# R Lab 3 - IPTW & the Positivity Assumption

#### Laura B. Balzer

### Biostat 683- Intro. to Causal Inference

### Goals:

- 1. Implement IPTW for a binary exposure.
- 2. Understand how the IPTW estimator is affected by "near" positivity violations and weight stabilization.
- 3. Extend IPTW to control for missingness on the outcome.

#### Next lab:

Code discrete Super Learner to select the estimator with the lowest cross-validated risk. Use R SuperLearner package to build the best convex combination of candidate algorithms and to evaluate the performance of Super Learner.

## 1 Background Story

You have been hired by the Queen to estimate the causal effect of scurvy on mortality among pirates. The data available on n pirates are

- W1: possession of at least one quintessential pirate characteristic (e.g., peg leg, eye patch, beard, parrot). This is a binary covariate measured when leaving port (yes:1, no:0).
- W2: a summary measure of route danger, including voyage length, hurricane season, travel through enemy waters, Bermuda triangle, etc. This is a categorical variable ranging from 0 as least dangerous to 3 as most dangerous.
- A: whether the pirate suffered from scurvy during the voyage. This is a binary exposure (yes:1, no:0).
- Y: pirate's mortality status upon returing to port. This is a binary outcome with 1 for deceased and 0 for alive.

### 1.1 Causal Roadmap Rundown

Please note: We are doing a very fast review here. In practice, each step of the road map requires very careful thinking.

## 1. Specify the Question:

What is the causal effect of scurvy on mortality among pirates?

### 2. Specify the causal model:

- Endogenous nodes: X = (W1, W2, A, Y), where W1 is an indicator for having an "awesome" pirate characteristic, W2 is a summary measure of route danger, A is scurvy status, and Y is mortality status.
- Background variables:  $U = (U_{W1}, U_{W2}, U_A, U_Y) \sim \mathbb{P}_U$ . We make no assumptions about the distribution  $\mathbb{P}_U$ .





Image 1: https://www.flickr.com/photos/talklikeapirateday/3933458622/in/set-990505 Image 2: http://www.huffingtonpost.com/2013/09/24/sir-stuffington-one-eyed-kitten\_n\_3982907.html

#### - Structural equations F:

$$W1 = f_{W1}(U_{W1})$$

$$W2 = f_{W2}(W1, U_{W2})$$

$$A = f_A(W1, W2, U_A)$$

$$Y = f_Y(W1, W2, A, U_Y)$$

There are no exclusion restrictions or assumptions about functional form.

#### 3. Specify the causal parameter of interest:

We are interested in the causal risk of death due to scurvy (i.e., the average treatment effect):

$$\Psi^*(\mathbb{P}^*) = \mathbb{E}^*(Y_1) - \mathbb{E}^*(Y_0) = \mathbb{P}^*(Y_1 = 1) - \mathbb{P}^*(Y_0 = 1)$$

where  $Y_a$  denotes the counterfactual outcome (mortality), if possibly contrary to fact, the pirate had scurvy status A = a.

#### 4. Specify the link between the SCM and the observed data:

The observed data were generated by sampling n independent times from a data generating system compatible with the structural causal model  $\mathcal{M}^*$ . This yields n i.i.d. copies of random variable  $O = (W1, W2, A, Y) \sim \mathbb{P}_0$ . The statistical model  $\mathcal{M}$  for the set of allowed distributions of the observed data is non-parametric.

#### 5. Assess identifiability:

The target causal parameter is not identified from the observed data distribution. There are several unblockable backdoor paths from the outcome Y into the exposure A. For identifiability to hold, we would need  $U_A \perp \!\!\!\perp U_Y$  and (i)  $U_A \perp \!\!\!\perp U_{W1}$ ,  $U_A \perp \!\!\!\perp U_{W2}$  or (ii)  $U_Y \perp \!\!\!\perp U_{W1}$ ,  $U_Y \perp \!\!\!\perp U_{W2}$ .

#### 6. Specify the target parameter of the observed data distribution:

Despite lack of identifiability, we can still "commit" to an interesting statistical estimand inspired by our scientific/causal question. Let W = (W1, W2) denote our adjustment set; then our statistical estimand is

$$\Psi(\mathbb{P}_0) = \mathbb{E}_0 \left[ \mathbb{E}_0(Y|A=1,W) \right] - \mathbb{E}_0 \left[ \mathbb{E}_0(Y|A=0,W) \right]$$

where the outer expectation is over the distribution of adjustment variables W. This can equivalently be expressed as the IPW estimand:

$$\Psi(\mathbb{P}_0) = \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=1)}{\mathbb{P}_0(A=1|W)} Y \right] - \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=0)}{\mathbb{P}_0(A=0|W)} Y \right]$$

where the second equality holds due to the linearity of expectations.

For identifiability, we also need the positivity assumption to hold:

$$min_{a \in \mathcal{A}} \mathbb{P}_0(A = a|W = w) > 0$$

for all w for which  $\mathbb{P}_0(W=w) > 0$ . In words, we need that each possible exposure level is represented in every covariate strata. This condition on data support ensures that our statistical estimand is well-defined. In our example, there must be a positive probability of having scurvy or not, within strata of "awesome" pirate status and route danger.

We have not changed our statistical model  $\mathcal{M}$ , which remains non-parametric.

#### 7. Estimate the chosen parameter of the observed data distribution:

We have discussed two estimators of the statistical parameter. They rely on estimating different parts of the observed data distribution  $\mathbb{P}_0$ :

(a) Simple substitution estimator based on the G-Computation formula:

$$\hat{\Psi}_{SS}(\mathbb{P}_n) = \frac{1}{n} \sum_{i=1}^n \hat{\mathbb{E}}(Y|A=1, W_i) - \frac{1}{n} \sum_{i=1}^n \hat{\mathbb{E}}(Y|A=0, W_i)$$

where  $\mathbb{P}_n$  is the empirical distribution and  $\hat{\mathbb{E}}(Y|A,W)$  is the estimate of the conditional mean outcome given the exposure (scurvy) and adjustment variables  $\mathbb{E}_0(Y|A,W)$ .

(b) Inverse probability weighted estimator (IPTW):

$$\begin{split} \hat{\Psi}_{IPTW}(\mathbb{P}_n) &= \frac{1}{n} \sum_{i=1}^n \left( \frac{\mathbb{I}(A_i = 1)}{\hat{\mathbb{P}}(A = 1 | W_i)} - \frac{\mathbb{I}(A_i = 0)}{\hat{\mathbb{P}}(A = 0 | W_i)} \right) Y_i \\ &= \frac{1}{n} \sum_{i=1}^n \left( \frac{\mathbb{I}(A_i = 1)}{\hat{\mathbb{P}}(A = 1 | W_i)} \right) Y_i - \frac{1}{n} \sum_{i=1}^n \left( \frac{\mathbb{I}(A_i = 0)}{\hat{\mathbb{P}}(A = 0 | W_i)} \right) Y_i \end{split}$$

where  $\hat{\mathbb{P}}(A=1|W_i)$  is an estimate of the conditional probability of scurvy, given pirate *i*'s baseline characteristics  $\mathbb{P}_0(A=1|W_i)$ . This conditional probability is often referred to as the "propensity score". IPTW is the focus of today's lab.

- (c) TMLE: coming soon:)
- 8. Inference and interpret results: Coming soon.

## 2 Import and explore data set RLab3.IPTW.csv.

- 1. Use the read.csv function to import the data set and assign it to data frame ObsData.
- 2. Use the nrow function to count the number of pirates in the data set.
- 3. Use the names, tail and summary functions to explore the data.
- 4. With the table function, check the number of pirates in each covariate strata without scurvy A = 0 and the number of pirates in each adjustment strata with scurvy A = 1. Note: these tables are simply counting the number of observations within each strata of (W1, W2, A) in a single sample of size n; we are not formally evaluating the structural positivity assumption, which is a statistical assumption on the true data generating process  $\mathbb{P}_0$ .

```
Solution:
> # 1. read in data
> ObsData<- read.csv('RLab3.IPTW.csv')</pre>
> # 2. number of pirates
> nrow(ObsData)
[1] 5000
> # 3. get the column names
> names(ObsData)
[1] "W1" "W2" "A" "Y"
> # show the obsv data on the last six pirates
> tail(ObsData)
    W1 W2 A Y
4995 0 1 1 1
4996 1 2 1 1
4997 0 0 0 0
4998 1 2 1 1
4999 0 1 1 1
5000 1 1 0 0
> # recall: W1-awesomeness, W2-danger, A-scurvy, Y-mortality
> summary(ObsData)
                                                     Y
      W1
                      W2
Min. :0.0000 Min. :0.000 Min. :0.0000 Min. :0.000
1st Qu.:0.0000 1st Qu.:1.000 1st Qu.:0.0000 1st Qu.:1.000
Median :1.0000 Median :2.000 Median :1.0000 Median :1.000
Mean :0.5002 Mean :1.494 Mean :0.6788 Mean :0.751
3rd Qu.:1.0000 3rd Qu.:2.000 3rd Qu.:1.0000
                                               3rd Qu.:1.000
Max. :1.0000 Max. :3.000 Max. :1.0000 Max. :1.000
> # 4. use the table function to examine support in exposure-covariate strata
> table(ObsData[,c('W1', 'W2', 'A')])
, , A = 0
  W2
W1 0 1
             2
                3
 0 238 145
             6
 1 290 758 167
, A = 1
  W2
```

W1 0 1 2 3 0 69 801 967 273 1 2 189 806 287

- > # top table corresponds to no scurvy A=0, and bottom table corresponds to scurvy A=1 > # the rows correspond to W1 (awesomeness) and columns to W2 (danger)
- There are certain covariate combinations with little or no variability in the exposure (scurvy). For example, there are zero "not-awesome" pirates (W1 = 0) without scurvy (A = 0) in the highest level of route danger (W2 = 3). Likewise, there are only 2 "awesome" pirates (W1 = 1) with scurvy (A = 1) in the lowest level of route danger (W2 = 0). In the following subsection, we will explore how sparsity (i.e., lack of data support) can affect estimator performance.

## 3 Implement IPTW for a binary exposure

1. Estimate the propensity score  $\mathbb{P}_0(A=1|W)$ , which is the conditional probability of scurvy, given the pirate's characteristics. Use the following *a priori*-specified parametric regression model:

$$\mathbb{P}_{0}(A = 1|W) = logit^{-1} [\beta_{0} + \beta_{1}W1 + \beta_{2}W2]$$

*Hint:* Run glm with specifications family='binomial' for logistic regression and data=ObsData. In practice, we would generally use a machine learning algorithm, such as Super Learner (coming next).

- 2. Predict each pirate's probability of having and not having scurvy, given their covariates:  $\hat{\mathbb{P}}(A=1|W_i)$  and  $\hat{\mathbb{P}}(A=0|W_i)$ .
  - (a) Obtain the predicted probability of having scurvy, given the baseline covariates prob.1W.

    Hint: Apply the predict function to the above fitted logistic regression function with type='response'.
  - (b) Also obtain the predicted probability of not having scurvy, given the baseline covariates prob.OW:

$$\hat{\mathbb{P}}(A=0|W_i) = 1 - \hat{\mathbb{P}}(A=1|W_i)$$

- (c) Use the summary function to examine the distribution of the predicted probabilities prob.1W and prob.0W. Any cause for concern?
- 3. Create the weights for each pirate as an indicator of receiving the exposure of interest, divided by their predicted probability of receiving that exposure
  - (a) Create a vector wt1 as an indicator of having scurvy, divided by the estimated probability of having scurvy given the adjustment set:  $\mathbb{I}(A_i = 1)/\hat{\mathbb{P}}(A = 1|W_i)$ .
    - > wt1 <- as.numeric(ObsData\$A==1)/prob.1W</pre>

where we are coding the indicator function with as.numeric applied to a logical statement.

- (b) Create a vector wt0 as an indicator of noth having scurvy, divided by the estimated probability of not having scurvy given the adjustment set:  $\mathbb{I}(A_i = 0)/\hat{\mathbb{P}}(A = 0|W_i)$ .
  - > wt0 <- as.numeric(ObsData\$A==0)/prob.0W
- (c) Comment on the distribution of the weights.
- 4. Evaluate the IPTW estimand by taking the difference of the empirical means of the weighted outcomes:

$$\hat{\Psi}_{IPTW}(\mathbb{P}_n) = \frac{1}{n} \sum_{i=1}^{n} \frac{\mathbb{I}(A_i = 1)}{\hat{\mathbb{P}}(A = 1|W_i)} Y_i - \frac{1}{n} \sum_{i=1}^{n} \frac{\mathbb{I}(A_i = 0)}{\hat{\mathbb{P}}(A = 0|W_i)} Y_i$$

```
> mean(wt1*ObsData$Y) - mean(wt0*ObsData$Y)
>
This is equivalent to
> mean( (wt1-wt0)*ObsData$Y)
>
```

- 5. Comment on the results.
- 6. Arbitrarily truncate the weights at 10 and evaluate the IPTW estimand.

Hint: The following code copies the weight vector (wt1) into a new vector (wt1.trunc) and truncates the weights at 10.

```
> wt1.trunc<- wt1
> wt1.trunc[wt1.trunc > 10]<- 10</pre>
```

7. Implement the stabilized IPTW estimator (a.k.a., the modified Horvitz-Thompson estimator):

$$\hat{\Psi}_{St.IPTW}(\mathbb{P}_n) = \frac{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 1)}{\hat{\mathbb{P}}(A = 1|W_i)} Y_i}{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 1)}{\hat{\mathbb{P}}(A = 1|W_i)}} - \frac{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 0)}{\hat{\mathbb{P}}(A = 0|W_i)} Y_i}{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 0)}{\hat{\mathbb{P}}(A = 0|W_i)}}$$

Dividing by the mean of the weights ensures that the IPTW estimator is bounded.

```
Solution:
```

```
> # 1. Estimate the treatment mechanism P(A|W)
> # run the logistic regression to estimate the treatment mechanism P(A/W)
> prob.AW.reg<- glm(A ~ W1 +W2, family="binomial", data=ObsData)
> # 2. # predicted probability of having scurvy, given the obs cov P(A=1/W)
> prob.1W <- predict(prob.AW.reg, type= "response")</pre>
> # predicted probability of not having scurvy, given the obs cov P(A=O|W)
> prob.0W <- 1 - prob.1W
> # look at the distribution of predicted probabilities
> summary(prob.1W)
  Min. 1st Qu. Median
                           Mean 3rd Qu.
0.01171 0.21860 0.83212 0.67880 0.99023 0.99958
> summary(prob.0W)
    Min.
            1st Qu.
                       Median
                                          3rd Qu.
                                   Mean
0.0004178 0.0097692 0.1678808 0.3212000 0.7813972 0.9882867
```

IPTW is extremely sensitive to theoretical and practical positivity violations. From above summaries, we see that there are certain covariate combinations with little variability in the exposure (scurvy status).

```
> # 3. Create the weights
> wt1 <- as.numeric(ObsData$A==1)/prob.1W
> wt0 <- as.numeric(ObsData$A==0)/prob.OW
> summary(wt1)
   Min. 1st Qu. Median
                             Mean 3rd Qu.
                                               Max.
 0.0000 0.0000 1.0086 0.9806 1.1748 85.3733
> summary(wt0)
    Min. 1st Qu.
                     Median
                                  Mean 3rd Qu.
                                                     Max.
  0.0000
            0.0000
                     0.0000
                               0.8843
                                         1.2424 117.9966
"Near" violations of the positivity assumptions often yield poor finite sample performance. Here, at least
one pirate is being up-weighted by 118.
> # 4. Point estimate:
> IPTW<- mean( wt1*ObsData$Y) - mean( wt0*ObsData$Y)
> IPTW
[1] 0.4450127
> mean( (wt1-wt0)*ObsData$Y)
[1] 0.4450127
5. The IPTW estimate of \Psi(\mathbb{P}_0) is 44.5%. The true value is \psi_0 = 28.7\%. We can interpret the statistical
estimand \psi_0 as the marginal difference in the mortality risk associated with scurvy, after controlling for
pirate characteristics and route danger. If the identifiability assumptions (i.e., randomization and positivity)
held, we could then interpret \psi_0 in terms of the average treatment effect (a.k.a., the causal risk difference).
> # 6. truncate weights ARBITRARILY at 10
> # first let's see how many weights under the exposure are greater than 10
> sum(wt1>10)
[1] 2
> wt1.trunc<- wt1
> wt1.trunc[ wt1.trunc>10] <-10
> # same for weights under no exposure
> sum(wt0>10)
[1] 8
> wt0.trunc<- wt0
> wt0.trunc[ wt0.trunc>10] <-10
> # evaluate the IPTW estimand with the truncated weights
> mean( wt1.trunc*ObsData$Y) - mean( wt0.trunc*ObsData$Y)
```

#### [1] 0.5814045

By bounding the predicted probabilities (weights), we are ensuring that the estimator of the propensity score  $\mathbb{P}_0(A=1|W)$  is not consistent and thereby the resulting IPTW estimator will be biased. The point estimate from the IPTW estimator, using truncated weights, was 58.1%; the true value of the statistical estimand was  $\psi_0 = 28.7\%$ .

```
> # 7. Stabilized IPTW estimator - Modified Horvitz-Thompson estimator
> mean( wt1*ObsData$Y)/mean( wt1) - mean( wt0*ObsData$Y)/mean( wt0)
```

[1] 0.3963289

```
> # this is equivalent to
> sum( wt1*ObsData$Y)/sum( wt1) - sum( wt0*ObsData$Y)/sum( wt0)
```

[1] 0.3963289

For this single sample of n pirates, the modified Horvitz-Thompson estimator yielded a point estimate 39.6% that was closest to the true value.

As shown in the Appendix, we were using the correctly specified regression to estimate the propensity score - and still saw a big between the true value of 28.7% and our estimates. To evaluate the performance of these 3 IPTW estimators, we could draw another independent sample of size n, implement the three estimators, and repeat 500+ times. Then we could evaluate the bias, variance, and mean squared error of these estimators for this data generating process. See R Lab2 and R homework 2.

## 4 Extensions to handle missingness

In the previous dataset, all pirates were magically followed to death or conclusion of their voyage - regardless of their characteristics or the route danger (e.g., Bermuda Triangle). More "realistically", we have to deal with incomplete measurement of outcome. One approach to handle missingness is to modify our scientific question: "what would be the causal effect of scurvy on mortality among pirates under complete measurement of the outcome?". As extra practice, you can work through the Causal Roadmap for this modified question. For the purposes of the R lab, we are going to skip to estimation.

In the following, let  $\Delta$  be an indicator that the outcome was observed, and redefine the outcome Y equal to 1 if the pirate was observed/reported to have died and 0 otherwise (either did not die or had a missing outcome). Again let W = (W1, W2) denote our adjustment set. We are now focused on estimating the following statistical estimand, corresponding to this causal question if the identifiability assumptions held:

$$\begin{split} &\Psi(\mathbb{P}_0) = \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=1,\Delta=1)}{\mathbb{P}_0(A=1,\Delta=1\mid W)} Y \right] - \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=0,\Delta=1)}{\mathbb{P}_0(A=0,\Delta=1\mid W)} Y \right] \\ &= \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=1,\Delta=1)}{\mathbb{P}_0(A=1\mid W)\mathbb{P}(\Delta=1\mid A,W)} Y \right] - \mathbb{E}_0 \left[ \frac{\mathbb{I}(A=0,\Delta=1)}{\mathbb{P}_0(A=0\mid W)\mathbb{P}_0(\Delta=1\mid A,W)} Y \right] \end{split}$$

where in the second equality, we factored the denominator of the weights according to the assumed time-ordering: the exposure of scurvy happens before measurement/missingness on the outcome.

1. Use the read.csv function to import the modified data RLab3.IPTW.missing.csv, and assign it to object ObsData.

- 2. Use the summary function to explore the dataset. How many pirates have their outcome measured?
- 3. Estimate the propensity score  $\mathbb{P}_0(A=1|W)$ , which is the conditional probability of scurvy, given the pirate's characteristics. Use the following *a priori*-specified parametric regression model:

$$\mathbb{P}_0(A=1|W) = logit^{-1} \left[ \beta_0 + \beta_1 W 1 + \beta_2 W 2 \right]$$

Hint: We already did this. Skip to the next step.

4. Predict each pirate's probability of having and not having scurvy, given their personal characteristics, and route danger:  $\hat{\mathbb{P}}(A=1|W_i)$  and  $\hat{\mathbb{P}}(A=0|W_i)$ .

Hint: We already did this; the vectors of the corresponding probabilies are given by prob. 1W and prob. OW, respectively. Skip to the next step.

5. Estimate the probability of being measured, given the exposure, personal characteristics, and route danger:  $\mathbb{P}_0(\Delta = 1|A, W)$ . Use the following *a priori*-specified parametric regression model:

$$\mathbb{P}_0(\Delta = 1|A, W1, W2) = logit^{-1} [\beta_0 + \beta_1 W1 + \beta_2 W2 + \beta_3 A]$$

6. Predict each pirate's probability of being measured, given their observed past  $\hat{\mathbb{P}}(\Delta = 1 | A_i, W_i)$ . Name the vector of the resulting probabilities probabilities probabilities to examine the distribution of probabilities. Any cause for concern?

Hint: Apply the predict function to the above fitted logistic regression function with type='response'.

- 7. Create the weights for each pirate as an indicator of receiving the exposure of interest and being measured, divided by their predicted probability of receiving that exposure and being measured.
  - (a) Create a vector wt1 with numerator as an indicator of having scurvy and being measured, and with denominator as the estimated probability of having scurvy, given the adjustment set, times the estimated probability of being measured, given the observed past:

$$wt1_i = \frac{\mathbb{I}(A_i = 1, \Delta_i = 1)}{\hat{\mathbb{P}}(A = 1|W_i) \times \hat{\mathbb{P}}(\Delta = 1|A_i, W_i)}$$

- > wt1 <- as.numeric(ObsData\$A==1 & ObsData\$Delta==1)/(prob.1W\*prob.delta)
- (b) Create a vector wt0 with numerator as an indicator of not having scurvy and being measured, and with denominator as the estimated probability of not having scurvy, given the adjustment set, times the estimated probability of being measured, given the observed past:

$$wt0_i = \frac{\mathbb{I}(A_i = 0, \Delta_i = 1)}{\hat{\mathbb{P}}(A = 0|W_i) \times \hat{\mathbb{P}}(\Delta = 1|A_i, W_i)}$$

- > wt0 <- as.numeric(ObsData\$A==0 & ObsData\$Delta==1)/(prob.OW\*prob.delta)
- (c) Comment on the distribution of the weights.
- 8. Evaluate the IPTW estimand by taking the difference of the empirical means of the weighted outcomes:

$$\hat{\Psi}_{IPTW}(\mathbb{P}_n) = \frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 1, \Delta_i = 1)}{\hat{\mathbb{P}}(A = 1|W_i)\hat{\mathbb{P}}(\Delta = 1|A_i, W_i)} Y_i - \frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 0, \Delta_i = 1)}{\hat{\mathbb{P}}(A = 0|W_i)\hat{\mathbb{P}}(\Delta = 1|A_i, W_i)} Y_i$$

9. Arbitrarily truncate the weights at 10 and evaluate the IPTW estimand.

10. Implement the stabilized IPTW estimator (a.k.a., the modified Horvitz-Thompson estimator).

11. Comment on the results.

```
Solution:
> # 1. read in data
> ObsData <- read.csv('RLab3.IPTW.missing.csv')</pre>
> # 2. summarize
> summary(ObsData)
      W1
                      W2
                                                   Delta
Min. :0.0000 Min. :0.000 Min. :0.0000 Min. :0.000
1st Qu.:0.0000 1st Qu.:1.000 1st Qu.:0.0000 1st Qu.:0.000
Median: 1.0000 Median: 2.000 Median: 1.0000 Median: 0.000
Mean :0.5002 Mean :1.494 Mean :0.6788
                                               Mean :0.353
3rd Qu.:1.0000 3rd Qu.:2.000 3rd Qu.:1.0000
                                                3rd Qu.:1.000
Max. :1.0000 Max. :3.000 Max. :1.0000 Max. :1.000
      Y
Min. :0.000
 1st Qu.:0.000
Median : 0.000
Mean :0.135
 3rd Qu.:0.000
Max. :1.000
> # how many were measured
> sum(ObsData$Delta==1)
[1] 1765
> # 3. skip
> prob.AW.reg
Call: glm(formula = A ~ W1 + W2, family = "binomial", data = ObsData)
Coefficients:
                                W2
(Intercept)
                    W1
                -3.161
                              3.018
Degrees of Freedom: 4999 Total (i.e. Null); 4997 Residual
Null Deviance:
                        6278
Residual Deviance: 3100
                             AIC: 3106
> # 4. skip
> summary(prob.1W); summary(prob.0W)
  Min. 1st Qu. Median
                         Mean 3rd Qu.
0.01171 0.21860 0.83212 0.67880 0.99023 0.99958
```

```
1st Qu.
                       Median
                                   Mean
                                          3rd Qu.
0.0004178\ 0.0097692\ 0.1678808\ 0.3212000\ 0.7813972\ 0.9882867
> # 5. estimate the probability of being measured, given the past
> prob.Delta.reg <- glm(Delta ~ W1 + W2 + A, family='binomial', data=ObsData)
> # 6. predicted probability of measurement, given the past
> prob.delta <- predict(prob.Delta.reg, type='response')</pre>
> summary(prob.delta)
            1st Qu. Median
                                   Mean
                                          3rd Qu.
     Min.
0.0000832 0.0152631 0.2727635 0.3530000 0.7498011 0.9982111
> # Eeek! Some small probabilities
> # 7 create the weights
> wt1 <- as.numeric(ObsData$A==1 & ObsData$Delta==1)/(prob.1W*prob.delta)
> wt0 <- as.numeric(ObsData$A==0 & ObsData$Delta==1)/(prob.0W*prob.delta)
> summary(wt1)
   Min. 1st Qu. Median Mean 3rd Qu.
                                                 Max.
  0.0000 0.0000 0.0000 0.9946 0.0000 501.6879
> summary(wt0)
  Min. 1st Qu. Median
                           Mean 3rd Qu.
0.0000 0.0000 0.0000 0.7531 1.0137 60.9922
"Near" violations of the positivity assumptions often yield poor finite sample performance. Here, at least
one pirate is being up-weighted by 502.
> # 8. Point estimate:
> IPTW<- mean( wt1*ObsData$Y) - mean( wt0*ObsData$Y)</pre>
> IPTW
[1] 0.5702749
> # 9. truncate weights ARBITRARILY at 10
> sum(wt1>10)
[1] 40
> wt1.trunc<- wt1
> wt1.trunc[ wt1.trunc>10] <-10
> # same for weights under no exposure
> sum(wt0>10)
```

```
[1] 103

> wt0.trunc<- wt0
> wt0.trunc[ wt0.trunc>10] <-10
> # evaluate the IPTW estimand with the truncated weights
> mean( wt1.trunc*ObsData$Y) - mean( wt0.trunc*ObsData$Y)

[1] 0.3084572

> # 10 Stabilized IPTW estimator - Modified Horvitz-Thompson estimator
> mean( wt1*ObsData$Y)/mean( wt1) - mean( wt0*ObsData$Y)/mean( wt0)

[1] 0.4430749

11. The new IPTW estimate of Ψ(P<sub>0</sub>) is 57.0%. The true value is still ψ<sub>0</sub> = 28.7%. The point estimate
```

Again, to evaluate the performance of these 3 IPTW estimators, we should draw another independent sample of size n, implement the three estimators, and repeat 500+ times. Then we could evaluate the bias, variance, and mean squared error of these estimators for this data generating process. See R Lab2 and R homework 2.

from the IPTW estimator, using truncated weights, is now closest to the truth: 30.8%. Now the stabilied

## Solution:

IPW yielded a point estimate 44.3%.

# Appendix A: a specific data generating process

The following code was used to generate the data set RLab3.IPTW.csv. In this data generating process (one of many compatible with the structural causal model  $\mathcal{M}^*$ ), all background errors are independent.

```
Y.1<- get.outcome(W1, W2, A=1, U.Y)
   Y.0<- get.outcome(W1, W2, A=0, U.Y)
   # outcome if no missingness
+ Y <- get.outcome(W1, W2, A, U.Y)
  # outcome if allow for missingess
   Y.na <- Y
    Y.na[Delta==0] <- 0
    data.frame(W1, W2, A, Y.1, Y.0, Delta, Y, Y.na)
> # create the RLab3.IPTW.csv & RLab3.IPTW.missing.csv
> set.seed(252)
> Full<- genData(n=5000)
> ObsData<- subset(Full, select=c(W1,W2,A,Y))</pre>
> write.csv(ObsData, file="RLab3.IPTW.csv", row.names=F)
> ObsData<- subset(Full, select=c(W1, W2, A, Delta, Y.na))
> colnames(ObsData) <- c('W1', 'W2', 'A', 'Delta','Y')
> write.csv(ObsData, file="RLab3.IPTW.missing.csv", row.names=F)
   • Given this specific data generating process, we could estimate the true value of the average treatment
effect by drawing a huge number of observations and taking the mean difference in the counterfactual
outcomes.
> set.seed(252)
> # calculate true ATE by drawing a huge number of observations
> nTot=100000
> Full <- genData(n=nTot)</pre>
> Psi.star <- mean(Full$Y.1) - mean(Full$Y.0)</pre>
> Psi.star
[1] 0.28653
```

The counterfactual risk of mortality would  $\Psi^*(\mathbb{P}^*)=28.7\%$  higher if all pirates had scurvy than if all pirates did not have scurvy. Avast! Pirates need their vitamin C!