## A guided tour in targeted learning territory

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## 1 Introduction

```
set.seed(54321) ## because reproducibility matters...
suppressMessages(library(R.utils)) ## make sure it is installed
suppressMessages(library(tidyverse)) ## make sure it is installed
suppressMessages(library(caret)) ## make sure it is installed
suppressMessages(library(ggdag)) ## make sure it is installed
expit <- plogis
logit <- qlogis
redo_fixed <- c(TRUE, FALSE)[2]</pre>
redo_varying <- c(TRUE, FALSE)[2]</pre>
## if 'redo_$' then recompute 'learned_features_$_sample_size', otherwise
## upload it if it is not already in the environment.
if (!redo_fixed) {
  if (!exists("learned_features_fixed_sample_size")) {
   learned_features_fixed_sample_size <-</pre>
      loadObject("data/learned_features_fixed_sample_size_new.xdr")
 }
```

```
if (!redo_varying) {
   if (!exists("learned_features_varying_sample_size")) {
     learned_features_varying_sample_size <-
        loadObject("data/learned_features_varying_sample_size_new.xdr")
   }
}</pre>
```

Function expit implements the link function expit :  $\mathbb{R} \to ]0,1[$  given by  $\exp it(x) \equiv (1+e^{-x})^{-1}$ . Function  $\log it$  implements its inverse function  $\log it: ]0,1[\to \mathbb{R}$  given by  $\log it(p) \equiv \log[p/(1-p)]$ .

## 2 A simulation study

blabla

2.1 Reproducible experiment as a law. We are interested in a reproducible experiment. The generic summary of how one realization of the experiment unfolds, our observation, is called O. We view O as a random variable drawn from what we call the law  $P_0$  of the experiment. The law  $P_0$  is viewed as an element of what we call the model. Denoted by  $\mathcal{M}$ , the model is the collection of all laws from which O can be drawn and that meet some constraints. The constraints translate the knowledge we have about the experiment. The more we know about the experiment, the smaller is  $\mathcal{M}$ . In all our examples, model  $\mathcal{M}$  will put very few restrictions on the candidate laws.

Consider the following chunk of code:

```
run_experiment <- function(n, ideal = FALSE) {</pre>
  ## preliminary
  n <- Arguments$getInteger(n, c(1, Inf))</pre>
  ideal <- Arguments$getLogical(ideal)</pre>
  ## ## 'Gbar' and 'Qbar' factors
  Gbar <- function(W) {</pre>
    expit(1 + 2 * W - 4 * sqrt(abs((W - 5/12))))
  }
  Qbar <- function(AW) {
    A \leftarrow AW[, 1]
    W \leftarrow AW[, 2]
    ## A * (\cos((1 + W) * pi / 4) + (1/3 \le W \& W \le 1/2) / 5) +
    ## (1 - A) * (\sin(4 * W^2 * pi) / 4 + 1/2)
    A * (cos((-1/2 + W) * pi) * 2/5 + 1/5 + (1/3 <= W & W <= 1/2) / 5 +
          (W >= 3/4) * (W - 3/4) * 2) +
      (1 - A) * (\sin(4 * W^2 * pi) / 4 + 1/2)
  }
  ## sampling
  ## ## context
  mixture_weights <-c(1/10, 9/10, 0)
  mins <-c(0, 11/30, 0)
  maxs <- c(1, 14/30, 1)
  latent <- findInterval(runif(n), cumsum(mixture_weights)) + 1</pre>
  W <- runif(n, min = mins[latent], max = maxs[latent])</pre>
  ## ## counterfactual rewards
```

```
zeroW <- cbind(A = 0, W)
oneW \leftarrow cbind(A = 1, W)
Qbar.zeroW <- Qbar(zeroW)
Qbar.oneW <- Qbar(oneW)
Yzero <- rbeta(n, shape1 = 2, shape2 = 2 * (1 - Qbar.zeroW) / Qbar.zeroW)
Yone <- rbeta(n, shape1 = 3, shape2 = 3 * (1 - Qbar.oneW) / Qbar.oneW)
## ## action undertaken
A <- rbinom(n, size = 1, prob = Gbar(W))
## ## actual reward
Y \leftarrow A * Yone + (1 - A) * Yzero
## ## observation
if (ideal) {
  obs <- cbind(W = W, Yzero = Yzero, Yone = Yone, A = A, Y = Y)
} else {
  obs \leftarrow cbind(W = W, A = A, Y = Y)
}
attr(obs, "Gbar") <- Gbar</pre>
attr(obs, "Qbar") <- Qbar</pre>
attr(obs, "QW") <- function(W) {</pre>
  out <- sapply(1:length(mixture_weights),</pre>
                 function(ii){
                   mixture_weights[ii] *
                      dunif(W, min = mins[ii], max = maxs[ii])
                 })
  return(rowSums(out))
}
attr(obs, "qY") <- function(AW, Y, Qbar){</pre>
  A \leftarrow AW[, 1]
  W \leftarrow AW[, 2]
  Qbar.AW <- do.call(Qbar, list(AW)) # is call to 'do.call' necessary?
  shape1 <- ifelse(A == 0, 2, 3)
  dbeta(Y, shape1 = shape1, shape2 = shape1 * (1 - Qbar.AW) / Qbar.AW)
}
##
return(obs)
```

We can interpret run\_experiment as a law  $P_0$  since we can use the function to sample observations from a common law. It is even a little more than that, because we can tweak the experiment, by setting its ideal argument to TRUE, in order to get what appear as intermediary (counterfactual) variables in the regular experiment. The next chunk of code runs the (regular) experiment five times independently and outputs the resulting observations:

```
(five_obs <- run_experiment(5))</pre>
```

```
## W A Y
## [1,] 0.4533028 0 0.8979460
## [2,] 0.3716077 0 0.9905312
## [3,] 0.3875802 0 0.8080567
## [4,] 0.4008279 1 0.9954100
## [5,] 0.4038325 0 0.9772926
## attr(,"Gbar")
## function (W)
```

```
## {
##
                       expit(1 + 2 * W - 4 * sqrt(abs((W - 5/12))))
## }
## <bytecode: 0x7fd1fd7e7d00>
## <environment: 0x7fd1fd1df720>
## attr(,"Qbar")
## function (AW)
## {
##
                       A \leftarrow AW[, 1]
##
                       W \leftarrow AW[, 2]
                       A * (cos((-1/2 + W) * pi) * 2/5 + 1/5 + (1/3 \le W \& W \le
                                    1/2)/5 + (W >= 3/4) * (W - 3/4) * 2) + (1 - A) * (sin(4 * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) * 3/4) 
##
##
                                   W^2 * pi)/4 + 1/2
## }
## <bytecode: 0x7fd1fe50d3f8>
## <environment: 0x7fd1fd1df720>
## attr(,"QW")
## function (W)
##
##
                       out <- sapply(1:length(mixture weights), function(ii) {
##
                                   mixture_weights[ii] * dunif(W, min = mins[ii], max = maxs[ii])
##
                       return(rowSums(out))
##
## }
        <br/>
<br/>
<br/>
de: 0x7fd2606e2ad8>
## <environment: 0x7fd1fd1df720>
## attr(,"qY")
## function (AW, Y, Qbar)
## {
##
                       A \leftarrow AW[, 1]
##
                       W \leftarrow AW[, 2]
##
                       Qbar.AW <- do.call(Qbar, list(AW))</pre>
##
                       shape1 <- ifelse(A == 0, 2, 3)
##
                       dbeta(Y, shape1 = shape1, shape2 = shape1 * (1 - Qbar.AW)/Qbar.AW)
## }
## <bytecode: 0x7fd1ffa9e5e0>
## <environment: 0x7fd1fd1df720>
```

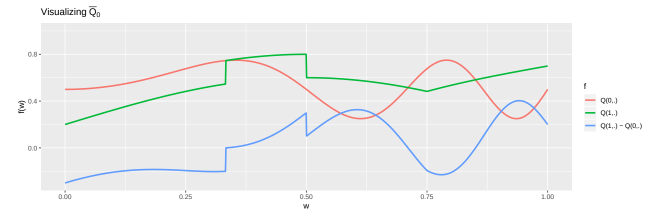
We can view the attributes of object five\_obs because, in this section, we act as oracles, i.e., we know completely the nature of the experiment. In particular, we have included several features of  $P_0$  that play an important role in our developments. The attribute QW describes the density of W, of which the law  $Q_{0,W}$  is a mixture of the uniform laws over [0,1] (weight 1/10) and [11/30,14/30] (weight 9/10).\* The attribute Gbar describes the conditional probability of action A=1 given W. For each  $a\in\{0,1\}$ , we denote  $\bar{G}_0(W)\equiv \Pr_{P_0}(A=1|W)$  and  $\ell\bar{G}_0(a,W)\equiv \Pr_{P_0}(A=a|W)$ . Obvisously,  $\ell\bar{G}_0(A,W)\equiv A\bar{G}_0(W)+(1-A)(1-\bar{G}_0(W))$ . The attribute qY describes the conditional density of Y given A and W. For each  $y\in ]0,1[$ , we denote by  $q_{0,Y}(y,A,W)$  the conditional density evaluated at y of Y given A and W. Similarly, the attribute Qbar describes the conditional mean of Y given A and W, and we denote  $\bar{Q}_0(A,W)=\mathrm{E}_{P_0}(Y|A,W)$  the conditional mean of Y given A and W.



<sup>\*</sup>We fine-tuned (or tweaked, or something else?) the marginal law of W to make it easier later on to drive home important messages. Specifically, . . . (do we explain what happens?)

1. Run the following chunk of code. It visualizes the conditional mean  $\bar{Q}_0$ .

```
Gbar <- attr(five_obs, "Gbar")</pre>
Qbar <- attr(five_obs, "Qbar")</pre>
QW <- attr(five_obs, "QW")
features <- tibble(w = seq(0, 1, length.out = 1e3)) %>%
  mutate(Qw = QW(w),
         Gw = Gbar(w),
         Q1w = Qbar(cbind(A = 1, W = w)),
         QOw = Qbar(cbind(A = 0, W = w)),
         blip_Qw = Q1w - Q0w)
features %>% select(-Qw, -Gw) %>%
  rename("Q(1,.)" = Q1w,
         "Q(0,.)" = Q0w,
         Q(1,.) - Q(0,.) = blip_Qw %
  gather("f", "value", -w) %>%
  ggplot() +
  geom_line(aes(x = w, y = value, color = f), size = 1) +
  labs(y = "f(w)", title = bquote("Visualizing" ~ bar(Q)[0])) +
 ylim(NA, 1)
```



- 2. Adapt the above chunk of code to visualize the marginal density  $Q_{0,W}$  and conditional probability  $\bar{G}_0$ .
- **2.2** The parameter of interest, first pass. It happens that we especially care for a finite-dimensional feature of  $P_0$  that we denote by  $\psi_0$ . Its definition involves two of the aforementioned infinite-dimensional features:

$$\psi_0 \equiv \int \left( \bar{Q}_0(1, w) - \bar{Q}_0(0, w) \right) dQ_{0,W}(w)$$

$$= \mathcal{E}_{P_0} \left( \mathcal{E}_{P_0}(Y \mid A = 1, W) - \mathcal{E}_{P_0}(Y \mid A = 0, W) \right).$$
(1)

Acting as oracles, we can compute explicitly the numerical value of  $\psi_0$ .

```
integrand <- function(w) {
   Qbar <- attr(five_obs, "Qbar")
   QW <- attr(five_obs, "QW")
   ( Qbar(cbind(1, w)) - Qbar(cbind(0, w)) ) * QW(w)
}
(psi_zero <- integrate(integrand, lower = 0, upper = 1)$val)</pre>
```

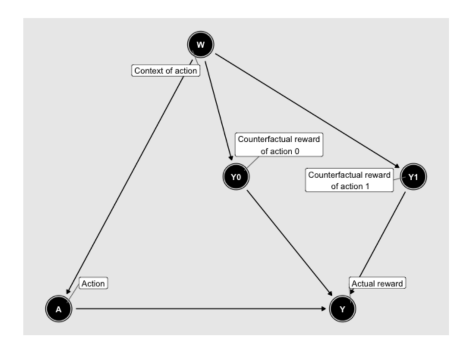


Figure 1: Causal graph summarizing the inner causal mechanism at play in 'run\_experiment'.

#### ## [1] 0.08317711

Our interest in  $\psi_0$  is of causal nature. Taking a closer look at run\_experiment reveals indeed that the random making of an observation O drawn from  $P_0$  can be summarized by the following causal graph and nonparametric system of structural equations:

and, for some deterministic functions  $f_w$ ,  $f_a$ ,  $f_y$  and independent sources of randomness  $U_w$ ,  $U_a$ ,  $U_y$ ,

- 1. sample the context where the counterfactual rewards will be generated, the action will be undertaken and the actual reward will be obtained,  $W = f_w(U_w)$ ;
- 2. sample the two counterfactual rewards of the two actions that can be undertaken,  $Y_0 = f_y(0, W, U_y)$  and  $Y_1 = f_y(1, W, U_y)$ ;

- 3. sample which action is carried out in the given context,  $A = f_a(W, U_a)$ ;
- 4. define the corresponding reward,  $Y = AY_1 + (1 A)Y_0$ ;
- 5. summarize the course of the experiment with the observation O = (W, A, Y), thus concealing  $Y_0$  and  $Y_1$ .

The above description of the experiment run\_experiment is useful to reinforce what it means to run the "ideal" experiment by setting argument ideal to TRUE in a call to run\_experiment. Doing so triggers a modification of the nature of the experiment, enforcing that the counterfactual rewards  $Y_0$  and  $Y_1$  be part of the summary of the experiment eventually. In light of the above enumeration,  $\mathbb{O} \equiv (W, Y_0, Y_1, A, Y)$  is output, as opposed to its summary measure O. This defines another experiment and its law, that we denote  $\mathbb{P}_0$ .

It is straightforward to show<sup>†</sup> that

$$\psi_0 = \mathcal{E}_{\mathbb{P}_0}(Y_1 - Y_0) = \mathcal{E}_{\mathbb{P}_0}(Y_1) - \mathcal{E}_{\mathbb{P}_0}(Y_0). \tag{2}$$

Thus,  $\psi_0$  describes the average difference of the two counterfactual rewards. In other words,  $\psi_0$  quantifies the difference in average of the reward one would get in a world where one would always enforce action a=1 with the reward one would get in a world where one would always enforce action a=0. This said, it is worth emphasizing that  $\psi_0$  is a well-defined parameter beyond its causal interpretation, and that it describes a standardized association between the action A and reward Y.

To conclude this subsection, we use our position as oracles to sample observations from the ideal experiment. We call run\_experiment with its argument ideal set to TRUE in order to numerically approximate  $\psi_0$ . By the law of large numbers, the following code approximates  $\psi_0$  and shows it approximate value.

```
B <- 1e6
ideal_obs <- run_experiment(B, ideal = TRUE)
(psi_approx <- mean(ideal_obs[, "Yone"] - ideal_obs[, "Yzero"]))</pre>
```

#### ## [1] 0.08332233

The object  $psi_approx$  contains an approximation to  $\psi_0$  based on B observations from the ideal experiment. The random sampling of observations results in uncertainty in the numerical approximation of  $\psi_0$ . This uncertainty can be quantified by constructing a 95% confidence interval for  $\psi_0$ . The central limit theorem and Slutsky's lemma<sup>‡</sup> allow us to build such an interval as follows.

```
sd_approx <- sd(ideal_obs[, "Yone"] - ideal_obs[, "Yzero"])
alpha <- 0.05
(psi_approx_CI <- psi_approx + c(-1, 1) * qnorm(1 - alpha / 2) * sd_approx / sqrt(B))</pre>
```

†For a = 0, 1,

$$\begin{split} \mathbf{E}_{\mathbb{P}_{0}}(Y_{a}) &= \int \mathbf{E}_{\mathbb{P}_{0}}(Y_{a} \mid W = w)dQ_{0,W}(w) = \int \mathbf{E}_{\mathbb{P}_{0}}(Y_{a} \mid A = a, W = w)dQ_{0,W}(w) \\ &= \int \mathbf{E}_{P_{0}}(Y \mid A = a, W = w)dQ_{0,W}(w) = \int \bar{Q}_{0}(a, W)dQ_{0,W}(w). \end{split}$$

The second equality follows from the conditional independence of the counterfactual rewards  $(Y_0,Y_1)$  and action A given W (in words, the randomization assumption " $(Y_0,Y_1) \perp A \mid W$ " is met under  $\mathbb{P}_0$ ). The third equality results from the facts that the observed reward Y equals the counterfactual reward  $Y_a$  when A=a (in words, the consistency assumption " $Y_a=Y \mid A=a$ " is met under  $\mathbb{P}_0$ ) and that  $\Pr_{P_0}(\ell \bar{G}_0(a,W)>0)=1$  (in words, the positivity assumption is met under  $P_0$  — this is needed for  $\operatorname{E}_{P_0}(Y\mid A=a,W)$  to be well-defined).

†Let  $X_1,\ldots,X_n$  be independently drawn from a law such that  $\sigma^2 \equiv \operatorname{Var}(X_1)$  is finite. Let  $m \equiv \operatorname{E}(X_1)$  and  $\bar{X}_n \equiv n^{-1} \sum_{i=1}^n X_i$ 

 $^{\ddagger}$ Let  $X_1, \ldots, X_n$  be independently drawn from a law such that  $\sigma^2 \equiv \text{Var}(X_1)$  is finite. Let  $m \equiv \mathrm{E}(X_1)$  and  $\bar{X}_n \equiv n^{-1} \sum_{i=1}^n X_i$  be the empirical mean. It holds that  $\sqrt{n}(\bar{X}_n - m)$  converges in law as n grows to the centered Gaussian law with variance  $\sigma^2$ . Moreover, if  $\sigma_n^2$  is a (positive) consistent estimator of  $\sigma^2$ , then  $\sqrt{n}/\sigma_n(\bar{X}_n - m)$  converges in law to the standard normal law. The empirical variance  $n^{-1} \sum_{i=1}^n (X_i - \bar{X}_n)^2$  is such an estimator. In conclusion, denoting by  $\Phi$  the standard normal distribution function,  $[\bar{X}_n \pm \Phi^{-1}(1-\alpha)\sigma_n/\sqrt{n}]$  is a confidence interval for m with asymptotic level  $(1-2\alpha)$ .

#### ## [1] 0.08271472 0.08392994

We note that the interpretation of this confidence interval is that in 95% of draws of size B from the ideal data generating experiment, the true value of  $\psi_0$  will be contained in the generated confidence interval.



As discussed above, parameter  $\psi_0$  (2) is the difference in average rewards if we enforce action a=1 rather than a=0. An alternative way to describe the rewards under different actions involves quantiles as opposed to averages.

Let  $Q_{0,Y}(y,A,W) = \int_0^y q_{0,Y}(u,A,W)du$  be the conditional cumulative distribution of reward Y given A and W, evaluated at  $y \in ]0,1[$ , that is implied by  $P_0$ . For each action  $a \in \{0,1\}$  and  $c \in ]0,1[$ , introduce

$$\gamma_{0,a,c} \equiv \inf \left\{ y \in ]0,1[: \int Q_{0,Y}(y,a,w)dQ_{0,W}(w) \ge c \right\}.$$
(3)

It is not difficult to check (see Problem 1 below) that

$$\gamma_{0,a,c} = \inf \{ y \in ]0,1[ : \Pr_{\mathbb{P}_0}(Y_a \le y) \ge c \}.$$
 (4)

Thus,  $\gamma_{0,a,c}$  can be interpreted as a covariate-adjusted c-th quantile reward when action a is enforced. The difference

$$\delta_{0,c} \equiv \gamma_{0,1,c} - \gamma_{0,0,c}$$

is the c-th quantile counterpart to parameter  $\psi_0$  (2).

- 1. Prove (4).
- 2. Compute the numerical value of  $\gamma_{0,a,c}$  for each  $(a,c) \in \{0,1\} \times \{1/4,1/2,3/4\}$  using the appropriate attributes of five\_obs. Based on these results, report the numerical value of  $\delta_{0,c}$  for each  $c \in \{1/4,1/2,3/4\}$ .
- 3. Approximate the numerical values of  $\gamma_{0,a,c}$  for each  $(a,c) \in \{0,1\} \times \{1/4,1/2,3/4\}$  by drawing a large sample from the "ideal" data experiment and using empirical quantile estimates. Deduce from these results a numerical approximation to  $\delta_{0,c}$  for  $c \in \{1/4,1/2,3/4\}$ . Confirm that your results closely match those obtained in the previous problem.
- **2.3** The parameter of interest, second pass. Suppose we know beforehand that O drawn from  $P_0$  takes its values in  $\mathcal{O} \equiv [0,1] \times \{0,1\} \times [0,1]$  and that  $\bar{G}(W) = P_0(A=1|W)$  is bounded away from zero and one  $Q_{0,W}$ -almost surely (this is the case indeed). Then we can define model  $\mathcal{M}$  as the set of all laws P on  $\mathcal{O}$  such that  $\bar{G}(W) \equiv P(A=1|W)$  is bounded away from zero and one  $Q_W$ -almost surely, where  $Q_W$  is the marginal law of W under P.

Let us also define generically  $\bar{Q}$  as

$$\bar{Q}(A, W) \equiv E_P(Y|A, W)$$
,

where, for simplicity, we have suppressed the dependence of  $\bar{Q}$  on P. Central to our approach is viewing  $\psi_0$  as the value at  $P_0$  of the statistical mapping  $\Psi$  from  $\mathcal{M}$  to [0,1] characterized by

$$\Psi(P) \equiv \int \left(\bar{Q}(1, w) - \bar{Q}(0, w)\right) dQ_W(w)$$

$$= \mathbf{E}_P \left( \bar{Q}(1, W) - \bar{Q}(0, W) \right),\,$$

a clear extension of (1). For instance, although the law  $\Pi_0 \in \mathcal{M}$  encoded by default (*i.e.*, with h=0) in run\_another\_experiment defined below differs starkly from  $P_0$ ,

```
run_another_experiment <- function(n, h = 0) {</pre>
  ## preliminary
  n <- Arguments$getInteger(n, c(1, Inf))</pre>
  h <- Arguments$getNumeric(h)</pre>
  ## ## 'Gbar' and 'Qbar' factors
  Gbar <- function(W) {</pre>
    sin((1 + W) * pi / 6)
  Qbar <- function(AW, hh = h) {
    A \leftarrow AW[, 1]
    W \leftarrow AW[, 2]
    expit( logit( A * W + (1 - A) * W^2) +
            hh * 10 * sqrt(W) * A )
  }
  ## sampling
  ## ## context
  W \leftarrow runif(n, min = 1/10, max = 9/10)
  ## ## action undertaken
  A <- rbinom(n, size = 1, prob = Gbar(W))
  ## ## reward
  shape1 <- 4
  QAW <- Qbar(cbind(A, W))
  Y <- rbeta(n, shape1 = shape1, shape2 = shape1 * (1 - QAW) / QAW)
  ## ## observation
  obs \leftarrow cbind(W = W, A = A, Y = Y)
  attr(obs, "Gbar") <- Gbar</pre>
  attr(obs, "Qbar") <- Qbar</pre>
  attr(obs, "QW") <- function(x){dunif(x, min = 1/10, max = 9/10)}
  attr(obs, "shape1") <- shape1</pre>
  attr(obs, "qY") <- function(AW, Y, Qbar, shape1){</pre>
    A \leftarrow AW[,1]; W \leftarrow AW[,2]
    Qbar.AW <- do.call(Qbar, list(AW))</pre>
    dbeta(Y, shape1 = shape1, shape2 = shape1 * (1 - Qbar.AW) / Qbar.AW)
  }
  ##
  return(obs)
```

the parameter  $\Psi(\Pi_0)$  is well defined. Straightforward algebra confirms that  $\Psi(\Pi_0) = 59/300$ , which is confirmed by our numeric computation below.

```
five_obs_from_another_experiment <- run_another_experiment(5)
another_integrand <- function(w) {
    Qbar <- attr(five_obs_from_another_experiment, "Qbar")
    QW <- attr(five_obs_from_another_experiment, "QW")
    ( Qbar(cbind(1, w)) - Qbar(cbind(0, w)) ) * QW(w)
}
(psi_Pi_zero <- integrate(another_integrand, lower = 0, upper = 1)$val)</pre>
```

## [1] 0.1966687



As above, we define  $q_Y(y, A, W)$  to be the conditional density of Y given A and W, evaluated at y, that is implied by a generic  $P \in \mathcal{M}$ . Similarly, we use  $Q_Y$  to denote the corresponding cumulative distribution function. The covariate-adjusted c-th quantile reward for action  $a \in \{0, 1\}$  may be viewed as a mapping  $\Gamma_{a,c}$  from  $\mathcal{M}$  to [0, 1] characterized by

$$\Gamma_{a,c}(P) = \inf \left\{ y \in \left[ 0, 1 \right[ : \int Q_Y(y, a, w) dQ_W(w) \ge c \right\}.$$

The difference in c-th quantile rewards may similarly be viewed as a mapping  $\Delta_c$  from  $\mathcal{M}$  to [0, 1], characterized by  $\Delta_c(P) \equiv \Gamma_{1,c}(P) - \Gamma_{0,c}(P)$ .

- 1. Compute the numerical value of  $\Gamma_{a,c}(\Pi_0)$  for  $(a,c) \in \{0,1\} \times \{1/4,1/2,3/4\}$  using the appropriate attributes of five\_obs\_from\_another\_experiment. Based on these results, report the numerical value of  $\Delta_c(\Pi_0)$  for each  $c \in \{1/4,1/2,3/4\}$ .
- 2. Approximate the value of  $\Gamma_{0,a,c}(\Pi_0)$  for  $(a,c) \in \{0,1\} \times \{1/4,1/2,3/4\}$  by drawing a large sample from the "ideal" data experiment and using empirical quantile estimates. Deduce from these results a numerical approximation to  $\Delta_{0,c}(\Pi_0)$  for each  $c \in \{1/4,1/2,3/4\}$ . Confirm that your results closely match those obtained in the previous problem.
- 3. Building upon the code you wrote to solve the previous problem, construct a confidence interval with asymptotic level 95% for  $\Delta_{0,c}(\Pi_0)$ , with  $c \in \{1/4, 1/2, 3/4\}$ .
- **2.4** The parameter of interest, third pass. In the previous subsection, we reoriented our view of the target parameter to be that of a statistical functional of the law of the observed data. Specifically, we viewed the parameter as a function of specific features of the observed data law, namely  $Q_W$  and  $\bar{Q}$ . It is straightforward  $\P$  to show an equivalent representation of the parameter as

$$\psi_0 = \int \frac{2a - 1}{\ell \bar{G}_0(a, w)} y dP_0(w, a, y)$$

$$= \mathcal{E}_{P_0} \left( \frac{2A - 1}{\ell \bar{G}_0(A, W)} Y \right). \tag{5}$$

Viewing again the parameter as a statistical mapping from  $\mathcal{M}$  to [0,1], it also holds that

$$\Psi(P) = \int \frac{2a-1}{\ell \bar{G}(a,w)} y dP(w,a,y)$$

$$= \mathcal{E}_P \left( \frac{2A-1}{\ell \bar{G}_0(A,W)} Y \right). \tag{6}$$

§Let  $X_1,\ldots,X_n$  be independently drawn from a continuous distribution function F. Set  $p\in ]0,1[$  and, assuming that n is large, find  $k\geq 1$  and  $l\geq 1$  such that  $k/n\approx p-\Phi^{-1}(1-\alpha)\sqrt{p(1-p)/n}$  and  $l/n\approx p+\Phi^{-1}(1-\alpha)\sqrt{p(1-p)/n}$ , where  $\Phi$  is the standard normal distribution function. Then  $[X_{(k)},X_{(l)}]$  is a confidence interval for  $F^{-1}(p)$  with asymptotic level  $1-2\alpha$ .

¶We temporarily drop the subscript  $P_0$  to save space and note, for the same reason, that (2a-1) equals 1 if a=1 and -1 if a=0. Now, for each a=0,1,

$$\begin{split} \mathbf{E}\left(\frac{\mathbf{1}\{A=a\}Y}{\ell\bar{G}(a,W)}\right) &= \mathbf{E}\left(\mathbf{E}\left(\frac{\mathbf{1}\{A=a\}Y}{\ell\bar{G}(a,W)}\bigg|A,W\right)\right) = \mathbf{E}\left(\frac{\mathbf{1}\{A=a\}}{\ell\bar{G}(a,W)}\bar{Q}(A,W)\right) = \mathbf{E}\left(\frac{\mathbf{1}\{A=a\}}{\ell\bar{G}(a,W)}\bar{Q}(a,W)\right) \\ &= \mathbf{E}\left(\mathbf{E}\left(\frac{\mathbf{1}\{A=a\}}{\ell\bar{G}(a,W)}\bar{Q}(a,W)\bigg|W\right)\right) = \mathbf{E}\left(\frac{\ell\bar{G}(a,W)}{\ell\bar{G}(a,W)}\bar{Q}(a,W)\bigg|W\right) = \mathbf{E}\left(\bar{Q}(a,W)\right), \end{split}$$

where the first, fourth and sixth equalities follow from the tower rule, and the second and fifth hold by definition of the conditional expectation, This completes the proof.

Our reason for introducing this alternative view of the target parameter will become clear when we discuss estimation of the target parameter. Specifically, the representations (1) and (5) naturally suggest different estimation strategies for  $\psi_0$ . The former suggests building an estimator of  $\psi_0$  using estimators of  $\bar{Q}_0$  and of  $Q_{W,0}$ . The latter suggests building an estimator of  $\psi_0$  using estimators of  $\ell \bar{Q}_0$  and of  $\ell \bar{Q$ 



1. Show that for  $a' = 0, 1, \gamma_{0,a',c}$  as defined in (3) can be equivalently expressed as

$$\inf \left\{ z \in ]0,1[: \int \frac{\mathbf{1}\{a=a'\}}{\ell \bar{G}(a',W)} \mathbf{1}\{y \le z\} dP_0(w,a,y) \ge c \right\}.$$

2.5 Smooth parameters, first pass. Within our view of the target parameter as a statistical mapping, it is natural to inquire of properties this functional enjoys. For example, we may be interested in asking how the value of  $\Psi(P)$  changes as we consider laws that get nearer to P in  $\mathcal{M}$ . If small deviations from  $P_0$  result in large changes in  $\Psi(P_0)$ , then we might hypothesize that it will be difficult to produce stable estimators of  $\psi_0$ . Fortunately, this turns out not to be the case for the mapping  $\Psi$ , and so we say that  $\Psi$  is a smooth parameter mapping. We formalize this notion in Section 2.6, and here provide an informal description of smoothness.

To discuss how  $\Psi(P)$  changes for distributions near P in the model, we require a more concrete definition of nearness. To that end, consider the law encoded in run\_another\_experiment as a function of the input parameter h.

Let  $\Pi_h \in \mathcal{M}$  be the law encoded by run\_another\_experiment for a given  $h \in [-1,1]$ . Note that  $\mathcal{P} \equiv \{\Pi_h : h \in [-1,1]\}$  defines a collection of laws, that is, a statistical model. We say that  $\mathcal{P}$  is a submodel of  $\mathcal{M}$  because  $\mathcal{P} \subset \mathcal{M}$ . Moreover, we say that this submodel is through  $\Pi_0$  since  $\Pi_h \to \Pi_0$  as  $h \to 0$ . One could enumerate many possible submodels in  $\mathcal{M}$  through  $\Pi_0$ . It turns out that all that matters for our purposes is the form of the submodel in a neighborhood of  $\Pi_0$ . We informally say that this local behavior describes the direction of a submodel through  $\Pi_0$ . We formalize this notion in the next subsection.

We now have a notion of how to move through the model space  $P \in \mathcal{M}$  and can study how the value of the parameter changes as we move away from a law P. Above, we said that  $\Psi$  is a smooth parameter if it does not change "too much" as we move towards P in any particular direction. That is, we should hope that  $\Psi$  is differentiable along our submodel at P. This idea too is formalized in the next subsection, and we now turn to illustrating this idea numerically. The code below evaluates how the parameter changes for laws in  $\mathcal{P}$ , and approximates the derivative of the parameter along the submodel  $\mathcal{P}$  at  $\Pi_0$ .

```
approx <- seq(-1, 1, length.out = 1e2)
psi_Pi_h <- sapply(approx, function(t) {
   obs_from_another_experiment <- run_another_experiment(1, h = t)
   integrand <- function(w) {
      Qbar <- attr(obs_from_another_experiment, "Qbar")
      QW <- attr(obs_from_another_experiment, "QW")
      ( Qbar(cbind(1, w)) - Qbar(cbind(0, w)) ) * QW(w)
   }
   integrate(integrand, lower = 0, upper = 1)$val
})
slope_approx <- (psi_Pi_h - psi_Pi_zero) / approx
slope_approx <- slope_approx[min(which(approx > 0))]
ggplot() +
   geom_point(data = data.frame(x = approx, y = psi_Pi_h), aes(x, y),
```

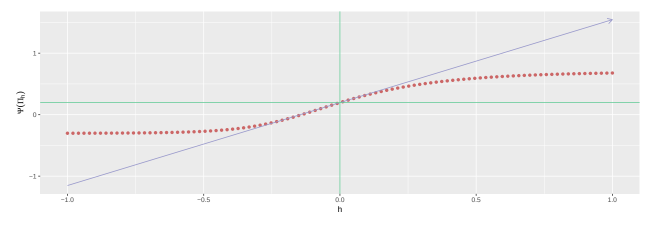


Figure 2: Evolution of statistical parameter  $\Psi$  along fluctuation  $\{\Pi_h : h \in H\}$ .

The dotted curve represents the function  $h \mapsto \Psi(\Pi_h)$ . The blue line represents the tangent to the previous curve at h = 0, which indeed appears to be differentiable around h = 0. In the next subsection, we derive a closed-form expression for the slope of the blue curve from the chunk of code where run\_another\_experiment is defined.

# Exercises

1. Adapt the code from problem 1 in Section ?? to visualize  $E_{\Pi_h}(Y \mid A=1, W)$ ,  $E_{\Pi_h}(Y \mid A=0, W)$ , and  $E_{\Pi_h}(Y \mid A=1, W) - E_{\Pi_h}(Y \mid A=0, W)$ , for  $h \in \{-1/2, 0, 1/2\}$ .

Define a new experiment with law  $\Pi'_0$  by adapting the code used to define run\_another\_experiment. Leave all aspects of the new experiment identical to  $\Pi_0$ , but set

- 2. Repeat the previous problem for this new experiment. Comment on the similarities and differences between  $\Pi_0$  and  $\Pi'_0$  for different values of h.
- 3. Re-produce Figure 1 for law  $\Pi'_0$ . Comment on the similarities and differences between this figure for  $\Pi_0$  and  $\Pi'_0$ . In particular, how does the behavior of the target parameter around h=0 compare between

laws  $\Pi_0$  and  $\Pi'_0$ ?

**2.6** Being smooth, second pass. Let us now formally define what it means for statistical mapping  $\Psi$  to be smooth at every  $P \in \mathcal{M}$ . For every  $h \in H \equiv ]-M^{-1}, M^{-1}[$ , we can define a law  $P_h \in \mathcal{M}$  by setting  $P_h \ll P^{\parallel}$  and

$$\frac{dP_h}{dP} \equiv 1 + hs,\tag{7}$$

where  $s: \mathcal{O} \to \mathbb{R}$  is a (measurable) function of O such that s(O) is not equal to zero P-almost surely,  $E_P(s(O)) = 0$ , and s bounded by M. We make the observation that

(i) 
$$P_h|_{h=0} = P$$
, (ii)  $\frac{d}{dh} \log \frac{dP_h}{dP}(O)\Big|_{h=0} = s(O)$ . (8)

Because of (i),  $\{P_h : h \in H\}$  is a submodel through P (also referred to as a fluctuation of P). As above note that the fluctuation is a one-dimensional submodel of  $\mathcal{M}$  with univariate parameter  $h \in H$ . We note that (ii) indicates that the score of this submodel at h = 0 is s. Thus, we say that the fluctuation is in the direction of s. Fluctuations of P do not necessarily take the same form as in (7). No matter how the fluctuation is built, for our purposes the most important feature of the fluctuation is its local shape in a neighborhood of P.

We are now prepared to provide a formal definition of smoothness of statistical mappings. We say that a statistical mapping  $\Psi$  is smooth at every  $P \in \mathcal{M}$  if for each  $P \in \mathcal{M}$ , there exists a (measurable) function  $D^*(P): \mathcal{O} \to \mathbb{R}$  such that  $E_P(D^*(P)(O)) = 0$ ,  $\operatorname{Var}_P(D^*(P)(O)) < \infty$ , and, for every fluctuation  $\{P_h : h \in H\}$  with score s at h = 0, the real-valued mapping  $h \mapsto \Psi(P_h)$  is differentiable at h = 0, with a derivative equal to

$$E_P(D^*(P)(O)s(O)). (9)$$

The object  $D^*(P)$  in (9) is called a gradient of  $\Psi$  at P.

This terminology has a direct parallel to directional derivatives in the calculus of Euclidean geometry. Recall that if f is a differentiable mapping from  $\mathbb{R}^p$  to  $\mathbb{R}$ , then the directional derivative of f at x (a point in  $\mathbb{R}^p$ ) in direction u (a unit vector in  $\mathbb{R}^p$ ) is the dot product of the gradient of f and u. In words, the directional derivative of f at x can be represented as an inner product of the direction that we approach x and the change of the function's value at x. In the present problem, the law P is the point at which we evaluate the function  $\Psi$ , the score x of the fluctuation is the "direction" in which we approach the point, and the gradient describes the change in the function's value at the point.

In general, it is possible for many gradients to exist\*\*. However, in the special case that the model is nonparametric, only a single gradient exists, which is sometimes referred to as the canonical gradient. In the more general setting, the canonical gradient may be defined as the minimizer of  $D \mapsto \operatorname{Var}_P(D(O))$  over the set of all gradients.

That is,  $P_h$  is dominated by P: if an event A satisfies P(A) = 0, then necessarily  $P_h(A) = 0$  too.

<sup>\*\*</sup>This may be at first surprising given the previous parallel drawn to Euclidean geometry. However, it is important to remember that the model dictates fluctuations of P that are valid submodels with respect to the full model. In turn, this determines the possible directions from which we may approach P. Thus, depending on the direction, (9) may hold with different choices of  $D^*$ .

**2.7** Revisiting Section 2.5. It is not difficult (though cumbersome) to verify that, up to a constant,  $\{\Pi_h : h \in [-1,1]\}$  is a fluctuation of  $\Pi_0$  in the direction (in the sense of (7)) of

$$\sigma_0(O) \equiv -10\sqrt{W}A \times \beta_0(A,W) \left(\log(1-Y) + \sum_{k=0}^3 \left(k + \beta_0(A,W)\right)^{-1}\right) + \text{constant},$$
 where  $\beta_0(A,W) \equiv \frac{1 - \bar{Q}_{\Pi_0}(A,W)}{\bar{Q}_{\Pi_0}(A,W)}$ .

Consequently, the slope of the dotted curve in Figure 2 is equal to

$$E_{\Pi_0}(D^*(\Pi_0)(O)\sigma_0(O)) \tag{10}$$

(since  $D^*(\Pi_0)$  is centered under  $\Pi_0$ , knowing  $\sigma_0$  up to a constant is not problematic).

In the following code, we check this numerically by implementing the direction  $\sigma_0$  with R function sigma0\_run\_another\_experiment, which we use to numerically approximate (10) (pointwise and with a confidence interval of asymptotic level 95%):

```
sigma0 run another experiment <- function(obs) {</pre>
  ## preliminary
  Qbar <- attr(obs, "Qbar")</pre>
  QAW <- Qbar(obs[, c("A", "W")])
  shape1 <- Arguments$getInteger(attr(obs, "shape1"), c(1, Inf))</pre>
  ## computations
  betaAW <- shape1 * (1 - QAW) / QAW
  out <- log(1 - obs[, "Y"])
  for (int in 1:shape1) {
    out <- out + 1/(int - 1 + betaAW)
  out <- - out * shape1 * (1 - QAW) / QAW * 10 * sqrt(obs[, "W"]) * obs[, "A"]
  ## no need to center given how we will use it
  return(out)
}
## DEBUGGING:
## 1) drawing 'obs_from_another_experiment' here (duplicated)
## 2) adding definition of 'eic' here (duplicated)
obs_from_another_experiment <- run_another_experiment(B)
eic <- function(obs, psi) {</pre>
  Qbar <- attr(obs, "Qbar")</pre>
  Gbar <- attr(obs, "Gbar")</pre>
  QAW <- Qbar(obs[, c("A", "W")])
  QoneW <- Qbar(cbind(A = 1, W = obs[, "W"]))
  QzeroW <- Qbar(cbind(A = 0, W = obs[, "W"]))</pre>
  GW <- Gbar(obs[, "W", drop = FALSE])</pre>
  1GAW \leftarrow obs[, "A"] * GW + (1 - obs[, "A"]) * (1 - GW)
  out <- (QoneW - QzeroW - psi) + (2 * obs[, "A"] - 1) / 1GAW * (obs[, "Y"] - QAW)
  out <- as.vector(out)</pre>
  return(out)
}
```

```
vars <- eic(obs_from_another_experiment, psi = 59/300) *
   sigma0_run_another_experiment(obs_from_another_experiment)
sd_hat <- sd(vars)
(slope_hat <- mean(vars))</pre>
```

## [1] 1.359158

```
(slope_CI <- slope_hat + c(-1, 1) * qnorm(1 - alpha / 2) * sd_hat / sqrt(B))
```

## [1] 1.353891 1.364425

Equal to 1.349 (rounded to three decimal places), the first numerical approximation slope\_approx is not too off.

2.8 Influence functions and the efficient influence function. If an estimator  $\psi_n$  of  $\Psi(P)$  can be written as

$$\psi_n = \Psi(P) + \frac{1}{n} \sum_{i=1}^n \text{IF}(O_i) + o_P(1/\sqrt{n})$$

for some function IF :  $\mathcal{O} \to \mathbb{R}$  such that  $E_P(\operatorname{IF}(O)) = 0$  and  $\operatorname{Var}_P(\operatorname{IF}(O)) < \infty$ , then we say that  $\psi_n$  is asymptotically linear with influence function IF. We note that weak convergence of asymptotically linear estimators is implied by the central limit theorem. That is, if  $\psi_n$  is asymptotically linear with influence function IF, then  $\sqrt(n)(\psi_n - \Psi(P)) = \frac{1}{\sqrt(n)} \sum_{i=1}^n \operatorname{IF}(O_i) + o_P(1)$ . The central limit theorem implies the the right-hand-side converges in law to a centered Gaussian distribution with variance  $\operatorname{Var}_P(\operatorname{IF}(O))$ .

As it happens, influence functions of regular<sup>††</sup> estimators are intimately related to gradients. In fact, if  $\psi_n$  is a regular, asymptotically linear estimator of  $\Psi(P)$  with influence function IF, then it must be true that  $\Psi$  is a smooth at P and that IF is a gradient of  $\Psi$  at P. Moreover, the converse is also true: given a gradient  $D^*$  of  $\Psi$  at P, under regularity conditions, it is possible to construct an estimator with influence function equal to  $D^*(P)$ .

These results, combined with the definition of the canonical gradient, imply that if  $\psi_n$  is a regular, asymptotically linear estimator of  $\Psi(P)$  built from n independent observations drawn from P, then the asymptotic variance of  $\sqrt{n}(\psi_n - \Psi(P))$  cannot be smaller than the variance of the canonical gradient of  $\Psi$  at P,

$$\operatorname{Var}_{P}(D^{*}(P)(O)). \tag{11}$$

That is, (11) is the lower bound on the asymptotic variance of any regular, asymptotically linear estimator of  $\Psi(P)$ . This bound is referred to as the *Cramér-Rao bound*. Any regular estimator that achieves this variance bound is said to be asymptotically efficient at P. Since, the canonical gradient is the influence function of an asymptotically efficient estimator, it is often referred to as the efficient influence function or the efficient influence curve.

<sup>&</sup>lt;sup>††</sup>We can view  $\psi_n$  as the by product of an algorithm  $\widehat{\Psi}$  trained on independent observations  $O_1,\ldots,O_n$  drawn from P. We say that the estimator is regular at P if, for any direction  $s \neq 0$  such that  $\mathrm{E}_P(s(O)) = 0$  and  $\mathrm{Var}_P(s(O)) < \infty$  and fluctuation  $\{P_h: h \in H\}$  satisfying (8), the estimator  $\psi_{n,1/\sqrt{n}}$  of  $\Psi(P_{1/\sqrt{n}})$  obtained by training  $\widehat{\Psi}$  on independent observations  $O_1,\ldots,O_n$  drawn from  $P_{1/\sqrt{n}}$  is such that  $\sqrt{n}(\psi_{n,1/\sqrt{n}} - \Psi(P_{1/\sqrt{n}}))$  converges in law to a limit that does not depend on s.

It is not difficult to check (do we give the proof?) that the efficient influence function of  $\Psi$  at  $P \in \mathcal{M}$  can be written as  $D^*(P) \equiv D_1^*(P) + D_2^*(P)$  where

$$D_1^*(P)(O) \equiv \bar{Q}(1, W) - \bar{Q}(0, W) - \Psi(P),$$
  
$$D_2^*(P)(O) \equiv \frac{2A - 1}{\ell \bar{G}(A, W)} (Y - \bar{Q}(A, W)).$$



- 1. Numerically approximate the Cramér-Rao bound at  $P_0$  and at  $\Pi_0$ . With a large sample and using regular estimators, can we more precisely estimate  $\Psi(P_0)$  or  $\Psi(\Pi_0)$ ?
- 2. Numerically approximate the Cramér-Rao bound at the law encoded for problems 2 and 3 of Section ??. Compare this bound with those computed in problem 1.
- **2.9 Double-robustness** The efficient influence curve  $D^*(P)$  at  $P \in \mathcal{M}$  enjoys another remarkable property: it is double-robust. Specifically, if we define for all  $P' \in \mathcal{M}$

$$\operatorname{Rem}_{P}(\bar{Q}', \bar{G}') \equiv \Psi(P') - \Psi(P) + \operatorname{E}_{P}(D^{*}(P')(O)),$$
 (12)

then the so called remainder term  $\operatorname{Rem}_{P}(\bar{Q}', \bar{G}')$  satisfies<sup>‡‡</sup>

$$\operatorname{Rem}_{P}(\bar{Q}', \bar{G}')^{2} \leq \|\bar{Q}' - \bar{Q}\|_{P}^{2} \times \|(\bar{G}' - \bar{G})/\ell\bar{G}'\|_{P}^{2}. \tag{13}$$

In particular, if

$$E_P(D^*(P')(O)) = 0, (14)$$

and either  $\bar{Q}' = \bar{Q}$  or  $\bar{G}' = \bar{G}$ , then  $\operatorname{Rem}_P(\bar{Q}', \bar{G}') = 0$  hence  $\Psi(P') = \Psi(P)$ . In words, if P' solves the so called P-specific efficient influence curve equation (14) and if, in addition, P' has the same  $\bar{Q}$ -component or  $\bar{G}$ -component as P, then  $\Psi(P') = \Psi(P)$  no matter how P' may differ from P otherwise. This property is useful to build consistent estimators of  $\Psi(P)$ .

However, there is much more to double-robustness than the above straightforward implication. Indeed, 12 is useful to build a consistent etimator of  $\Psi(P)$  that, in addition, satisfies a central limit theorem and thus lends itself to the construction of confidence intervals.

Let  $P_n^0 \in \mathcal{M}$  be an element of model  $\mathcal{M}$  of which the choice is data-driven, based on observing n independent draws from P. Equality 12 reveals that the statistical behavior of the corresponding substitution estimator  $\psi_n^0 \equiv \Psi(P_n^0)$  is easier to analyze when the remainder term  $\operatorname{Rem}_P(\bar{Q}_n^0, \bar{G}_n^0)$  goes to zero at a fast (relative to n) enough rate. In light of 13, this happens if the features  $\bar{Q}_n^0$  and  $\bar{G}_n^0$  of  $P_n^0$  converge to their counterparts under P at rates of which the product is fast enough.

<sup>&</sup>lt;sup>‡‡</sup>For any (measurable)  $f: \mathcal{O} \to \mathbb{R}$ , we denote  $||f||_P = \mathcal{E}_P(f(O)^2)^{1/2}$ .

**2.10** Inference assuming  $\bar{G}_0$  known, or not, first pass. Let  $O_1, \ldots, O_n$  be a sample of independent observations drawn from  $P_0$ . Let  $P_n$  be the corresponding empirical measure, *i.e.*, the law consisting in drawing one among  $O_1, \ldots, O_n$  with equal probabilities  $n^{-1}$ .

Let us assume for a moment that we know  $\bar{G}_0$ . This may be the case indeed if  $P_0$  was a controlled experiment. Note that, on the contrary, assuming  $\bar{Q}_0$  known would be difficult to justify.

```
## Debug -- couldn't find obs when I tried to compile
obs <- run_experiment(1e3)
Gbar <- attr(obs, "Gbar")
iter <- 1e3</pre>
```

Then, the alternative expression 5 suggests to estimate  $\psi_0$  with

$$\psi_n^b \equiv \mathcal{E}_{P_n} \left( \frac{2A - 1}{\ell \bar{G}_0(A, W)} Y \right) = \frac{1}{n} \sum_{i=1}^n \left( \frac{2A_i - 1}{\ell \bar{G}_0(A_i, W_i)} Y_i \right). \tag{15}$$

Note how  $P_n$  is substituted for  $P_0$  in (15) relative to (5).

It is easy to check that  $\psi_n^b$  estimates  $\psi_0$  consistently, but this is too little to request from an estimator of  $\psi_0$ . Better,  $\psi_n^b$  also satisfies a central limit theorem:  $\sqrt{n}(\psi_n^b - \psi_0)$  converges in law to a centered Gaussian law with asymptotic variance

$$v^b \equiv \operatorname{Var}_{P_0} \left( \frac{2A - 1}{\ell \bar{G}_0(A, W)} Y \right),$$

where  $v^b$  can be consistently estimated by its empirical counterpart

$$v_n^b \equiv \text{Var}_{P_n} \left( \frac{2A - 1}{\ell \bar{G}_0(A, W)} Y \right) = \frac{1}{n} \sum_{i=1}^n \left( \frac{2A_i - 1}{\ell \bar{G}_0(A_i, W_i)} Y_i - \psi_n^b \right)^2.$$
 (16)

Let us investigate how  $\psi_n^b$  behaves based on obs. Because we are interested in the law of  $\psi_n^b$ , the next chunk of code constitutes iter = 1000 independent samples of independent observations drawn from  $P_0$ , each consisting of n equal to nrow(obs)/iter = 1 data points, and computes the realization of  $\psi_n^b$  on all samples.

Before proceeding, let us introduce

$$\begin{split} \psi_n^a &\equiv \mathbf{E}_{P_n} \left( Y | A = 1 \right) - \mathbf{E}_{P_n} \left( Y | A = 0 \right) \\ &= \frac{1}{n_1} \sum_{i=1}^n \mathbf{1} \{ A_i = 1 \} Y_i - \frac{1}{n_0} \sum_{i=1}^n \mathbf{1} \{ A_i = 0 \} Y_i \\ &= \frac{1}{n_1} \sum_{i=1}^n A_i Y_i - \frac{1}{n_0} \sum_{i=1}^n (1 - A_i) Y_i, \end{split}$$

where  $n_1 = \sum_{i=1}^n A_i = n - n_0$  is the number of observations  $O_i$  such that  $A_i = 1$ . It is an estimator of

$$E_{P_0}(Y|A=1) - E_{P_0}(Y|A=0).$$

We seize this opportunity to demonstrate numerically the obvious fact that  $\psi_n^a$  does not estimate  $\psi_0$ .

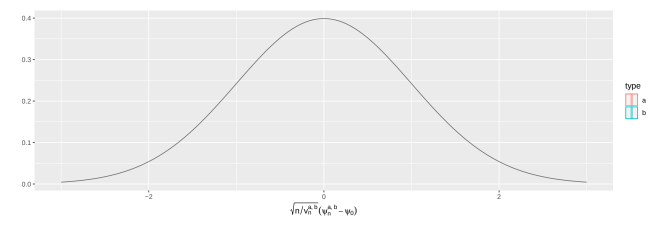


Figure 3: Kernel density estimators of the law of two estimators of  $\psi_0$  (recentered with respect to  $\psi_0$ , and renormalized), one of them misconceived (a), the other assuming that  $\bar{G}_0$  is known (b). Built based on iter independent realizations of each estimator.

```
std_b = sd(Y * (2 * A - 1) / 1GAW) / sqrt(n()),
            clt_b = (est_b - psi_approx) / std_b) %>%
  mutate(std_a = sd(est_a),
         clt_a = (est_a - psi_approx) / std_a) %>%
  gather("key", "value", -id) %>%
  extract(key, c("what", "type"), "([^_]+)_([ab])") %>%
  spread(what, value)
(bias_ab <- psi_hat_ab %>% group_by(type) %>% summarise(bias = mean(clt)))
## # A tibble: 2 x 2
##
            bias
     type
##
     <chr> <dbl>
## 1 a
             NaN
## 2 b
fig <- ggplot() +
  geom_line(aes(x = x, y = y),
            data = tibble(x = seq(-3, 3, length.out = 1e3),
                         y = dnorm(x)),
            linetype = 1, alpha = 0.5) +
  geom_density(aes(clt, fill = type, colour = type),
               psi_hat_ab, alpha = 0.1) +
  geom_vline(aes(xintercept = bias, colour = type),
             bias_ab, size = 1.5, alpha = 0.5)
fig +
  labs(y = "",
       x = \exp(\sqrt{n/v[n]^{\{list(a, b)\}})*(psi[n]^{\{list(a, b)\}}) - psi[0])))}
```

Let  $v_n^a$  be n times the empirical variance of the iter realizations of  $\psi_n^a$ . By the above chunk of code, the averages of  $\sqrt{n/v_n^a}(\psi_n^a-\psi_0)$  and  $\sqrt{n/v_n^b}(\psi_n^b-\psi_0)$  computed across the realizations of the two estimators are respectively equal to NaN and NA (both rounded to three decimal places — see bias\_ab). Interpreted as amounts of bias, those two quantities are represented by vertical lines in Figure 3. The red and blue bell-shaped curves represent the empirical laws of  $\psi_n^a$  and  $\psi_n^b$  (recentered with respect to

 $psi_0$ , and renormalized) as estimated by kernel density estimation. The latter is close to the black curve, which represents the standard normal density.

**2.11 Inference assuming**  $\bar{G}_0$  known, or not, second pass. At the beginning of Section 2.10, we assumed that  $\bar{G}_0$  was known. Let us suppose now that it is not. The definition of  $\psi_n^b$  can be adapted to overcome this difficulty, by substituting an estimator of  $\ell\bar{G}_0$  for  $\ell\bar{G}_0$  in (15).

For simplicity, we consider the case that  $\bar{G}_0$  is estimated by minimizing a loss function on a single working model, both fine-tune-parameter-free. By adopting this stance, we exclude estimating procedures that involve penalization (e.g. the LASSO) or aggregation of competing estimators (via stacking/super learning) – see Section ??. Defined in the next chunk of code, the generic function estimate\_G fits a user-specified working model by minimizing the empirical risk associated to the user-specified loss function and provided data.

Comment on new structure of estimate\_G and say a few words about compute\_lGhatAW.

```
estimate_G <- function(dat, algorithm, ...) {</pre>
  if (!is.data.frame(dat)) {
    dat <- as.data.frame(dat)</pre>
  }
  if (!attr(algorithm, "ML")) {
    fit <- algorithm[[1]](formula = algorithm[[2]], data = dat)</pre>
  } else {
    fit <- algorithm[[1]](dat, ...)</pre>
  fit$type_of_preds <- algorithm$type_of_preds
  return(fit)
}
compute_lGhatAW <- function(A, W, Ghat, threshold = 0.05) {</pre>
  dat \leftarrow data.frame(A = A, W = W)
  Ghat_W <- predict(Ghat, newdata = dat, type = Ghat$type_of_preds)</pre>
  IGAW <- A * Ghat_W + (1 - A) * (1 - Ghat_W)</pre>
  pred <- pmin(1 - threshold, pmax(lGAW, threshold))</pre>
  return(pred)
}
```

Note how the prediction of any  $\ell \bar{G}_0(A,W)$  is manually bounded away from 0 and 1 at the last but one line of compute\_lGhatAW. This is desirable because the *inverse* of each  $\ell \bar{G}_0(A_i,W_i)$  appears in the definition of  $\psi_n^b$  (15).

For sake of illustration, we choose argument working\_model\_G\_one of function estimate\_G as follows:

```
attr(working_model_G_one, "ML") <- FALSE
working_model_G_one$formula</pre>
```

```
## A ~ I(W^0.5) + I(abs(W - 5/12)^0.5) + I(W^1) + I(abs(W - 5/12)^1) +
## I(W^1.5) + I(abs(W - 5/12)^1.5)
```

In words, we choose the so called logistic (or negative binomial) loss function  $L_a$  given by

$$-L_a(f)(A, W) \equiv A \log f(W) + (1 - A) \log(1 - f(W))$$
(17)

for any function  $f:[0,1] \to [0,1]$  paired with the working model  $\mathcal{F} \equiv \{f_{\theta}: \theta \in \mathbb{R}^5\}$  where, for any  $\theta \in \mathbb{R}^5$ , logit  $f_{\theta}(W) \equiv \theta_0 + \sum_{j=1}^4 \theta_j W^{j/2}$ . The working model is well specified: it happens that  $\bar{G}_0$  is the unique minimizer of the risk entailed by  $L_a$  over  $\mathcal{F}$ :

$$\bar{G}_0 = \operatorname*{arg\,min}_{f_{\theta} \in \mathcal{F}} \mathrm{E}_{P_0} \left( L_a(f_{\theta})(A, W) \right).$$

Therefore, the estimator  $\bar{G}_n$  output by estimate\_G and obtained by minimizing the empirical risk

$$E_{P_n}(L_a(f_\theta)(A, W)) = \frac{1}{n} \sum_{i=1}^n L_a(f_\theta)(A_i, W_i)$$

over  $\mathcal{F}$  consistently estimates  $\bar{G}_0$ .

In light of (15), introduce

$$\psi_n^c \equiv \frac{1}{n} \sum_{i=1}^n \left( \frac{2A_i - 1}{\ell \bar{G}_n(A_i, W_i)} Y_i \right). \tag{18}$$

Because  $\bar{G}_n$  minimizes the empirical risk over a finite-dimensional and well-specified working model,  $\sqrt{n}(\psi_n^c - \psi_0)$  converges in law to a centered Gaussian law. Let us compute  $\psi_n^c$  on the same iter = 1000 independent samples of independent observations drawn from  $P_0$  as in Section 2.10:

```
if (redo fixed) {
  learned_features_fixed_sample_size <-</pre>
    obs %>% as tibble() %>%
   mutate(id = (seq_len(n()) - 1) %% iter) %>%
   nest(-id, .key = "obs") %>%
   mutate(Ghat = map(obs, ~ estimate_G(., algorithm = working_model_G_one))) %>%
    mutate(lGAW = map2(Ghat, obs, ~ compute_lGhatAW(.y$A, .y$W, .x)))
}
psi_hat_abc <-
  learned_features_fixed_sample_size %>%
  unnest(obs, 1GAW) %>%
  group_by(id) %>%
  summarize(est = mean(Y * (2 * A - 1) / 1GAW)) %>%
  mutate(std = sd(est),
         clt = (est - psi_approx) / std,
         type = "c") %>%
  full_join(psi_hat_ab)
## DEBUG : This was breaking when I compiled.
(bias_abc <- psi_hat_abc %>% group_by(type) %>% summarise(bias = mean(clt)))
```

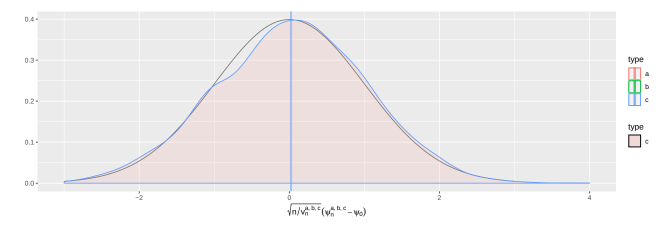


Figure 4: Kernel density estimators of the law of three estimators of  $\psi_0$  (recentered with respect to  $\psi_0$ , and renormalized), one of them misconceived (a), one assuming that  $\bar{G}_0$  is known (b) and one that hinges on the estimation of  $\bar{G}_0$  (c). The present figure includes Figure 3 (but the colors differ). Built based on iter independent realizations of each estimator.

```
## # A tibble: 3 x 2
## type bias
## <chr> <dbl>
## 1 a NaN
## 2 b NA
## 3 c 0.0201
```

Note how we exploit the independent realizations of  $\psi_n^c$  to estimate the asymptotic variance of the estimator with  $v_n^c/n$ . By the above chunk of code, the average of  $\sqrt{n/v_n^c}(\psi_n^c-\psi_0)$  computed across the realizations is equal to 0.02 (rounded to three decimal places — see bias\_abc). We represent the empirical laws of the recentered (with respect to

 $psi_0$ ) and renormalized  $\psi_n^a$ ,  $\psi_n^b$  and  $\psi_n^c$  in Figures 4 (kernel density estimators) and 5 (quantile-quantile plots).

Figures 4 and 5 reveal that  $\psi_n^c$  behaves as well as  $\psi_n^b$  — but remember that we did not discuss how to estimate its asymptotic variance.

# Exercises

geom\_qq(alpha = 1)

- 1. Compute a numerical approximation of  $E_{P_0}(Y|A=1) E_{P_0}(Y|A=0)$ . How accurate is it?
- 2. Building upon the piece of code devoted to the repeated computation of  $\psi_n^b$  and its companion quantities,

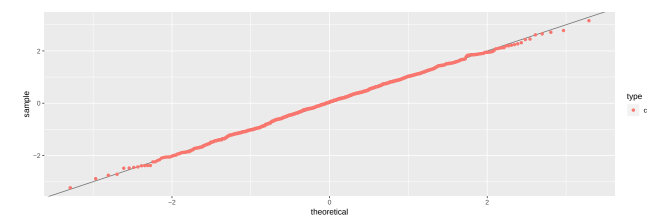


Figure 5: Quantile-quantile plot of the standard normal law against the empirical laws of three estimators of  $\psi_0$ , one of them misconceived (a), one assuming that  $\bar{G}_0$  is known (b) and one that hinges on the estimation of  $\bar{G}_0$  (c). Built based on iter independent realizations of each estimator.

construct confidence intervals for  $\psi_0$  of (asymptotic) level 95%, and check if the empirical coverage is satisfactory. Note that if the coverage was exactly 95%, then the number of confidence intervals that would contain  $\psi_0$  would follow a binomial law with parameters iter and 0.95, and recall that function binom.test performs an exact test of a simple null hypothesis about the probability of success in a Bernoulli experiment against its three one-sided and two-sided alternatives.

- 3. The call to compute\_1GhatAW makes predictions on the same data points as those exploited to learn  $\bar{G}_0$  by fitting the user-supplied working model. Why could that be problematic? Can you think of a simple workaround, implement and test it?
- 4. Discuss what happens when the dimension of the (still well-specified) working model grows. You could use the following chunk of code

play around with argument powers (making sure that 1/2 and 1 belong to it), and plot graphics similar to those presented in Figures 4 and 5.

5. Discuss what happens when the working model is mis-specified. You could use the following chunk of code:

```
transform <- c("cos", "sin", "sqrt", "log", "exp")
working_model_G_three <- list(</pre>
```

```
## A ~ I(cos(W)) + I(sin(W)) + I(sqrt(W)) + I(log(W)) + I(exp(W))
```

- 6. Prawing inspiration from (16), one may consider estimating the asymptotic variance of  $\psi_n^c$  with the counterpart of  $v_n^b$  obtained by substituting  $\ell \bar{G}_n$  for  $\ell \bar{G}_0$  in (16). By adapting the piece of code devoted to the repeated computation of  $\psi_n^b$  and its companion quantities, discuss if that would be legitimate.
- 2.12 Inference based on the estimation of  $\bar{Q}_0$ . Comment on structure of estimate\_Q, similar to that of estimate\_G.

Demonstrating the inference of  $\psi_0$  based on the estimation of  $\bar{Q}_0$  (and of the marginal law of W). Once based on a (mis-specified) working model, and once based on a non-parametric algorithm.

```
estimate_Q <- function(dat, algorithm, ...) {</pre>
  if (!is.data.frame(dat)) {
    dat <- as.data.frame(dat)</pre>
  if (!attr(algorithm, "ML")) {
    fit <- algorithm[[1]](formula = algorithm[[2]], data = dat)</pre>
  } else {
    fit <- algorithm[[1]](dat, ...)</pre>
  fit$type_of_preds <- algorithm$type_of_preds</pre>
  return(fit)
}
compute_QhatAW <- function(Y, A, W, Qhat, blip = FALSE) {</pre>
  if (!blip) {
    dat \leftarrow data.frame(Y = Y, A = A, W = W)
    pred <- predict(Qhat, newdata = dat, type = Qhat$type_of_preds)</pre>
  } else {
    pred <- predict(Qhat, newdata = data.frame(A = 1, W = W),</pre>
                     type = Qhat$type_of_preds) -
      predict(Qhat, newdata = data.frame(A = 0, W = W),
               type = Qhat$type_of_preds)
  }
  return(pred)
working_model_Q_one <- list(</pre>
  model = function(...) {trim_glm_fit(glm(family = binomial(), ...))},
 formula = as.formula(
```

```
paste("Y ~ A * (",
          paste("I(W^*), seq(1/2, 3/2, by = 1/2), sep = "", collapse = ") + "),
          "))")
  ),
  type_of_preds = "response"
attr(working_model_Q_one, "ML") <- FALSE</pre>
working model Q one $formula
## Y ~ A * (I(W^0.5) + I(W^1) + I(W^1.5))
## k-NN
kknn_algo <- list(
  algo = function(dat, ...) {
    args <- list(...)</pre>
    if ("Subsample" %in% names(args)) {
      keep <- sample.int(nrow(dat), args$Subsample)</pre>
      dat <- dat[keep, ]</pre>
    }
    fit <- caret::train(Y ~ I(10*A) + W, ## a tweak
                         data = dat,
                         method = "kknn",
                         verbose = FALSE,
    fit$finalModel$fitted.values <- NULL</pre>
    ## nms <- names(fit$finalModel$data)</pre>
    ## for (ii in match(setdiff(nms, ".outcome"), nms)) {
    ## fit$finalModel$data[[ii]] <- NULL</pre>
    ## }
    fit$trainingData <- NULL</pre>
    return(fit)
  },
  type_of_preds = "raw"
attr(kknn algo, "ML") <- TRUE</pre>
kknn_grid <- expand.grid(kmax = 5, distance = 2, kernel = "gaussian")
control <- trainControl(method = "cv", number = 2,</pre>
                         predictionBounds = c(0, 1),
                         trim = TRUE,
                         allowParallel = TRUE)
if(redo_fixed) {
  learned_features_fixed_sample_size <-</pre>
    learned_features_fixed_sample_size %>% # head(n = 100) %>%
    mutate(Qhat_d = map(obs, ~ estimate_Q(., algorithm = working_model_Q_one)),
           Qhat_e = map(obs, ~ estimate_Q(., algorithm = kknn_algo,
                                            trControl = control,
                                            tuneGrid = kknn_grid))) %>%
    mutate(blip_QW_d = map2(Qhat_d, obs,
                              compute_QhatAW(.y$Y, .y$A, .y$W, .x, blip = TRUE)),
           blip_QW_e = map2(Qhat_e, obs,
                              compute_QhatAW(.y$Y, .y$A, .y$W, .x, blip = TRUE)))
}
```

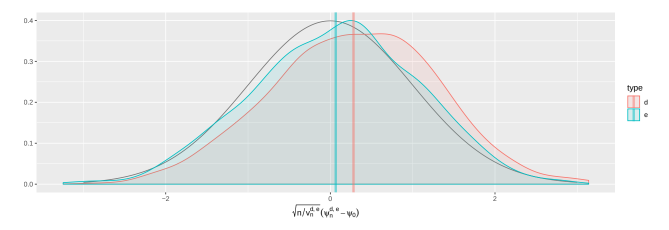


Figure 6: Write caption.

```
psi_hat_de <- learned_features_fixed_sample_size %>%
  unnest(blip_QW_d, blip_QW_e) %>%
  group_by(id) %>%
  summarize(est_d = mean(blip_QW_d),
           est_e = mean(blip_QW_e)) %>%
  mutate(std_d = sd(est_d),
        std_e = sd(est_e),
         clt_d = (est_d - psi_approx) / std_d,
         clt_e = (est_e - psi_approx) / std_e) %>%
  gather("key", "value", -id) %>%
  extract(key, c("what", "type"), "([^_]+)_([de])") %>%
  spread(what, value)
(bias_de <- psi_hat_de %>% group_by(type) %>% summarize(bias = mean(clt)))
## # A tibble: 2 x 2
     type
            bias
     <chr> <dbl>
          0.281
## 1 d
## 2 e
          0.0655
fig <- ggplot() +
  geom_line(aes(x = x, y = y),
            data = tibble(x = seq(-3, 3, length.out = 1e3),
                         y = dnorm(x)),
           linetype = 1, alpha = 0.5) +
  geom_density(aes(clt, fill = type, colour = type),
              psi_hat_de, alpha = 0.1) +
  geom_vline(aes(xintercept = bias, colour = type),
            bias_de, size = 1.5, alpha = 0.5)
fig +
 labs(y = "",
       x = \exp(\sqrt{n/v} [n]^{(i,j)} (psi[n]^{(i,j)} - psi[0])))
```

No that bad! Yet, we know that  $\sqrt{n}$  times bias is bound to increase with sample size. To see this, check out the next chunks of code.

```
sample_size <- c(4e3, 9e3)
block_size <- sum(sample_size)

label <- function(xx, sample_size = c(1e3, 2e3)) {
   by <- sum(sample_size)
        xx <- xx[seq_len((length(xx) %/% by) * by)] - 1
   prefix <- xx %/% by
   suffix <- findInterval(xx %% by, cumsum(sample_size))
   paste(prefix + 1, suffix + 1, sep = "_")
}

if (redo_varying) {
   learned_features_varying_sample_size <- obs %>% as.tibble %>%
        head(n = (nrow(.) %/% block_size) * block_size) %>%
        mutate(block = label(1:nrow(.), sample_size)) %>%
        nest(-block, .key = "obs")
}
```

First, we cut the data set into independent sub-data sets of sample size n in { 4000, 9000 }. Second, we infer  $\psi_0$  as shown two chunks earlier. We thus obtain 0 independent realizations of each estimator derived on data sets of 2, increasing sample sizes.

```
if(redo_varying) {
  learned_features_varying_sample_size <-</pre>
    learned_features_varying_sample_size %>%
    mutate(Qhat_d = map(obs, ~ estimate_Q(., algorithm = working_model_Q_one)),
           Qhat_e = map(obs, ~ estimate_Q(., algorithm = kknn_algo,
                                          trControl = control,
                                          tuneGrid = kknn_grid))) %>%
    mutate(blip QW d = map2(Qhat d, obs,
                            ~ compute_QhatAW(.y$Y, .y$A, .y$W, .x, blip = TRUE)),
           blip_QW_e = map2(Qhat_e, obs,
                            compute_QhatAW(.y$Y, .y$A, .y$W, .x, blip = TRUE)))
}
root_n_bias <- learned_features_varying_sample_size %>%
  unnest(blip QW d, blip QW e) %>%
  group_by(block) %>%
  summarize(clt_d = sqrt(n()) * (mean(blip_QW_d) - psi_approx),
            clt_e = sqrt(n()) * (mean(blip_QW_e) - psi_approx)) %>%
  gather("key", "value", -block) %>%
  extract(key, c("what", "type"), "([^_]+)_([de])") %>%
  spread(what, value) %>%
  mutate(block = unlist(map(strsplit(block, "_"), ~.x[2])),
         sample_size = sample_size[as.integer(block)])
```

The tibble called root\_n\_bias reports root-n times bias for all combinations of estimator and sample size. The next chunk of code presents visually our findings, see Figure 7. Note how we include the realizations of the estimators derived earlier and contained in psi\_hat\_de (thus breaking the independence between components of root\_n\_bias, a small price to pay in this context).

```
root_n_bias <- learned_features_fixed_sample_size %>%
  mutate(sample_size = B/iter) %>%  # because *fixed* sample size
  unnest(blip_QW_d, blip_QW_e) %>%
```

```
group_by(id) %>%
  summarize(clt_d = sqrt(n()) * (mean(blip_QW_d) - psi_approx),
            clt_e = sqrt(n()) * (mean(blip_QW_e) - psi_approx),
            sample_size = sample_size[1]) %>%
  gather("key", "clt", -id, -sample_size) %>%
  extract(key, c("what", "type"), "([^_]+)_([de])") %>%
  mutate(block = "0") %>% select(-id, -what) %>%
  full join(root n bias)
root_n_bias %>%
  ggplot() +
  stat_summary(aes(x = sample_size, y = clt,
                   group = interaction(sample_size, type),
                   color = type),
               fun.data = mean_se, fun.args = list(mult = 2),
               position = position_dodge(width = 250), cex = 1) +
  stat_summary(aes(x = sample_size, y = clt,
                   group = interaction(sample_size, type),
                   color = type),
               fun.data = mean_se, fun.args = list(mult = 2),
               position = position_dodge(width = 250), cex = 1,
               geom = "errorbar", width = 750) +
  stat_summary(aes(x = sample_size, y = clt,
                   color = type),
               fun.y = mean,
               position = position dodge(width = 250),
               geom = "polygon", fill = NA) +
  geom_point(aes(x = sample_size, y = clt,
                 group = interaction(sample_size, type),
                 color = type),
             position = position_dodge(width = 250),
             alpha = 0.1) +
  scale_x_continuous(breaks = unique(c(B / iter, sample_size))) +
  labs(x = "sample size n",
       y = expression(paste(sqrt(n) * (psi[n]^{list(d, e)} - psi[0]))))
## execute
## rm(learned_features_fixed_sample_size)
## as soon as possible!
```

2.13 One-step estimation. Function set\_Qbar\_Gbar implements the change of the Qbar and Gbar attributes of obs (which are accessible only by oracles).

```
set_Qbar_Gbar <- function(obs, Qhat, Ghat) {
  attr(obs, "Qbar") <- function(newdata) {
    if (!is.data.frame(newdata)) {
        newdata <- as.data.frame(newdata)
    }
    predict(Qhat, newdata = newdata, type = Qhat$type_of_preds)
}
attr(obs, "Gbar") <- function(newdata) {
    if (!is.data.frame(newdata)) {
        newdata <- as.data.frame(newdata)</pre>
```

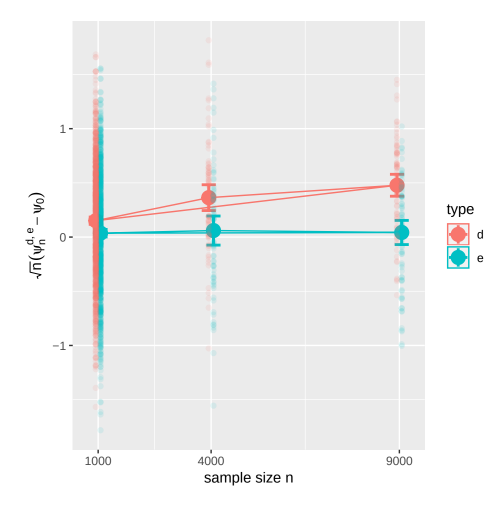


Figure 7: Evolution of root-n times bias versus sample size for two inference methodology of  $\psi_0$  based on the estimation of  $\bar{Q}_0$ . Big dots represent the average biases and vertical lines represent twice the standard error.

```
predict(Ghat, newdata = newdata, type = Ghat$type_of_preds)
  return(obs)
eic_hat <- function(obs, Qhat, Ghat, psi_hat) {</pre>
  Qbar <- function(newdata) {
    if (!is.data.frame(newdata)) {
      newdata <- as.data.frame(newdata)</pre>
    predict(Qhat, newdata = newdata, type = Qhat$type_of_preds)
  Gbar <- function(newdata) {</pre>
    if (!is.data.frame(newdata)) {
      newdata <- as.data.frame(newdata)</pre>
    }
    predict(Ghat, newdata = newdata, type = Ghat$type_of_preds)
  QAW <- Qbar(obs[, c("A", "W")])
  QoneW <- Qbar(cbind(A = 1, W = obs[, "W"]))</pre>
  QzeroW <- Qbar(cbind(A = 0, W = obs[, "W"]))</pre>
  GW <- Gbar(obs[, "W", drop = FALSE])</pre>
  1GAW \leftarrow obs[, "A"] * GW + (1 - obs[, "A"]) * (1 - GW)
  out <- (QoneW - QzeroW - psi_hat) + (2 * obs[, "A"] - 1) / 1GAW * (obs[, "Y"] - QAW)
  out <- out[[1]]
  return(out)
}
```

We first call function eic\_hat to compute the values of the estimated efficient influence at the observations in obs. Constructing the one-step estimators is then straightforward.

```
psi_hat_de_one_step <- learned_features_fixed_sample_size %>%
  mutate(est_d = map(blip_QW_d, mean),
         est_e = map(blip_QW_e, mean)) %>%
  mutate(eic_obs_d = pmap(list(obs, Qhat_d, Ghat, est_d),
                         eic_hat),
         eic_obs_e = pmap(list(obs, Qhat_e, Ghat, est_e),
                          unnest(blip_QW_d, eic_obs_d,
         blip_QW_e, eic_obs_e) %>%
  group_by(id) %>%
  summarize(est_d = mean(blip_QW_d) + mean(eic_obs_d),
           std_d = sd(eic_obs_d),
           clt_d = sqrt(n()) * (est_d - psi_approx) / std_d,
           est_e = mean(blip_QW_e) + mean(eic_obs_e),
           std_e = sd(eic_obs_e),
            clt_e = sqrt(n()) * (est_e - psi_approx) / std_e) %>%
  gather("key", "value", -id) %>%
  extract(key, c("what", "type"), "([^_]+)_([de])") %>%
  spread(what, value) %>%
  mutate(type = paste0(type, "_one_step"))
(bias_de_one_step <- psi_hat_de_one_step %>%
   group_by(type) %>% summarize(bias = mean(clt)))
```

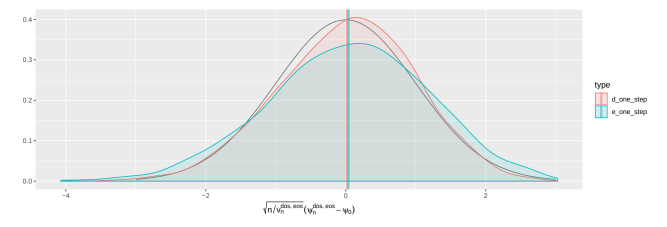


Figure 8: Write caption.

```
## # A tibble: 2 x 2
##
    type
                  bias
     <chr>
                 <dbl>
## 1 d_one_step 0.0189
## 2 e_one_step 0.0379
ggplot() +
  geom_line(aes(x = x, y = y),
            data = tibble(x = seq(-3, 3, length.out = 1e3),
                          y = dnorm(x)),
            linetype = 1, alpha = 0.5) +
  geom_density(aes(clt, fill = type, colour = type),
               psi_hat_de_one_step, alpha = 0.1) +
  geom_vline(aes(xintercept = bias, colour = type),
             bias_de_one_step, size = 1.5, alpha = 0.5) +
  labs(y = "",
       x = expression(
         paste(sqrt(n/v[n]^{list(dos, eos)}) * (psi[n]^{list(dos, eos)} - psi[0]))))
```

It seems that the one-step correction is quite good (in particular, compare bias\_de with bias\_de\_one\_step): bind\_rows(bias\_de, bias\_de\_one\_step)

```
## # A tibble: 4 x 2
## type bias
## <chr> <dbl>
## 1 d 0.281
## 2 e 0.0655
## 3 d_one_step 0.0189
## 4 e_one_step 0.0379
```

What about the estimation of the asymptotic variance, and of the mean-square errors of the estimators?

```
psi_hat_de %>%
full_join(psi_hat_de_one_step) %>% group_by(type) %>%
summarize(sd = mean(std * ifelse(str_detect(type, "one_step"), 1, NA),
```

```
se = sd(est) * sqrt(n()),
mse = mean((est - psi_approx)^2) * n()))
```

The sd (estimator of the asymptotic standard deviation) and se (empirical standard deviation) entries for type d\_one\_step are very similar: this indicates that the inference of the asymptotic variance of the d-variant of the one-step estimator based on the influence function is accurate. This comes as a surprise, since theory suggests that estimation should be conservative, that is, that the estimator should produce an upper bound to the actual asymptotic variance. On the contrary, the influence function-based estimator of the asymptotic variance is clearly very conservative for type e\_one-step. As for the mean square error, it is diminished by the one-step update for type d and enlarged for type e, the d\_one\_step estimator exhibiting the smallest mean square error.

#### 2.14 Targeted inference.

#### 2.15 Appendix. For later...

```
working_model_Q_two <- list(</pre>
  model = function(...) {trim_glm_fit(glm(family = binomial(), ...))},
  formula = as.formula(
    paste("Y ~ A * (",
          paste("I(W^{-}", seq(1/2, 3, by = 1/2), sep = "", collapse = ") + "),
  ),
  type_of_preds = "response"
attr(working_model_Q_two, "ML") <- FALSE</pre>
## xgboost based on trees
xgb_tree_algo <- list(</pre>
  algo = function(dat, ...) {
    caret::train(Y ~ I(10*A) + W,
                  data = dat,
                  method = "xgbTree",
                  trControl = control,
                  tuneGrid = grid,
                  verbose = FALSE)
  },
  type_of_preds = "response"
attr(xgb_tree_algo, "ML") <- TRUE</pre>
xgb_tree_grid <- expand.grid(nrounds = 350,</pre>
                               \max_{depth} = c(4, 6),
                               eta = c(0.05, 0.1),
```

```
gamma = 0.01,
                              colsample_bytree = 0.75,
                              subsample = 0.5,
                              min child weight = 0)
## nonparametric kernel smoothing regression
npreg <- list(</pre>
 label = "Kernel regression",
 type = "Regression",
 library = "np",
  parameters = data.frame(parameter =
                             c("subsample", "regtype",
                               "ckertype", "ckerorder"),
                           class = c("integer", "character",
                                      "character", "integer"),
                           label = c("#subsample", "regtype",
                                     "ckertype", "ckerorder")),
  grid = function(x, y, len = NULL, search = "grid") {
    if (!identical(search, "grid")) {
      stop("No random search implemented.\n")
    } else {
      out <- expand.grid(subsample = c(50, 100),
                          regtype = c("lc", "ll"),
                          ckertype =
                            c("gaussian",
                              "epanechnikov",
                              "uniform"),
                          ckerorder = seq(2, 8, 2))
    }
    return(out)
  },
  fit = function(x, y, wts, param, lev, last, classProbs, ...) {
    ny <- length(y)</pre>
    if (ny > param$subsample) {
      ## otherwise far too slow for what we intend to do here...
      keep <- sample.int(ny, param$subsample)</pre>
      x \leftarrow x[keep,]
      y <- y[keep]
    bw <- np::npregbw(xdat = as.data.frame(x), ydat = y,</pre>
                      regtype = param$regtype,
                       ckertype = param$ckertype,
                       ckerorder = param$ckerorder,
                       remin = FALSE, ftol = 0.01, tol = 0.01,
                       ...)
   np::npreg(bw)
  },
  predict = function (modelFit, newdata, preProc = NULL, submodels = NULL) {
    if (!is.data.frame(newdata)) {
      newdata <- as.data.frame(newdata)</pre>
    }
    np:::predict.npregression(modelFit, se.fit = FALSE, newdata)
  },
```

```
sort = function(x) {
    x[order(x$regtype, x$ckerorder), ]
 },
 loop = NULL, prob = NULL, levels = NULL
npreg_algo <- list(</pre>
 algo = function(dat, ...) {
    caret::train(working_model_Q_one$formula,
                 data = dat,
                 method = npreg, # no quotes!
                 verbose = FALSE,
 },
 type_of_preds = "response"
attr(npreg_algo, "ML") <- TRUE</pre>
npreg_grid <- data.frame(subsample = 100,</pre>
                          regtype = "lc",
                          ckertype = "gaussian",
                          ckerorder = 4,
                          stringsAsFactors = FALSE)
```