lm(cov ~ data$linps))[data$exposure == 1])
      den <- var(resid(lm(cov ~ data$linps))[data$exposure == 0])
      res[i] <- round(num/den, 3)
   }
   final <- data_frame(name = names(covlist), resid.var.ratio = res)
   return(final)
}</pre>
```

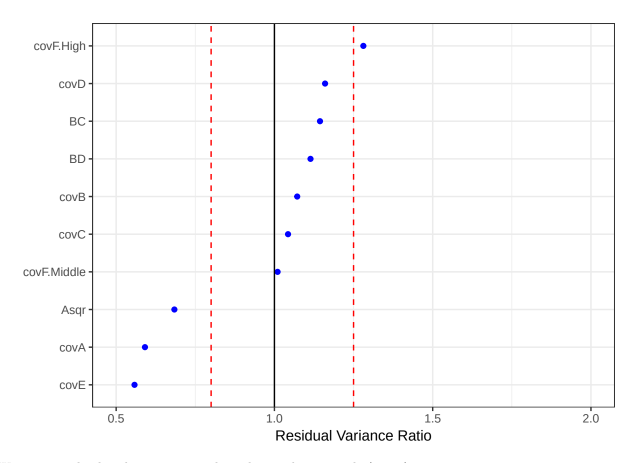
Now, then, applying the rule to our sample prior to propensity score adjustment, we get the following result. Note that I'm using the indicator variable forms for the covF information.

```
# A tibble: 10 x 2
   name
               resid.var.ratio
   <chr>>
                          <dbl>
 1 covA
                          0.591
2 covB
                          1.07
3 covC
                          1.04
 4 covD
                          1.16
5 covE
                          0.558
6 covF.Middle
                          1.01
7 covF.High
                          1.28
8 Asgr
                          0.684
9 BC
                          1.14
10 BD
                          1.11
```

Some of these covariates look to have residual variance ratios near 1, while others are further away, but all are within the (0.5, 2.0) range. So we'd pass Rule 3 here, although we'd clearly like to see some covariates (A and E, in particular) with ratios closer to 1.

7.3.1 A Cleveland Dot Chart of the Rubin's Rule 3 Results

```
ggplot(rubin3.unadj, aes(x = resid.var.ratio, y = reorder(name, resid.var.ratio))) +
    geom_point(col = "blue") +
    theme_bw() +
    xlim(0.5, 2.0) +
    geom_vline(aes(xintercept = 1)) +
    geom_vline(aes(xintercept = 4/5), linetype = "dashed", col = "red") +
    geom_vline(aes(xintercept = 5/4), linetype = "dashed", col = "red") +
    labs(x = "Residual Variance Ratio", y = "")
```



We see several values between 0.5 and 0.8, but nothing outside (0.5, 2).

8 Task 4. Use 1:1 greedy matching on the linear PS, then check post-match balance

As requested, we'll do 1:1 greedy matching on the linear propensity score without replacement and breaking ties randomly. To start, we won't include an outcome variable in our call to the Match function within the Matching package We'll wind up with a match including 70 treated and 70 control subjects.

```
X <- toy$linps ## matching on the linear propensity score
Tr <- as.logical(toy$treated)
match1 <- Match(Tr=Tr, X=X, M = 1, replace=FALSE, ties=FALSE)
summary(match1)</pre>
```

```
Estimate... 0
SE..... 0
T-stat.... NaN
p.val..... NA

Original number of observations..... 20
Original number of treated obs..... 70
Matched number of observations..... 70
Matched number of observations (unweighted). 70
```

8.1 Balance Assessment (Semi-Automated)

Next, we'll assess the balance imposed by this greedy match on our covariates, and their transformations (A^2 and B*C and B*D) as well as the raw and linear propensity scores. The default output from the MatchBalance function is extensive...

**** (V1) covA ****		
	Before Matching	After Matching
mean treatment		0.59071
mean control	0.45046	0.55443
std mean diff	63.684	16.476
mean raw eQQ diff	0.15014	0.054286
med raw eQQ diff	0.175	0.05
max raw eQQ diff	0.28	0.14
mean eCDF diff	0.15106	0.059127
med eCDF diff	0.18352	0.042857
max eCDF diff	0.28352	0.2
var ratio (Tr/Co)	0.58886	0.83347
T-test p-value		0.29257
KS Bootstrap p-value		0.084
KS Naive p-value		0.12159
KS Statistic		0.2
No bodolbolo	0.20002	0.2
***** (V2) covB ****		
(12) 3312	Before Matching	After Matching
mean treatment	0.48571	0.48571
mean control	0.30769	0.38571
std mean diff		19.865
Bod medir dili	00.001	10.000
mean raw eQQ diff	0.18571	0.1
med raw eQQ diff	0	0
max raw eQQ diff	1	1
mean eCDF diff	0.089011	0.05
med eCDF diff	0.089011	0.05
max eCDF diff		0.1
var ratio (Tr/Co)	1.1805	1.0543
T-test p-value		0.22225
<u> </u>	· · · · · ·	· -
***** (V3) covC *****		
•	Before Matching	After Matching
mean treatment	9.6257	9.6257
mean control	9.7623	9.7329

std mean diff	-6.7767	-5.3156
mean raw eQQ diff	0.18714	0.26429
med raw eQQ diff	0.1	0.2
max raw eQQ diff	1.7	1.7
mean eCDF diff	0.021871	0.031027
med eCDF diff	0.012088	0.028571
max eCDF diff	0.078022	0.1
var ratio (Tr/Co)	0.99092	0.98676
T-test p-value	0.64881	0.76698
KS Bootstrap p-value	0.87	0.804
KS Naive p-value	0.94461	0.8752
KS Statistic	0.078022	0.1
AD DUALIBUTE	0.070022	0.1
**** (V4) covD ****		
	Before Matchin	ng After Matching
mean treatment	10.096	10.096
mean control	9.7377	10.103
std mean diff	19.074	-0.38055
mean raw eQQ diff	0.43143	0.23286
med raw eQQ diff	0.4	0.1
max raw eQQ diff.	1	1.5
mean eCDF diff	0.053676	0.024654
med eCDF diff	0.050549	0.024004
max eCDF diff	0.10989	0.028371
max ecur dili	0.10969	0.071429
var ratio (Tr/Co)	1.1249	1.2306
T-test p-value	0.19162	0.9803
KS Bootstrap p-value	0.482	0.98
KS Naive p-value	0.64191	0.99407
KS Statistic	0.10989	0.071429
No bodolbole	0.10303	0.011425
**** (V5) covE ****		
	Before Matchin	ng After Matching
mean treatment	9.6714	9.6714
mean control	10.5	10.029
std mean diff	-30.199	-13.017
mean raw eQQ diff	1.0429	0.87143
med raw eQQ diff	1	1
max raw eQQ diff	3	3
mean eCDF diff	0.058242	0.058095
med eCDF diff	0.05	0.057143
max eCDF diff	0.12857	0.11429
var ratio (Tr/Co)	0.60225	0.61843
T-test p-value	0.068094	0.48672

KS Bootstrap p-value KS Naive p-value KS Statistic	0.258 0.43948 0.12857	0.504 0.75053 0.11429
***** (V6) covF2-Middle	· ****	
	Before Matching	After Matching
mean treatment	0.42857	0.42857
mean control	0.41538	0.38571
std mean diff	2.6456	8.5982
stu mean ulli	2.0430	0.0902
mean raw eQQ diff	0.014286	0.042857
med raw eQQ diff	0	0
max raw eQQ diff	1	1
mean eCDF diff	0.0065024	0.001400
	0.0065934	0.021429
med eCDF diff	0.0065934	0.021429
max eCDF diff	0.013187	0.042857
var ratio (Tr/Co)	1.0152	1.0336
T-test p-value	0.85825	0.59098
r		
(177)		
***** (V7) covF3-High *		After Motahing
	Before Matching	•
mean treatment	0.3	0.3
mean control	0.21538	0.27143
std mean diff	18.332	6.1901
mean raw eQQ diff	0.085714	0.028571
med raw eQQ diff	0	0
max raw eQQ diff	1	1
mean eCDF diff	0.042308	0.014286
med eCDF diff	0.042308	0.014286
max eCDF diff	0.084615	0.028571
var ratio (Tr/Co)	1.251	1.0619
T-test p-value	0.20201	0.65566
-		
**** (V8) Asqr ****		
May That	Before Matching	After Matching
mean treatment	0.39675	0.39675
mean control		0.36475
	0.28465	
std mean diff	45.623	13.023
mean raw eQQ diff	0.12648	0.063506
med raw eQQ diff	0.1308	0.06825
max raw eQQ diff	0.2544	0.1568
	0.2011	3.1000
mean eCDF diff	0.15106	0.059127
med eCDF diff	0.18352	0.042857
max eCDF diff	0.28352	0.2

var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value KS Statistic	< 2.22e-16 0.0013313	0.77694 0.43078 0.084 0.12159 0.2
***** (V9) BC ****		
	Before Matching	After Matching
mean treatment	4.5843	4.5843
${\tt mean control}$	2.8569	3.5614
std mean diff	34.494	20.426
mean raw eQQ diff	1.78	1.0229
med raw eQQ diff		0
max raw eQQ diff	8.8	7.8
mean eCDF diff	0.099473	0.062458
med eCDF diff	0.099451	0.071429
max eCDF diff.	0.18681	0.12857
var ratio (Tr/Co)	1.2672	1.1369
T-test p-value	0.017031	0.19412
KS Bootstrap p-value	0.026	0.348
KS Naive p-value	0.083513	0.60929
KS Statistic	0.18681	0.12857
***** (V10) BD ****	Before Matching	After Matching
***** (V10) BD ***** mean treatment	Before Matching 4.8157	After Matching 4.8157
mean treatment	4.8157	4.8157
mean treatment mean control	4.8157 3 35.245	4.8157 3.7786
mean treatment mean control std mean diff	4.8157 3 35.245	4.8157 3.7786 20.132
mean treatment mean control std mean diff mean raw eQQ diff	4.8157 3 35.245 1.8629	4.8157 3.7786 20.132 1.0371
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff	4.8157 3 35.245 1.8629 0 9.5	4.8157 3.7786 20.132 1.0371 0 8.3
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011	4.8157 3.7786 20.132 1.0371 0 8.3
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff	4.8157 3 35.245 1.8629 0 9.5	4.8157 3.7786 20.132 1.0371 0 8.3
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co)	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967 0.03	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157 0.6
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967 0.03	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157 0.6
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967 0.03 0.10039	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157 0.6 0.8752
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value KS Statistic	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967 0.03 0.10039	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157 0.6 0.8752
mean treatment mean control std mean diff mean raw eQQ diff med raw eQQ diff max raw eQQ diff mean eCDF diff med eCDF diff var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value KS Statistic	4.8157 3 35.245 1.8629 0 9.5 0.10274 0.1011 0.18132 1.2538 0.014967 0.03 0.10039 0.18132	4.8157 3.7786 20.132 1.0371 0 8.3 0.065306 0.071429 0.1 1.1086 0.2157 0.6 0.8752 0.1

std mean diff	101.45	37.975
mean raw eQQ diff med raw eQQ diff max raw eQQ diff	0.19569 0.21857 0.26218	0.073618 0.081407 0.1385
mean eCDF diff med eCDF diff max eCDF diff	0.26132 0.25165 0.47253	0.1249 0.14286 0.25714
var ratio (Tr/Co) T-test p-value KS Bootstrap p-value KS Naive p-value KS Statistic	1.7258e-10 < 2.22e-16 9.8491e-10	1.4126 2.591e-06 0.014 0.019182 0.25714

***** (V12) linps *****

(VIZ) IIIpo		
	Before Matching	g After Matching
mean treatment	-0.13912	-0.13912
mean control	-1.2152	-0.45294
std mean diff	117.3	34.211
mean raw eQQ diff	1.0997	0.3201
med raw eQQ diff	1.0798	0.33541
max raw eQQ diff	1.9834	0.73393
mean eCDF diff	0.26132	0.1249
med eCDF diff	0.25165	0.14286
max eCDF diff	0.47253	0.25714
var ratio (Tr/Co)	0.58354	1.4158
T-test p-value	3.3878e-11	3.1952e-06
KS Bootstrap p-value	< 2.22e-16	0.014
KS Naive p-value	9.8491e-10	0.019182
KS Statistic	0.47253	0.25714

```
Before Matching Minimum p.value: < 2.22e-16
```

Variable Name(s): covA Asqr ps linps Number(s): 1 8 11 12

```
After Matching Minimum p.value: 2.591e-06 Variable Name(s): ps Number(s): 11
```

The cobalt package has some promising tools for taking this sort of output and turning it into something useful. We'll look at that approach soon. For now, some old-school stuff...

8.2 Extracting, Tabulating and Plotting Standardized Differences (without cobalt)

We'll start by naming the covariates that the MatchBalance output contains...

```
"A^2","B*C", "B*D", "raw PS", "linear PS")
```

The next step is to extract the standardized differences (using the pooled denominator to estimate, rather than the treatment-only denominator used in the main output above.)

```
pre.szd <- NULL; post.szd <- NULL
for(i in 1:length(covnames)) {
  pre.szd[i] <- mb1$BeforeMatching[[i]]$sdiff.pooled
  post.szd[i] <- mb1$AfterMatching[[i]]$sdiff.pooled
}</pre>
```

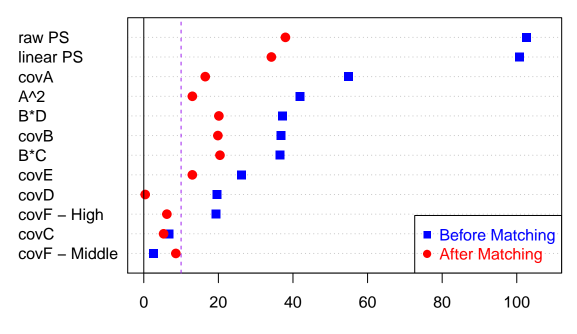
Now, we can build a table of the standardized differences:

```
temp <- data.frame(pre.szd, post.szd, row.names=covnames)
print(temp, digits=3)</pre>
```

```
pre.szd post.szd
                54.83
                       16.476
COVA
covB
                36.80
                       19.865
covC
                -6.76
                        -5.316
covD
                19.63
                        -0.381
               -26.18 -13.017
covE
covF - Middle
                 2.66
                         8.598
                19.33
                         6.190
covF - High
A^2
                41.90
                       13.023
                36.47
                        20.426
B*C
B*D
                37.18
                        20.132
raw PS
               102.60
                        37.975
linear PS
               100.70
                        34.211
```

And then, more usefully, we can plot the absolute values of the standardized differences...

Absolute Standardized Difference Plot

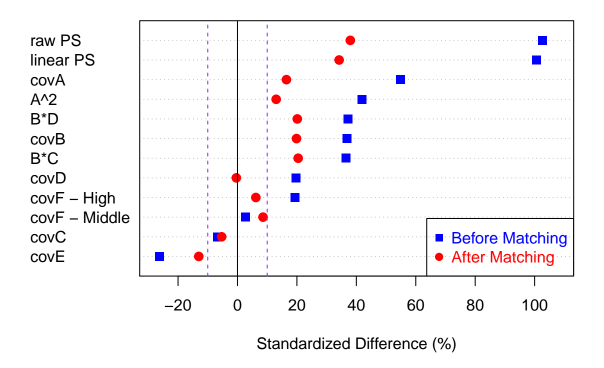


Absolute Standardized Difference (%)

Or, we can plot the standardized differences with their signs intact...

```
temp <- data.frame(pre.szd, post.szd, row.names=covnames)</pre>
tempsort <- temp[with(temp, order(pre.szd)), ]</pre>
low <- min(min(pre.szd), min(post.szd), -0.1)</pre>
high <- max(max(pre.szd), max(post.szd), 0.1)
dotchart(tempsort$pre.szd, xlim=c(1.05*low, 1.05*high), pch="",
         labels=row.names(tempsort), main="Standardized Difference Plot",
         xlab="Standardized Difference (%)")
points(tempsort$pre.szd, seq(1:length(tempsort$pre.szd)),
       pch=15, col="blue", cex=1.2)
points(tempsort$post.szd, seq(1:length(tempsort$post.szd)),
       pch=19, col="red", cex=1.2)
abline(v=0, lty=1)
abline(v=10, lty=2, col="purple")
abline(v=-10, lty=2, col="purple")
legend("bottomright", legend = c("Before Matching", "After Matching"),
       col=c("blue", "red"), text.col=c("blue", "red"), bty="o", pch = c(15, 19))
```

Standardized Difference Plot



8.3 Using cobalt to build a "Love Plot" after Matching

```
b <- bal.tab(match1, treated ~ covA + covB + covC + covD + covE + covF + Asqr + BC + BD + ps + linps, data=toy, un = TRUE)
b
```

Balance Measures:

	Туре	Diff.Un	Diff.Adj
covA	Contin.	0.6368	0.1648
covB	Binary	0.1780	0.1000
covC	${\tt Contin.}$	-0.0678	-0.0532
covD	${\tt Contin.}$	0.1907	-0.0038
covE	${\tt Contin.}$	-0.3020	-0.1302
covF_1-Low	Binary	-0.0978	-0.0714
$covF_2$ -Middle	Binary	0.0132	0.0429
covF_3-High	Binary	0.0846	0.0286
Asqr	Contin.	0.4562	0.1302
BC	${\tt Contin.}$	0.3449	0.2043
BD	${\tt Contin.}$	0.3525	0.2013
ps	${\tt Contin.}$	1.0145	0.3798
linps	Contin.	1.1730	0.3421

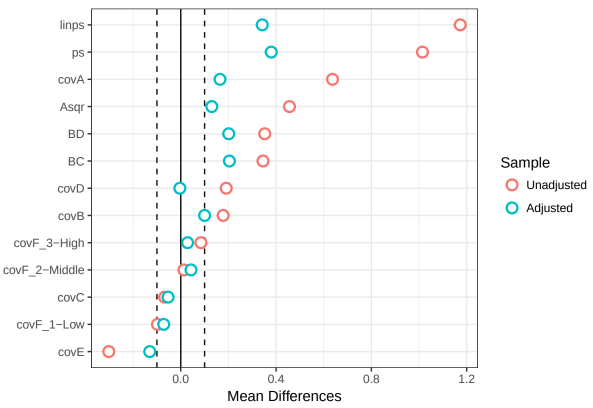
Sample sizes:

Control Treated All 130 70

```
Matched 70 70
Unmatched 60 0
```

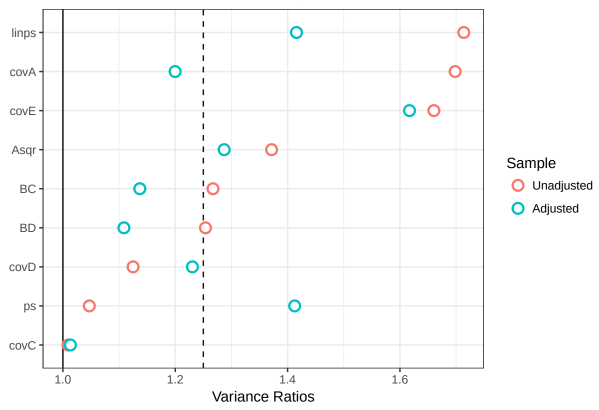
8.3.1 Building a Plot of Standardized Differences, with cobalt

Standardized Differences and 1:1 Matching



8.3.2 Building a Plot of Variance Ratios, with cobalt

Variance Ratios and 1:1 Matching



8.4 Extracting, Tabulating and Plotting Variance Ratios (without cobalt)

Next, we extract the variance ratios, build a table and then plot them.

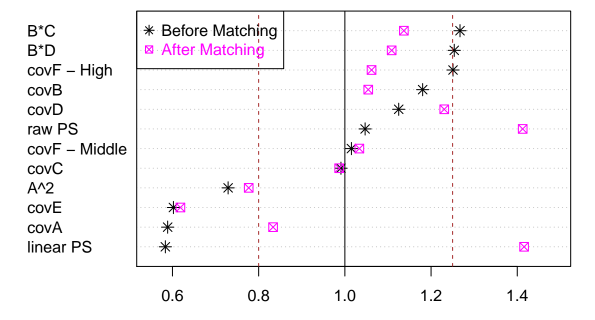
```
pre.vratio <- NULL; post.vratio <- NULL
for(i in 1:length(covnames)) {
   pre.vratio[i] <- mb1$BeforeMatching[[i]]$var.ratio
   post.vratio[i] <- mb1$AfterMatching[[i]]$var.ratio
}

## Table of Variance Ratios
temp <- data.frame(pre.vratio, post.vratio, row.names=covnames)
print(temp, digits=2)</pre>
```

	nro uratio	nost uratio
	pre.vracio	post.vratio
covA	0.59	0.83
covB	1.18	1.05
covC	0.99	0.99
covD	1.12	1.23
covE	0.60	0.62
covF - Middle	1.02	1.03
covF - High	1.25	1.06
A^2	0.73	0.78
B*C	1.27	1.14
B*D	1.25	1.11
raw PS	1.05	1.41

```
linear PS
                    0.58
                                 1.42
## Variance Ratio Plot
temp <- data.frame(pre.vratio, post.vratio, row.names=covnames)</pre>
tempsort <- temp[with(temp, order(pre.vratio)), ]</pre>
low <- min(min(pre.vratio), min(post.vratio))</pre>
high <- max(max(pre.vratio), max(post.vratio))</pre>
dotchart(tempsort$pre.vratio, xlim=c(0.95*low, 1.05*high),
         pch="", labels=row.names(tempsort), main="Plot of Variance Ratios",
         xlab="Treatment Variance / Control Variance")
points(tempsort$pre.vratio, seq(1:length(tempsort$pre.vratio)),
       pch=8, col="black", cex=1.2)
points(tempsort$post.vratio, seq(1:length(tempsort$post.vratio)),
       pch=7, col="magenta", cex=1.2)
abline(v=1, lty=1)
abline(v=4/5, lty=2, col="brown")
abline(v=5/4, lty=2, col="brown")
legend("topleft", legend = c("Before Matching", "After Matching"),
       col=c("black", "magenta"), text.col=c("black", "magenta"),
       bty="o", pch = c(8, 7))
```

Plot of Variance Ratios



Treatment Variance / Control Variance

8.5 Creating a New Data Frame, Containing the Matched Sample (without cobalt)

Now, we build a new matched sample data frame in order to do some of the analyses to come. This will contain only the 140 matched subjects (70 treated and 70 control).

```
matches <- factor(rep(match1$index.treated, 2))
toy.matchedsample <- cbind(matches, toy[c(match1$index.control, match1$index.treated),])</pre>
```

Some sanity checks:

```
table(toy.matchedsample$treated_f)
```

Treated Control 70 70

head(toy.matchedsample)

```
matches subject treated covA covB covC covD covE
                                                           covF out1.cost
                44
                         0 0.76
                                       8.5 10.7
1
      131
                                    0
                                                    11
                                                         3-High
                                                                        32
2
      132
               100
                          0 0.60
                                    1
                                        4.6
                                            7.3
                                                    11
                                                         3-High
                                                                        38
3
      133
                72
                          0 0.68
                                    0
                                        8.1 10.4
                                                    10
                                                          1-Low
                                                                        24
4
      134
                84
                          0 0.61
                                    1
                                        9.5
                                            7.4
                                                    13
                                                         3-High
                                                                        35
5
      135
               108
                          0 0.22
                                    0 11.1 10.1
                                                     7
                                                      2-Middle
                                                                        49
                          0 0.58
                                                                        49
      136
                18
                                    1 10.1 11.5
                                                     9
                                                          1-Low
  out2.event out3.time treated_f covB_f
                                             out2 f out2 covF.Low covF.Middle
1
         Yes
                     94
                           Control
                                     No B
                                              Event
                                                        1
                                                                  0
                                                                               0
2
         Yes
                     98
                           Control
                                    Has B
                                                                  0
                                                                               0
                                              Event
                                                        1
3
                                                                               0
          No
                     95
                           Control
                                     No B No Event
                                                        0
                                                                  1
                                                                  0
                                                                               0
4
                                    Has B No Event
                                                        0
          No
                    122
                           Control
5
         Yes
                    104
                           Control
                                     No B
                                              Event
                                                                  0
                                                                               1
                                                        1
6
         Yes
                    101
                           Control
                                    Has B
                                              Event
                                                        1
                                                                  1
                                                                               0
  covF.High
                      BC
                            BD
                                               linps exposure
               Asqr
                                       ps
1
          1 0.5776
                     0.0
                           0.0 0.5414953
                                           0.1663638
                                                             0
2
           1 0.3600
                     4.6
                           7.3 0.5587964
                                                             0
                                           0.2362787
3
          0 0.4624
                     0.0
                           0.0 0.4074310 -0.3745958
                                                             0
4
          1 0.3721
                    9.5
                          7.4 0.4739821 -0.1041658
                                                             0
5
          0 0.0484 0.0 0.0 0.1605378 -1.6542318
                                                             0
6
          0 0.3364 10.1 11.5 0.6940724 0.8192280
                                                             0
```

8.6 Rubin's Rules to Check Balance After Matching

8.6.1 Rubin's Rule 1

Rubin's Rule 1 states that the absolute value of the standardized difference of the linear propensity score, comparing the treated group to the control group, should be close to 0, ideally below 10%, and in any case less than 50%. If so, we may move on to Rule 2.

Recall that our result without propensity matching (or any other adjustment) was

rubin1.unadj

[1] 88.11531

To run this for our matched sample, we use:

[1] 36.54225

Here, we've at least got this value down below 50%, so we would pass Rule 1, although perhaps a different propensity score adjustment (perhaps by weighting or subclassification, or using a different matching approach) might improve this result by getting it closer to 0.

8.6.2 Rubin's Rule 2

Rubin's Rule 2 states that the ratio of the variance of the linear propensity score in the treated group to the variance of the linear propensity score in the control group should be close to 1, ideally between 4/5 and 5/4, but certainly not very close to or exceeding 1/2 and 2. If so, we may move on to Rule 3.

Recall that our result without propensity matching (or any other adjustment) was

```
rubin2.unadj
```

[1] 0.5835438

To run this for our matched sample, we use:

```
rubin2.match <- with(toy.matchedsample, var(linps[treated==1])/var(linps[treated==0]))
rubin2.match</pre>
```

[1] 1.415833

This is moderately promising - a substantial improvement over our unadjusted result, though not yet within our desired range of 4/5 to 5/4, but clearly within 1/2 to 2. We pass Rule 2, as well.

8.6.3 Rubin's Rule 3

For Rubin's Rule 3, we begin by calculating regression residuals for each covariate of interest (usually, each of those included in the propensity model) regressed on a single predictor - the linear propensity score. We then look to see if the ratio of the variance of the residuals of this model for the treatment group divided by the variance of the residuals of this model for the control group is close to 1. Again, ideally this will fall between 4/5 and 5/4 for each covariate, but certainly between 1/2 and 2. If so, then the use of regression models seems well justified.

Recall that our result without propensity matching (or any other adjustment) was

rubin3.unadj

A tibble: 10 x 2

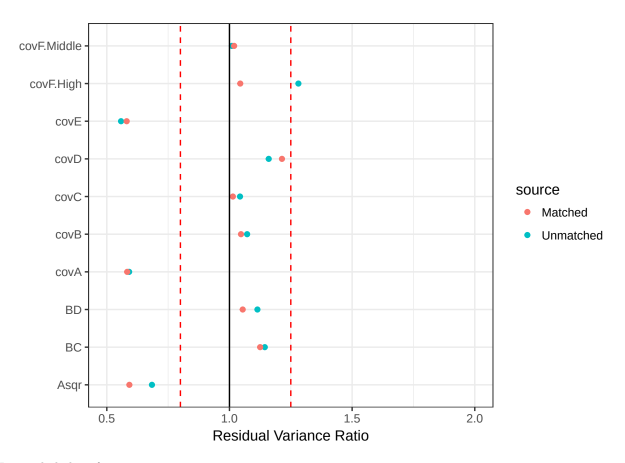
	name	resid.var.ratio
	<chr></chr>	<dbl></dbl>
1	covA	0.591
2	covB	1.07
3	covC	1.04
4	covD	1.16
5	covE	0.558
6	${\tt covF.Middle}$	1.01
7	covF.High	1.28
8	Asqr	0.684
9	BC	1.14
10	BD	1.11

After propensity matching, we use this code to assess Rubin's 3rd Rule in our matched sample.

A tibble: 10 x 2 nameresid.var.ratio <chr> <dbl> 0.583 1 covA 2 covB 1.05 3 covC 1.01 4 covD 1.21 5 covE 0.581 6 covF.Middle 1.02 7 covF.High 1.04 8 Asqr 0.592 9 BC 1.12 10 BD 1.05

It looks like the results are basically unchanged, except that covF.High is improved. The dotplot of these results comparing pre- to post-matching is shown below.

8.6.4 A Cleveland Dot Chart of the Rubin's Rule 3 Results Pre vs. Post-Match



Not a whole lot of improvement to report.

9 Task 5. After matching, estimate the causal effect of treatment on ...

9.1 Outcome 1 (a continuous outcome)

9.1.1 Approach 1. Automated Approach from the Matching Library - ATT Estimate

First, we'll look at the essentially automatic answer which can be obtained when using the Matching package and inserting an outcome Y. For a continuous outcome, this is often a reasonable approach.

```
X <- toy$linps ## matching on the linear propensity score
Tr <- as.logical(toy$treated)
Y <- toy$out1.cost
match1 <- Match(Y=Y, Tr=Tr, X=X, M = 1, replace=FALSE, ties=FALSE)
summary(match1)</pre>
```

Estimate... 15.557 SE..... 2.0397 T-stat.... 7.6273 p.val..... 2.3981e-14

Original number of observations..... 200

We can obtain an approximate 95% confidence interval by adding and subtracting 1.96 times (or just double) the standard error (SE) to the point estimate, which is 15.6 here. Here, using the 1.96 figure, that would yields an approximate 95% CI of (11.6, 19.6).

9.1.2 Approach 2. Automated Approach from the Matching Library - ATE Estimate

```
match1.ATE <- Match(Y=Y, Tr=Tr, X=X, M = 1, replace=FALSE, ties=FALSE, estimand="ATE")
Warning in Match(Y = Y, Tr = Tr, X = X, M = 1, replace = FALSE, ties =
FALSE, : replace==FALSE, but there are more (weighted) control obs than
treated obs. Some control obs will not be matched. You may want to estimate
ATT instead.
summary(match1.ATE)
Estimate... 16.014
SE..... 1.5568
T-stat..... 10.287
p.val..... < 2.22e-16
Original number of observations.....
Original number of treated obs.....
                                            70
Matched number of observations..... 140
Matched number of observations (unweighted).
rm(match1.ATE)
```

And our 95% CI for this ATE estimate would be $16.014 \pm 1.96(1.5614)$, or (13.0, 19.1), but we'll stick with the ATT estimate for now.

9.1.3 ATT vs. ATE: Definitions

- Informally, the average treatment effect on the treated (ATT) estimate describes the difference in potential outcomes (between treated and untreated subjects) summarized across the population of people who actually received the treatment.
 - In our initial match, we identified a unique and nicely matched control patient for each of the 70 people in the treated group. We have a 1:1 match on the treated, and thus can describe subjects across that set of treated patients reasonably well.
- On the other hand the **average treatment effect** (ATE) refers to the difference in potential outcomes summarized across the entire population, including those who did not receive the treatment.
 - In our ATE match, we have less success, in part because if we match to the treated patients in a 1:1 way, we'll have an additional 60 unmatched control patients, about whom we can describe results only vaguely. We could consider matching up control patients to treated patients, perhaps combined with a willingness to re-use some of the treated patients to get a better estimate across the whole population.

9.1.4 Approach 3. Mirroring the Paired T test in a Regression Model

We can mirror the paired t test result in a regression model that treats the match identifier as a fixed factor in a linear model, as follows. This takes the pairing into account, but treating pairing as a fixed, rather than random, factor, isn't really satisfactory as a solution, although it does match the paired t test.

```
adj.m.out1 <- lm(out1.cost ~ treated + factor(matches), data=toy.matchedsample)
coef(adj.m.out1)["treated"] # point estimate for treated effect

treated
15.55714
confint(adj.m.out1)["treated",1] # lower limit of 95% CI

[1] 11.45873
confint(adj.m.out1)["treated",2] # lower limit of 95% CI

[1] 19.65556</pre>
```

So, this regression approach produces an estimate that is exactly the same as the paired t $test^2$, but this isn't something I'm completely comfortable with.

9.1.5 Approach 4. A Mixed Model to account for 1:1 Matching

What I think of as a more appropriate result comes from a mixed model where the matches are treated as a random factor, but the treatment group is treated as a fixed factor. This is developed like this, using the lme4 package. Note that we have to create a factor variable to represent the matches, since that's the only thing that lme4 understands.

```
toy.matchedsample$matches.f <- as.factor(toy.matchedsample$matches)</pre>
## Need to use matches as a factor in R here
mixedmodel.out1 <- lmer(out1.cost ~ treated + (1 | matches.f), data=toy.matchedsample)
summary(mixedmodel.out1); confint(mixedmodel.out1)
Linear mixed model fit by REML ['lmerMod']
Formula: out1.cost ~ treated + (1 | matches.f)
   Data: toy.matchedsample
REML criterion at convergence: 1122.5
Scaled residuals:
             1Q Median
   Min
                             3Q
                                    Max
-2.3102 -0.5639 0.1046 0.6678 1.9395
Random effects:
Groups
           Name
                       Variance Std.Dev.
                                 6.727
matches.f (Intercept) 45.25
Residual
                       147.72
                                12.154
Number of obs: 140, groups: matches.f, 70
Fixed effects:
            Estimate Std. Error t value
(Intercept)
             45.057
                          1.660 27.137
```

²I'll leave checking that this is true as an exercise for the curious.

```
treated
              15.557
                           2.054
                                   7.573
Correlation of Fixed Effects:
        (Intr)
treated -0.619
Computing profile confidence intervals ...
                2.5 %
                          97.5 %
.sig01
             0.503528 9.895039
.sigma
            10.313755 14.378709
(Intercept) 41.802672 48.311613
            11.503977 19.610308
treated
```

9.1.6 Practically, does any of this matter in this example?

Not in this example, no, as long as you stick to the ATT approaches.

Approach	Effect Estimate	Standard Error	95% CI
"Automated" ATT via Match	15.56	2.04	11.56, 19.55
Linear Model (pairs as fixed factor)	15.56	2.05	11.46, 19.66
Mixed Model (pairs as random factor)	15.57	2.05	11.50, 19.61

9.2 Outcome 2 (a binary outcome)

9.2.1 Approach 1. Automated Approach from the Matching Library (ATT)

First, we'll look at the essentially automatic answer which can be obtained when using the Matching package and inserting an outcome Y. For a binary outcome, this is often a reasonable approach, especially if you don't wish to adjust for any other covariate, and the result will be expressed as a risk difference, rather than as a relative risk or odds ratio. Note that I have used the 0-1 version of Outcome 2, rather than a factor version. The estimate produced is the difference in risk associated with $\mathtt{out2} = 1$ (Treated subjects) minus $\mathtt{out2} = 0$ (Controls.)

```
X <- toy$linps ## matching on the linear propensity score
Tr <- as.logical(toy$treated)
Y <- toy$out2
match1 <- Match(Y=Y, Tr=Tr, X=X, M = 1, replace=FALSE, ties=FALSE)
summary(match1)</pre>
```

```
Estimate... 0.1
SE...... 0.075997
T-stat.... 1.3158
p.val..... 0.18823

Original number of observations...... 200
Original number of treated obs...... 70
Matched number of observations....... 70
Matched number of observations (unweighted). 70
```

As in the continuous case, we obtain an approximate 95% confidence interval by adding and subtracting 1.96 times (or just double) the standard error (SE) to the point estimate, which is 0.1 (i.e. 10 percentage points)

here.

- Here, using the 1.96 figure, that would yields an approximate 95% CI of (-0.05, 0.25).
- Again, the estimated average causal effect is not statistically significant here, since 0 is contained in this confidence interval.

9.2.2 Approach 2. Using the matched sample to perform a conditional logistic regression

Since we have the matched sample available, we can simply perform a conditional logistic regression to estimate the treatment effect in terms of a log odds ratio (or, by exponentiating, an odds ratio.) Again, I use the 0/1 version of both the outcome and treatment indicator. The key modeling function clogit is part of the survival package.

```
adj.m.out2 <- clogit(out2 ~ treated + strata(matches), data=toy.matchedsample)</pre>
summary(adj.m.out2)
Call:
coxph(formula = Surv(rep(1, 140L), out2) ~ treated + strata(matches),
    data = toy.matchedsample, method = "exact")
  n= 140, number of events= 71
          coef exp(coef) se(coef)
                                       z Pr(>|z|)
treated 0.4925
                  1.6364
                            0.3827 1.287
                                            0.198
        exp(coef) exp(-coef) lower .95 upper .95
                                 0.7729
treated
            1.636
                       0.6111
                                            3.465
Rsquare= 0.012
                 (max possible= 0.25 )
Likelihood ratio test= 1.71
                              on 1 df,
                                         p=0.1914
Wald test
                      = 1.66
                              on 1 df,
                                         p=0.1982
Score (logrank) test = 1.69
                              on 1 df,
                                         p=0.1936
```

The odds ratio in the exp(coef) section above is the average causal effect estimate - it describes the odds of having an event (out2) occur associated with being a treated subject, as compared to the odds of the event when a control subject.

- Again, the result, though nominally fairly far from 1, is nowhere near statistically significant, according to our 95% confidence interval.
- Our estimate is 1.64, with 95% CI (0.77, 3.47).
- I'll use this conditional logistic regression approach to summarize the findings with regard to an odds ratio in my summary of matching results to come.

9.3 Outcome 3 (a time-to-event outcome)

9.3.1 Approach 1. Automated Approach from the Matching Library

Again, we'll start by thinking about the essentially automatic answer which can be obtained when using the Match function. The problem here is that this approach doesn't take into account the right censoring at all, and assumes that all of the specified times in Outcome 3 are observed. This causes the result (or the ATE version) to be non-sensical, given what we know about the data. So I don't recommend you use this approach when dealing with a time-to-event outcome.

And as a result, I won't even shos it here.

9.3.2 Approach 2. A stratified Cox proportional hazards model

rho chisq

treated 0.288

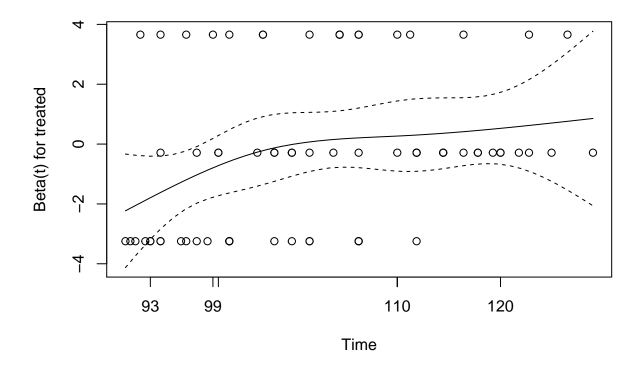
5.9 0.0152

plot(cox.zph(adj.m.out3), var="treated")

Since we have the matched sample, we can use a stratified Cox proportional hazards model to compare the treatment groups on our time-to-event outcome, while accounting for the matched pairs. The main results will be a relative hazard rate estimate, with 95% CI. Again, I use the 0/1 numeric version of the event indicator (out2), and of the treatment indicator (treated) here.

adj.m.out3 <- coxph(Surv(out3.time, out2) ~ treated + strata(matches), data=toy.matchedsample)

```
summary(adj.m.out3)
Call:
coxph(formula = Surv(out3.time, out2) ~ treated + strata(matches),
    data = toy.matchedsample)
 n= 140, number of events= 71
           coef exp(coef) se(coef)
                                         z Pr(>|z|)
treated -0.2877
                   0.7500
                             0.3118 -0.923
                                               0.356
        exp(coef) exp(-coef) lower .95 upper .95
             0.75
                        1.333
                                 0.4071
                                             1.382
treated
Concordance= 0.571 (se = 0.5)
Rsquare= 0.006
                 (max possible= 0.34)
Likelihood ratio test= 0.86 on 1 df,
                                         p=0.3537
Wald test
                      = 0.85
                              on 1 df,
                                         p=0.3562
Score (logrank) test = 0.86 on 1 df,
                                         p=0.3545
The relative hazard rate (from the exp(coef) section of the output) is estimated to be 0.75, with 95% CI
(0.41, 1.39) - again not statistically significant. Checking the proportional hazards assumption suggests a
possible issue, as well...
cox.zph(adj.m.out3) # Quick check for proportional hazards assumption
```



9.4 Results So Far (After Propensity Matching)

So, here's our summary again, now incorporating both our unadjusted results and the results after matching. Automated results and my favorite of our various non-automated approaches are shown. Note that I've left out the "automated" approach for a time-to-event outcome entirely, so as to discourage you from using it as presented above.

Est. Treatment Effect (95% CI)	Outcome 1 (Cost diff.)	Outcome 2 (Risk diff.)	Outcome 2 (Odds Ratio)	Outcome 3 (Relative Hazard Rate)
No covariate adjustment	15.7	+0.11	1.56	0.86
(unadjusted)	(12.0, 19.3)	(-0.03, +0.25)	(0.87, 2.82)	(0.57, 1.29)
After 1:1 PS Match	15.6	+0.11	N/A	N/A
(Match: Automated)	(11.6, 19.6)	(-0.05, +0.25)	N/A	N/A
After 1:1 PS Match	15.6	N/A	1.64	0.75
("Regression" Models)	(11.5, 19.6)	N/A	(0.77, 3.47)	(0.41, 1.38)

10 Task 6. Subclassify by PS quintile, then display postsubclassification balance

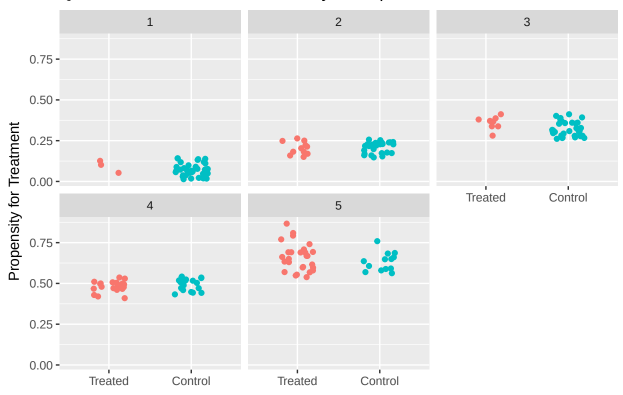
First, we divide the data by the propensity score into 5 strata of equal size using the cut2 function from the $\tt Hmisc$ package. Then we create a quintile variable which specifies $1 = \tt lowest$ propensity scores to $5 = \tt highest$.

```
toy$stratum <- cut2(toy$ps, g=5)</pre>
by(toy$ps, toy$stratum, summary) # sanity check
toy$stratum: [0.00777,0.147)
   Min. 1st Qu. Median
                           Mean 3rd Qu.
0.007769 0.045272 0.068558 0.072741 0.097041 0.144572
     _____
toy$stratum: [0.14670,0.260)
  Min. 1st Qu. Median Mean 3rd Qu.
0.1467  0.1736  0.2241  0.2089  0.2367  0.2597
______
toy$stratum: [0.25985,0.415)
  Min. 1st Qu. Median Mean 3rd Qu.
0.2598 \quad 0.2892 \quad 0.3243 \quad 0.3298 \quad 0.3663 \quad 0.4081
toy$stratum: [0.41456,0.543)
  Min. 1st Qu. Median Mean 3rd Qu.
0.4146 0.4694 0.4898 0.4872 0.5136 0.5415
toy$stratum: [0.54295,0.870]
  Min. 1st Qu. Median Mean 3rd Qu.
0.5430 0.5931 0.6485 0.6514 0.6865 0.8699
toy$quintile <- factor(toy$stratum, labels=1:5)</pre>
table(toy$stratum, toy$quintile) ## sanity check
                 1 2 3 4 5
  [0.00777,0.147) 40 0 0 0
  [0.14670,0.260) 0 40 0 0 0
  [0.25985,0.415) 0 0 40 0 0
  [0.41456,0.543) 0 0 0 40 0
 [0.54295,0.870] 0 0 0 40
```

10.1 Check Balance and Propensity Score Overlap in Each Quintile

We want to check the balance and propensity score overlap for each quintile. I'll start with a set of facetted, jittered plots to look at overlap.

Quintile Subclassification in the Toy Example



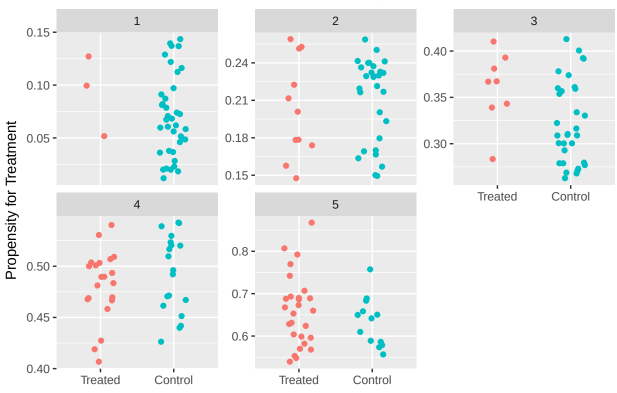
It can be helpful to know how many observations (by exposure group) are in each quintile.

```
addmargins(table(toy$quintile, toy$treated_f))
```

	Treated	Control	Sum
1	3	37	40
2	11	29	40
3	8	32	40
4	21	19	40
5	27	13	40
$\operatorname{\mathtt{Sum}}$	70	130	200

The overlap may show a little better in the plot if you free up the y axes...





10.2 Creating a Standardized Difference Calculation Function

We'll need to be able to calculate standardized differences in this situation so I've created a simple szd function to do this - using the average denominator method.

```
szd <- function(covlist, g) {
  covlist2 <- as.matrix(covlist)
  g <- as.factor(g)
  res <- NA
  for(i in 1:ncol(covlist2)) {
    cov <- as.numeric(covlist2[,i])
    num <- 100*diff(tapply(cov, g, mean, na.rm=TRUE))
    den <- sqrt(mean(tapply(cov, g, var, na.rm=TRUE)))
    res[i] <- round(num/den,2)
  }
  names(res) <- names(covlist)
  res
}</pre>
```

10.3 Creating the Five Subsamples, by PS Quintile

Next, we split the complete sample into the five quintiles.

```
## Divide the sample into the five quintiles
quin1 <- filter(toy, quintile==1)</pre>
```

```
quin2 <- filter(toy, quintile==2)
quin3 <- filter(toy, quintile==3)
quin4 <- filter(toy, quintile==4)
quin5 <- filter(toy, quintile==5)</pre>
```

10.4 Standardized Differences in Each Quintile, and Overall

Now, we'll calculate the standardized differences within each quintile, as well as overall.

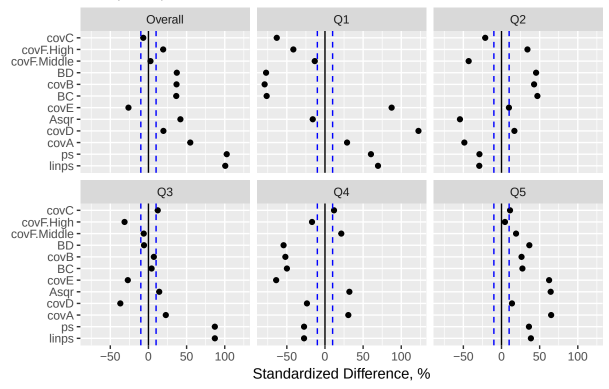
```
covs <- c("covA", "covB", "covC", "covD", "covE", "covF.Middle", "covF.High", "Asqr", "BC", "BD", "ps",
d.q1 <- szd(quin1[covs], quin1$treated)</pre>
d.q2 <- szd(quin2[covs], quin2$treated)</pre>
d.q3 <- szd(quin3[covs], quin3$treated)</pre>
d.q4 <- szd(quin4[covs], quin4$treated)</pre>
d.q5 <- szd(quin5[covs], quin5$treated)</pre>
d.all <- szd(toy[covs], toy$treated)</pre>
toy.szd <- data_frame(covs, Overall = d.all, Q1 = d.q1, Q2 = d.q2, Q3 = d.q3, Q4 = d.q4, Q5 = d.q5)
toy.szd <- gather(toy.szd, "quint", "sz.diff", 2:7)</pre>
toy.szd
# A tibble: 72 x 3
               quint sz.diff
   covs
               <chr>
                        <dbl>
   <chr>>
             Overall 54.8
 1 covA
             Overall 36.8
2 covB
3 covC
               Overall - 6.76
4 covD
               Overall 19.6
               Overall -26.2
5 covE
6 covF.Middle Overall 2.66
7 covF.High Overall 19.3
8 Asqr
               Overall 41.9
9 BC
               Overall 36.5
10 BD
               Overall 37.2
# ... with 62 more rows
```

10.5 Plotting the Standardized Differences

```
ggplot(toy.szd, aes(x = sz.diff, y = reorder(covs, -sz.diff), group = quint)) +
    geom_point() +
    geom_vline(xintercept = 0) +
    geom_vline(xintercept = c(-10,10), linetype = "dashed", col = "blue") +
    facet_wrap(~ quint) +
    labs(x = "Standardized Difference, %", y = "",
        title = "Comparing Standardized Differences by PS Quintile",
        subtitle = "The toy example")
```

Comparing Standardized Differences by PS Quintile

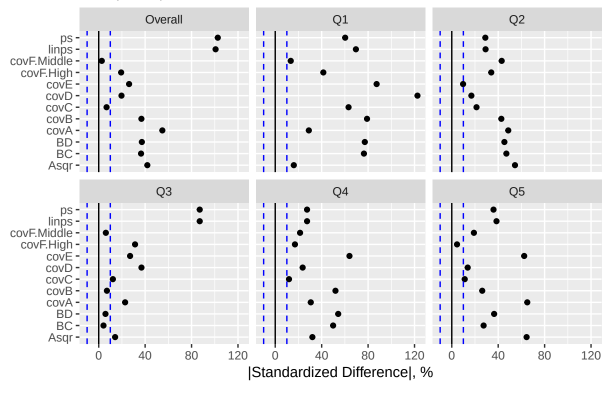
The toy example



```
ggplot(toy.szd, aes(x = abs(sz.diff), y = covs, group = quint)) +
    geom_point() +
    geom_vline(xintercept = 0) +
    geom_vline(xintercept = c(-10,10), linetype = "dashed", col = "blue") +
    facet_wrap(~ quint) +
    labs(x = "|Standardized Difference|, %", y = "",
        title = "Absolute Standardized Differences by PS Quintile",
        subtitle = "The toy example")
```

Absolute Standardized Differences by PS Quintile

The toy example



10.6 Checking Rubin's Rules Post-Subclassification

10.6.1 Rubin's Rule 1

As a reminder, prior to adjustment, Rubin's Rule 1 for the toy example was:

[1] 88.11531

After propensity score subclassification, we can obtain the same summary within each of the five quintiles...

```
rubin1.sub <- c(rubin1.q1, rubin1.q2, rubin1.q3, rubin1.q4, rubin1.q5)
names(rubin1.sub)=c("Q1", "Q2", "Q3", "Q4", "Q5")
rubin1.sub</pre>
```

```
Q1 Q2 Q3 Q4 Q5
60.58052 29.90587 80.45697 27.58516 35.77173
```

Each quintile shows (in some cases, only a slightly) better result than the full data set. With a small sample size like this, and some subclasses having very few subjects in one exposure group, it was always a long shot that subclassification alone would reduce all of these values below 10%, but I had hoped to get more than 3/5 below 50%.

10.6.2 Rubin's Rule 2

As a reminder, prior to adjustment, Rubin's Rule 2 for the toy example was:

```
rubin2.unadj <- with(toy, var(linps[treated==1])/var(linps[treated==0]))
rubin2.unadj</pre>
```

[1] 0.5835438

After Subclassification, we can obtain the same summary within each of the five quintiles...

```
rubin2.q1 <- with(quin1, var(linps[treated==1])/var(linps[treated==0]))
rubin2.q2 <- with(quin2, var(linps[treated==1])/var(linps[treated==0]))
rubin2.q3 <- with(quin3, var(linps[treated==1])/var(linps[treated==0]))
rubin2.q4 <- with(quin4, var(linps[treated==1])/var(linps[treated==0]))
rubin2.q5 <- with(quin5, var(linps[treated==1])/var(linps[treated==0]))
rubin2.sub <- c(rubin2.q1, rubin2.q2, rubin2.q3, rubin2.q4, rubin2.q5)
names(rubin2.sub)=c("Q1", "Q2", "Q3", "Q4", "Q5")</pre>
```

```
Q1 Q2 Q3 Q4 Q5
0.4780421 1.2041215 0.7905793 0.7950520 2.4852723
```

Some of these variance ratios are actually a bit further from 1 than the full data set. Again, with a small sample size like this, subclassification looks like a weak choice. At most, three of the quintiles (2-4) show OK variance ratios after propensity score subclassification.

10.6.3 Rubin's Rule 3

Prior to propensity adjustment, recall that Rubin's Rule 3 summaries were:

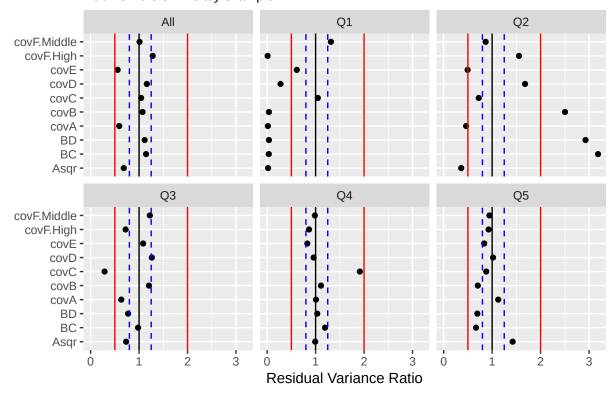
After subclassification, then, Rubin's Rule 3 summaries within each quintile are:

```
rubin3.q1 <- rubin3(data=quin1, covlist=quin1[covs])
rubin3.q2 <- rubin3(data=quin2, covlist=quin2[covs])
rubin3.q3 <- rubin3(data=quin3, covlist=quin3[covs])
rubin3.q4 <- rubin3(data=quin4, covlist=quin4[covs])
rubin3.q5 <- rubin3(data=quin5, covlist=quin5[covs])</pre>
```

```
toy.rubin3 <- data_frame(covs, All = rubin3.unadj$resid.var.ratio,</pre>
                           Q1 = rubin3.q1\$resid.var.ratio,
                          Q2 = rubin3.q2\$resid.var.ratio,
                          Q3 = rubin3.q3\(\frac{1}{2}\)resid.var.ratio,
                          Q4 = rubin3.q4\$resid.var.ratio,
                          Q5 = rubin3.q5\(\frac{1}{2}\)resid.var.ratio)
toy.rubin3 <- gather(toy.rubin3, "quint", "rubin3", 2:7)</pre>
ggplot(toy.rubin3, aes(x = rubin3, y = covs, group = quint)) +
    geom point() +
    geom vline(xintercept = 1) +
    geom_vline(xintercept = c(0.8, 1.25), linetype = "dashed", col = "blue") +
    geom_vline(xintercept = c(0.5, 2), col = "red") +
    facet_wrap(~ quint) +
    labs(x = "Residual Variance Ratio", y = "",
         title = "Residual Variance Ratios by PS Quintile",
         subtitle = "Rubin's Rule 3: The toy example")
```

Residual Variance Ratios by PS Quintile

Rubin's Rule 3: The toy example



Most of the residual variance ratios are in the range of (0.5, 2) but the results within quintiles vary widely. Quintiles 1 and 2 are especially problematic in this regard.

11 Task 7. After subclassifying, what is the estimated average treatment effect?

... on Outcome 1 [a continuous outcome]

First, we'll find the estimated average causal effect (and standard error) within each quintile via linear regression.

```
quin1.out1 <- lm(out1.cost ~ treated, data=quin1)
quin2.out1 <- lm(out1.cost ~ treated, data=quin2)
quin3.out1 <- lm(out1.cost ~ treated, data=quin3)
quin4.out1 <- lm(out1.cost ~ treated, data=quin4)
quin5.out1 <- lm(out1.cost ~ treated, data=quin5)
coef(summary(quin1.out1)); coef(summary(quin2.out1)); coef(summary(quin3.out1)); coef(summary(quin4.out
             Estimate Std. Error t value
                                                Pr(>|t|)
(Intercept) 45.45946
                        1.508909 30.127372 3.931690e-28
treated
            -12.45946
                        5.509756 -2.261345 2.954424e-02
             Estimate Std. Error
                                     t value
                                                 Pr(>|t|)
(Intercept) 44.206897 1.850820 23.8850290 1.727762e-24
                        3.529376 0.3019884 7.643075e-01
treated
             1.065831
                                 t value
            Estimate Std. Error
                                               Pr(>|t|)
(Intercept)
             44.375 2.035534 21.800171 4.412752e-23
              10.500 4.551593 2.306884 2.660602e-02
treated
                                 t value
            Estimate Std. Error
                                               Pr(>|t|)
(Intercept) 43.84211
                       2.895296 15.142532 1.105938e-17
treated
            16.91980
                      3.995888 4.234303 1.400593e-04
            Estimate Std. Error
                                  t value
                                               Pr(>|t|)
(Intercept) 48.07692
                       2.991778 16.069682 1.556770e-18
            23.44160
                       3.641476 6.437388 1.432788e-07
treated
We could probably figure out a cleverer way to accomplish this using the broom package.
Next, we find the mean of the five quintile-specific estimated regression coefficients
est.st <- (coef(quin1.out1)[2] + coef(quin2.out1)[2] + coef(quin3.out1)[2] +
               coef(quin4.out1)[2] + coef(quin5.out1)[2])/5
est.st
treated
7.893553
To get the combined standard error estimate, we do the following:
se.q1 <- summary(quin1.out1)$coefficients[2,2]</pre>
se.q2 <- summary(quin2.out1)$coefficients[2,2]</pre>
se.q3 <- summary(quin3.out1)$coefficients[2,2]</pre>
```

```
[1] 1.926223
```

se.st

se.q4 <- summary(quin4.out1)\$coefficients[2,2]</pre> se.q5 <- summary(quin5.out1)\$coefficients[2,2]</pre>

 $se.st \leftarrow sqrt((se.q1^2 + se.q2^2 + se.q3^2 + se.q4^2 + se.q5^2)*(1/25))$

The resulting 95% confidence Interval for the average causal treatment effect is then:

```
temp.result1 <- c(est.st, est.st - 1.96*se.st, est.st + 1.96*se.st)
names(temp.result1) <- c("Estimate", "Low 95% CI", "High 95% CI")
temp.result1</pre>
```

```
Estimate Low 95% CI High 95% CI 7.893553 4.118156 11.668950
```

11.2 ... on Outcome 2 [a binary outcome]

First, we find the estimated average causal effect (and standard error) within each quintile via logistic regression:

```
quin1.out2 <- glm(out2 ~ treated, data=quin1, family=binomial())
quin2.out2 <- glm(out2 ~ treated, data=quin2, family=binomial())</pre>
quin3.out2 <- glm(out2 ~ treated, data=quin3, family=binomial())</pre>
quin4.out2 <- glm(out2 ~ treated, data=quin4, family=binomial())</pre>
quin5.out2 <- glm(out2 ~ treated, data=quin5, family=binomial())
coef(summary(quin1.out2)); coef(summary(quin2.out2)); coef(summary(quin3.out2)); coef(summary(quin4.out
               Estimate Std. Error
                                      z value Pr(>|z|)
(Intercept) -0.05406722   0.3289181 -0.1643790   0.8694328
treated
            -0.63907996 1.2681430 -0.5039495 0.6142969
              Estimate Std. Error
                                    z value
                                              Pr(>|z|)
(Intercept) -0.7985077 0.4013863 -1.989374 0.04665989
treated
             0.2388919  0.7442903  0.320966  0.74823614
              Estimate Std. Error
                                     z value Pr(>|z|)
(Intercept) -0.2513144  0.3563439 -0.7052581  0.4806496
             0.2513144 0.7918213 0.3173878 0.7509494
treated
             Estimate Std. Error
                                   z value Pr(>|z|)
(Intercept) 0.3184537  0.4646602 0.6853476 0.4931246
            0.1670541 0.6463994 0.2584379 0.7960690
treated
              Estimate Std. Error
                                     z value Pr(>|z|)
(Intercept) -0.1541507  0.5563486 -0.2770757  0.7817220
treated
             Next, we find the mean of the five quintile-specific estimated logistic regression coefficients
est.st \leftarrow (coef(quin1.out2)[2] + coef(quin2.out2)[2] + coef(quin3.out2)[2] +
               coef(quin4.out2)[2] + coef(quin5.out2)[2])/5
est.st ## this is the estimated log odds ratio
  treated
0.1405919
## And we exponentiate this to get the overall odds ratio estimate
exp(est.st)
```

1.150955

treated

To get the combined standard error estimate across the five quintiles, we do the following:

```
se.q1 <- summary(quin1.out2)$coefficients[2,2]
se.q2 <- summary(quin2.out2)$coefficients[2,2]
se.q3 <- summary(quin3.out2)$coefficients[2,2]
se.q4 <- summary(quin4.out2)$coefficients[2,2]
se.q5 <- summary(quin5.out2)$coefficients[2,2]
se.st <- sqrt((se.q1^2 + se.q2^2 + se.q3^2 + se.q4^2 + se.q5^2)*(1/25))
se.st</pre>
```

[1] 0.3834222

Wald test

```
## Of course, this standard error is also on the log odds ratio scale
```

Now, we obtain a 95% Confidence Interval for the Average Causal Effect of our treatment (as an Odds Ratio) through combination and exponentiation, as follows:

```
temp.result2 <- c(exp(est.st), exp(est.st - 1.96*se.st), exp(est.st + 1.96*se.st))
names(temp.result2) <- c("Estimate", "Low 95% CI", "High 95% CI")
temp.result2</pre>
```

```
Estimate Low 95% CI High 95% CI 1.1509548 0.5428536 2.4402474
```

11.3 ... on Outcome 3 [a time to event]

Subjects with out2.event = "Yes" are truly observed events, while those with out2.event == "No" are censored before an event can happen to them.

The Cox model comparing treated to control, stratifying on quintile, is...

```
adj.s.out3 <- coxph(Surv(out3.time, out2) ~ treated + strata(quintile), data=toy)
summary(adj.s.out3) ## exp(coef) gives relative hazard associated with treatment
Call:
coxph(formula = Surv(out3.time, out2) ~ treated + strata(quintile),
   data = toy)
 n= 200, number of events= 97
           coef exp(coef) se(coef)
                                        z Pr(>|z|)
                   0.7858
treated -0.2411
                           0.2436 - 0.989
        exp(coef) exp(-coef) lower .95 upper .95
treated
           0.7858
                       1.273
                                0.4874
                                           1.267
Concordance= 0.538 (se = 0.056)
Rsquare= 0.005
                 (max possible= 0.946 )
Likelihood ratio test= 0.98 on 1 df,
                                        p=0.3212
```

11.3.1 Checking the Proportional Hazards Assumption

= 0.98 on 1 df,

The proportional hazards assumption looks fairly reasonable.

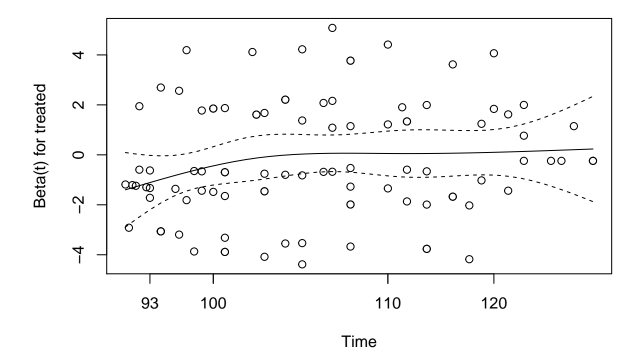
Score (logrank) test = 0.98 on 1 df,

p=0.3224

p=0.3219

```
cox.zph(adj.s.out3) ## checking the proportional hazards assumption
```

```
rho chisq p
treated 0.161 2.39 0.122
plot(cox.zph(adj.s.out3), var="treated")
```



11.4 Results So Far (After Matching and Subclassification)

These subclassification results describe the average treatment effect, while the previous analyses we have completed describe the average treatment effect on the treated. This is one reason for the meaningful difference between the estimates.

Est. Treatment Effect (95% CI)	Outcome 1 (Cost diff.)	Outcome 2 (Risk diff.)	Outcome 2 (Odds Ratio)	Outcome 3 (Relative Hazard Rate)
No covariate	15.7	+0.11	1.56	0.86
adjustment (unadjusted)	(12.0, 19.3)	(-0.03, +0.25)	(0.87, 2.82)	(0.57, 1.29)
After 1:1 PS Match	15.6	+0.11	N/A	N/A
(Match: Automated)	(11.6, 19.6)	(-0.05, +0.25)	N/A	N/A
After 1:1 PS Match	15.6	N/A	1.64	0.75
("Regression" models)	(11.5, 19.6)	N/A	(0.77, 3.47)	(0.41, 1.38)
After PS	7.9	N/A	1.15	0.79
Subclassification				

				Outcome 3
Est. Treatment Effect	Outcome 1 (Cost	Outcome 2 (Risk	Outcome 2	(Relative Hazard
(95% CI)	diff.)	diff.)	(Odds Ratio)	Rate)
("Regression" models, ATE)	(4.1, 11.7)	N/A	(0.54, 2.44)	(0.49, 1.27)

12 Task 8. Execute weighting by the inverse PS, then assess covariate balance

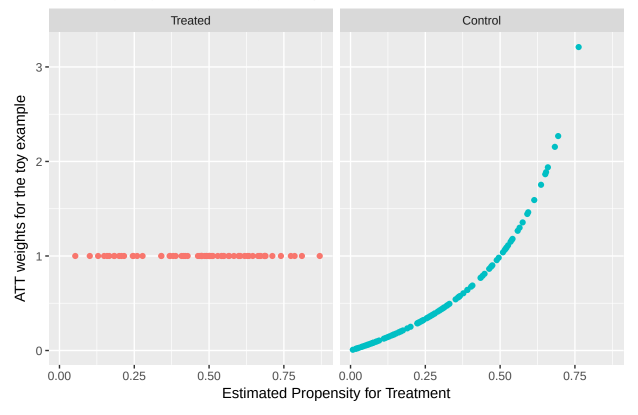
12.1 ATT approach: Weight treated subjects as 1; control subjects as ps/(1-ps)

```
toy$wts1 <- ifelse(toy$treated==1, 1, toy$ps/(1-toy$ps))</pre>
```

Here is a plot of the resulting ATT (average treatment effect on the treated) weights:

```
ggplot(toy, aes(x = ps, y = wts1, color = treated_f)) +
    geom_point() +
    guides(color = FALSE) +
    facet_wrap(~ treated_f) +
    labs(x = "Estimated Propensity for Treatment",
        y = "ATT weights for the toy example",
        title = "ATT weighting structure: Toy example")
```

ATT weighting structure: Toy example



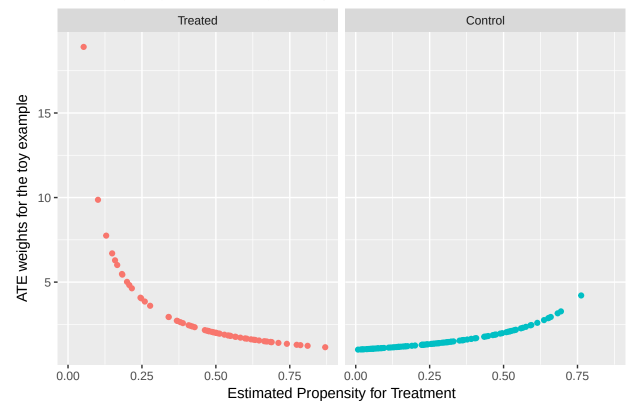
12.2 ATE Approach: Weight treated subjects by 1/ps; Control subjects by 1/(1-PS)

```
toy$wts2 <- ifelse(toy$treated==1, 1/toy$ps, 1/(1-toy$ps))</pre>
```

Here's a plot of the ATE (average treatment effect) weights...

```
ggplot(toy, aes(x = ps, y = wts2, color = treated_f)) +
    geom_point() +
    guides(color = FALSE) +
    facet_wrap(~ treated_f) +
    labs(x = "Estimated Propensity for Treatment",
        y = "ATE weights for the toy example",
        title = "ATE weighting structure: Toy example")
```

ATE weighting structure: Toy example



12.3 Assessing Balance after Weighting

The twang package provides several functions for assessing balance after weighting, in addition to actually doing the weighting using more complex propensity models. For this example, we'll demonstrate balance assessment for our two (relatively simple) weighting schemes. In other examples, we'll use twang to do more complete weighting work.

12.3.1 Reminder of ATT vs. ATE Definitions

- Informally, the average treatment effect on the treated (ATT) estimate describes the difference in potential outcomes (between treated and untreated subjects) summarized across the population of people who actually received the treatment. This is usually the estimate we work with in making causal estimates from observational studies.
- On the other hand, the **average treatment effect** (ATE) refers to the difference in potential outcomes summarized across the entire population, including those who did not receive the treatment.

12.3.2 For ATT weights (wts1)

```
toy_df <- data.frame(toy) # twanq doesn't react well to tibbles
covlist <- c("covA", "covB", "covC", "covD", "covE", "covF", "Asqr", "BC", "BD", "ps", "linps")
# for ATT weights
bal.wts1 <- dx.wts(x=toy_df$wts1, data=toy_df, vars=covlist,
                   treat.var="treated", estimand="ATT")
bal.wts1
  type n.treat n.ctrl ess.treat ess.ctrl
                                              max.es
                                                        mean.es
                                                                   max.ks
                             70 130.00000 1.1730321 0.40948475 0.4725275
1
  unw
            70
                  130
2
            70
                  130
                             70 62.42659 0.1259544 0.07031615 0.1998033
     mean.ks iter
1 0.19771767
               NΑ
2 0.08785673
bal.table(bal.wts1)
$unw
               tx.mn tx.sd ct.mn ct.sd std.eff.sz
                                                      stat
                                                               р
covA
               0.591 0.220 0.450 0.287
                                              0.637
                                                     3.863 0.000 0.284
               0.486 0.503 0.308 0.463
                                              0.354
                                                    2.461 0.015 0.178
covB
               9.626 2.016 9.762 2.025
                                             -0.068 -0.458 0.647 0.078
COVC
              10.096 1.877
                            9.738 1.770
covD
                                              0.191
                                                     1.317 0.189 0.110
covE
               9.671 2.744 10.500 3.536
                                             -0.302 -1.842 0.067 0.129
covF:1-Low
               0.271 0.445 0.369 0.483
                                             -0.220
                                                    1.317 0.269 0.098
covF:2-Middle
               0.429 0.495
                            0.415 0.493
                                              0.027
                                                        NA
                                                              NA 0.013
covF:3-High
               0.300 0.458
                            0.215 0.411
                                              0.185
                                                        NA
                                                              NA 0.085
               0.397 0.246
                            0.285 0.288
                                              0.456
                                                     2.904 0.004 0.284
Asqr
BC
               4.584 5.008
                           2.857 4.449
                                              0.345
                                                     2.427 0.016 0.187
BD
               4.816 5.152 3.000 4.601
                                              0.352
                                                     2.476 0.014 0.181
               0.476 0.191 0.282 0.187
                                              1.014
                                                     6.922 0.000 0.473
ps
              -0.139 0.917 -1.215 1.201
                                              1.173 7.100 0.000 0.473
linps
              ks.pval
                0.001
covA
                0.098
covB
                0.921
covC
covD
                0.598
covE
                0.401
covF:1-Low
                0.269
covF:2-Middle
                0.269
covF:3-High
                0.269
                0.001
Asqr
```

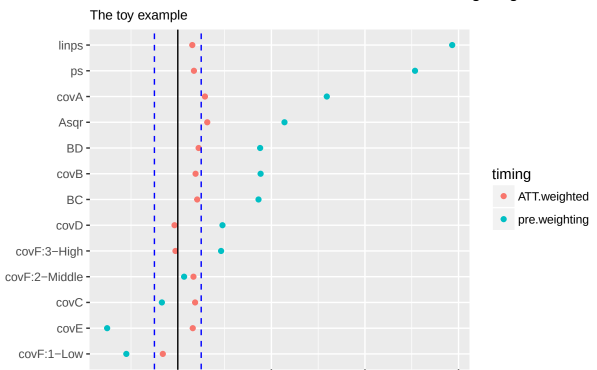
```
BC
                0.072
BD
                0.087
ps
                0.000
                0.000
linps
[[2]]
               tx.mn tx.sd ct.mn ct.sd std.eff.sz
                                                      stat
                                                               р
covA
               0.591 0.220 0.565 0.216
                                              0.116
                                                     0.740 0.460 0.200
               0.486 0.503 0.447 0.499
covB
                                              0.076 0.440 0.660 0.038
covC
               9.626 2.016 9.477 1.960
                                              0.074 0.442 0.659 0.086
covD
              10.096 1.877 10.122 1.711
                                             -0.014 -0.086 0.931 0.078
               9.671 2.744
                           9.495 3.430
                                                     0.323 0.747 0.125
covE
                                              0.064
covF:1-Low
               0.271 0.445 0.300 0.458
                                             -0.064
                                                     0.090 0.913 0.028
covF:2-Middle 0.429 0.495 0.396 0.489
                                             0.067
                                                        NA
                                                              NA 0.033
               0.300 0.458 0.305 0.460
                                             -0.010
                                                              NA 0.005
covF:3-High
                                                        NA
Asqr
               0.397 0.246
                            0.366 0.246
                                              0.126
                                                     0.775 0.439 0.200
BC
                                                     0.482 0.630 0.081
               4.584 5.008 4.170 4.844
                                              0.083
BD
               4.816 5.152 4.359 4.960
                                              0.089
                                                     0.517 0.606 0.081
                                              0.069
               0.476 0.191 0.463 0.177
                                                     0.410 0.682 0.093
ps
linps
              -0.139 0.917 -0.196 0.838
                                              0.062 0.380 0.705 0.093
              ks.pval
                0.121
covA
                1.000
covB
                0.943
covC
COVD
                0.972
covE
                0.619
covF:1-Low
                0.913
covF:2-Middle
                0.913
covF:3-High
                0.913
                0.121
Asqr
BC
                0.963
BD
                0.963
                0.903
ps
                0.903
linps
```

The std.eff.sz shows the standardized difference, but as a proportion, rather than as a percentage. We'll create a data frame (tibble) so we can plot the data more easily.

OK - here is the plot of standardized differences before and after ATT weighting.

```
ggplot(balance.att.weights, aes(x = szd, y = reorder(names, szd), color = timing)) +
    geom_point() +
    geom_vline(xintercept = 0) +
    geom_vline(xintercept = c(-10,10), linetype = "dashed", col = "blue") +
    labs(x = "Standardized Difference", y = "",
        title = "Standardized Difference before and after ATT Weighting",
        subtitle = "The toy example")
```

Standardized Difference before and after ATT Weighting



40

Standardized Difference

0

0.397 0.246 0.285 0.288

80

120

12.3.3 For ATE weights (wts2)

Asqr

```
bal.wts2 <- dx.wts(x=toy_df$wts2, data=toy_df, vars=covlist,
                   treat.var="treated", estimand="ATE")
bal.wts2
  type n.treat n.ctrl ess.treat ess.ctrl
                                            max.es
                                                      mean.es
                                                                 max.ks
1
  unw
                  130 70.00000 130.0000 0.9256094 0.36753949 0.4725275
            70
                  130 38.96583 115.2132 0.1866526 0.05313253 0.1436468
     mean.ks iter
1 0.19771767
              NA
2 0.07716551
bal.table(bal.wts2)
$unw
               tx.mn tx.sd ct.mn ct.sd std.eff.sz
                                                     stat
                                                              р
covA
               0.591 0.220 0.450 0.287
                                             0.513 3.863 0.000 0.284
covB
               0.486 0.503 0.308 0.463
                                             0.368
                                                    2.461 0.015 0.178
               9.626 2.016 9.762 2.025
                                            -0.068 -0.458 0.647 0.078
covC
                           9.738 1.770
                                             0.198 1.317 0.189 0.110
covD
              10.096 1.877
              9.671 2.744 10.500 3.536
                                            -0.251 -1.842 0.067 0.129
covE
covF:1-Low
              0.271 0.445 0.369 0.483
                                            -0.220 1.317 0.269 0.098
covF:2-Middle 0.429 0.495 0.415 0.493
                                             0.027
                                                       NA
                                                             NA 0.013
covF:3-High
              0.300 0.458 0.215 0.411
                                             0.185
                                                       NA
                                                             NA 0.085
```

0.403 2.904 0.004 0.284

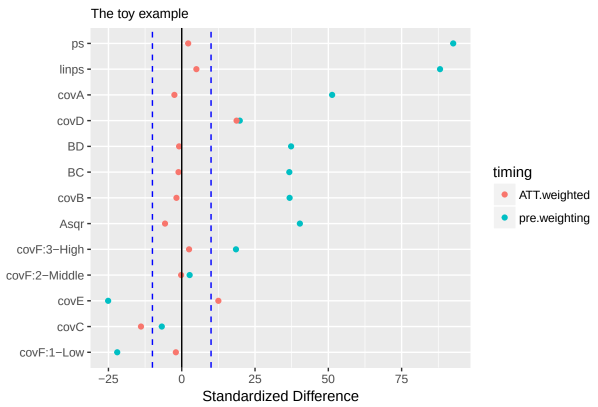
```
4.584 5.008 2.857 4.449
BC
                                              0.367 2.427 0.016 0.187
BD
               4.816 5.152 3.000 4.601
                                              0.373 2.476 0.014 0.181
                                              0.926 6.922 0.000 0.473
ps
               0.476 0.191 0.282 0.187
              -0.139 0.917 -1.215 1.201
                                              0.881 7.100 0.000 0.473
linps
              ks.pval
                0.001
covA
covB
                0.098
                0.921
covC
covD
                0.598
covE
                0.401
covF:1-Low
                0.269
covF:2-Middle
                0.269
covF:3-High
                0.269
Asqr
                0.001
BC
                0.072
BD
                0.087
                0.000
ps
linps
                0.000
[[2]]
               tx.mn tx.sd ct.mn ct.sd std.eff.sz
                                                      stat
                                                                     ks
                                                                р
covA
               0.483 0.254 0.490 0.270
                                             -0.025 -0.130 0.897 0.114
covB
               0.347 0.479 0.356 0.481
                                             -0.018 -0.109 0.913 0.009
covC
               9.395 1.860 9.664 2.007
                                             -0.139 -0.764 0.446 0.144
              10.208 1.860 9.870 1.759
                                              0.187 1.087 0.278 0.133
covD
covE
              10.566 3.063 10.154 3.532
                                              0.125
                                                     0.657 0.512 0.087
covF:1-Low
               0.336 0.472 0.345 0.475
                                             -0.020
                                                     0.011 0.988 0.010
covF:2-Middle 0.407 0.491 0.409 0.492
                                             -0.002
                                                        NA
                                                               NA 0.001
covF:3-High
               0.257 0.437
                            0.246 0.431
                                              0.025
                                                        NA
                                                               NA 0.011
                                             -0.057 -0.322 0.748 0.114
Asqr
               0.297 0.260 0.313 0.277
BC
               3.258 4.673
                            3.309 4.631
                                             -0.011 -0.066 0.948 0.050
BD
               3.426 4.852 3.468 4.772
                                             -0.009 -0.051 0.959 0.048
               0.349 0.212 0.344 0.203
                                              0.022 0.105 0.916 0.141
ps
              -0.806 1.133 -0.864 1.193
                                              0.050 0.225 0.822 0.141
linps
              ks.pval
                0.802
covA
covB
                1.000
COVC
                0.541
covD
                0.641
covE
                0.963
covF:1-Low
                0.988
covF:2-Middle
                0.988
covF:3-High
                0.988
Asqr
                0.802
BC
                1.000
BD
                1.000
ps
                0.566
                0.566
linps
bal.before.wts2 <- bal.table(bal.wts2)[1]</pre>
bal.after.wts2 <- bal.table(bal.wts2)[2]</pre>
balance.ate.weights <- data_frame(names = rownames(bal.before.wts2$unw),
                               pre.weighting = 100*bal.before.wts2$unw$std.eff.sz,
```

```
ATT.weighted = 100*bal.after.wts2[[1]]$std.eff.sz)
balance.ate.weights <- gather(balance.ate.weights, timing, szd, 2:3)
```

Here is the plot of standardized differences before and after ATE weighting.

```
ggplot(balance.ate.weights, aes(x = szd, y = reorder(names, szd), color = timing)) +
    geom_point() +
    geom_vline(xintercept = 0) +
    geom_vline(xintercept = c(-10,10), linetype = "dashed", col = "blue") +
    labs(x = "Standardized Difference", y = "",
        title = "Standardized Difference before and after ATE Weighting",
        subtitle = "The toy example")
```

Standardized Difference before and after ATE Weighting



12.4 Rubin's Rules after ATT weighting

For our weighted sample, our summary statistic for Rules 1 and 2 may be found from the bal.table output.

12.4.1 Rubin's Rule 1

We can read off the standardized effect size after weighting for the linear propensity score as 0.062. Multiplying by 100, we get 6.2%, so we would pass Rule 1.

12.4.2 Rubin's Rule 2

We can read off the standard deviations within the treated and control groups. We can then square each, to get the relevant variances, then take the ratio of those variances. Here, we have $0.917^2 / 0.838^2 = 1.1974314$, which is a substantial improvement over our unadjusted result, and now within our desired range of 4/5 to 5/4, as well as clearly within 1/2 to 2. Pass Rule 2, also.

12.4.3 Rubin's Rule 3

Rubin's Rule 3 requires some more substantial manipulation of the data. I'll skip that for now.

12.5 Rubin's Rules after ATE weighting

Again, our summary statistic for Rules 1 and 2 may be found from the bal.table output.

12.5.1 Rubin's Rule 1

The standardized effect size after ATE weighting for the linear propensity score is 0.050. Multiplying by 100, we get 5.0%, so we would pass Rule 1.

12.5.2 Rubin's Rule 2

We can read off the standard deviations within the treated and control groups from the ATE weights, then square to get the variances, then take the ratio. Here, we have $1.133^2 / 1.193^2 = 0.9019427$, which is also a substantial improvement over our unadjusted result, and within our desired range of 4/5 to 5/4. Pass Rule 2, also.

12.5.3 Rubin's Rule 3

Again, for now, I'm skipping Rubin's Rule 3 after weighting.

13 Using TWANG for Alternative PS Estimation and ATT Weighting

Here, I'll demonstrate the use of the twang package's functions to fit the propensity model and then perform ATT weighting, mostly using default options.

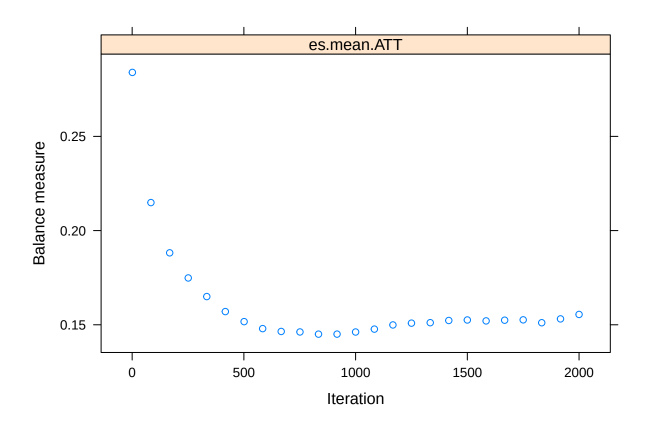
13.1 Estimate the Propensity Score using Generalized Boosted Regression, and then perfom ATT Weighting

We can directly use the twang (toolkit for weighting and analysis of nonequivalent groups) package to weight our results, and even to re-estimate the propensity score using generalized boosted regression rather than a logistic regression model. The twang vignette is very helpful and found at this link.

To begin, we'll estimate the propensity score using the twang function ps. This uses a *generalized boosted* regression approach to estimate the propensity score and produce material for checking balance.

13.1.1 Did we let the simulations run long enough to stabilize estimates?

```
plot(ps.toy)
```



13.1.2 What is the effective sample size of our weighted results?

```
n.treat n.ctrl ess.treat ess.ctrl max.es mean.es
unw 70 130 70 130.0000 0.6368393 0.2850744
es.mean.ATT 70 130 70 73.3784 0.2340580 0.1448418
```

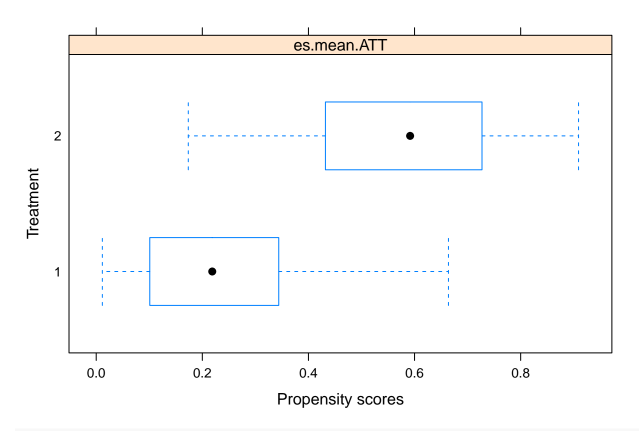
```
        max.ks
        max.ks.p
        mean.ks
        iter

        unw
        0.2835165
        NA 0.14775225
        NA

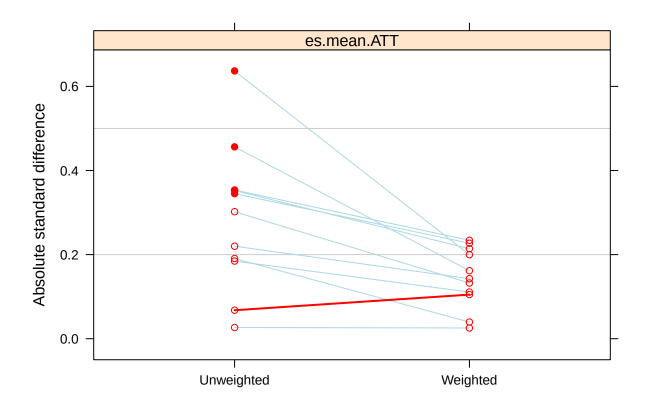
        es.mean.ATT
        0.1524293
        NA 0.09974725
        870
```

13.1.3 How is the balance?

```
plot(ps.toy, plots = 2)
```



plot(ps.toy, plots = 3)



13.1.4 Assessing Balance with cobalt

```
bal.tab(ps.toy, full.stop.method = "es.mean.att")
```

Call:

```
ps(formula = treated ~ covA + covB + covC + covD + covE + covF +
   Asqr + BC + BD, data = toy_df, n.trees = 2000, interaction.depth = 2,
   verbose = FALSE, estimand = "ATT", stop.method = c("es.mean"))
```

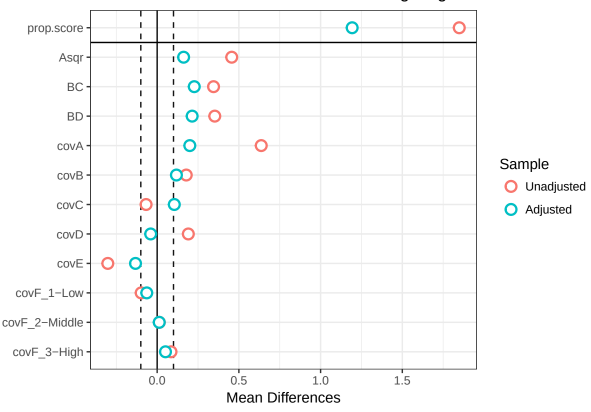
Balance Measures:

Balance Measures:				
	Туре	Diff.Adj		
prop.score	${\tt Distance}$	1.1932		
covA	Contin.	0.1998		
covB	Binary	0.1178		
covC	Contin.	0.1049		
covD	Contin.	-0.0396		
covE	Contin.	-0.1325		
covF_1-Low	Binary	-0.0635		
${\tt covF_2-Middle}$	Binary	0.0127		
covF_3-High	Binary	0.0509		
Asqr	Contin.	0.1618		
BC	Contin.	0.2270		
BD	Contin.	0.2142		

```
Effective sample sizes:
Control Treated
Unadjusted 130.00 70
Adjusted 73.38 70
```

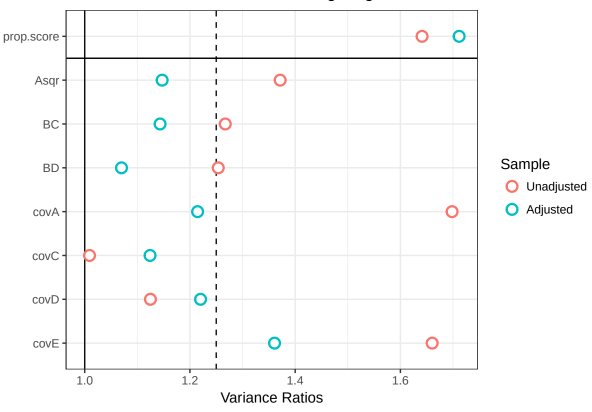
13.2 Semi-Automated Love plot of Standardized Differences

Standardized Diffs and TWANG ATT weighting



13.3 Semi-Automated Love plot of Variance Ratios

Variance Ratios: TWANG ATT weighting



14 Task 9. After weighting, what is the estimated average average causal effect of treatment?

14.1 ... on Outcome 1 [a continuous outcome]

14.1.1 with ATT weights

```
The relevant regression approach uses the svydesign and svyglm functions from the survey package.
```

```
toywt1.design <- svydesign(ids=~1, weights=~wts1, data=toy) # using ATT weights
adjout1.wt1 <- svyglm(out1.cost ~ treated, design=toywt1.design)
summary(adjout1.wt1); confint(adjout1.wt1)</pre>
```

```
15.221
treated
                         2.323 6.553 4.78e-10 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 189.3912)
Number of Fisher Scoring iterations: 2
               2.5 %
                      97.5 %
(Intercept) 42.98170 47.80528
           10.66821 19.77339
treated
14.1.2 with ATE weights
toywt2.design <- svydesign(ids=~1, weights=~wts2, data=toy) # using ATE weights
adjout1.wt2 <- svyglm(out1.cost ~ treated, design=toywt2.design)</pre>
summary(adjout1.wt2); confint(adjout1.wt2)
Call:
svyglm(formula = out1.cost ~ treated, design = toywt2.design)
Survey design:
svydesign(ids = ~1, weights = ~wts2, data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 45.0951
                        0.9245 48.78
                                         <2e-16 ***
treated
             7.0474
                        3.5236
                                  2.00
                                         0.0469 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 223.8246)
Number of Fisher Scoring iterations: 2
                 2.5 %
                       97.5 %
(Intercept) 43.2831607 46.90713
treated
            0.1412766 13.95348
14.1.3 with TWANG ATT weights
toywt3.design <- svydesign(ids=~1,
                          weights=~get.weights(ps.toy,
                                               stop.method = "es.mean"),
```

Call:

data=toy) # using twang ATT weights

adjout1.wt3 <- svyglm(out1.cost ~ treated, design=toywt3.design)</pre>

summary(adjout1.wt3); confint(adjout1.wt3)

```
svyglm(formula = out1.cost ~ treated, design = toywt3.design)
svydesign(ids = ~1, weights = ~get.weights(ps.toy, stop.method = "es.mean"),
   data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 45.055
                         1.109 40.616 < 2e-16 ***
             15.560
                         2.261
                                 6.882 7.6e-11 ***
treated
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 202.799)
Number of Fisher Scoring iterations: 2
              2.5 %
                      97.5 %
(Intercept) 42.88061 47.22893
treated
           11.12823 19.99080
```

14.2 ... on Outcome 2 [a binary outcome]

For a binary outcome, we build the outcome model using the quasibinomial, rather than the usual binomial family. We use the same svydesign information as we built for outcome 1.

14.2.1 Using ATT weights

treated

```
adjout2.wt1 <- svyglm(out2 ~ treated, design=toywt1.design, family=quasibinomial())
summary(adjout2.wt1)
Call:
svyglm(formula = out2 ~ treated, design = toywt1.design, family = quasibinomial())
Survey design:
svydesign(ids = ~1, weights = ~wts1, data = toy)
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.1611
                        0.2538 -0.635
                                          0.526
treated
             0.3907
                        0.3502 1.116
                                          0.266
(Dispersion parameter for quasibinomial family taken to be 1.005025)
Number of Fisher Scoring iterations: 4
exp(summary(adjout2.wt1)$coef)
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.8512082 1.288958 0.5301153 1.692797
```

1.4779751 1.419314 3.0515633 1.304634

```
exp(confint(adjout2.wt1))
               2.5 % 97.5 %
(Intercept) 0.5175732 1.399909
           0.7440438 2.935863
treated
14.2.2 Using ATE weights
adjout2.wt2 <- svyglm(out2.event ~ treated, design=toywt2.design, family=quasibinomial())</pre>
summary(adjout2.wt2)
Call:
svyglm(formula = out2.event ~ treated, design = toywt2.design,
   family = quasibinomial())
Survey design:
svydesign(ids = ~1, weights = ~wts2, data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.1972 0.1877 -1.051 0.295
treated
             0.2545
                        0.3713 0.685
                                         0.494
(Dispersion parameter for quasibinomial family taken to be 1.005025)
Number of Fisher Scoring iterations: 3
exp(summary(adjout2.wt2)$coef)
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.821019 1.206466 0.3496972 1.342700
treated
           exp(confint(adjout2.wt2))
               2.5 %
                     97.5 %
(Intercept) 0.5683120 1.186095
treated
        0.6229616 2.670553
14.2.3 with TWANG ATT weights
adjout2.wt3 <- svyglm(out2 ~ treated, design=toywt3.design,</pre>
                     family=quasibinomial())
summary(adjout2.wt3)
svyglm(formula = out2 ~ treated, design = toywt3.design, family = quasibinomial())
Survey design:
svydesign(ids = ~1, weights = ~get.weights(ps.toy, stop.method = "es.mean"),
   data = toy)
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.2581 0.2335 -1.105 0.270
treated 0.4877 0.3358 1.452 0.148
```

(Dispersion parameter for quasibinomial family taken to be 1.005025)

Number of Fisher Scoring iterations: 4

```
exp(summary(adjout2.wt3)$coef)
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.7725293   1.263056  0.331168  1.310547
treated   1.6285006  1.398989  4.273565  1.159466
exp(confint(adjout2.wt3))
```

```
2.5 % 97.5 % (Intercept) 0.4887992 1.220954 treated 0.8433287 3.144698
```

14.3 ... on Outcome 3 [a time to event]

As before, subjects with out2.event = "Yes" are truly observed events, while those with out2.event == "No" are censored before an event can happen to them.

14.3.1 Using ATT weights

The Cox model comparing treated to control, weighting by ATT weights (wts1), is...

```
adjout3.wt1 <- coxph(Surv(out3.time, out2) ~ treated, data=toy, weights=wts1)
summary(adjout3.wt1)</pre>
```

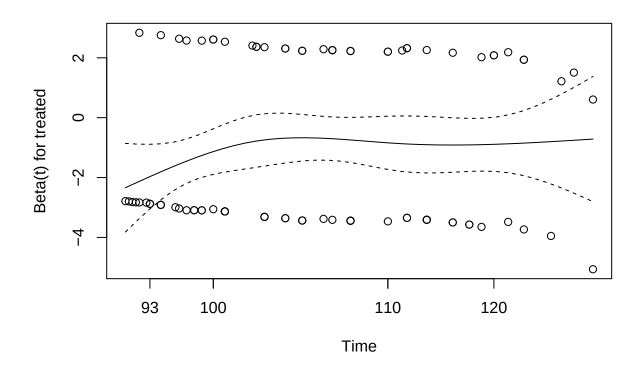
Call:

```
coxph(formula = Surv(out3.time, out2) ~ treated, data = toy,
   weights = wts1)
 n= 200, number of events= 97
          coef exp(coef) se(coef)
                                       z Pr(>|z|)
treated -0.1975
                  0.8208
                           0.2417 - 0.817
                                            0.414
        exp(coef) exp(-coef) lower .95 upper .95
          0.8208
                       1.218
                                0.511
                                          1.318
treated
Concordance= 0.549 (se = 0.033)
Rsquare= 0.003
                (max possible= 0.951)
Likelihood ratio test= 0.66 on 1 df,
                                       p=0.4158
Wald test
                    = 0.67
                            on 1 df,
                                       p=0.4139
Score (logrank) test = 0.67 on 1 df,
                                       p=0.4132
```

The exp(coef) output gives the relative hazard of the event comparing treated subjects to control subjects. And here's the check of the proportional hazards assumption...

```
cox.zph(adjout3.wt1); plot(cox.zph(adjout3.wt1), var="treated")
```

 $\begin{array}{ccc} & \text{rho chisq} & \text{p} \\ \text{treated 0.111} & 1.57 & 0.21 \end{array}$



14.3.2 Using ATE weights

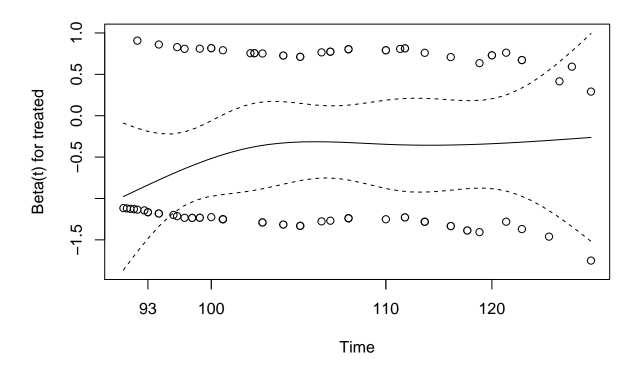
```
adjout3.wt2 <- coxph(Surv(out3.time, out2) ~ treated, data=toy, weights=wts2)</pre>
summary(adjout3.wt2)
coxph(formula = Surv(out3.time, out2) ~ treated, data = toy,
    weights = wts2)
 n= 200, number of events= 97
           coef exp(coef) se(coef)
                                       z Pr(>|z|)
                  0.8341 0.1451 -1.251
treated -0.1814
        exp(coef) exp(-coef) lower .95 upper .95
treated
          0.8341
                       1.199
                               0.6276
Concordance= 0.547 (se = 0.02)
Rsquare= 0.008 (max possible= 1)
Likelihood ratio test= 1.56 on 1 df,
                                       p=0.2124
```

```
Wald test = 1.56 on 1 df, p=0.2111 Score (logrank) test = 1.57 on 1 df, p=0.2105
```

And here's the check of the proportional hazards assumption...

```
cox.zph(adjout3.wt2); plot(cox.zph(adjout3.wt2), var="treated")
```

rho chisq p treated 0.151 1.06 0.303



14.3.3 with TWANG ATT weights

```
wts3 <- get.weights(ps.toy, stop.method = "es.mean")
adjout3.wt3 <- coxph(Surv(out3.time, out2) ~ treated, data=toy, weights=wts3)
summary(adjout3.wt3)

Call:
coxph(formula = Surv(out3.time, out2) ~ treated, data = toy,
    weights = wts3)

n= 200, number of events= 97

    coef exp(coef) se(coef)    z Pr(>|z|)
treated -0.1056    0.8997    0.2758 -0.383    0.702
    exp(coef) exp(-coef) lower .95 upper .95
```

```
treated 0.8997 1.111 0.524 1.545

Concordance= 0.539 (se = 0.034)

Rsquare= 0.001 (max possible= 0.913)

Likelihood ratio test= 0.15 on 1 df, p=0.7033

Wald test = 0.15 on 1 df, p=0.7017

Score (logrank) test = 0.15 on 1 df, p=0.7016
```

14.4 Results So Far (After Matching, Subclassification and Weighting)

Est. Treatment Effect (95% CI)	Outcome 1 (Cost diff.)	Outcome 2 (Risk diff.)	Outcome 2 (Odds Ratio)	Outcome 3 (Relative Hazard Rate)
No covariate	15.7	+0.11	1.56	0.86
$\operatorname{adjustment}$				
(unadjusted, ATT)	(12.0, 19.3)	(-0.03, +0.25)	(0.87, 2.82)	(0.57, 1.29)
1:1 PS Match	15.6	+0.11	N/A	N/A
(Match: ATT)	(11.6, 19.6)	(-0.05, +0.25)	N/A	N/A
1:1 PS Match	15.6	N/A	1.64	0.75
("Regression", ATT)	(11.5, 19.6)	N/A	(0.77, 3.47)	(0.41, 1.38)
PS Subclassification	7.9	N/A	1.15	0.79
(ATE)	(4.1, 11.7)	N/A	(0.54, 2.44)	(0.49, 1.27)
ATT Weighting	15.2	N/A	1.48	0.82
(ATT)	(10.7, 19.8)	N/A	(0.74, 2.94)	(0.51, 1.32)
ATE Weighting	7.1	N/A	1.29	0.83
(ATE)	(0.1, 14.0)	N/A	(0.62, 2.67)	(0.63, 1.11)
twang ATT weights	15.6	N/A	1.63	0.90
(ATT)	(11.1, 20.0)	N/A	(0.84, 3.14)	(0.52, 1.55)

15 Task 10. After direct adjustment for the linear PS, what is the estimated average causal treatment effect?

15.1 ... on Outcome 1 [a continuous outcome]

Here, we fit a linear regression model with linps added as a covariate.

```
adj.reg.out1 <- lm(out1.cost ~ treated + linps, data=toy)
summary(adj.reg.out1); confint(adj.reg.out1)</pre>
```

```
treated
            12.4629
                        1.9782
                                  6.300 1.9e-09 ***
              2.9859
                        0.7746 3.855 0.000157 ***
linps
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 12.1 on 197 degrees of freedom
Multiple R-squared: 0.3165,
                               Adjusted R-squared: 0.3095
F-statistic: 45.6 on 2 and 197 DF, p-value: < 2.2e-16
                2.5 %
                        97.5 %
(Intercept) 45.769157 51.364410
            8.561701 16.364085
treated
linps
            1.458316 4.513435
## provides treated effect and confidence interval estimates
      ... on Outcome 2 [a binary outcome]
15.2
Here, fit a logistic regression with linps added as a covariate
adj.reg.out2 <- glm(out2 ~ treated + linps, data=toy, family=binomial())</pre>
summary(adj.reg.out2)
Call:
glm(formula = out2 ~ treated + linps, family = binomial(), data = toy)
Deviance Residuals:
   Min
             1Q
                 Median
                                       Max
                                3Q
-1.4003 -1.1256 -0.9414
                          1.1915
                                     1.4281
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.05119
                        0.23531 -0.218
                                           0.828
                                0.916
                                           0.360
treated
            0.30064
                        0.32835
linps
            0.13716
                        0.13027
                                  1.053
                                           0.292
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 277.08 on 199 degrees of freedom
Residual deviance: 273.71 on 197 degrees of freedom
AIC: 279.71
Number of Fisher Scoring iterations: 4
exp(coef(adj.reg.out2)) # produces odds ratio estimate
(Intercept)
                treated
                              linps
 0.9501023
              1.3507214
                          1.1470102
exp(confint(adj.reg.out2)) # produces 95% CI for odds ratio
Waiting for profiling to be done...
                2.5 %
                       97.5 %
(Intercept) 0.5977473 1.508760
treated
           0.7097588 2.580375
```

15.3 ... on Outcome 3 [a time-to-event outcome]

Again, subjects with out2.event No are right-censored, those with Yes for out2.event have their times to event observed.

We fit a Cox proportional hazards model predicting time to event (with event=Yes indicating non-censored cases) based on treatment group (treated) and now also the linear propensity score.

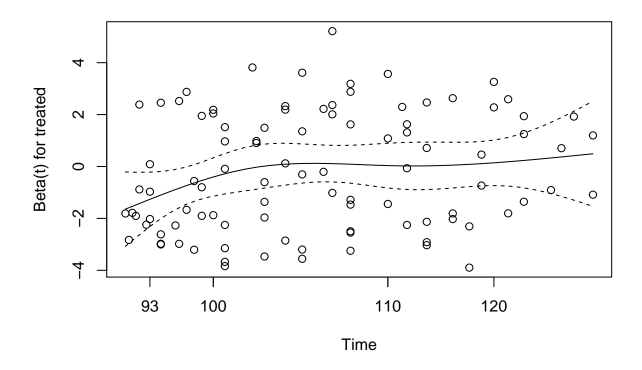
```
adj.reg.out3 <- coxph(Surv(out3.time, out2) ~ treated + linps, data=toy)</pre>
summary(adj.reg.out3)
coxph(formula = Surv(out3.time, out2) ~ treated + linps, data = toy)
 n= 200, number of events= 97
           coef exp(coef) se(coef)
                                        z Pr(>|z|)
treated -0.2274
                   0.7966
                            0.2343 -0.97
                                             0.332
         0.0684
                   1.0708
                            0.1005 0.68
                                             0.496
linps
        exp(coef) exp(-coef) lower .95 upper .95
           0.7966
                      1.2553
                                0.5032
                                            1.261
treated
linps
           1.0708
                      0.9339
                                0.8793
                                            1.304
Concordance= 0.559 (se = 0.033)
Rsquare= 0.005
                 (max possible= 0.988 )
Likelihood ratio test= 1.01
                             on 2 df,
                                        p=0.6022
Wald test
                     = 1.01
                             on 2 df,
                                        p=0.6025
                                        p=0.6021
Score (logrank) test = 1.01 on 2 df,
```

The exp(coef) section indicates the relative hazard estimates and associated 95% CI.

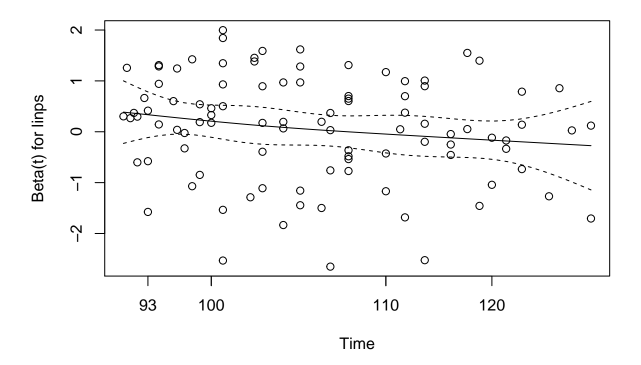
15.3.1 Check proportional hazards assumption

Not the best of news here. The results are close to being statistically significant.

```
rho chisq p
treated 0.193 3.40 0.0651
linps -0.174 3.17 0.0751
GLOBAL NA 4.52 0.1045
plot(cox.zph(adj.reg.out3), var="treated")
```



plot(cox.zph(adj.reg.out3), var="linps")



15.4 Results So Far (After Matching, Subclassification, Weighting, Adjustment)

Outcome 3 (Relative Hazard Rate)	Outcome 2 (Odds Ratio)	Outcome 2 (Risk diff.)	Outcome 1 (Cost diff.)	Est. Treatment Effect (95% CI)
0.86	1.56	+0.11	15.7	No covariate adjustment
(0.57, 1.29)	(0.87, 2.82)	(-0.03, +0.25)	(12.0, 19.3)	(unadjusted, ATT)
N/A	N/A	+0.11	15.6	1:1 PS Match
N/A	N/A	(-0.05, +0.25)	(11.6, 19.6)	(Match: ATT)
0.75	1.64	N/A	15.6	1:1 PS Match
(0.41, 1.38)	(0.77, 3.47)	N/A	(11.5, 19.6)	("Regression", ATT)
0.79	1.15	N/A	7.9	PS Subclassification
(0.49, 1.27)	(0.54, 2.44)	N/A	(4.1, 11.7)	("Regression", ATE)
0.82	1.48	N/A	15.2	ATT Weighting
	(0.51, 1.32)	(0.74, 2.94)	(10.7, 19.8)	(ATT)
0.83	1.29	N/A	7.1	ATE Weighting
(0.63, 1.11)	(0.62, 2.67)	N/A	(0.1, 14.0)	(ATE)
0.90	1.63	N/A	15.6	twang ATT weights
(0.52, 1.55)	(0.84, 3.14)	N/A	(11.1, 20.0)	(ATT)
0.80	1.35	N/A	12.5	Direct Adjustment
(0.50, 1.26)	(0.71, 2.58)	N/A	(8.56, 16.36)	(with linps, ATT)

16 Task 11. "Double Robust" Approach - Weighting + Adjustment, what is the estimated average causal effect of treatment?

This approach is essentially identical to the weighting analyses done in Task 9. The only change is to add linps to treated in the outcome models.

16.1 ... on Outcome 1 [a continuous outcome]

16.1.1 with ATT weights

```
The relevant regression approach uses the svydesign and svyglm functions from the survey package.
```

```
toywt1.design <- svydesign(ids=~1, weights=~wts1, data=toy) # using ATT weights
dr.out1.wt1 <- svyglm(out1.cost ~ treated + linps, design=toywt1.design)
summary(dr.out1.wt1); confint(dr.out1.wt1)</pre>
```

```
Call:
svyglm(formula = out1.cost ~ treated + linps, design = toywt1.design)
Survey design:
svydesign(ids = ~1, weights = ~wts1, data = toy)
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 46.817
                         1.534 30.528 < 2e-16 ***
             14.808
                         2.122 6.980 4.41e-11 ***
treated
                         1.230 5.906 1.51e-08 ***
linps
              7.264
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 148.853)
Number of Fisher Scoring iterations: 2
               2.5 %
                       97.5 %
(Intercept) 43.810924 49.82230
```

16.1.2 with ATE weights

10.649919 18.96660 4.853559 9.67487

treated

linps

```
toywt2.design <- svydesign(ids=~1, weights=~wts2, data=toy) # using ATE weights
dr.out1.wt2 <- svyglm(out1.cost ~ treated + linps, design=toywt2.design)
summary(dr.out1.wt2); confint(dr.out1.wt2)</pre>
```

```
Call:
svyglm(formula = out1.cost ~ treated + linps, design = toywt2.design)
```

```
Survey design:
svydesign(ids = ~1, weights = ~wts2, data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                       1.695 29.750 < 2e-16 ***
(Intercept) 50.420
treated
              6.691
                         2.581 2.593 0.0102 *
              6.162
                         1.356 4.545 9.57e-06 ***
linps
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 172.751)
Number of Fisher Scoring iterations: 2
               2.5 %
                        97.5 %
(Intercept) 47.098373 53.741875
treated
            1.633441 11.749311
linps
            3.504662 8.818882
16.1.3 with twang based ATT weights
wts3 <- get.weights(ps.toy, stop.method = "es.mean")</pre>
toywt3.design <- svydesign(ids=~1, weights=~wts3, data=toy) # twanq ATT weights
dr.out1.wt3 <- svyglm(out1.cost ~ treated + linps, design=toywt3.design)</pre>
summary(dr.out1.wt3); confint(dr.out1.wt3)
Call:
svyglm(formula = out1.cost ~ treated + linps, design = toywt3.design)
Survey design:
svydesign(ids = ~1, weights = ~wts3, data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 48.694
                         1.717 28.367 < 2e-16 ***
treated
             12.954
                         2.294 5.646 5.68e-08 ***
                         1.260
                                5.900 1.56e-08 ***
linps
              7.435
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 155.9117)
Number of Fisher Scoring iterations: 2
               2.5 %
                        97.5 %
(Intercept) 45.329842 52.058658
treated
            8.457203 17.451434
            4.964688 9.904405
linps
```

16.2 ... on Outcome 2 [a binary outcome]

For a binary outcome, we build the outcome model using the quasibinomial, rather than the usual binomial family. We use the same svydesign information as we built for outcome 1.

16.2.1 Using ATT weights

Call:

family = quasibinomial())

```
dr.out2.wt1 <- svyglm(out2 ~ treated + linps, design=toywt1.design,</pre>
                     family=quasibinomial())
summary(dr.out2.wt1)
Call:
svyglm(formula = out2 ~ treated + linps, design = toywt1.design,
   family = quasibinomial())
Survey design:
svydesign(ids = ~1, weights = ~wts1, data = toy)
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                        0.2781 -0.427
(Intercept) -0.1186
                                         0.670
treated
             0.3817
                        0.3559 1.072
                                         0.285
linps
             0.2262
                        0.2052 1.102
                                         0.272
(Dispersion parameter for quasibinomial family taken to be 1.005348)
Number of Fisher Scoring iterations: 4
exp(summary(dr.out2.wt1)$coef)
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.888133 1.320594 0.6527147 1.954483
treated
           1.464723
                    1.427472 2.9223339 1.329576
           linps
exp(confint(dr.out2.wt1))
                     97.5 %
               2.5 %
(Intercept) 0.5149615 1.531727
treated
          0.7291363 2.942405
linps
           0.8385347 1.874632
16.2.2 Using ATE weights
dr.out2.wt2 <- svyglm(out2.event ~ treated + linps, design=toywt2.design,</pre>
                     family=quasibinomial())
summary(dr.out2.wt2)
```

svyglm(formula = out2.event ~ treated + linps, design = toywt2.design,

```
Survey design:
svydesign(ids = ~1, weights = ~wts2, data = toy)
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.11246 0.27201 -0.413
                                      0.680
treated
                     0.38320 0.652
                                      0.515
           0.24972
linps
           0.09896
                     0.20309 0.487
                                      0.627
(Dispersion parameter for quasibinomial family taken to be 1.005037)
Number of Fisher Scoring iterations: 4
exp(summary(dr.out2.wt2)$coef)
           Estimate Std. Error t value Pr(>|t|)
treated
          linps
exp(confint(dr.out2.wt2))
              2.5 %
                   97.5 %
(Intercept) 0.5243509 1.522994
treated 0.6057234 2.720375
          0.7414934 1.643802
linps
16.2.3 Using twang ATT weights
dr.out2.wt3 <- svyglm(out2 ~ treated + linps, design=toywt3.design,</pre>
                   family=quasibinomial())
summary(dr.out2.wt3)
Call:
svyglm(formula = out2 ~ treated + linps, design = toywt3.design,
   family = quasibinomial())
Survey design:
svydesign(ids = ~1, weights = ~wts3, data = toy)
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.1707 0.2748 -0.621
                                      0.535
treated
            0.4271
                      0.3546 1.204
                                      0.230
                      0.1930 0.947
                                      0.345
linps
            0.1828
(Dispersion parameter for quasibinomial family taken to be 1.005653)
Number of Fisher Scoring iterations: 4
exp(summary(dr.out2.wt3)$coef)
           Estimate Std. Error t value Pr(>|t|)
```

```
(Intercept) 0.8430793
                        1.316301 0.5373532 1.707879
treated
            1.5327500
                        1.425640 3.3343637 1.258506
linps
            1.2006145
                        1.212918 2.5784289 1.411577
exp(confint(dr.out2.wt3))
                2.5 %
                        97.5 %
(Intercept) 0.4919674 1.444776
treated
           0.7649226 3.071321
linps
            0.8224259 1.752711
```

16.3 ... on Outcome 3 [a time to event]

As before, subjects with out2.event = "Yes" are truly observed events, while those with out2.event == "No" are censored before an event can happen to them.

16.3.1 Using ATT weights

```
The Cox model comparing treated to control, weighting by ATT weights (wts1), is...
dr.out3.wt1 <- coxph(Surv(out3.time, out2) ~ treated + linps, data=toy, weights=wts1)
```

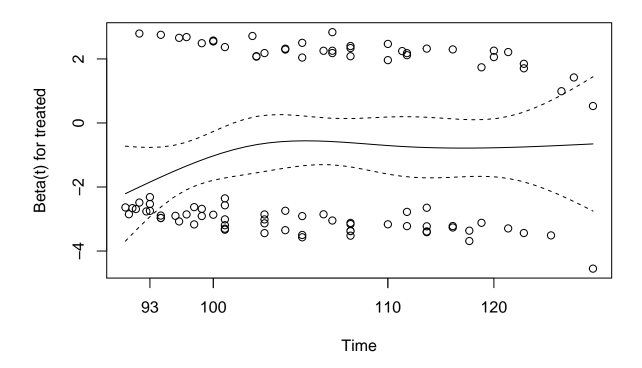
```
summary(dr.out3.wt1)
Call:
coxph(formula = Surv(out3.time, out2) ~ treated + linps, data = toy,
   weights = wts1)
 n= 200, number of events= 97
           coef exp(coef) se(coef)
                                        z Pr(>|z|)
treated -0.2086
                   0.8118
                            0.2424 -0.861
                                              0.39
         0.1045
                   1.1102
                            0.1445 0.723
                                              0.47
linps
        exp(coef) exp(-coef) lower .95 upper .95
                      1.2319
treated
           0.8118
                                0.5048
                                           1.305
linps
           1.1102
                      0.9007
                                0.8363
                                           1.474
Concordance= 0.564 (se = 0.039)
Rsquare= 0.006
                 (max possible= 0.951 )
Likelihood ratio test= 1.2 on 2 df,
                                       p=0.5501
Wald test
                     = 1.19 on 2 df,
                                        p=0.5514
Score (logrank) test = 1.19 on 2 df,
                                        p=0.5508
```

The exp(coef) output gives the relative hazard of the event comparing treated subjects to control subjects.

And here's the check of the proportional hazards assumption...

```
cox.zph(dr.out3.wt1); plot(cox.zph(dr.out3.wt1), var="treated")
```

```
rho chisq p
treated 0.1151 1.539 0.215
linps -0.0332 0.298 0.585
GLOBAL NA 1.759 0.415
```



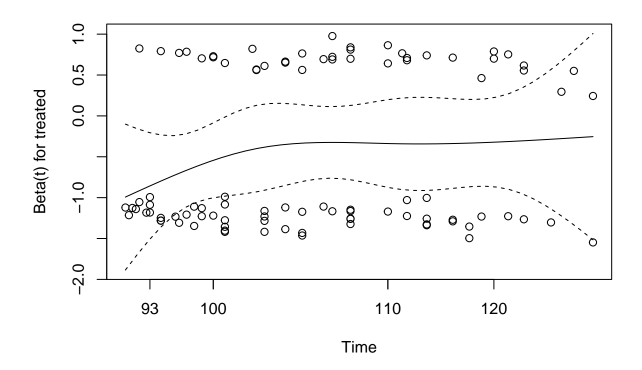
16.3.2 Using ATE weights

```
dr.out3.wt2 <- coxph(Surv(out3.time, out2) ~ treated + linps, data=toy, weights=wts2)</pre>
summary(dr.out3.wt2)
Call:
coxph(formula = Surv(out3.time, out2) ~ treated + linps, data = toy,
    weights = wts2)
  n= 200, number of events= 97
             coef exp(coef)
                             se(coef)
                                           z Pr(>|z|)
treated -0.181102 0.834351
                             0.145821 -1.242
                                                0.214
linps
        -0.001565 0.998436
                             0.065906 -0.024
        exp(coef) exp(-coef) lower .95 upper .95
           0.8344
treated
                       1.199
                                0.6269
                                           1.110
linps
           0.9984
                       1.002
                                0.8774
                                           1.136
Concordance= 0.533 (se = 0.023)
Rsquare= 0.008
                (max possible= 1 )
Likelihood ratio test= 1.56 on 2 df,
                                        p=0.4594
                     = 1.56
                             on 2 df,
                                        p=0.4574
Score (logrank) test = 1.57 on 2 df,
                                        p=0.4564
```

And here's the check of the proportional hazards assumption...

```
cox.zph(dr.out3.wt2); plot(cox.zph(dr.out3.wt2), var="treated")
```

```
rho chisq p
treated 0.173 1.27 0.260
linps -0.136 1.02 0.313
GLOBAL NA 2.08 0.353
```



16.3.3 Using twang ATT weights

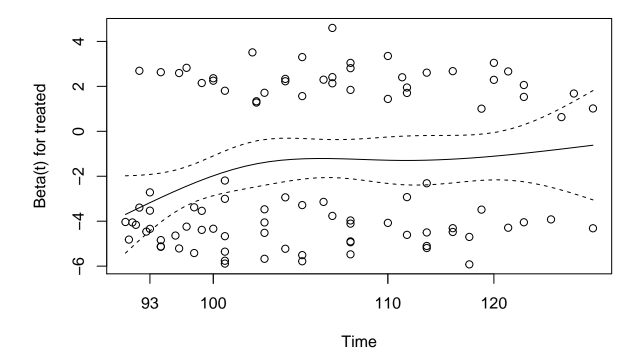
```
dr.out3.wt3 <- coxph(Surv(out3.time, out2) ~ treated + linps,</pre>
                     data=toy, weights=wts3)
summary(dr.out3.wt3)
Call:
coxph(formula = Surv(out3.time, out2) ~ treated + linps, data = toy,
   weights = wts3)
 n= 200, number of events= 97
            coef exp(coef) se(coef)
                                          z Pr(>|z|)
                   0.87874 0.28119 -0.460
treated -0.12927
                                               0.646
        0.06495
                   1.06711 0.15181 0.428
linps
        exp(coef) exp(-coef) lower .95 upper .95
```

```
0.8787
treated
                      1.1380
                                0.5064
                                            1.525
           1.0671
                      0.9371
linps
                                0.7925
                                            1.437
Concordance= 0.552 (se = 0.042)
Rsquare= 0.002
                 (max possible= 0.913)
Likelihood ratio test= 0.33
                             on 2 df,
                                        p=0.8478
Wald test
                     = 0.33
                             on 2 df,
                                        p=0.8476
Score (logrank) test = 0.33
                             on 2 df,
                                        p=0.8475
```

The exp(coef) output gives the relative hazard of the event comparing treated subjects to control subjects. And here's the check of the proportional hazards assumption...

```
cox.zph(dr.out3.wt3); plot(cox.zph(dr.out3.wt3), var="treated")
```

```
rho chisq p
treated 0.1931 5.26 0.0218
linps -0.0745 1.52 0.2177
GLOBAL NA 5.91 0.0521
```



17 Task 12. Results

17.1 Treatment Effect Estimates

We now can build the table of all of the outcome results we've obtained here.

				Outcome 3
Est. Treatment Effect	Outcome 1 (Cost	,	Outcome 2	(Relative Hazard
(95% CI)	diff.)	diff.)	(Odds Ratio)	Rate)
No covariate	15.7	+0.11	1.56	0.86
adjustment				
(unadjusted, ATT)	(12.0, 19.3)	(-0.03, +0.25)	(0.87, 2.82)	(0.57, 1.29)
1:1 PS Match	15.6	+0.11	N/A	N/A
(Match: ATT)	(11.6, 19.6)	(-0.05, +0.25)	N/A	N/A
1:1 PS Match	15.6	N/A	1.64	0.75
("Regression", ATT)	(11.5, 19.6)	N/A	(0.77, 3.47)	(0.41, 1.38)
PS Subclassification	7.9	N/A	1.15	0.79
("Regression", ATE)	(4.1, 11.7)	N/A	(0.54, 2.44)	(0.49, 1.27)
ATT Weighting	15.2	N/A	1.48	0.82
(ATT)	(10.7, 19.8)	N/A	(0.74, 2.94)	(0.51, 1.32)
ATE Weighting	7.1	N/A	1.29	0.83
(ATE)	(0.1, 14.0)	N/A	(0.62, 2.67)	(0.63, 1.11)
twang ATT weights	15.6	N/A	1.63	0.90
(ATT)	(11.1, 20.0)	N/A	(0.84, 3.14)	(0.52, 1.55)
Direct Adjustment	12.5	N/A	1.35	0.80
(with linps, ATT)	(8.6, 16.4)	N/A	(0.71, 2.58)	(0.50, 1.26)
Double Robust	14.8	N/A	1.46	0.81
(ATT wts + adj.)	(10.6, 19.0)	N/A	(0.73, 2.94)	(0.50, 1.31)
Double Robust	6.7	N/A	1.28	0.83
(ATE wts + adj.)	(1.6, 11.7)	N/A	(0.61, 2.72)	(0.63, 1.11)
Double Robust	13.0	N/A	1.53	0.88
(twang ATT $wts +$	(8.5, 17.5)	N/A	(0.76, 3.07)	(0.50, 1.53)
adj.)	. ,	, 		

So, for outcome 1, we have a significant result (indicating higher costs with the treatment) with every approach, and with outcomes 2 and 3, we do not.

17.2 Quality of Balance: Standardized Differences and Variance Ratios

We're looking at the balance across the following 10 covariates and transformations here: covA, covB, covC, covD, covE, covF[middle], covF[high], A squared, BxC and BxD...

Approach	Standardized Diffs	Variance Ratios
Most Desirable Values	Between -10 and $+10$	Between 0.8 and 1.25
No Adjustments	-30 to 63	0.59 to 1.27
Subclassification Quintile 1	-79 to 123	0 to 1.35
Quintile 2	-54 to 47	0.40 to 2.99
Quintile 3	-37 to 23	0.32 to 1.22
Quintile 4	-64 to 32	0.84 to 1.85
Quintile 5	5 to 65	0.80 to 1.32
1:1 Propensity Matching	-13 to 20	0.62 to 1.23
Propensity Weighting, ATT	-6 to 13	0.64 to 1.20
Propensity Weighting, ATE	-14 to 19	0.86 to 1.12

17.3 Quality of Balance: Rubin's Rules

Approach	Rubin 1	Rubin 2	Rubin 3
"Pass" Range, per Rubin	0 to 50	0.5 to 2.0	0.5 to 2.0
No Adjustments	88	0.58	0.59 to 1.28
Subclassification: Quintile 1	61	0.48	0.02 to 1.32
Quintile 2	30	1.20	0.36 to 3.19
Quintile 3	80	0.79	0.29 to 1.26
Quintile 4	28	0.80	0.83 to 1.91
Quintile 5	36	2.49	0.67 to 1.42
1:1 Propensity Matching	37	1.42	0.56 to 1.28
Propensity Weighting, ATT	6.2	1.20	Not evaluated
Propensity Weighting, ATE	5.0	0.90	Not evaluated

Clearly, the matching and propensity weighting show improvement over the initial (no adjustments) results, although neither is completely satisfactory in terms of all covariates. In practice, I would be comfortable with either a 1:1 match or a weighting approach, I think. It isn't likely that the subclassification will get us anywhere useful in terms of balance. Rubin's Rule 3 could also be applied after weighting on the propensity score.

18 What is a Sensitivity Analysis for Matched Samples?

We'll study a formal sensitivity analysis approach for **matched** samples. Note well that this specific approach is appropriate only when we have

- 1. a statistically significant conclusion
- 2. from a matched samples analysis using the propensity score.

18.1 Goal of a Formal Sensitivity Analysis for Matched Samples

To replace a general qualitative statement that applies in all observational studies, like ...

the association we observe between treatment and outcome does not imply causation

or

hidden biases can explain observed associations

... with a quantitative statement that is specific to what is observed in a particular study, such as ... to explain the association seen in a particular study, one would need a hidden bias of a particular magnitude.

If the association is strong, the hidden bias needed to explain it would be large.

- If a study is free of hidden bias (main example: a carefully randomized trial), this means that any two units (patients, subjects, whatever) that appear similar in terms of their observed covariates actually have the same chance of assignment to treatment.
- There is *hidden bias* if two units with the same observed covariates have different chances of receiving the treatment.

A sensitivity analysis asks: How would inferences about treatment effects be altered by hidden biases of various magnitudes? How large would these differences have to be to alter the qualitative conclusions of the study?

The methods for building such sensitivity analyses are largely due to Paul Rosenbaum, and as a result the methods are sometimes referred to as **Rosenbaum bounds**.

18.2 The Sensitivity Parameter, Γ

Suppose we have two units (subjects, patients), say, j and k, with the same observed covariate values \mathbf{x} but different probabilities p of treatment assignment (possibly due to some unobserved covariate), so that $\mathbf{x}_j = \mathbf{x}_k$ but that possibly $p_j \neq p_k$.

Units j and k might be matched to form a matched pair in our attempt to control overt bias due to the covariates \mathbf{x} .

• The odds that units j and k receive the treatment are, respectively, $\frac{p_j}{1-p_j}$ and $\frac{p_k}{1-p_k}$, and the odds ratio is thus the ratio of these odds.

Imagine that we knew that this odds ratio for units with the same \mathbf{x} was at most some number Γ , so that $\Gamma \geq 1$. That is,

$$\frac{1}{\Gamma} \le \frac{p_j(1-p_j)}{p_k(1-p_k)} \le \Gamma$$

We call Γ the **sensitivity parameter**, and it is the basis for our sensitivity analyses.

• If $\Gamma = 1$, then $p_j = p_k$ whenever $\mathbf{x}_j = \mathbf{x}_k$, so the study would be free of hidden bias, and standard statistical methods designed for randomized trials would apply.

If $\Gamma = 2$, then two units who appear similar in that they have the same set of observed covariates \mathbf{x} , could differ in their odds of receiving the treatment by as much as a factor of 2, so that one could be twice as likely as the other to receive the treatment.

So Γ is a value between 1 and ∞ where the size of Γ indicates the degree of a departure from a study free of hidden bias.

18.3 Interpreting the Sensitivity Parameter, Γ

Again, Γ is a measure of the degree of departure from a study that is free of hidden bias.

A sensitivity analysis will consider possible values of Γ and show how the inference for our outcomes might change under different levels of hidden bias, as indexed by Γ .

- A study is *sensitive* if values of Γ close to 1 could lead to inferences that are very different from those obtained assuming the study is free of hidden bias.
- A study is insensitive (a good thing here) if extreme values of Γ are required to alter the inference.

When we perform this sort of sensitivity analysis, we will specify different levels of hidden bias (different Γ values) and see how large a Γ we can have while still retaining the fundamental conclusions of the matched outcomes analysis.

19 Task 13. Sensitivity Analysis for Matched Samples, Outcome 1, using rbounds

In our matched sample analysis, for outcome 1 (cost) in the toy example, we saw a statistically significant result. A formal *sensitivity analysis* is called for, as a result, and we will accomplish one for this quantitative outcome, using the **rbounds** package.

The rbounds package is designed to work with the output from Matching, and can calculate Rosenbaum sensitivity bounds for the treatment effect, which help us understand the impact of hidden bias needed to invalidate our significant conclusions from the matched samples analysis.

19.1 Rosenbaum Bounds for the Wilcoxon Signed Rank test (Quantitative outcome)

We have already used the Match function from the Matching package to develop a matched sample. Given this, we need only run the psens function from the rbounds package to obtain sensitivity results.

```
X <- toy$linps ## matching on the linear propensity score
Tr <- as.logical(toy$treated)
Y <- toy$out1.cost
match1 <- Match(Tr=Tr, X=X, Y = Y, M = 1, replace=FALSE, ties=FALSE)
summary(match1)</pre>
```

```
Estimate... 15.557

SE...... 2.0397

T-stat.... 7.6273
p.val..... 2.3981e-14

Original number of observations....... 200
Original number of treated obs....... 70

Matched number of observations....... 70

Matched number of observations (unweighted). 70

psens(match1, Gamma = 5, GammaInc = 0.25)
```

Rosenbaum Sensitivity Test for Wilcoxon Signed Rank P-Value

Unconfounded estimate 0

Gamma	Lower	bound	Upper bound
1.00		0	0.0000
1.25		0	0.0000
1.50		0	0.0000
1.75		0	0.0001
2.00		0	0.0003
2.25		0	0.0008
2.50		0	0.0022
2.75		0	0.0046
3.00		0	0.0087
3.25		0	0.0148
3.50		0	0.0233
3.75		0	0.0343
4.00		0	0.0480
4.25		0	0.0644
4.50		0	0.0833
4.75		0	0.1046
5.00		0	0.1280

Note: Gamma is Odds of Differential Assignment To Treatment Due to Unobserved Factors

If the study were free of hidden bias, that is, if $\Gamma = 1$, then there would be **strong** evidence that the treated patients had higher costs, and the specific Wilcoxon signed rank test we're looking at here shows a p value < 0.0001. The sensitivity analysis we'll conduct now asks how this conclusion might be changed by hidden

biases of various magnitudes, depending on the significance level we plan to use in our test.

19.2 Specifying The Threshold Γ value

From the output above, find the Γ value where the upper bound for our p value slips from "statistically significant" to "not significant" territory.

 We're doing a two-tailed test, with a 95% confidence level, so the Γ statistic for this situation is between 3.50 and 3.75, since that is the point where the upper bound for the p value crosses the threshold of α/2 = 0.025.

So this study's conclusion (that treated patients had significantly higher costs) would still hold even in the face of a hidden bias with $\Gamma = 3.5$, but not with $\Gamma = 3.75$.

The tipping point for the sensitivity parameter is a little over 3.5. To explain away the observed association between treatment and this outcome (cost), a hidden bias or unobserved covariate would need to increase the odds of treatment by more than a factor of $\Gamma = 3.5$.

Returning to the output:

- If instead we were doing a one-tailed test, with a 95% confidence level, then the Γ statistic for this
 situation is between 4 and 4.25, since that is the point where the upper bound for the p value crosses
 α = 0.05.
- And if instead we were doing a one-tailed test with a 90% confidence level, then the Γ statistic would be between 4.75 and 5.0, since that is where the upper bound for the p value crosses $\alpha = 0.10$.

19.3 Interpreting Γ appropriately

 Γ tells you only how big a bias is needed to change the answer. By itself, it says NOTHING about the likelihood that a bias of that size is present in your study, except that, of course, smaller biases hide more effectively than large ones, on average.

In some settings, we'll think of Γ in terms of small (< 1.5), modest (1.5 - 2.5), moderate (2.5 - 4) and large (> 4) hidden bias requirements. But these are completely arbitrary distinctions, and I can provide no good argument for their use.

The **only** defense against hidden bias affecting your conclusions is to try to reduce the potential for hidden bias in the first place. We work on this via careful design of observational studies, especially by including as many different dimensions of the selection problem as possible in your propensity model.

19.4 An Alternate Approach - the Hodges-Lehman estimate

hlsens(match1)

Rosenbaum Sensitivity Test for Hodges-Lehmann Point Estimate

Unconfounded estimate 16.5

${\tt Gamma}$	Lower	bound	Upper	bound
1		16.5		16.5
2		11.0		21.6
3		7.5		24.1
4		5.5		26.1
5		3.5		27.6

```
6 2.0 28.6
```

Note: Gamma is Odds of Differential Assignment To Treatment Due to Unobserved Factors

If the Γ value is 3.0, then this implies that the Hodges-Lehmann estimate might be as low as 7.5 or as high as 24.1 (it is 16.5 in the absence of hidden bias in this case - when $\Gamma = 0$.)

19.5 What about other types of outcomes?

The rbounds package can evaluate binary outcomes using the binarysens and Fishersens functions.

Survival outcomes can be assessed, too, but not, I believe, using **rbounds** unless there is no censoring. Some time back, I built a spreadsheet for this task, which I'll be happy to share.

19.6 What about when we match 1:2 or 1:3 instead of 1:1?

The mcontrol function in the rbounds package can be helpful in such a setting.

20 Wrapup

If you run this script, you'll wind up with a version of the toy tibble that contains 200 observations on 28 variables, along with a codebook list.

You'll also have two new functions, called szd and rubin3, that, with some modification, may be useful elsewhere.

To drop everything else in the global environment created by this Markdown file, run the code that follows.

```
rm(list = c("adj.m.out1", "adj.m.out2", "adj.m.out3",
            "adj.reg.out1", "adj.reg.out2", "adj.reg.out3",
            "adj.s.out3", "adjout1.wt1", "adjout1.wt2",
            "adjout2.wt1", "adjout2.wt2", "adjout3.wt1",
            "adjout3.wt2", "bal.after.wts1", "bal.after.wts2",
            "bal.before.wts1", "bal.before.wts2", "bal.wts1",
            "bal.wts2", "balance.ate.weights", "balance.att.weights",
            "cov.sub", "covlist", "covnames", "covs",
            "d.all", "d.q1", "d.q2", "d.q3", "d.q4", "d.q5",
            "dr.out1.wt1", "dr.out1.wt2", "dr.out2.wt1",
            "dr.out2.wt2", "dr.out3.wt1", "dr.out3.wt2", "est.st",
            "factorlist", "high", "i", "low", "match1", "matches",
            "mb1", "mixedmodel.out1", "post.szd", "post.vratio",
            "pre.szd", "pre.vratio", "psmodel", "quin1",
            "quin1.out1", "quin1.out2", "quin2", "quin2.out1",
            "quin2.out2", "quin3", "quin3.out1", "quin3.out2",
            "quin4", "quin4.out1", "quin4.out2", "quin5",
            "quin5.out1", "quin5.out2", "rubin1.match", "rubin1.q1",
            "rubin1.q2", "rubin1.q3", "rubin1.q4", "rubin1.q5",
            "rubin1.sub", "rubin1.unadj", "rubin2.match",
            "rubin2.q1", "rubin2.q2", "rubin2.q3", "rubin2.q4",
            "rubin2.q5", "rubin2.sub", "rubin2.unadj", "rubin3.both",
            "rubin3.matched", "rubin3.q1", "rubin3.q2", "rubin3.q3",
```

```
"rubin3.q4", "rubin3.q5", "rubin3.unadj", "se.q1",
"se.q2", "se.q3", "se.q4", "se.q5", "se.st", "temp",
"temp.result1", "temp.result2", "tempsort",
"toy.matchedsample", "toy.rubin3", "toy.szd",
"toy_df", "toywt1.design", "toywt2.design", "Tr",
"unadj.out1", "unadj.out2", "unadj.out3", "varlist", "X", "Y"))
```