Inserm Workshop 282 - practical session Mediation analyses with the ltmle and medoutcon packages

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Chapter 1

Introduction



Figure 1.1: QR code towards the github repository of the workshop

We will use the following data set, you can download the data and import it in R.

For example, you can create a R project folder for this practical session, add a "data" folder and copy-paste the "df.csv" file in the data folder.

```
df <- read.csv2("data/df.csv")</pre>
```

1.1 Data generating system

The data generating mechanism is defined by the following set of structural equations, where:

- baseline confounders are sex $(L(0)_{sex}, 0 \text{ for women, 1 for men})$ and low parental education level $(L(0)_{low.par.edu}, 0 \text{ for no, 1 for yes})$,
- the exposure of interest is the individual's educational level $(A_{edu}, 0 \text{ for high}, 1 \text{ for low}),$
- 2 intermediate confounders affected by the exposure: physical activity $(L(1)_{phys}, 0 \text{ for no, } 1 \text{ for yes})$ and occupation $(L(1)_{occupation}, 0 \text{ for non-manual, } 1 \text{ for manual}),$
- the mediator of interest is smoking $(M_{smoking}, 0 \text{ for no, } 1 \text{ for yes}),$
- the outcome Y can be death (0 for no, 1 for yes) or a continuous functional score (higher values correspond to higher function).

```
\begin{array}{lll} L(0)_{sex} & = & f\left(U_{sex}\right) \\ L(0)_{low.par.edu} & = & f\left(L(0)_{sex}, U_{low.par.edu}\right) \\ A_{edu} & = & f\left(L(0)_{sex}, L(0)_{low.par.edu}, U_{edu}\right) \\ L(1)_{phys} & = & f\left(L(0)_{sex}, L(0)_{low.par.edu}, A_{edu}, U_{L(1)_{phys}}\right) \\ L(1)_{occupation} & = & f\left(L(0)_{sex}, L(0)_{low.par.edu}, A_{edu}, L(1)_{phys}, U_{L(1)_{occupation}}\right) \\ M_{smoking} & = & f\left(L(0)_{sex}, L(0)_{low.par.edu}, A_{edu}, L(1)_{phys}, L(1)_{occupation}, U_{M_{smoking}}\right) \\ Y & = & f\left(L(0)_{sex}, L(0)_{low.par.edu}, A_{edu}, L(1)_{phys}, L(1)_{occupation}, M_{smoking}, U_{Y}\right) \end{array}
```

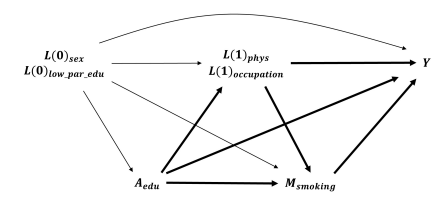


Figure 1.2: Causal model 1

Data were simulated using simple logistic and linear models.

Note that an "exposure - mediator" interaction term is included in the equations to simulate the binary and quantitative outcomes. This will have to be considered during the estimations (in case of exposure-mediator interaction, the results will depend on the choice of estimands: pure or total indirect and direct effects).

```
edu1 <- rep(1, N)
### intermediate counfounders L1
# physical activity: 1 = yes ; 0 = no
phys <- rexpit(qlogis(0.6) + log(1.5) * sex + log(0.8) * low_par_edu +
                 log(0.7) * edu
phys0 \leftarrow rexpit(qlogis(0.6) + log(1.5) * sex + log(0.8) * low par edu +
                  log(0.7) * edu0)
phys1 <- rexpit(qlogis(0.6) + log(1.5) * sex + log(0.8) * low_par_edu +
                  log(0.7) * edu1)
# occupation: 1 = manual; 0 = non-manual
occupation \leftarrow rexpit(qlogis(0.5) + log(1.3) * sex + log(1.2) * low_par_edu +
                 log(2.5) * edu + log(2) * phys)
occupation0 <- rexpit(qlogis(0.5) + log(1.3) * sex + log(1.2) * low_par_edu +
                         log(2.5) * edu0 + log(2) * phys0)
occupation1 <- rexpit(qlogis(0.5) + log(1.3) * sex + log(1.2) * low_par_edu +
                         log(2.5) * edu1 + log(2) * phys1)
### mediator
smoking \leftarrow rexpit(qlogis(0.3) + log(1.8) * sex + log(1.5) * low_par_edu +
                     log(2) * edu + log(0.7) * phys + log(1.8) * occupation)
smoking0 <- rep(0, N)</pre>
smoking1 <- rep(1, N)</pre>
smoking\_tot0 \leftarrow rexpit(qlogis(0.3) + log(1.8) * sex + log(1.5) * low_par_edu +
                          log(2) * edu0 + log(0.7) * phys0 + log(1.8) * occupation0)
smoking\_tot1 \leftarrow rexpit(qlogis(0.3) + log(1.8) * sex + log(1.5) * low_par_edu +
                          log(2) * edu1 + log(0.7) * phys1 + log(1.8) * occupation1)
### outcomes
death \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                  log(1.7) * edu + log(0.8) * phys + log(1.5) * occupation +
                  log(2.5) * smoking + log(1.5) * edu * smoking * inter)
death00 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                     log(1.7) * edu0 + log(0.8) * phys0 + log(1.5) * occupation0 +
                     log(2.5) * smoking0 + log(1.5) * edu0 * smoking0 * inter)
death01 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                     log(1.7) * edu0 + log(0.8) * phys0 + log(1.5) * occupation0 +
                     log(2.5) * smoking1 + log(1.5) * edu0 * smoking1 * inter)
death10 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low par edu +
                     log(1.7) * edu1 + log(0.8) * phys1 + log(1.5) * occupation1 +
                     log(2.5) * smoking0 + log(1.5) * edu1 * smoking0 * inter)
death11 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                     log(1.7) * edu1 + log(0.8) * phys1 + log(1.5) * occupation1 +
                     log(2.5) * smoking1 + log(1.5) * edu1 * smoking1 * inter)
death_tot0 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                        log(1.7) * edu0 + log(0.8) * phys0 + log(1.5) * occupation0 +
                        log(2.5) * smoking_tot0 + log(1.5) * edu0 * smoking_tot0 * inter)
death_tot1 \leftarrow rexpit(qlogis(0.05) + log(1.5) * sex + log(1.6) * low_par_edu +
                        log(1.7) * edu1 + log(0.8) * phys1 + log(1.5) * occupation1 +
                        log(2.5) * smoking_tot1 + log(1.5) * edu1 * smoking_tot1 * inter)
score \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                 -10 * edu + 8 * phys -7 * occupation +
                 -15 * smoking + -8 * edu * smoking * inter,
```

```
sd = 15
score00 \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                   -10 * edu0 + 8 * phys0 -7 * occupation0 +
                   -15 * smoking0 + -8 * edu0 * smoking0 * inter,
                 sd = 15)
score01 \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                   -10 * edu0 + 8 * phys0 -7 * occupation0 +
                   -15 * smoking1 + -8 * edu0 * smoking1 * inter,
                 sd = 15)
score10 \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                   -10 * edu1 + 8 * phys1 - 7 * occupation1 +
                   -15 * smoking0 + -8 * edu1 * smoking0 * inter,
                 sd = 15)
score11 \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                   -10 * edu1 + 8 * phys1 - 7 * occupation1 +
                   -15 * smoking1 + -8 * edu1 * smoking1 * inter,
                 sd = 15
score_{tot0} \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                      -10 * edu0 + 8 * phys0 -7 * occupation0 +
                      -15 * smoking_tot0 + -8 * edu0 * smoking_tot0 * inter,
                    sd = 15)
score_{tot1} \leftarrow rnorm(N, mean = 50 + 5 * sex -5 * low_par_edu +
                      -10 * edu1 + 8 * phys1 - 7 * occupation1 +
                      -15 * smoking tot1 + -8 * edu1 * smoking tot1 * inter,
                    sd = 15)
if (psi == FALSE) {
 return(data.sim = data.frame(subjid = 1:N,
                                sex = sex,
                                low_par_edu = low_par_edu,
                                edu = edu,
                                phys = phys,
                                occupation = occupation,
                                smoking = smoking,
                                death = death,
                                score = score))
} else {
  return(Psi = list(EY00_death = mean(death00),
                    EY01_death = mean(death01),
                    EY10 death = mean(death10),
                    EY11_death = mean(death11),
                    EYO death = mean(death tot0),
                    EY1_death = mean(death_tot1),
                    ATE_death = mean(death_tot1) - mean(death_tot0),
                    CDEO_death = mean(death10) - mean(death00),
                    CDE1_death = mean(death11) - mean(death01),
                    EY00_score = mean(score00),
                    EY01_score = mean(score01),
                    EY10_score = mean(score10),
                    EY11_score = mean(score11),
                    EYO_score = mean(score_tot0),
                    EY1_score = mean(score_tot1),
                    ATE_score = mean(score_tot1) - mean(score_tot0),
                    CDE0_score = mean(score10) - mean(score00),
                    CDE1_score = mean(score11) - mean(score01)))
```

```
## Simulate the data.frame df
set.seed(1234)
df <- GenerateData.CDE(N = 10000, inter = rep(1, 10000), psi = FALSE)
summary(df)
                                 low_par_edu
##
      subjid
                      sex
                                                    edu
               Min. :0.0000
##
  Min. : 1
                               Min. :0.0000
                                                      :0.0000
                                               Min.
  1st Qu.: 2501 1st Qu.:0.0000 1st Qu.:0.0000 1st Qu.:0.0000
## Median : 5000 Median :0.0000
                               Median :1.0000
                                               Median :1.0000
## Mean : 5000 Mean :0.4488
                               Mean :0.7329 Mean :0.5953
## 3rd Qu.: 7500
                 3rd Qu.:1.0000 3rd Qu.:1.0000 3rd Qu.:1.0000
##
   Max. :10000
                 Max. :1.0000 Max. :1.0000 Max. :1.0000
##
       phys
                  occupation
                                smoking
                                                   death
##
  Min. :0.0000 Min. :0.000 Min. :0.0000 Min.
                                                     :0.0000
  ## Median: 1.0000 Median: 1.000 Median: 1.0000 Median: 0.0000
## Mean :0.5579 Mean :0.744 Mean :0.5842
                                               Mean :0.2561
## 3rd Qu.:1.0000 3rd Qu.:1.000 3rd Qu.:1.0000 3rd Qu.:1.0000
## Max. :1.0000
                 Max. :1.000 Max. :1.0000 Max. :1.0000
##
      score
## Min. :-49.40
## 1st Qu.: 15.18
## Median: 30.37
## Mean : 30.48
## 3rd Qu.: 45.67
## Max. : 97.00
## Calculate the "true" estimands for:
## - the Average total effect: ATE = E(Y_0) - E(Y_1)
## - the controlled direct effect CDE(M=m), setting the mediator to M=m
     CDE(M=0) = E(Y_10) - E(Y_00)
     CDE(M=1) = E(Y_11) - E(Y_01)
set.seed(1234)
true <- GenerateData.CDE(N = 10000, inter = rep(1, 1e6), psi = TRUE) # CHANGE N TO 1e6 +++++++++
true
## $EY00 death
## [1] 0.096781
## $EY01 death
## [1] 0.209633
##
## $EY10_death
## [1] 0.163479
##
## $EY11_death
## [1] 0.416236
##
## $EYO_death
## [1] 0.153527
##
```

```
## $EY1_death
## [1] 0.332711
##
## $ATE_death
## [1] 0.179184
## $CDEO_death
## [1] 0.066698
##
## $CDE1_death
## [1] 0.206603
##
## $EY00_score
## [1] 48.78223
##
## $EY01_score
## [1] 33.57381
## $EY10_score
## [1] 36.85815
##
## $EY11_score
## [1] 13.95586
##
## $EYO_score
## [1] 41.77577
## $EY1_score
## [1] 21.84661
##
## $ATE_score
## [1] -19.92916
##
## $CDEO_score
## [1] -11.92408
##
## $CDE1_score
```

[1] -19.61794

Chapter 2

Reminders - Estimate the ATE

2.1 Estimation of the Average Total Effect (ATE) by g-computation

G-computation can be used for the estimation of the total effect and two-way decomposition (CDE, randomized direct and indirect effects). Analogs of the 3-way and 4-way decompositions are also given by the CMAverse package.

The following steps describe the implementation of the g-computation estimator of the average total effect ATE = $\mathbb{E}(Y_{A=1}) - \mathbb{E}(Y_{A=0})$.

0. The g-formula for the ATE is:

$$\begin{split} \Psi^{\text{ATE}} &= \sum_{l(0) \in L(0)} \mathbb{E}\left(Y \mid A = 1, L(0) = l(0)\right) P(L(0) = l(0)) - \sum_{l(0) \in L(0)} \mathbb{E}\left(Y \mid A = 0, L(0) = l(0)\right) P(L(0) = l(0)) \\ \Psi^{\text{ATE}} &= \sum_{l(0) \in L(0)} \overline{Q}(A = 0) \times P(L(0) = l(0)) - \sum_{l(0) \in L(0)} \overline{Q}(A = 1) \times P(L(0) = l(0)) \end{split}$$

- 1. Fit a logistic or a linear regression to estimate $\overline{Q} = \mathbb{E}(Y \mid A, L(0))$
- 2. Use this estimate to predict an outcome for each subject $\widehat{\overline{Q}}(A=0)_i$ and $\widehat{\overline{Q}}(A=1)_i$, by evaluating the regression fit \overline{Q} at A=0 and A=1 respectively
- 3. Plug the predicted outcomes in the g-formula and use the sample mean to estimate Ψ_{ATE} .

We can estimate its components and plug them directly in the g-formula:

$$\hat{\Psi}_{\text{gcomp}}^{\text{ATE}} = \frac{1}{n} \sum_{i=1}^{n} \left[\hat{\overline{Q}}(A=1)_i - \hat{\overline{Q}}(A=0)_i \right]$$
 (2.1)

For continuous outcomes, $\overline{Q}(A=a)$ functions can be estimated using linear regressions. For binary outcomes, they can be estimated using logistic regressions.

```
## 2. Predict an outcome for each subject, setting A=0 and A=1
# prepare data sets used to predict the outcome under the counterfactual
# scenarios setting A=O and A=1
data_Ais1 <- data_Ais0 <- df</pre>
data_Ais1$edu <- 1
data_AisO$edu <- 0
# predict values
Y1_death_pred <- predict(Q_tot_death, newdata = data_Ais1, type = "response")
YO_death_pred <- predict(Q_tot_death, newdata = data_AisO, type = "response")
Y1_score_pred <- predict(Q_tot_score, newdata = data_Ais1, type = "response")
YO_score_pred <- predict(Q_tot_score, newdata = data_AisO, type = "response")
## 3. Plug the predicted outcome in the gformula and use the sample mean
     to estimate the ATE
ATE_death_gcomp <- mean(Y1_death_pred) - mean(Y0_death_pred)
ATE_death_gcomp
## [1] 0.1867172
ATE_score_gcomp <- mean(Y1_score_pred) - mean(Y0_score_pred)
ATE_score_gcomp
## [1] -19.75274
```

A 95% confidence interval can be estimated applying a bootstrap procedure. An example is given in the following code.

```
## set seed for reproducibility
set.seed(1234)
B <- 10 # use a large number here (at least 200 to 1000)
## we will store estimates from each bootstrap sample in a data.frame:
bootstrap_estimates <- data.frame(matrix(NA, nrow = B, ncol = 2))</pre>
colnames(bootstrap_estimates) <- c("boot_death_est", "boot_score_est")</pre>
for (b in 1:B){
  # sample the indices 1 to n with replacement
  bootIndices <- sample(1:nrow(df), replace=T)</pre>
  bootData <- df[bootIndices,]</pre>
  if (round(b/100, 0) == b/100 ) print(paste0("bootstrap number ",b))
  Q_tot_death <- glm(death ~ edu + sex + low_par_edu,
                   family = "binomial", data = bootData)
  Q_tot_score <- glm(score ~ edu + sex + low_par_edu,
                   family = "gaussian", data = bootData)
  boot_Ais1 <- boot_Ais0 <- bootData</pre>
  boot_Ais1$edu <- 1
  boot_AisO$edu <- 0
  Y1_death_boot <- predict(Q_tot_death, newdata = boot_Ais1, type = "response")
  YO_death_boot <- predict(Q_tot_death, newdata = boot_AisO, type = "response")
```

2.2 Estimation of the ATE by IPTW

2.2.1 "Classic" Horvitz Thompson estimator

If the average total effect (ATE) is identifiable, $\Psi_{ATE} = \mathbb{E}(Y_{A=1}) - \mathbb{E}(Y_{A=0})$ can be expressed using Inverse probability of treatment weighting (IPTW), denoting $\mathbb{P}(A=a\mid L(0))=g(A=a\mid L(0))$:

$$\Psi_{ATE} = \mathbb{E}\left(\frac{\mathbb{I}(A=1)}{g(A=1\mid L(0))}Y\right) - \mathbb{E}\left(\frac{\mathbb{I}(A=0)}{g(A=0\mid L(0))}Y\right) \tag{2.2}$$

The following steps describe the implementation of the IPTW estimator

- 1. Estimate the treatment mechanism $g(A = 1 \mid L(0))$
- 2. Predict each individual's probability of being exposed to her own exposure
- 3. Apply weights corresponding to the inverse of the predicted probability $w_i = \frac{1}{\hat{g}(A=a_i|L(0)_i)}$
- 4. Use the empirical mean of the weighted outcome Y: $\hat{\mathbb{E}}(Y_a) = \frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = a)}{\hat{a}(A = a, |L(0)|)} Y_i$

```
gA_L[df$edu == 0] <- pred_g0_L[df$edu == 0]

## 3. Apply weights corresponding to the inverse of the predicted probability
wt <- 1 / gA_L

## 4. Use the empirical mean of the weighted outcome
# point estimates:
IPTW_death <- mean(wt * as.numeric(df$edu == 1) * df$death) -
    mean(wt * as.numeric(df$edu == 0) * df$death)
IPTW_death</pre>
```

[1] 0.1865118

```
IPTW_score <- mean(wt * as.numeric(df$edu == 1) * df$score) -
   mean(wt * as.numeric(df$edu == 0) * df$score)
IPTW_score</pre>
```

```
## [1] -19.76854
```

The ATE estimates using IPTW for death probability and mean quality of life are respectively +18.65% and -19.77.

2.2.2 Stabilized IPTW for the ATE

If the average total effect (ATE) is identifiable, Ψ_{ATE} can be estimated using a stabilized IPTW estimator:

$$\widehat{\mathbb{E}}(Y_1) - \widehat{\mathbb{E}}(Y_0) = \frac{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 1) \widehat{g}^*(A_i = 1)}{\widehat{g}(A_i = 1|L(0)_i)} Y_i}{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 1) \widehat{g}^*(A_i = 1)}{\widehat{g}(A_i = 1|L(0)_i)}} - \frac{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 0) \widehat{g}^*(A_i = 0)}{\widehat{g}(A_i = 0|L(0)_i)} Y_i}{\frac{1}{n} \sum_{i=1}^n \frac{\mathbb{I}(A_i = 0) \widehat{g}^*(A_i = 0)}{\widehat{g}(A_i = 0|L(0)_i)}}$$
(2.3)

The estimation algorithm is the same as for IPTW, but taking into account any non-null function of A ($g^*(A_i = a)$) in the denominator of the weight in step 3, and applying the stabilized estimator in step 4.

[1] 0.1865687

```
## [1] -19.75943
```

The ATE estimates using stabilized IPTW for death probability and mean quality of life are respectively +18.66% and -19.76.

2.2.3 Using an MSM estimated by IPTW

We can also express the ATE using coefficients of a MSM. For a continuous or binary outcome, we can use the following MSM to summarize the relationship between the counterfactual outcome (Y_a) and the exposure(s) A:

$$\mathbb{E}(Y_a) = \alpha_0 + \alpha_A a \tag{2.4}$$

The Average Total Effect ATE = $\mathbb{E}(Y_{A=1}) - \mathbb{E}(Y_{A=0})$ can then be expressed using the coefficients of this MSM (2.4):

$$\text{ATE} := (\alpha_0 + \alpha_A \times 1) - (\alpha_0 + \alpha_A \times 0) = \alpha_A$$

In this example, the coefficient α_A corresponds to the ATE.

Such a model is not very useful for a binary exposure. It would be much more useful for higher-dimensional exposures, for example with a continuous exposure, where the relationship between all the possible continuous values of the exposure A=a and the corresponding outcomes Y_a is summarized (and arbitrarily simplified) by a single line and the slope coefficient α_A .

MSM coefficients can be easily estimated using an Inverse Probability of Treatment (IPTW) approach based on weighted regressions.

For example, in order to fit the MSM (2.4) described above, we can use a linear regression of the (observed) outcome Y on the exposure, weighted by individual weights w_i or sw_i :

$$\mathbb{E}(Y) = \alpha_0 + \alpha_A a \tag{2.5}$$

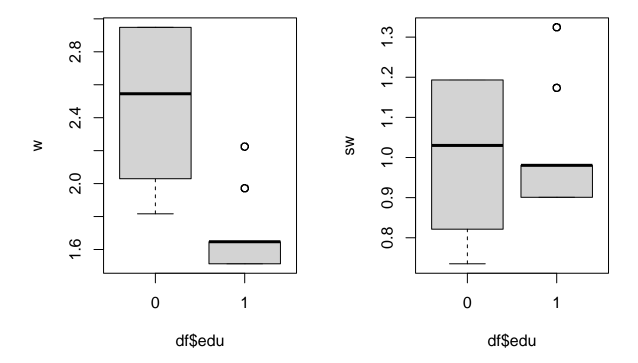
where
$$w_i=\frac{1}{P(A=a_i|L(0)=l(0)_i)}$$
 or $sw_i=\frac{P(A=a_i)}{P(A=a_i|L(0)=l(0)_i)}.$

The "no-unmeasured confounding" assumption is addressed by the application of weights w_i or sw_i , which balance confounders L(0) relative to the exposure A.

Below, we give an example where the parameters of an MSM are estimated using a weighted regression.

```
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
## 1. Denominator of the weight
# 1a. Estimate q(A=a_i|L(0)) (denominator of the weight)
g_A_L <- glm(edu ~ sex + low_par_edu,</pre>
              family = "binomial", data = df)
# 1b. Predict each individual's probability of being exposed to her own exposure
# predict the probabilities P(edu = 1) \& P(edu = 0)
pred_g1_L <- predict(g_A_L, type="response")</pre>
pred_g0_L <- 1 - pred_g1_L</pre>
# the predicted probability of the observed treatment P(A = a_i \mid L(0)) is :
gAi L <- rep(NA, nrow(df))
gAi_L[df$edu==1] <- pred_g1_L[df$edu==1]
gAi_L[df$edu==0] <- pred_g0_L[df$edu==0]</pre>
## 2. Numerator of the weight
# The numerator of the weight can be 1 for simple weights,
# or q(A=a_i/V) to obtain stabilized weights which put less weight to individuals
# with less observation. Stabilized weights enable a weaker positivity assumption.
# 2a. Estimate g(A=a_i) (numerator of the stabilized weight)
g_A \leftarrow glm(edu \sim 1,
           family = "binomial", data = df)
```

```
# 2b. Predict each individual's probability of being exposed to her own exposure
# predict the probabilities P(edu = 1) \& P(edu = 0)
pred_g1 <- predict(g_A, type="response")</pre>
pred_g0 <- 1 - pred_g1</pre>
# the predicted probability of the observed treatment P(A = a_i) is :
gAi <- rep(NA, nrow(df))
gAi[df$edu==1] <- pred_g1[df$edu==1]
gAi[df$edu==0] <- pred_g0[df$edu==0]
## 3. Define individual weights:
# We can use simple weights w = 1 / g(A=a_i / L(0))
w <- 1 / gAi_L
# Or alternatively, we can use stabilized weights :
# sw = g(A=a_i | sex) / g(A=a_i | L(0))
sw <- gAi / gAi_L
# we can see that stabilized weights have less extreme values
par(mfcol = c(1,2))
boxplot(w ~ df$edu)
boxplot(sw ~ df$edu)
```



```
par(mfcol = c(1,1))

## applying these weights creates a pseudo-population were the baseline
## confounders are balanced, relative to the exposure:
## before applying weights to the individuals:
prop.table(table(df$sex, df$edu, deparse.level = 2),
```

```
margin = 2)
##
         df$edu
## df$sex
                  0
##
        0 0.5245861 0.5692928
##
        1 0.4754139 0.4307072
prop.table(table(df$low_par_edu, df$edu, deparse.level = 2),
          margin = 2)
##
                 df$edu
## df$low_par_edu
                          0
##
                0 0.3397578 0.2177054
##
                1 0.6602422 0.7822946
## after applying weights to the individuals:
library(questionr) # The questionr package enables to describe weighted populations
prop.table(wtd.table(x = df$sex, y = df$edu, weights = w), margin = 2)
##
             0
## 0 0.5518912 0.5516449
## 1 0.4481088 0.4483551
prop.table(wtd.table(x = df$low_par_edu, y = df$edu, weights = w), margin = 2)
## 0 0.2668761 0.2669119
## 1 0.7331239 0.7330881
## 4. Estimate coefficients of the MSM using a weighted regression E(Y | A, sex)
 \hbox{\it\# a GLM with gaussian family can be applied to estimate risk differences }
# (for relative risk or rate ratios, we can apply a Poisson family;
# for OR, we can apply a binomial family)
msm1 <- glm(death ~ edu,</pre>
           weights = w,
           family = "gaussian",
           data = df
coef(msm1)
## (Intercept)
                       edu
## 0.1441017 0.1865687
msm2 <- glm(death ~ edu,
            weights = sw,
            family = "gaussian",
            data = df)
coef(msm2)
## (Intercept)
## 0.1441017 0.1865687
```

0.1865687

```
## 5. Estimate the ATE stratified by sex
# According to MSM1 (with simple weights)
coef(msm1)["edu"]

## edu
## 0.1865687

# According to the MSM2 (with stabilized weights)
coef(msm2)["edu"]
## edu
```

The ATE estimates of death probability using an MSM estimated by IPTW is +18.65%.

The results are the same with unstabilized or stabilized weights because there is no violation of the positivity assumption. In case of positivity violation, stabilized weights would give more accurate estimates.

95% confidence intervals can be calculated by bootstrap.

Chapter 3

Applying the CMAverse

If we use the CMAverse package with the df data set, we can estimate the Average Total Effect (ATE) and the Controlled direct effects (setting the mediator to 0) (CDE(M=0)).

$$\begin{split} ATE &= \mathbb{E}(Y_{A=1}) - \mathbb{E}(Y_{A=0}) \\ CDE(M=0) &= \mathbb{E}(Y_{A=1,M=0}) - \mathbb{E}(Y_{A=0,M=0}) \end{split}$$

In case of intermediate confounders affected by the exposure, the estimation based on regression coefficients cannot be used to estimate CDE and other direct and indirect effects. We need to use a g-computation, an IPTW estimator or a double-robust estimator.

3.1 Estimation by "parametric" g-computation

For example, we can estimate the two estimands ATE and CDE(M=0) on a risk difference scale as described below. In order to get estimates on the risk difference scale, the yreg argument needs to be set to "linear".

In order to estimate CDE(M=0) by parametric g-computation, we need:

- a model of the outcome (note that the exposure*mediator interaction is correctly included)
- a model of each of the intermediate confounder (2 models in our example).

Using those 3 models, we can simulated counterfactual values (under $\{A = 1, M = 0\}$ and $\{A = 0, M = 0\}$) exactly as what we did in the introduction chapter to simulate the data set df.

In the results, the model of the mediator is not needed for the CDE, but it is needed to estimate the interventional (stochastic) direct and indirect effects.

Note that for the model of the intermediate confounder occupation, physical activity (phys) was not included as a predictor whereas it was present in the corresponding equation of the data-generating system: Can this "misspecification" result in some bias? (I don't know the answer, it might be interesting to test on simulations).

|

```
summary(gformula_death_RD)
```

```
## Causal Mediation Analysis
##
## # Outcome regression:
##
## Call:
## glm(formula = death ~ edu + smoking + edu * smoking + sex + low_par_edu +
      phys + occupation, family = gaussian(), data = getCall(x$reg.output$yreg)$data,
##
      weights = getCall(x$reg.output$yreg)$weights)
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.0006107 0.0128884 -0.047 0.9622
## edu
              0.1122164 0.0130604 8.592 < 2e-16 ***
## smoking
## sex
               0.0633703 0.0084086
                                   7.536 5.25e-14 ***
## low_par_edu 0.0652140 0.0094112
                                    6.929 4.49e-12 ***
             -0.0168476 0.0084636 -1.991
## phys
                                            0.0466 *
## occupation 0.0413486 0.0097641
                                    4.235 2.31e-05 ***
## edu:smoking 0.1484815 0.0170875
                                   8.689 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 0.1671299)
##
##
      Null deviance: 1905.1 on 9999 degrees of freedom
## Residual deviance: 1670.0 on 9992 degrees of freedom
## AIC: 10499
## Number of Fisher Scoring iterations: 2
##
##
## # Mediator regressions:
##
## Call:
## glm(formula = smoking ~ edu + sex + low_par_edu + phys + occupation,
      family = binomial(), data = getCall(x$reg.output$mreg[[1L]])$data,
```

```
##
      weights = getCall(x$reg.output$mreg[[1L]])$weights)
##
## Coefficients:
##
             Estimate Std. Error z value Pr(>|z|)
0.70047
                        0.04402 15.911
                                         <2e-16 ***
## sex
              0.62499
                      0.04365 14.318
                                        <2e-16 ***
## low_par_edu 0.41552
                        0.04772
                                 8.707
                                         <2e-16 ***
                      0.04418 -8.761
            -0.38703
                                       <2e-16 ***
## phys
## occupation 0.59325
                      0.04946 11.993 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 13578 on 9999 degrees of freedom
## Residual deviance: 12685 on 9994 degrees of freedom
## AIC: 12697
##
## Number of Fisher Scoring iterations: 4
##
##
## # Regressions for mediator-outcome confounders affected by the exposure:
##
## Call:
## glm(formula = phys ~ edu + sex + low_par_edu, family = binomial(),
      data = getCall(x$reg.output$postcreg[[1L]])$data, weights = getCall(x$reg.output$postcreg[[1L]])$we
##
## Coefficients:
             Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 0.38815 0.04765 8.145 3.78e-16 ***
                      0.04205 -8.215 < 2e-16 ***
             -0.34545
                      0.04124 10.599 < 2e-16 ***
## sex
             0.43714
## low_par_edu -0.19185
                        0.04689 -4.092 4.28e-05 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 13729 on 9999 degrees of freedom
## Residual deviance: 13519 on 9996 degrees of freedom
## AIC: 13527
##
## Number of Fisher Scoring iterations: 4
##
##
##
## glm(formula = occupation ~ edu + sex + low_par_edu, family = binomial(),
      data = getCall(x$reg.output$postcreg[[2L]])$data, weights = getCall(x$reg.output$postcreg[[2L]])$we
##
##
## Coefficients:
             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.37498 0.05083 7.378 1.61e-13 ***
## edu
             0.85924 0.04734 18.150 < 2e-16 ***
## sex
```

```
## low_par_edu 0.09333
                         0.05217
                                 1.789
                                           0.0736 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 11377 on 9999 degrees of freedom
## Residual deviance: 10977 on 9996 degrees of freedom
  AIC: 10985
## Number of Fisher Scoring iterations: 4
##
##
## # Effect decomposition on the mean difference scale via the g-formula approach
##
## Direct counterfactual imputation estimation with
## bootstrap standard errors, percentile confidence intervals and p-values
##
                                     95% CIL 95% CIU P.val
##
                 Estimate Std.error
## cde
                0.0715898 0.0102149 0.0648080
                                              0.079 <2e-16 ***
## rpnde
                0.1415395 0.0124769 0.1284949
                                              0.145 <2e-16 ***
                0.1735670 0.0166478 0.1519051 0.174 <2e-16 ***
## rtnde
                0.0242051 0.0002929 0.0222740
                                              0.023 <2e-16 ***
## rpnie
                0.0562325 0.0038779 0.0460777 0.051 <2e-16 ***
## rtnie
## te
                0.0699497 0.0022620 0.0636869 0.067 <2e-16 ***
## rintref
## rintmed
                0.0320275 0.0041709 0.0234102
                                              0.029 <2e-16 ***
## cde(prop)
                0.3619816 0.0210889 0.3711416
                                              0.399 <2e-16 ***
## rintref(prop) 0.3536883 0.0188551 0.3395706
                                              0.365 <2e-16 ***
## rintmed(prop) 0.1619413 0.0100660 0.1340541
                                              0.148 <2e-16 ***
## rpnie(prop)
                0.1223887 0.0122998 0.1133772
                                              0.130 <2e-16 ***
## rpm
                0.2843301 0.0022338 0.2609549
                                              0.264 <2e-16 ***
## rint
                0.5156296 0.0087890 0.4871483
                                              0.499 <2e-16 ***
                0.6380184 0.0210889 0.6005255
                                              0.629 <2e-16 ***
## rpe
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (cde: controlled direct effect; rpnde: randomized analogue of pure natural direct effect; rtnde: random
##
## Relevant variable values:
## $a
## [1] 1
##
## $astar
## [1] 0
## $mval
## $mval[[1]]
## [1] 0
```

The ATE = 0.1978 and the CDE(M=0) = 0.0716.

3.2 Estimation by MSM estimated by IPTW

The CDE(M=0) can also by estimated using Marginal structural models (MSM) estimated by IPTW.

In order to get estimations by IPTW, the model argument needs to be set to msm.

The cmest function will estimate:

• a MSM for the outcome (depending only on the exposure and the mediator). Confounding is handled by weighting. This MSM can be used to estimate the controlled direct effect.

$$\begin{split} \mathbb{E}(Y_{A=a,M=m}) &= \beta_0 + \beta_{A_{educ}} \times a + \beta_{M_{smoking}} \times m + \beta_{A\star M} \times (a \times m) \\ CDE(M=m) &= \mathbb{E}(Y_{A=1,M=m}) - \mathbb{E}(Y_{A=0,M=m}) = \beta_{A_{educ}} + \beta_{A\star M} \times m \end{split}$$

• the weights $sw = sw_A \times sw_M$ is needed to handle confounding in the MSM of the outcome, where sw_A is a weight balancing parents of the exposure A and sw_M is a weight balancing parents of the mediator. To calculate those weights, we need to specify the numerator and denominator for the mediator's weight with the wmnomreg and wmdenomreg arguments. The denominator for the exposure's weight is specified with the ereg argument.

$$sw = sw_A \times sw_M = \frac{P(A_i)}{P(A_i \mid L(0))} \times \frac{P(M_i \mid A)}{P(M_i \mid L(0), A, L(1))}$$

An MSM of the mediator is also estimated (depending only on the exposure, confounding is handled by weighting). This MSM is not useful to estimate the CDE, but is needed to estimated the "interventional" or "stochastic" natural direct and indirect effects.

```
set.seed(1234)
iptw death RD <- cmest(data = df,
                       model = "msm", # using MSM estimated by IPTW
                       outcome = "death", # outcome variable
                       exposure = "edu", # exposure variable
                       mediator = "smoking", # mediator
                       basec = c("sex",
                                 "low_par_edu"), # baseline confounders
                       postc = c("phys",
                                 "occupation"), # intermediate confounders
                       EMint = TRUE, # exposures*mediator interaction
                       ereg = "logistic", # exposure regression model g(A=1/L(0))
                       mreg = list("logistic"), # q(M=1/L1,A,L0)
                       yreg = "linear",# Qbar.L2 = P(Y=1/M,L1,A,L0)
                       postcreg = list("logistic", "logistic"), # Qbar.L1 = P(L1=1/A,L0)
                       wmnomreg = list("logistic"), #g(M=1/A) wgt nominator
                       wmdenomreg = list("logistic"), # g(M=1/L1,A,L(0)) wgt denominator
                       astar = 0,
                       a = 1,
                       mval = list(0), # do(M=0) to estimate CDE_m
                       estimation = "imputation", # if model= gformula
                       inference = "bootstrap",
                       boot.ci.type = "per", # for percentile, other option: "bca"
                       nboot = 2) # we should use a large number of bootstrap samples
```

| | | |

```
summary(iptw_death_RD)
## Causal Mediation Analysis
##
## # Outcome regression:
##
## Call:
## glm(formula = death ~ edu + smoking + edu * smoking, family = gaussian(),
       data = getCall(x$reg.output$yreg)$data, weights = getCall(x$reg.output$yreg)$weights)
##
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                        0.008865
                                    9.355 < 2e-16 ***
## (Intercept) 0.082933
                                    5.514 3.59e-08 ***
## edu
              0.070447
                         0.012775
## smoking
              0.119740
                        0.012960
                                    9.239 < 2e-16 ***
## edu:smoking 0.142064
                         0.017184
                                    8.267 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 0.1690611)
##
##
      Null deviance: 1880.9 on 9999 degrees of freedom
## Residual deviance: 1689.9 on 9996 degrees of freedom
## AIC: 10969
## Number of Fisher Scoring iterations: 2
##
##
## # Mediator regressions:
##
## glm(formula = smoking ~ edu, family = binomial(), data = getCall(x$reg.output$mreg[[1L]])$data,
##
       weights = getCall(x$reg.output$mreg[[1L]])$weights)
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -0.11692
                          0.03149 -3.713 0.000205 ***
               0.78869
                          0.04174 18.895 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 13576 on 9999 degrees of freedom
## Residual deviance: 13214 on 9998 degrees of freedom
## AIC: 13197
##
## Number of Fisher Scoring iterations: 4
##
##
## # Mediator regressions for weighting (denominator):
##
## Call:
## glm(formula = smoking ~ edu + sex + low_par_edu + phys + occupation,
```

```
family = binomial(), data = getCall(x$reg.output$wmdenomreg[[1L]])$data,
##
##
      weights = getCall(x$reg.output$wmdenomreg[[1L]])$weights)
##
## Coefficients:
##
             Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.85996
                         0.06143 -13.998 <2e-16 ***
                         0.04402 15.911
              0.70047
                                          <2e-16 ***
              0.62499
                         0.04365 14.318
                                          <2e-16 ***
## low_par_edu 0.41552
                         0.04772
                                  8.707
                                          <2e-16 ***
## phys
            -0.38703
                         0.04418 -8.761
                                          <2e-16 ***
## occupation 0.59325
                         0.04946 11.993 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 13578 on 9999 degrees of freedom
## Residual deviance: 12685 on 9994 degrees of freedom
## AIC: 12697
## Number of Fisher Scoring iterations: 4
##
##
## # Mediator regressions for weighting (nominator):
##
## Call:
## glm(formula = smoking ~ edu, family = binomial(), data = getCall(x$reg.output$wmnomreg[[1L]])$data,
      weights = getCall(x$reg.output$wmnomreg[[1L]])$weights)
##
##
## Coefficients:
##
             Estimate Std. Error z value Pr(>|z|)
0.04178 19.493 < 2e-16 ***
## ed11
              0.81446
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 13578 on 9999 degrees of freedom
##
## Residual deviance: 13192 on 9998 degrees of freedom
## AIC: 13196
##
## Number of Fisher Scoring iterations: 4
##
##
## # Exposure regression for weighting:
##
  glm(formula = edu ~ sex + low_par_edu, family = binomial(), data = getCall(x$reg.output$ereg)$data,
##
      weights = getCall(x$reg.output$ereg)$weights)
##
## Coefficients:
             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.02910 0.04186 0.695
                                           0.487
             -0.23178
                       0.04154 -5.580 2.41e-08 ***
## low_par_edu 0.63793
                         0.04599 13.871 < 2e-16 ***
```

```
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
    Null deviance: 13497 on 9999 degrees of freedom
## Residual deviance: 13285 on 9997 degrees of freedom
## AIC: 13291
##
## Number of Fisher Scoring iterations: 4
##
##
## # Effect decomposition on the mean difference scale via the marginal structural model
## Direct counterfactual imputation estimation with
## bootstrap standard errors, percentile confidence intervals and p-values
##
##
           Estimate Std.error 95% CIL 95% CIU P.val
           0.070447 0.007889 0.067239 0.078 <2e-16 ***
## cde
## rpnde
           ## rtnde
           ## rpnie
           ## rtnie
## te
           ## rintref
## rintmed
           0.067440 0.002075 0.067636 0.070 <2e-16 ***
           ## cde(prop)
           ## rintref(prop) 0.360098 0.020701 0.345418 0.373 <2e-16 ***
## rintmed(prop) 0.143118 0.018617 0.136462 0.161 <2e-16 ***
## rpnie(prop)
           ## rpm
## rint
           ## rpe
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (cde: controlled direct effect; rpnde: randomized analogue of pure natural direct effect; rtnde: random
##
## Relevant variable values:
## $a
## [1] 1
##
## $astar
## [1] 0
##
## $mval
## $mval[[1]]
## [1] 0
```

Applying an IPTW estimator using the CMAverse, the ATE = 0.1873 and the CDE(M=0) = 0.0704.

Chapter 4

Estimate the ATE by TMLE

When estimating a mean counterfactual outcome using g-computation methods, we have to estimate some \bar{Q} functions (functions of the outcome conditional on the exposures and confounders, $\bar{Q} = \mathbb{E}(Y \mid A, L(0))$). For example, the Average Total Effect (ATE) is defined as a marginal effect, estimated using the empirical mean of such \bar{Q} functions:

$$\hat{\Psi}_{\mathrm{gcomp}}^{\mathrm{ATE}} = \frac{1}{n} \sum_{i=1}^{n} \left[\hat{\overline{Q}} (A=1)_i - \hat{\overline{Q}} (A=0)_i \right]$$

Unless the \bar{Q} functions are not misspecified, its estimate is expected to be biased (and \bar{Q} are expected to be misspecified, especially if the set of baseline confounders L(0) is high dimensional, for example if it includes is a large number of variables or continuous variables). In order to improve the estimation of $\bar{Q}(A,L)$, it is possible to use data-adaptive methods (machine learning algorithms) in order to optimize the bias-variance trade-off. However, this bias-variance trade-off would be optimized for the \bar{Q} functions, not for the ATE estimate $\hat{\Psi}_{\text{gcomp}}^{\text{ATE}}$. If the \bar{Q} function is unknown and has to be estimated (preferably by data-adaptive methods), it can be shown that the g-computation estimate of Ψ^{ATE} is asymptotically biased.

The Targeted Maximum Likelihood Estimation (TMLE) method has been developed as an asymptotically linear estimator, so that the estimation of any target parameter in any semiparametric statistical model is unbiased and efficient. In order to estimate a parameter $\Psi(P_0)$, where P_0 is an unknown probability distribution among a set \mathcal{M} of possible statistical models, the TMLE is described as a two-step procedure (Laan and Rose 2011):

- The first step is to obtain an initial estimate of the relevant part (\bar{Q}_0 in our applications) of the probability distribution P_0 . Data adaptive methods (machine learning algorithms) can be used to optimize this first step.
- The second step is to update the initial fit in order to "target toward making an optimal bias-variance tradeoff for the parameter of interest" $\Psi(\bar{Q})$.

Several R packages have been developed in order to carry out TMLE estimation of causal effects. We will begin using the ltmle package, as it can be used to estimate ATE or CDE. More generally, this package can be used to estimate the counterfactual effects of repeated exposure in time-to-event settings. In the setting of mediation analysis, a controlled direct effect (CDE) corresponds to a sequence of counterfactual interventions on 2 "exposure variables": the initial exposure A and the mediator of interest M. The package can also be used in simpler settings with only one binary or continuous outcome, measure only once at the end a the study.

4.1 TMLE for the ATE

In order to illustrate the TMLE procedure, the estimation of a mean counterfactual outcome, denoted $\Psi(A=1) = \mathbb{E}\left[\bar{Q}(A=1,L(0))\right]$, will be described in detail, following the algorithm implemented in the ltmle package.

The basic steps of the procedure are the following (Laan and Rose 2011):

- 1. Estimate \bar{Q}_0 . Data-adaptive methods can be used here, the ltmle package relies on the SuperLearner package to fit and predict $\hat{\bar{Q}}(A=1)$.
- 2. Estimate the treatment mechanism (propensity score) $g(A = 1 \mid L(0))$. Once again, data-adaptive methods can be used to improve the estimation.
- 3. The initial estimator of $\bar{Q}_0(A=1)$ will be slightly modified using a parametric fluctuation model, in order to reduce the bias when estimating the ATE. For example, the following parametric model of $\bar{Q}_0(A=1)$ and a "clever covariate" $H=\frac{I(A=1)}{\hat{g}(A=1|L(0))}$ can be applied:

$$logitP(Y \mid \hat{\bar{Q}}, H) = logit\bar{Q} + \varepsilon H$$

This parametric fluctuation model is chosen so that the derivative of its log-likelihood loss function is equal to the appropriate component of the efficient influence curve of the target parameter $\Psi(A=1)$.

- 4. Modify the initial estimator of $\bar{Q}_0(A=1)$ with the parametric fluctuation model (using the estimation $\hat{\varepsilon}$ from the previous step). We denote $\hat{\bar{Q}}^*(A=1)$ the updated value of $\hat{\bar{Q}}(A=1)$
- 5. Use the updated values $\hat{\bar{Q}}^*(A=1)$ in the substitution estimator to estimate the target parameter $\Psi(A=1)$:

$$\hat{\Psi}(A=1)_{\rm TMLE} = \frac{1}{n} \sum_{i=1}^{n} \hat{\bar{Q}}^{*}(A=1,L(0))$$

6. Estimate the efficient influence curve $D^*(Q_0, g_0)$:

$$D^*(Q_0,g_0) = \frac{I(A=1)}{g_0(A=1 \mid L(0))} (Y - \bar{Q}_0(A,L(O))) + \bar{Q}_0(A=1,L(0)) - \Psi(A=1)$$

The variance of the target parameter can then be calculated using the variance of the efficient influence curve:

$$\operatorname{var}\hat{\Psi}(A=1)_{\mathrm{TMLE}} = \frac{\operatorname{var}\hat{D}^*}{n}$$

1 2 3 4 5 6 ## 0.6608369 0.6071248 0.6071248 0.4495011 0.6071248 0.6071248

```
# It is useful to check the distribution of qA_L, as values close to 0 or 1 are
# indicators of near positivity violation and can result in large variance for the
# estimation.
# In case of near positivity violation, gA_{-}L values can be truncated to decrease
# the variance (at the cost a increased bias).
summary(g1_L)
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
## 0.4495 0.5073 0.6071 0.5953 0.6608 0.6608
# there is no positivity issues in this example.
## 3) Determine a parametric family of fluctuations of Qbar.
# The fluctuation model is a model of logitQbar and g(A=1/L(0))
# The clever covariate H(A,L(0)) depends on g(A=1/L(0)):
H \leftarrow (df \cdot = 1) / g1_L
# Update the initial fit Qbar from step 1.
# This is achieved by holding Qbar fixed (as intercept) while estimating the
# coefficient epsilon for H
# for example we could use the following fluctuation model (from the "Targeted
# Learning" book)
update_fit <- glm(df$death ~ -1 + offset(logitQ) + H,
                  family = "quasibinomial")
coef(update_fit)
## -1.658129e-05
Qstar <- predict(update_fit, data = data.frame(logitQ, H), type = "response")</pre>
# In the ltmle package, the fluctuation parametric model is slightly different
# (but with the same purpose). The "clever covariate" H is scaled and used as a
# weight in the parametric quasi-logistic regression
S1 <- rep(1, nrow(df))</pre>
update_fit_ltmle <- glm(df$death ~ -1 + S1 + offset(logitQ),
                        family = "quasibinomial",
                        weights = scale(H, center = FALSE))
coef(update_fit_ltmle)
##
             S1
## -2.80861e-05
## 4) Update the initial estimate of Qbar using the fluctuation parametric model
Qstar_ltmle <- predict(update_fit_ltmle,</pre>
                       data = data.frame(logitQ, H),
                       type = "response")
head(Qstar_ltmle)
##
                               3
                                                    5
```

0.3073922 0.4243074 0.4243074 0.3015693 0.4243074 0.4243074

```
1 2 3
# 0.2872412 0.3441344 0.3441344 0.2591356 0.3441344 0.3441344
## 5) Obtain the substition estimator of Psi Ais1
Psi_Ais1 <- mean(Qstar_ltmle)</pre>
Psi_Ais1
## [1] 0.3306626
## 5) Calculate standard errors based on the influence curve of the TMLE
IC <- H * (df$death - Qstar_ltmle) + Qstar_ltmle - Psi_Ais1</pre>
head(IC)
##
                                    3
                                                                        6
                        2
                                                4
                                                            5
## -0.48842639 0.09364473 0.09364473 1.52469745 0.09364473 1.04187255
# the influence curve has a mean of O
summary(IC)
##
      Min. 1st Qu.
                      Median
                                 Mean 3rd Qu.
                                                   Max.
## -0.69999 -0.48843 -0.02909 0.00000 0.09364 1.52470
# The standard error of the target parameter Psi(A=1) can be estimated by :
sqrt(var(IC)/nrow(df))
## [1] 0.006090826
```

We can see that we can get the same output using the ltmle package (cf. ?ltmle to see how the function works):

```
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
library(ltmle)
# The Qform and gform arguments are defined from the DAG
Qform <- c(death="Q.kplus1 ~ sex + low_par_edu + edu")
gform <- c("edu ~ sex + low_par_edu")</pre>
# in the ltmle package, the data set should be formated so that the order of the
# columns corresponds to the time-ordering of the model
data ltmle <- subset(df,</pre>
                      select = c(sex, low_par_edu, edu, death))
# the counterfactual intervention is defined in the abar argument
abar <- 1
Psi_Ais1 <- ltmle(data_ltmle,</pre>
                   Anodes = "edu",
                   Ynodes = "death",
                   Qform = Qform,
                   gform = gform,
                   gbounds = c(0.01, 1), # by default, g function truncated at 0.01
                   abar = abar,
```

In practice, it is recommended to apply data-adaptive algorithms to estimate Q and g functions: the ltmle package relies on the SuperLearner package. As indicated in the Guide to SuperLearner, The SuperLearner is "an algorithm that uses cross-validation to estimate the performance of multiple machine learning models, or the same model with different settings. It then creates an optimal weighted average of those models (ensemble learning) using the test data performance."

Here is an example for our estimation of the Average Total Effect (ATE).

The SuperLearner package includes a set of algorithms with default parameters (showed by listWrappers()). Because the simulated data set only have 2 binary baseline variables, the set \mathcal{M} of possible statistical models is limited. In order to estimate the ATE, we will include a library with:

- SL.mean, the null-model which only predict the marginal mean (it can be used as a reference for a bad model);
- SL.glm, a glm using the main terms from the Qform and gform argument;
- SL.interaction.back, a step-by-step backward GLM prodecure (based on the AIC), starting with all 2 × 2 interactions between main terms. This function is customized from the SL.step.interaction available with the ltmle and SuperLearner packages, where the direction argument is set to both by default.
- SL.hal9001 fit the "Highly Adaptive Lasso (HAL) algorithm. See the vignette for more information on the HAL algorithm. One of its advantage is a very fast rate of convergence.

```
library(SuperLearner)
library(hal9001)
# Below, we use the same ltmle() function than previously,
# and specify a family of algorithms to be used with the SuperLearner
## we can change the default argument of the SL.xgboost algorithm and the
## SL.step.interaction algorithm
# We can check how arguments are used in the pre-specified algorithms
SL.step.interaction
# function (Y, X, newX, family, direction = "both", trace = 0,
      k = 2, ....
#
# {
#
      fit.glm \leftarrow glm(Y \sim ., data = X, family = family)
#
      fit.step <- step(fit.glm, scope = Y ~ .~2, direction = direction,
          trace = trace, k = k)
```

```
#
      pred <- predict(fit.step, newdata = newX, type = "response")</pre>
#
      fit <- list(object = fit.step)</pre>
#
      out <- list(pred = pred, fit = fit)</pre>
#
      class(out$fit) <- c("SL.step")</pre>
#
      return(out)
# }
# <bytecode: 0x000001b965ed0dc0>
# <environment: namespace:SuperLearner>
# the SL.step.interaction can be adapted, changing some arguments:
SL.interaction.back = function(...) {
  SL.step.interaction(..., direction = "backward")
## The HAL algorithm implemented by default does not deal correctly with
## continuous outcome,
## However, we can define your own learner algorithm, following the template:
# function (Y, X, newX, family, obsWeights, id, ...)
# {
#
      if (family$family == "gaussian") {
#
      if (family$family == "binomial") {
#
#
#
     pred <- numeric()</pre>
#
     fit <- vector("list", length = 0)
#
     class(fit) <- c("SL.template")</pre>
      out <- list(pred = pred, fit = fit)</pre>
#
      return(out)
# }
# <bytecode: 0x00000291b737f2e0>
# <environment: namespace:SuperLearner>
## We define your own HAL algorithm that can run on both continuous and binary outcomes
SL.hal9001.Qbar <- function (Y, X, newX, family, obsWeights, id, max_degree = 2,
                              smoothness_orders = 1, num_knots = 5, ...) {
  if (!is.matrix(X))
    X <- as.matrix(X)</pre>
  if (!is.null(newX) & !is.matrix(newX))
    newX <- as.matrix(newX)</pre>
  if (length(unique(Y)) == 2) { # for binomial family
    hal_fit <- hal9001::fit_hal(Y = Y, X = X, family = "binomial",
                                 weights = obsWeights, id = id, max_degree = max_degree,
                                 smoothness_orders = smoothness_orders, num_knots = num_knots,
                                  ...)
  }
  if (length(unique(Y)) > 2) { # for quasibinomial family
    hal_fit <- hal9001::fit_hal(Y = Y, X = X, family = "gaussian",
                                 weights = obsWeights, id = id, max_degree = max_degree,
                                 smoothness_orders = smoothness_orders, num_knots = num_knots,
                                 ...)
  }
  if (!is.null(newX)) {
```

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```
pred <- stats::predict(hal_fit, new_data = newX)</pre>
  else {
    pred <- stats::predict(hal_fit, new_data = X)</pre>
  fit <- list(object = hal_fit)</pre>
  class(fit) <- "SL.hal9001"</pre>
  out <- list(pred = pred, fit = fit)</pre>
  return(out)
}
environment(SL.hal9001.Qbar) <-asNamespace("SuperLearner")</pre>
## the algorithms we would like to use can be specified separately for the Q and
# q functions
SL.library <- list(Q=c("SL.mean", "SL.glm", "SL.interaction.back", "SL.hal9001"),
                   g=c("SL.mean","SL.glm","SL.interaction.back", "SL.hal9001"))
set.seed(1234)
Psi_ATE_tmle <- ltmle(data = data_ltmle,</pre>
                      Anodes = "edu",
                      Ynodes = "death",
                      Qform = Qform,
                      gform = gform,
                      gbounds = c(0.01, 1),
                      abar = list(1,0), # vector of the counterfactual treatment
                      SL.library = SL.library,
                      variance.method = "ic")
# The estimation is more computer intensive
# The function give the ATE on the difference scale (as well, as RR and OR)
summary(Psi_ATE_tmle, estimator = "tmle")
# Additive Treatment Effect:
#
    Parameter Estimate: 0.18646
#
     Estimated Std Err: 0.0082369
                p-value: <2e-16
#
#
      95% Conf Interval: (0.17031, 0.2026)
## We can see how the SuperLearner used the algorithms for the g function
Psi_ATE_tmle$fit$g[[1]]
# we see that the Risk is high for the bad model (SL.mean)
# and very similar for the 3 other models
                                            Coef
# SL.mean_All
                           0.2409651 0.004074661 # risk is higher for the bad model
# SL.qlm All
                          0.2358587 0.000000000
# SL.interaction.back_All 0.2358587 0.995925339
# SL.hal9001_All
                         0.2358824 0.000000000
# [[1]]$AO_PM2.5
# for the g function, the SuperLearner predicts the treatment mechanism
# base on a mix between the glm and the customized xgboost algorithm.
## We can see how the SuperLearner used the algorithms for the g function
Psi_ATE_tmle$fit$Q
                                Risk
                                           Coef
                                Risk
                                          Coef
# SL.mean All
                           0.1905554 0.0000000 # risk is higher for the bad model
# SL.glm_All
                           0.1775983 0.6619501
```

```
# SL.interaction.back_All 0.1775983 0.0000000
# SL.hal9001 All
                        0.1776157 0.3380499
# The SuperLearner predicts both the treatment mechanism q and the Q function
# from a mix between the glm (or the main term glm) and the
# HAL algorithm.
# However, the choice between the SL.qlm and the SL.interaction.back
# procedure was arbitrary: as we can see the Risk is exactly the same for both
# algorithms. The final model from the step-by-step procedure was much probably
# a main term qlm.
## The `ltmle` package can also be used to estimate the effect of binary exposures
## on continous outcomes
Qform <- c(score="Q.kplus1 ~ sex + low_par_edu + edu")
gform <- c("edu ~ sex + low_par_edu")</pre>
SL.library <- list(Q=c("SL.mean", "SL.glm", "SL.interaction.back", "SL.hal9001.Qbar"),
                   g=c("SL.mean","SL.glm","SL.interaction.back", "SL.hal9001"))
set.seed(1234)
Psi_ATE_tmle_score <- ltmle(data = subset(df,</pre>
                                           select = c(sex, low_par_edu,
                                                      edu,
                                                      score)),
                      Anodes = "edu",
                      Ynodes = "score",
                      Qform = Qform,
                      gform = gform,
                      gbounds = c(0.01, 1),
                      abar = list(1,0), # vector of the counterfactual treatment
                      SL.library = SL.library,
                      variance.method = "ic")
summary(Psi_ATE_tmle_score, estimator = "tmle")
# Additive Treatment Effect:
   Parameter Estimate: -19.76
#
    Estimated Std Err: 0.37746
#
               p-value: <2e-16
      95% Conf Interval: (-20.5, -19.021)
```

The TMLE estimation of the ATE from the ltmle package for death probability and mean quality of life is +18.65% (95% CI=[17.03%, +20.26%]) and -19.76 [-20.5, -19.021].

Note that the ltmle package can also be used to calculate the IPTW estimation of the ATE and the CDE.

```
# using the output from the previous ltmle() procedure
summary(Psi_ATE_tmle, estimator = "iptw")
# Additive Treatment Effect:
# Parameter Estimate: 0.18659
# Estimated Std Err: 0.0083201
# p-value: <2e-16
# 95% Conf Interval: (0.17029, 0.2029)

summary(Psi_ATE_tmle_score, estimator = "iptw")
# Additive Treatment Effect:
# Parameter Estimate: -19.763</pre>
```

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```
# Estimated Std Err: 0.38341

# p-value: <2e-16

# 95% Conf Interval: (-20.515, -19.012)
```

The IPTW estimation of the ATE from the ltmle package for death probability and mean quality of life is +18.66% (95% CI=[+17.03%, +20.29%]) and -19.76 [-20.52, -19.01].

Chapter 5

Estimate the CDE using the ltmle package

The ltmle package can be used to estimate controlled direct effects by:

- g-computation by iterative conditional expectation (ICE),
- IPTW,
- or TMLE.

5.1 G-computation by iterative conditional expectation

5.1.1 Algorithm and manual calculation

The following steps describe the implementation of the g-computation estimator by iterative conditional expectation (ICE) for the component $\mathbb{E}(Y_{A=a',M=m})$ used in the definition of CDE $\Psi^{\text{CDE}_m} = \mathbb{E}(Y_{A=1,M=m}) - \mathbb{E}(Y_{A=0,M=m})$. Interestingly, there is no need to estimate or simulate L(1) density with this method.

- 1. Fit a logistic or a linear regression of the final outcome, conditional on the exposure A, the mediator M and all the parents of Y preceding M, to estimate $\overline{Q}_{L(2)} = \mathbb{E}(Y \mid L(0), A, L(1), M);$
- 2. Use this estimate to predict an outcome for each subject $\widehat{\overline{Q}}_{L(2)}(A=a',M=m)_i$, by evaluating the regression fit $\overline{Q}_{L(2)}$ at the chosen value for the exposure A=a' and the mediator M=m;
- 3. Fit a quasibinomial or a linear regression of the predicted values $\overline{Q}_{L(2)}(A=a',M=m)_i$ conditional on the exposure A and baseline confounders L(0) to estimate $\overline{Q}_{L(1)}=\mathbb{E}\left(\widehat{\overline{Q}}_{L(2)}(A=a',M=m)\Big|L(0),A\right)$;
- 4. Use this estimate to predict the outcome $\hat{\overline{Q}}_{L(1)}(A=a')_i$ for each subject, by evaluating the regression fit $\overline{Q}_{L(1)}$ at A=a';
- 5. Use the sample mean to estimate $\Psi_{\text{gcomp}}^{\text{CDE}_m}$

$$\hat{\Psi}_{\text{gcomp}}^{\text{CDE}_m} = \frac{1}{n} \sum_{i=1}^n \left[\hat{\overline{Q}}_{L(1)} (A=1)_i - \hat{\overline{Q}}_{L(1)} (A=0)_i \right]$$
 (5.1)

Note that G-computation by iterative expectation is preferable if the set of intermediate confounders L(1) is high-dimensional as we only need to fit 1 model by counterfactual scenario (for a whole set of L(1) variables) in the

procedure described below, whereas at least 1 model by L(1) variable and by counterfactual scenario are needed with parametric g-computation.

In the following example, we will focus on the estimand $CDE(M=0) = \mathbb{E}(Y_{A=1,M=0}) - \mathbb{E}(Y_{A=0,M=0})$.

```
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
## 1) Regress the outcome on LO, A, L1 and M (and the A*M interaction if appropriate)
death_model <- glm(death ~ sex + low_par_edu + edu + phys + occupation +
                      smoking + edu:smoking,
                   family = "binomial", data = df)
score_model <- glm(score ~ sex + low_par_edu + edu + phys + occupation +</pre>
                      smoking + edu:smoking,
                   family = "gaussian", data = df)
# 2) Generate predicted values by evaluating the regression setting the exposure
     and the mediator at exposure history of interest:
     {A=1, M=0}, {A=0, M=0}
data_Ais0_Mis0 <- data_Ais1_Mis0 <- df</pre>
data_AisO_MisO$edu <- 0
data_Ais0_Mis0$smoking <- 0</pre>
data_Ais1_Mis0$edu <- 1
data_Ais1_Mis0$smoking <- 0</pre>
Q_L2_death_AOMO <- predict(death_model,
                            newdata = data_AisO_MisO, type="response")
Q_L2_death_A1MO <- predict(death_model,</pre>
                            newdata = data_Ais1_Mis0, type="response")
Q_L2_score_AOMO <- predict(score_model,
                            newdata = data_Ais0_Mis0, type="response")
Q_L2_score_A1MO <- predict(score_model,</pre>
                            newdata = data_Ais1_Mis0, type="response")
## 3) Regress the predicted values conditional on the exposure A
      and baseline confounders L(0)
L1_death_AOMO_model <- glm(Q_L2_death_AOMO ~ sex + low_par_edu + edu,
                            family = "quasibinomial", data = df)
L1_death_A1MO_model <- glm(Q_L2_death_A1MO ~ sex + low_par_edu + edu,
                            family = "quasibinomial", data = df)
L1_score_AOMO_model <- glm(Q_L2_score_AOMO ~ sex + low_par_edu + edu,
                            family = "gaussian", data = df)
L1_score_A1MO_model <- glm(Q_L2_score_A1MO ~ sex + low_par_edu + edu,
                            family = "gaussian", data = df)
## 4) generate predicted values by evaluating the regression at exposure
      of interest: {A=1} & {A=0}
Q_L1_death_AOMO <- predict(L1_death_AOMO_model,</pre>
                            newdata = data_Ais0_Mis0, type="response")
Q_L1_death_A1MO <- predict(L1_death_A1MO_model,</pre>
                            newdata = data_Ais1_Mis0, type="response")
```

[1] -12.04431

Applying g-computation by iterative expectation, the CDE setting the mediator to 0 is +7.20% for death and -12.04 for the quantitative score.

95% confidence intervals can be estimated by bootstrap methods.

5.1.2 G-computation by ICE using the ltmle package

The ltmle package can be used to estimate Controlled Direct Effects by g-computation, as shown below.

```
library(ltmle)
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
# the data set should be composed of continuous or binary variables,
# ordered following the cause-effect sequence of each variables.
# Note that within a set of exposures or intermediate confounders measured at a
# single discrete time t, any causal sequence can be applied (for example,
# with several L1 variable, it can be {L1.1, L1.2, L1.3} or {L1.2,L1.3,L1.1},
# without any consequences on the estimation.
df_death <- subset(df, select = -c(X, subjid, score))</pre>
df score <- subset(df, select = -c(X, subjid, death))</pre>
## 1) Define Q formulas (Qbar_L1 and Qbar_Y functions)
Q_formulas_death <- c(phys = "Q.kplus1 ~ sex + low_par_edu + edu",
                      death = "Q.kplus1 ~ sex + low_par_edu + phys + occupation +
                                edu * smoking") # add interaction
Q_formulas_score <- c(phys = "Q.kplus1 ~ sex + low_par_edu + edu",
                    score = "Q.kplus1 ~ sex + low_par_edu + phys + occupation +
                              edu * smoking") # add interaction
## 2) Define g formulas (needed for the ltmle package) but they are not used
     with the g-computation estimator
g_formulas <- c("edu ~ sex + low_par_edu",</pre>
                "smoking ~ sex + low_par_edu + edu + phys + occupation")
## 3) Use the ltmle() function
# arguments:
# - Anodes: indicate the exposure and the mediator variables
```

```
# - Lnodes: indicate the intermediate confounders (+/- baseline confounders)
# - Cnodes: censoring nodes, useless in our example
# - Ynodes: outcome variable
# - survivalOutcome = FALSE in our example
# - abar: list of the two values used to define counterfactual outcomes
#
           for the contrast of interest. For example, setting M=O,
#
           CDE(M=0) = E(Y \{A=1, M=0\}) - E(Y \{A=0, M=0\})
# - rule: to define dynamic rules (useless in our example)
# - gbounds = c(0.01, 1) by default. This parameter is not used with g-computation
# - Yrange = NULL, can be used to define range (min, max) for continuous outcomes
# - SL.library = "glm", will apply main terms glm models.
                  The argument can be used to specify SuperLearner libraries.
#
#
                  However, simple qlm models might be preferable as data.adaptive
#
                  algorithms rely on cross-validation, which is difficult and long to
                  implement with the bootstrap procedure needed for 95% confidence
#
#
                  intervals
# - stratify = FALSE by default. If TRUE, glm estimations are stratified for
                each counterfactual scenario defined in abar.
# - estimate.time = FALSE. If TRUE, print a rough estimate of computation time
# - iptw.only = FALSE, useless with g-computation
# - variance.method = "ic", computation is faster than with "tmle" which
#
                        is useless with g-comp: variance estimates rely on
                        influence curves which cannot be used with g-comp because
#
                        q-computation is not a asymptotically efficient estimator.
# - observation.weights = NULL, can be used to specify individual weights
# - id = subject identifiers, useful in case of clustered structure in the data
## With a binary outcome, CDE(M=1) = P(Y_{A=1,M=0} = 1) - P(Y_{A=0,M=0} = 1)
ltmle_gcomp_CDE_MO <- ltmle(data = df_death,</pre>
                            Anodes = c("edu", "smoking"),
                            Lnodes = c("phys", "occupation"),
                            Ynodes = c("death"), # binary outcome
                            survivalOutcome = FALSE,
                            Qform = Q_formulas_death, # Q formulas
                            gform = g_formulas, # q formulas
                            abar = list(c(1,0),
                                        c(0,0), # EY_{A=1,M=0} vs EY_{A=0,M=0}
                            rule = NULL,
                            gbounds = c(0.01, 1), # truncation of g, by default
                            Yrange = NULL,
                            deterministic.g.function = NULL,
                            stratify = FALSE,
                            SL.library = "glm"
                            SL.cvControl = list(),
                            estimate.time = FALSE,
                            gcomp = TRUE, # should be TRUE for g-computation
                            iptw.only = FALSE,
                            deterministic.Q.function = NULL,
                            variance.method = "ic",
                            observation.weights = NULL,
                            id = NULL)
summary(ltmle_gcomp_CDE_MO)
```

```
## Estimator: gcomp
## Warning: inference for gcomp is not accurate! It is based on TMLE influence curves.
## Call:
```

```
## ltmle(data = df_death, Anodes = c("edu", "smoking"), Lnodes = c("phys",
##
       "occupation"), Ynodes = c("death"), survivalOutcome = FALSE,
##
       Qform = Q_formulas_death, gform = g_formulas, abar = list(c(1,
           0), c(0, 0), rule = NULL, gbounds = c(0.01, 1), Yrange = NULL,
##
##
       deterministic.g.function = NULL, stratify = FALSE, SL.library = "glm",
##
       SL.cvControl = list(), estimate.time = FALSE, gcomp = TRUE,
       iptw.only = FALSE, deterministic.Q.function = NULL, variance.method = "ic",
##
##
       observation.weights = NULL, id = NULL)
##
## Treatment Estimate:
      Parameter Estimate: 0.15656
##
       Estimated Std Err: 0.0085093
##
##
                 p-value: <2e-16
##
       95% Conf Interval: (0.13988, 0.17324)
##
## Control Estimate:
##
      Parameter Estimate: 0.084535
##
      Estimated Std Err: 0.0063349
##
                 p-value: <2e-16
##
      95% Conf Interval: (0.072118, 0.096951)
##
## Additive Treatment Effect:
##
      Parameter Estimate: 0.072027
##
      Estimated Std Err: 0.010602
##
                 p-value: 1.0947e-11
       95% Conf Interval: (0.051247, 0.092808)
##
##
## Relative Risk:
      Parameter Estimate: 1.852
##
##
    Est Std Err log(RR): 0.092521
                 p-value: 2.7185e-11
##
##
       95% Conf Interval: (1.5449, 2.2203)
##
## Odds Ratio:
##
      Parameter Estimate: 2.0102
##
    Est Std Err log(OR): 0.10412
##
                 p-value: 1.9986e-11
##
      95% Conf Interval: (1.6391, 2.4653)
## With a continuous outcome, CDE(M=1) = E(Y_{A=1,M=1}) - E(Y_{A=0,M=1})
ltmle_gcomp_CDE_MO <- ltmle(data = df_score,</pre>
                            Anodes = c("edu", "smoking"),
                            Lnodes = c("phys", "occupation"),
                            Ynodes = c("score"), # continous outcome
                            survivalOutcome = FALSE,
                            Qform = Q_formulas_score, # Q formulas
                            gform = g_formulas, # g formulas
                            abar = list(c(1,0),
                                        c(0,0), # Y_{A=1,M=0} vs Y_{A=0,M=0}
                            rule = NULL,
                            gbounds = c(0.01, 1), # by default
                            Yrange = NULL,
                            deterministic.g.function = NULL,
                            stratify = FALSE,
                            SL.library = "glm",
                            SL.cvControl = list(),
```

```
estimate.time = FALSE,
                            gcomp = TRUE, # should be TRUE for q-computation
                            iptw.only = FALSE,
                            deterministic.Q.function = NULL,
                            variance.method = "ic",
                            observation.weights = NULL,
                            id = NULL)
## Some Ynodes are not in [0, 1], and Yrange was NULL, so all Y nodes are being
## transformed to (Y-min.of.all.Ys)/range.of.all.Ys
summary(ltmle_gcomp_CDE_MO)
## Estimator: gcomp
## Warning: inference for gcomp is not accurate! It is based on TMLE influence curves.
## Call:
## ltmle(data = df_score, Anodes = c("edu", "smoking"), Lnodes = c("phys",
##
       "occupation"), Ynodes = c("score"), survivalOutcome = FALSE,
##
       Qform = Q_formulas_score, gform = g_formulas, abar = list(c(1,
           0), c(0, 0), rule = NULL, gbounds = c(0.01, 1), Yrange = NULL,
##
##
       deterministic.g.function = NULL, stratify = FALSE, SL.library = "glm",
##
       SL.cvControl = list(), estimate.time = FALSE, gcomp = TRUE,
##
       iptw.only = FALSE, deterministic.Q.function = NULL, variance.method = "ic",
##
       observation.weights = NULL, id = NULL)
##
## Treatment Estimate:
      Parameter Estimate: 37.537
##
##
       Estimated Std Err: 0.35058
##
                 p-value: <2e-16
##
       95% Conf Interval: (36.849, 38.224)
##
##
  Control Estimate:
##
      Parameter Estimate: 49.669
##
       Estimated Std Err: 0.34282
##
                 p-value: <2e-16
##
       95% Conf Interval: (48.997, 50.341)
##
## Additive Treatment Effect:
##
      Parameter Estimate: -12.132
##
       Estimated Std Err: 0.48796
##
                 p-value: <2e-16
       95% Conf Interval: (-13.089, -11.176)
##
# in order to apply quasibinomial regressions, ltmle automatically transformed
# the continuous outcome by Y_{transformed} = (Y - min(Y)) / range(Y)
# so that the range of Y_transformed is [0,1],
# Then, the results are back-transformed.
ltmle_gcomp_CDE_MO$transformOutcome
## [1] TRUE
## attr(,"Yrange")
```

We can see that the results are exactly the same as the previous manual calculation by ICE for the binary outcome. It is slightly different for the continuous outcome (because we did not apply a quasibinomial model on the min-max transformation in the manual calculation).

[1] -49.40414 97.00414

5.2 IPTW estimator of the CDE

We can express the CDE using coefficients of an MSM, where the MSM's coefficients are estimated by IPTW.

The controlled direct effect is defined as $CDE_m = \mathbb{E}(Y_{am}) - \mathbb{E}(Y_{a^*m})$.

Using the following MSM

$$\mathbb{E}(Y_{am}) = \alpha_0 + \alpha_A a + \alpha_M m + \alpha_{A*M} a \times m \tag{5.2}$$

the controlled direct effect (keeping the mediator constant to the value M=m) can be expressed using the coefficients of the MSM (5.2):

$$\begin{split} \text{CDE}_m &= (\alpha_0 + \alpha_A a + \alpha_M m + \alpha_{A*M} a \times m) - (\alpha_0 + \alpha_A a^* + \alpha_M m + \alpha_{A*M} a^* \times m) \\ \text{CDE}_m &= \alpha_A (a - a^*) + \alpha_{A*M} \times (a - a^*) \times m \end{split}$$

For a binary exposure A, we have $CDE_m = \alpha_A + \alpha_{A*M} \times m$.

5.2.1 Estimation of the MSM coefficients by IPTW

MSM coefficients can be easily estimated using an Inverse Probability of Treatment (IPTW) approach based on weighted regressions.

In order to fit the MSM (5.2), we can use a linear regression of the (observed) outcome Y on the exposure and mediator, weighted by individual stabilized weights sw_i (VanderWeele 2009):

$$\mathbb{E}\left(Y\mid A,M\right) = \alpha_0 + \alpha_A a + \alpha_M m + \alpha_{A*M} a \times m \tag{5.3}$$

where sw_i is the product of two weights $sw_i = sw_{A,i} \times sw_{M,i}$,

$$sw_{A,i} = \tfrac{P(A=a_i)}{P(A=a_i|L(0)=l(0)_i)} \text{ and } sw_{M,i} = \tfrac{P(M=m_i|A=a_i)}{P(M=m_i|A=a_i,L(0)=l(0)_i),L(1)=l(1)_i}.$$

The "no-unmeasured confounding" assumption is addressed by the application of weights sw_i , which balances confounders L(0) relative to the exposure-outcome A-Y relationship, and balance the set of confounders $\{L(0),A,L(1)\}$ relative to the mediator-outcome M-Y relationship.

Below, we estimate the CDE by hand:

```
### MSM of CDE, estimated by IPTW
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
## 1. Stabilized weight for the exposure sw_{A,i}
# 1a. Estimate g(A=a_i|L(0)) (denominator of the weight)
g_A_L <- glm(edu ~ sex + low_par_edu,</pre>
             family = "binomial", data = df)
# 1b. Predict each individual's probability of being exposed to her own exposure
# the predicted probability of the observed treatment g(A = a_i \mid L(0)) is :
gAi_L <- rep(NA, nrow(df))</pre>
gAi_L[df$edu == 1] <- predict(g_A_L, type="response")[df$edu == 1]
gAi_L[df$edu == 0] <- (1 - predict(g_A_L, type="response"))[df$edu == 0]
# 1c. Estimate q(A=a_i) (numerator of the weight)
g_A <- glm(edu ~ 1, family = "binomial", data = df)</pre>
# 1d. Predict each individual's probability of being exposed to her own exposure
# the predicted probability of the observed treatment g(A = a_i) is :
gAi <- rep(NA, nrow(df))
gAi[df$edu == 1] <- predict(g_A, type="response")[df$edu == 1]
gAi[df$edu == 0] <- (1 - predict(g_A, type="response"))[df$edu == 0]
```

binomial!

```
# 1e. Calculate the stabilized weight for the exposure A: sw_{A,i}
sw_Ai <- gAi / gAi_L
## 2. Stabilized weight for the mediator sw_{M,i}
# 2a. Estimate g(M=m_i/L(0),A,L(1)) (denominator of the weight)
g_M_L <- glm(smoking ~ sex + low_par_edu + edu + phys + occupation,
             family = "binomial", data = df)
# 2b. Predict each individual's probability of being exposed to her own exposure
# the predicted probability of the observed treatment g(A = a_i \mid L(0)) is :
gMi_L <- rep(NA, nrow(df))</pre>
gMi_L[df$smoking == 1] <- predict(g_M_L,</pre>
                                   type="response")[df$smoking == 1]
gMi_L[df$smoking == 0] <- (1 - predict(g_M_L,
                                        type="response"))[df$smoking == 0]
# 2c. Estimate g(M=m_i/A) (numerator of the weight)
g_M_A <- glm(smoking ~ edu, family = "binomial", data = df)</pre>
# 2d. Predict each individual's probability of being exposed to her own exposure
# the predicted probability of the observed treatment q(M = m_i/A) is :
gMi_A <- rep(NA, nrow(df))
gMi_A[df$smoking==1] <- predict(g_M_A, type="response")[df$smoking==1]
gMi_A[df$smoking==0] <- (1 - predict(g_M_A, type="response"))[df$smoking==0]
# 2e. Calculate the stabilized weight for the mediator M: sw_{M,i}
sw_Mi <- gMi_A / gMi_L
## 3. Define the individual stabilized weight for the CDE_m
sw_cde <- sw_Ai * sw_Mi</pre>
## 4. Estimate coefficients of the MSM using a weighted regression E(Y \mid A, sex)
# a GLM with qaussian family can be applied to estimate risk differences
msm_cde <- glm(death ~ edu + smoking + edu:smoking,</pre>
               weights = sw_cde,
               family = "gaussian",
               data = df
coef (msm_cde)
## (Intercept)
                       edu
                                smoking edu:smoking
## 0.08293313 0.07044677 0.11974047 0.14206356
## 5. Estimate CDE for m=0 and for m=1 using the MSM's coefficients
CDE_mis0 <- coef(msm_cde)["edu"] + 0 * coef(msm_cde)["edu:smoking"]
CDE_mis0
          edu
## 0.07044677
## Note: we will see below that the Itmle package use a logistic model for the MSM
msm_cde_logit <- glm(death ~ edu + smoking + edu:smoking,</pre>
                     weights = sw_cde,
                     family = "binomial",
                     data = df
```

Warning in eval(family\$initialize): nombre de succès non entier dans un glm

```
coef(msm_cde_logit)

## (Intercept) edu smoking edu:smoking
## -2.4031458 0.6948115 1.0334784 0.3322800
```

5.2.2 IPTW estmation using the ltmle package

The ltmle package can be used to estimate Controlled Direct Effects by IPTW, as shown below.

```
library(ltmle)
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
df_death <- subset(df, select = -c(X, subjid, score))</pre>
df_score <- subset(df, select = -c(X, subjid, death))</pre>
## 1) Define Q formulas (Qbar_L1 and Qbar_Y functions)
## (not needed if you only want to apply an IPTW estimation)
Q_formulas_death <- c(phys = "Q.kplus1 ~ sex + low_par_edu + edu",
                       death = "Q.kplus1 ~ sex + low_par_edu + phys + occupation +
                                edu * smoking") # add interaction
## 2) Define g formulas (needed for the ltmle package) but they are not used
     with the g-computation estimator
g_formulas <- c("edu ~ sex + low_par_edu",</pre>
                 "smoking ~ sex + low_par_edu + edu + phys + occupation")
## 3) Use the ltmle() function
## With a binary outcome, CDE(M=1) = P(Y_{A=1,M=0} = 1) - P(Y_{A=0,M=0} = 1)
ltmle_iptw_CDE_MO <- ltmle(data = df_death,</pre>
                            Anodes = c("edu", "smoking"),
                            Lnodes = c("phys", "occupation"),
                            Ynodes = c("death"), # binary outcome
                            survivalOutcome = FALSE,
                            Qform = Q_formulas_death, # Q formulas
                            gform = g_formulas, # g formulas
                            abar = list(c(1,0),
                                        c(0,0)), # EY_{A=1,M=0} vs EY_{A=0,M=0}
                            rule = NULL,
                            gbounds = c(0.01, 1), # truncation of g, by default
                            Yrange = NULL,
                            deterministic.g.function = NULL,
                            stratify = FALSE,
                            SL.library = "glm", # better to used the SuperLearner
                            SL.cvControl = list(),
                            estimate.time = FALSE,
                            gcomp = FALSE,
                            iptw.only = TRUE, # to use only the IPTW estimator
                            deterministic.Q.function = NULL,
                            variance.method = "ic",
                            observation.weights = NULL,
                            id = NULL)
summary(ltmle_iptw_CDE_MO, estimator = "iptw")
```

Estimator: iptw

```
## Call:
## ltmle(data = df_death, Anodes = c("edu", "smoking"), Lnodes = c("phys",
##
       "occupation"), Ynodes = c("death"), survivalOutcome = FALSE,
##
       Qform = Q_formulas_death, gform = g_formulas, abar = list(c(1,
##
           0), c(0, 0), rule = NULL, gbounds = c(0.01, 1), Yrange = NULL,
##
       deterministic.g.function = NULL, stratify = FALSE, SL.library = "glm",
##
       SL.cvControl = list(), estimate.time = FALSE, gcomp = FALSE,
##
       iptw.only = TRUE, deterministic.Q.function = NULL, variance.method = "ic",
##
       observation.weights = NULL, id = NULL)
##
  Treatment Estimate:
##
##
      Parameter Estimate: 0.15338
##
       Estimated Std Err: 0.008554
                 p-value: <2e-16
##
##
       95% Conf Interval: (0.13661, 0.17015)
##
##
  Control Estimate:
##
      Parameter Estimate: 0.082933
##
       Estimated Std Err: 0.0063406
##
                 p-value: <2e-16
##
       95% Conf Interval: (0.070506, 0.09536)
##
  Additive Treatment Effect:
##
##
      Parameter Estimate: 0.070447
##
       Estimated Std Err: 0.010648
##
                 p-value: 3.6865e-11
##
       95% Conf Interval: (0.049578, 0.091316)
##
  Relative Risk:
##
##
      Parameter Estimate: 1.8494
##
     Est Std Err log(RR): 0.094633
##
                 p-value: 8.1651e-11
       95% Conf Interval: (1.5363, 2.2263)
##
##
## Odds Ratio:
##
      Parameter Estimate: 2.0033
##
     Est Std Err log(OR): 0.10625
##
                 p-value: 6.1821e-11
      95% Conf Interval: (1.6267, 2.4671)
##
ltmle_iptw_CDE_MO$beta.iptw
## (Intercept)
## -2.4031458
                 0.6948115
```

5.3 TMLE estimator of the CDE

Of course, the main interest of the ltmle package is to apply a TMLE estimator.

As with the G-computation method by iterative conditional expectation, the TMLE procedure relies on the estimation of 2 \bar{Q} functions:

```
• \bar{Q}_{L(2)}(A=a,M=m) = \mathbb{E}(Y\mid L(0),A,L(1),M)
• and \bar{Q}_{L(1)} = \mathbb{E}(\hat{\bar{Q}}_{L(2)}(A=a,M=m)\mid L(0),A);
```

And as with the IPTW method, the TMLE procedure relies also on the estimation of the 2 treatment mechanisms g:

```
 \begin{array}{l} \bullet \ \ g_A(L(0)) = P(A=1 \mid L(0)) \\ \bullet \ \ \text{and} \ g_M(L(0),A,L(1)) = P(M=1 \mid L(0),A,L(1)). \end{array}
```

5.3.1 For binary outcomes

```
rm(list = ls())
df <- read.csv2("data/df.csv")</pre>
library(ltmle)
library(SuperLearner)
library(hal9001)
## 1) Define the formulas for the estimation of the 2 barQ functions and 2 g functions
# Note that it is possible to specify the A*M interaction, if we really want to
# take it into account.
## Define Q formulas (Qbar_L1 and Qbar_Y functions)
Qform <- c(phys = "Q.kplus1 ~ sex + low_par_edu + edu",
           death = "Q.kplus1 ~ sex + low_par_edu + phys + occupation +
                    edu * smoking")
## Define the formulas for the estimation of the 2 g function
gform <- c("edu ~ sex + low_par_edu",
           "smoking ~ sex + low_par_edu + edu + phys + occupation")
## The data frame should follow the time-ordering of the nodes
data_binary <- subset(df, select = c(sex, low_par_edu,</pre>
                                      edu, phys, occupation,
                                      smoking, death))
## Choose a family of data-adaptive algorithms from the SuperLearner package
# Define the SL.interaction.back learner from the SL.step.interaction
SL.interaction.back = function(...) {
  SL.step.interaction(..., direction = "backward")
# Define an ad-hoc hal learner that can be applied with continous outcomes
SL.hal9001.Qbar <- function (Y, X, newX, family, obsWeights, id, max_degree = 2,
                              smoothness_orders = 1, num_knots = 5, ...) {
  if (!is.matrix(X))
    X <- as.matrix(X)</pre>
  if (!is.null(newX) & !is.matrix(newX))
   newX <- as.matrix(newX)</pre>
  if (length(unique(Y)) == 2) { # for binomial family
    hal_fit <- hal9001::fit_hal(Y = Y, X = X, family = "binomial",
                                 weights = obsWeights, id = id, max_degree = max_degree,
                                 smoothness_orders = smoothness_orders, num_knots = num_knots,
                                 ...)
  }
  if (length(unique(Y)) > 2) { # for quasibinomial family
    hal_fit <- hal9001::fit_hal(Y = Y, X = X, family = "gaussian",
```

```
weights = obsWeights, id = id, max_degree = max_degree,
                                  smoothness orders = smoothness orders, num knots = num knots,
                                  ...)
  }
  if (!is.null(newX)) {
    pred <- stats::predict(hal_fit, new_data = newX)</pre>
  }
  else {
    pred <- stats::predict(hal_fit, new_data = X)</pre>
  fit <- list(object = hal_fit)</pre>
  class(fit) <- "SL.hal9001"</pre>
  out <- list(pred = pred, fit = fit)</pre>
  return(out)
}
environment(SL.hal9001.Qbar) <-asNamespace("SuperLearner")</pre>
## Define the SuperLearner library
SL.library <- list(Q=c("SL.mean", "SL.glm", "SL.interaction.back", "SL.hal9001.Qbar"),
                    g=c("SL.mean","SL.glm","SL.interaction.back","SL.hal9001"))
## CDE, setting M=0
set.seed(1234) # for reproducibility with the SuperLearner
CDE_ltmle_MO_death <- ltmle(data = data_binary,</pre>
                             Anodes = c("edu", "smoking"),
                             Lnodes = c("phys",
                                         "occupation"), # intermediate L(1) +/- L(0)
                             Ynodes = c("death"),
                             survivalOutcome = FALSE, # TRUE for time-to-event outcomes Y
                             Qform = Qform,
                             gform = gform,
                             abar = list(c(1,0), \# counterfactual intervention do(A=1,M=0)
                                          c(0,0), # counterfactual intervention do(A=0,M=0)
                             SL.library = SL.library,
                             estimate.time = FALSE, # estimate computation time?
                             gcomp = FALSE,
                             variance.method = "ic") # a more robust variance can
                                                      # be estimated with
                                                       # variance.method = "tmle"
summary(CDE ltmle MO death, estimator = "tmle")
```

The controlled direct effect of education on the probability of death, had the mediator been set to 0 for every participant, estimated by TMLE, is 7.03%, 95%CI=[4.96%, 9.10%].

Note that an estimation by IPTW is also available (because it relies on the g-functions that have been estimated in the process)

```
summary(CDE_ltmle_MO_death, estimator = "iptw")
```

You can check which learner has been used:

```
## for the propensity scores g
CDE_ltmle_MO_death$fit$g
```

```
## for the models of the outcome (Qbar)
CDE_ltmle_MO_death$fit$Q
## In practice, the 1tmle function estimate an MSM with 2 parameters
## (enough to compare 2 counterfactual scenarios)
CDE_ltmle_MO_death$msm
## You can check the distribution of the g-functions to check the
## positivity assumption
boxplot(CDE_ltmle_MO_death$cum.g[,,1])
boxplot(CDE_ltmle_MO_death$cum.g[,,2])
# Note: cum.q indicates the distribution before truncation at 0.01
        cum.q.unbounded indicates the distribution after truncation at 0.01
## The influence curve of the 2 parameter of the MSM are givent in the IC vector
# this is used to apply a delta method to calculate the confidence intervals
summary(CDE_ltmle_MO_death$IC)
## the fit$Qstar vector is a list of the fluctuation models
## used to update the initial Qbar function estimated by g-computation
# We can see that each Qbar model have been updated in the TMLE process.
CDE_ltmle_MO_death$fit$Qstar
```

5.3.2 For continuous outcomes

As previously, for continuous outcomes, the ltmle package transforms the outcome on a 0 to 1 continuous scale, $Y_{\text{transformed}} = \frac{Y - \min(Y)}{\max(Y) - \min(Y)}$, so that quasi-binomial parametric models can be used in the computation procedure. Mean predictions are then back-transformed on the original scale.

```
## Define the data set with the continuous outcome score
data_continuous <- subset(df, select = c(sex, low_par_edu,</pre>
                                          edu, phys, occupation,
                                          smoking, score))
## Replace the Qbar function
## (the 2d formula should be named score instead of death)
Qform <- c(phys = "Q.kplus1 ~ sex + low_par_edu + edu",
           score = "Q.kplus1 ~ sex + low_par_edu + phys + occupation +
                                edu * smoking") # add interaction
## SuperLearner library
SL.library <- list(Q=c("SL.mean", "SL.glm", "SL.interaction.back", "SL.hal9001.Qbar"),
                   g=c("SL.mean","SL.glm","SL.interaction.back","SL.hal9001"))
set.seed(1234)
## CDE, setting M=0
CDE_ltmle_MO_score <- ltmle(data = data_continuous,</pre>
                             Anodes = c("edu", "smoking"),
                            Lnodes = c("phys", "occupation"),
                            Ynodes = c("score"),
                             survivalOutcome = FALSE,
                            Qform = Qform,
                             gform = gform,
                             abar = list(c(1,0), # do(A=1, M=0)
                                         c(0,0)), # do(A=0,M=0)
                            SL.library = SL.library,
                             estimate.time = FALSE, # estimate computation time?
```

The controlled direct effect of education on the score outcome, had the mediator been set to 0 for every participant, estimated by TMLE is -12.12, 95%CI=[-13.08, -11.17].

Chapter 6

Estimate rNDE and rNIE by TMLE

Laan, Mark J. van der, and Sherri Rose. 2011. Targeted Learning: Causal Inference for Observational and Experimental Data. 1st ed. Springer Series in Statistics. New York, NY: Springer.

VanderWeele, Tyler J. 2009. "Marginal Structural Models for the Estimation of Direct and Indirect Effects." Epidemiology 20: 18–26.