

---

# Causal Inference

a summary

---

# Contents

<b>1</b>	<b>General</b>	<b>3</b>		
<b>2</b>	<b>Models</b>	<b>5</b>		
2.1	Traditional Methods . . . . .	5	2.2.1	Time-varying A . . . . . 6
2.2	G-Methods . . . . .	5	2.3	Advanced Methods . . . . . 7
			<b>3</b>	<b>Longitudinal Data</b> <b>9</b>

# 1 General

**Causal Roadmap** (Petersen and van der Laan, 2014)  
systematic approach linking causality to statistical procedures

**1. Specifying Knowledge.** structural causal model (unifying counterfactual language, structural equations, & causal graphs): a set of possible data-generating processes, expresses background knowledge and its limits

**2. Linking Data.** specifying measured variables and sampling specifics (latter can be incorporated into the model)

**3. Specifying Target.** define hypothetical experiment: decide

1. variables to intervene on: one (point treatment), multiple (longitudinal, censoring/missing, (in)direct effects)
2. intervention scheme: static, dynamic, stochastic
3. counterfactual summary of interest: absolute or relative, marginal structural models, interaction, effect modification
4. population of interest: whole, subset, different population

**4. Assessing Identifiability.** are knowledge and data sufficient to derive estimand and if not, what else is needed?

**5. Select Estimand.** current best answer: knowledge-based assumptions + which minimal convenience-based assumptions (transparency) gets as close as possible

**6. Estimate.** choose estimator by statistical properties, nothing causal here

**7. Interpret.** hierarchy: statistical, counterfactual, feasible intervention, randomized trial

**Average Causal Effect**  $E[Y^{a=1}] \neq E[Y^{a=0}]$

$$E[Y^a] = \sum_y y p_{Y^a}(y) \quad (\text{discrete})$$

$$= \int y f_{Y^a}(y) dy \quad (\text{continuous})$$

individual causal effect  $Y_i^{a=1} \neq Y_i^{a=0}$  generally unidentifiable

*null hypothesis:* no average causal effect

*sharp null hypothesis:* no causal effect for any individual

**notation**  $A, Y$ : random variables (differ for individuals);  $a, y$ : particular values; counterfactual  $Y^{a=1}$ :  $Y$  under treatment  $a = 1$

**stable unit treatment value assumption (SUTVA)**  $Y_i^a$  is well-defined: no interference between individuals, no multiple versions of treatment (weaker: treatment variation irrelevance)

**causal effect measures** typically based on means

*risk difference:*  $\Pr[Y^{a=1} = 1] - \Pr[Y^{a=0} = 1]$

*risk ratio:*  $\frac{\Pr[Y^{a=1}=1]}{\Pr[Y^{a=0}=1]}$

*odds ratio:*  $\frac{\Pr[Y^{a=1}=1]/\Pr[Y^{a=1}=0]}{\Pr[Y^{a=0}=1]/\Pr[Y^{a=0}=0]}$

*number needed to treat (NNT)* to save 1 life:  $-1/\text{risk difference}$

**sources of random error:** sampling variability (use consistent estimators), nondeterministic counterfactuals

**association** compares  $E[Y|A = 1]$  and  $E[Y|A = 0]$ , **causation** compares  $E[Y^{a=1}]$  and  $E[Y^{a=0}]$  (whole population)

**Target Trial** emulating an ideal randomized experiment explicitly formulate target trial & show how it is emulated → less vague causal question, helps spot issues

**missing data problem** unknown counterfactuals

*randomized experiments:* missing completely at random →

exchangeability (= exogeneity as treatment is exogenous)

*ideal randomized experiment:* no censoring, double-blind,

well-defined treatment, & adherence → association is causation

*pragmatic trial:* no placebo/blindness, realistic monitoring

**PICO** (population, intervention, comparator, outcome): some components of target trial

**three types of causal effects:**

*intention-to-treat effect* (effect of treatment assignment)

*per-protocol effect* (usually dynamic when toxicity arises)

*other intervention effect* (strategy changed during follow-up)

**controlled direct effects:** effect of  $A$  on  $Y$  not through  $B$

*natural direct effect*  $A$  on  $Y$  if  $B^{a=0}$  (cross-world quantity)

*principal stratum effect*  $A$  on  $Y$  for subset with  $B^{a=0} = B^{a=1}$

**crossover experiment:** sequential treatment & outcome  $t=0, 1$   
individual causal effect  $Y_{it}^{a_t=1} - Y_{it}^{a_t=0}$  only identifiable if: no carryover effect, effect  $\perp$  time, outcome  $\perp$  time

**time zero** if eligibility at multiple  $t$  (observational data):

earliest, random  $t$ , all  $t$  (adjust variance with bootstrapping)

**grace periods:** usually treatment starts  $x$  months after first

eligible, if death before: randomly assign strategy/copy into both

**Identifiability Conditions** hold in ideal experiments

**consistency** counterfactuals correspond to data  $Y = Y^A$ :

if  $A = a$ , then  $Y^a = Y$  for each individual

- precise definition of  $Y^a$  via specifying  $a$  (sufficiently well-defined  $a$  maybe impossible (effect of DNA before it was discovered), relies on expert consensus)
- linkage of counterfactuals to data ( $a$  must be seen in data)

**positivity**  $\Pr[A = a|L = l] > 0 \ \forall l$  with  $\Pr[L = l] > 0$ ;

$$f_L(l) \neq 0 \Rightarrow f_{A|L}(a|l) > 0 \ \forall a, l$$

- structural violations (inference not on full population)
- random variability (smooth over with parametric models)

can sometimes be empirically verified (if all is seen in data)

**exchangeability** unverifiable without randomization

- *marginal:*  $Y^a \perp\!\!\!\perp A \triangleq$  randomized experiment, counterfactuals are missing completely at random (MCAR)
- *conditional:*  $Y^a \perp\!\!\!\perp A|L \triangleq$  conditionally randomized, counterfactuals are missing at random (MAR)

alternative definition:  $\Pr[A = 1|Y^{a=0}, L] = \Pr[A = 1|L]$

**additional conditions:**

*correct measurement* mismeasurement of  $A, Y, L$  results in bias

*correct model specification* models  $\xrightarrow{\text{may}}$  misspecification bias

**Effect Modification**  $A$  on  $Y$  varies across levels of  $V$

null average causal effect  $\neq$  null causal effect per subgroup

**population characteristics:** causal effect measure is actually “effect in a population with a particular mix of effect modifiers”

**transportability:** extrapolation of effect to another population (issues: effect modification, versions of treatment, interference)

effects conditional on  $V$  may be more transportable

**types:** additive/multiplicative scale, qualitative (effect in opposite directions)/quantitative, surrogate/causal

**calculation:**

- *stratify* by  $V$  then standardize/IP weight for  $L$ ,
- $L$  as *matching* factor (ensures positivity, difficult if high-dimensional  $L$ )

**collapsibility:** causal risk difference and ratio are weighted

averages of stratum-specific risks, can not be done for odds ratio

**Interaction** effects of joint interventions  $A$  and  $E$

$$\Pr[Y^{1,1}=1] - \Pr[Y^{0,1}=1] \neq \Pr[Y^{1,0}=1] - \Pr[Y^{0,0}=1]$$

$A$  and  $E$  have equal status and could also be considered a combined treatment  $AE$ , exchangeability for both is needed  
*additive scale* (above): “>” superadditive and “<” subadditive;  
*multiplicative scale*: “>” super- and “<” submultiplicative

**difference to effect modification**: if  $E$  is randomly assigned methods coincide, but  $V$  can not be intervened on as  $E$  can  
**monotonicity** effect is either nonnegative or nonpositive  $\forall i$   
**sufficient component-cause framework** pedagogic model  
*response types* for binary  $A$ : helped, immune, hurt, doomed;  
for binary  $A$  and  $E$ : 16 types

(minimal) sufficient causes:

- (minimal)  $U_1$  together with  $A = 1$  ensure  $Y = 1$
- (minimal)  $U_2$  together with  $A = 0$  ensure  $Y = 1$

sufficient cause interaction:  $A$  and  $E$  appear together in a minimal sufficient cause

**NPSEM** nonparametric structural equation model

$$V_m = f_m(pa_m, \epsilon_m)$$

counterfactuals are obtained recursively, e.g.  $V_3^{v_1} = V_3^{v_1, V_2^{v_1}}$   
implies any variable can be intervened on

aka finest causally interpreted structural tree graph (FCISTG)

**additional assumption**  $\cap$  FCISTG  $\Rightarrow$  causal Markov condition:

- independent errors (NPSEM-IE): all  $\epsilon_m$  mutually independent
- fully randomized (FFRCISTG):  $V_m^{\bar{v}_{m-1}} \perp\!\!\!\perp V_j^{\bar{v}_{j-1}}$  if  $\bar{v}_{j-1}$  subvector of  $\bar{v}_{m-1}$

NPSEM-IE  $\Rightarrow$  FFRCISTG (assume DAGs represent latter)

NPSEM-IE assume crossworld independencies  $\rightarrow$  unverifiable

**Causal DAG** draw assumptions before conclusions

*rules*: arrow means direct causal effect for at least one  $i$ , absence means sharp null holds, all common causes are on the graph

*neglects*: direction of cause (harmful/protective), interactions

*convention*: time flows from left to right

**causal Markov assumption**: any variable ( $v$ ) | its direct causes ( $pa_j$ )  $\perp\!\!\!\perp$  its non-descendants ( $\neg v_j$ )  $\Leftrightarrow$  Markov factorization

$$f(v) = \prod_{j=1}^M f(v_j|pa_j)$$

**d-separation** (d for directional): a pathway in a DAG is ...

- blocked if collider or conditioned on non-collider
- opened if conditioned on collider or descendent of collider

2 variables are d-separated if all connecting paths are blocked

under causal Markov: d-separation  $\Rightarrow$  independence

under faithfulness: independence  $\Rightarrow$  d-separation

**faithfulness**: effects don't cancel out perfectly

*discovery*: process of learning the causal structure; requires faithfulness, but even with it is often impossible

**SWIGs** single world intervention graphs

**counterfactual graphic approach**:  $A$  turns into  $A|a$ , the left (right) side inherits incoming (outgoing) arrows (intervention with  $A = a$ ); all outcomes of  $A$  get a superscript  $a$ , e.g.  $Y^a$ ; more than one intervention possible, dynamic strategies require additional arrows from  $L$  to  $a$

$A$  and  $Y^a$  are d-separated  $\rightarrow Y^a \perp\!\!\!\perp A|L$  (for FFRCISTG)

**Confounding** bias due to common cause of  $A$  &  $Y$  *not in*  $L$   
randomization prevents confounding

**backdoor path**: noncausal path  $A$  to  $Y$  with arrow into  $A$

**backdoor criterion**: all backdoor paths are blocked by  $L$  & no descendants of  $A$  in  $L \Rightarrow$  conditional exchangeability

$Y^a \perp\!\!\!\perp A|L \Rightarrow L$  fulfills backdoor criterion if faithful (FFRCISTG)

**confounders in observational studies**: occupational factors (*healthy worker bias*), clinical decisions (*confounding by indication/channeling*), lifestyle, genetic factors (*population stratification*), social factors, environmental exposures

given a DAG, confounding is an absolute, confounder is relative  
surrogate confounders in  $L$  may reduce confounding bias

**negative outcome controls**: if  $A$  and  $Y$  share a common cause  $U$ : measure effect for  $Y_0$  (before treatment) and  $Y_1$  (after), subtract (assumption of additive equi-confounding)

**front door criterion** using the full mediator  $M$ :  $\Pr[Y^a = 1] = \sum_m \Pr[M = m|A = a] \sum_{a'} \Pr[Y = 1|M = m, A = a'] \Pr[A = a']$

**Selection Bias** bias due to common effect of  $A$  &  $Y$  *in*  $L$   
 $=$  conditioning on collider (can't be fixed by randomization)

**examples**: informative censoring, nonresponse bias, healthy worker bias, volunteer bias; often M-bias ( $A \leftarrow U_1 \rightarrow L \leftarrow U_2 \rightarrow Y$ )

**solution**: target  $Y^{A,C}$ ,  $AC$  fulfills identifiability conditions, if competing events, interventions may not be well-defined

**multiplicative survival model**:  $\Pr[Y=0|E=e, A=a] = g(e)h(a) \rightarrow$  no interaction between  $E$  and  $A$  on the multiplicative scale; if  $Y = 0$  is conditionally independent, then  $Y = 1$  can't be as  $\Pr[Y=1|E=e, A=a] = 1 - g(e)h(a) \rightarrow$  conditioning on a collider could be unbiased if restricted to certain levels ( $Y = 0$ )

**measurement bias** chapter 9

**random variability** chapter 10

## 2 Models

**Modeling** data are a sample from the target population

*estimand*: quantity of interest, e. g.  $E[Y|A = a]$   
*estimator*: function to use, e. g.  $\hat{E}[Y|A = a]$   
*estimate*: apply function to data, e. g. 4.1

**model**: a priori restriction of joint distribution/dose-response curve; *assumption*: no model misspecification (usually wrong)

**non-parametric estimator**: no restriction (saturated model) = *Fisher consistent estimator* (entire population data  $\rightarrow$  true value)

**parsimonious model**: few parameters estimate many quantities

**bias-variance trade-off**:

wiggleness  $\uparrow \rightarrow$  misspecification bias  $\downarrow$ , CI width  $\uparrow$

**Variable Selection** can induce bias if  $L$  includes:

(descendant of) collider: *selection bias under the null*  
 noncollider effect of  $A$ : *selection bias under the alternative*  
 mediator: *overadjustment for mediators*

temporal ordering is not enough to conclude anything

**bias amplification**: e.g. by adjusting for an instrument  $Z$  (can also reduce bias)

**Machine Learning**  $L$  is high-dimensional  
 use lasso or ML for IP weighting/standardization

**but**: ML does not guarantee elimination of confounding and has largely unknown statistical properties

$\rightarrow$  **doubly robust estimator**: consistent if bias  $< \frac{1}{\sqrt{n}}$

*sample splitting*: train estimators on training sample, use resulting estimators for doubly robust method on estimation sample (CIs on estimation sample are valid, but  $n$  halved)

*cross-fitting*: do again the other way round, average the two estimates, get CI via bootstrapping

**problems**: unclear choice of algorithm, is bias small enough?

**Super Learning** (Van der Laan et al., 2007, 2011)

**oracle selector**: select best estimator of set of learners  $Z_i$

**discrete super learner**: select algorithm with smallest cross-validated error (converges to oracle for large sample size)

**super learner**: improves asymptotically on discrete version

$\text{logit}(Y = 1|Z) = \sum_i \alpha_i Z_i$ , with  $0 < \alpha_i < 1$  and  $\sum \alpha_i = 1$

weights  $\alpha_i$  are determined inside the cross-validation; for the prediction,  $Z_i$  trained on the full data set are used

can be cross-validated itself to check for overfitting (unlikely)

### 2.1 Traditional Methods

**Stratification** calculate risk for each stratum of  $L$

only feasible if enough data per stratum

**instrumental variable estimation** chapter 16

**Outcome regression** chapter 15

**causal survival analysis** chapter 17 (and technical point 22.3)

### 2.2 G-Methods

**G-Methods** generalized treatment contrasts: adjust for (surrogate) confounders  $L$

- **standardization** two types of g-formula
- **IP weighting** also g-formula
- **g-estimation**: not needed unless longitudinal

**Standardization** plug-in (or parametric if so) g-formula

$$E[Y^a] = \overbrace{E[E[Y|A=a, L=l]]}^{\text{conditional expectation}} = \overbrace{\int E[Y|L=l, A=a] f_L[l] dl}^{\text{joint density estimator}}$$

weighted average of stratum-specific risks; unknowns can be estimated non-parametrically or modeled

**no need to estimate  $f_L[l]$ /integrate** as empirical distribution

can be used: estimate outcome model  $\rightarrow$  predict counterfactuals on whole dataset  $\rightarrow$  average the results ( $\rightarrow$  CI by bootstrapping)

**for discrete  $L$**   $E[Y|A=a] = \sum_l E[Y|L=l, A=a] \Pr[L=l]$

**time-varying** standardize over all possible  $\bar{L}$ -histories  
simulates joint distribution of counterfactuals  $(Y^{\bar{a}}, \bar{L}^{\bar{a}})$  for  $\bar{a}$   
**joint density estimator (jde)**

$$\text{discrete: } E[Y^{\bar{a}}] = \sum_{\bar{l}} E[Y|\bar{A} = \bar{a}, \bar{L} = \bar{l}] \prod_{k=0}^K f(l_k|\bar{a}_{k-1}, \bar{l}_{k-1})$$

$$\text{continuous: } \int f(y|\bar{a}, \bar{l}) \prod_{k=0}^K f(l_k|\bar{a}_{k-1}, \bar{l}_{k-1}) dl$$

for *stochastic strategies* multiply with  $\prod_{k=0}^K f^{int}(a_k|\bar{a}_{k-1}, \bar{l}_k)$

**estimation** (Young et al., 2011; Schomaker et al., 2019)

1. model  $f(l_k|\bar{a}_{k-1}, \bar{l}_{k-1})$  and  $E[Y|\bar{A} = \bar{a}, \bar{L} = \bar{l}]$
2. simulate data forward in time:  
at  $k = 0$ : use empirical distribution of  $L_0$  (observed data)  
at  $k > 0$ : set  $\bar{A} = \bar{a}$ , draw from models estimated in 1.
3. calculate mean of  $\hat{Y}_{K,i}^{\bar{a}}$  (bootstrap for CI)

**iterated conditional expectation (ice)**

$$E[Y_T^{\bar{a}}] = E[E[E[...E[Y_T|\bar{A}_{T-1}=\bar{a}_{T-1}, \bar{L}_T]...|\bar{A}_0=a_0, L_1]|L_0]]$$

**estimation** (Schomaker et al., 2019)

1. model inside out:  $Q_T = E[Y_T|\bar{A}_{T-1}, \bar{L}_T]$  to  $Q_0 = E[Q_1|\bar{L}_0]$ ,  
predict  $Q_t$  with  $\bar{A} = \bar{a}$  in each step
2. calculate mean of  $\hat{Q}_{0,i}^{\bar{a}}$  (bootstrap for CI)

**g-null paradox** even if the sharp null holds, model misspecification can lead to it being falsely rejected

Proof: for  $L_0 \rightarrow A_0 \rightarrow Y_0 \rightarrow L_1 \rightarrow A_1 \rightarrow Y_1$ ,  $\bar{a} = (a_0, a_1)$

$$E[Y_1^{\bar{a}}] \stackrel{\text{CE}}{=} E[E[Y_1^{\bar{a}}|A_0=a_0, L_0]]$$

$$(\text{ice}) \stackrel{\text{CE}^*}{=} E[E[E[Y_1|\bar{L}, \bar{A}=\bar{a}, Y_0]|A_0=a_0, L_0]]$$

$$\stackrel{\text{LTP}}{=} E\left[\sum_{l_1} E[Y_1|A_0=a_0, \bar{L}, Y_0] \Pr[l_1|a_0, l_0, y_0]\right]$$

$$\stackrel{\text{LTP}}{=} \sum_{l_0} \left[\sum_{l_1} E[Y_1|A_0=a_0, \bar{L}, Y_0] \Pr[l_1|a_0, l_0, y_0]\right] \Pr[l_0]$$

$$(\text{jde}) \stackrel{\text{sum}}{=} \sum_{\bar{l}} E[Y_1|A_0=a_0, \bar{L}, Y_0] \Pr[l_1|a_0, l_0] \Pr[l_0]$$

CE: conditional expectation; \*: exchangeability;

LTP: law of total probability

**Marginal Structural Models** association is causation  
in the IP weighted pseudo-population

$$\text{associational model } E[Y|A] = \text{causal model } E[Y^a]$$

step 1: estimate/model  $f[A|L]$  (and  $f[A]$ )  $\rightarrow$  get  $(S)W^A$

step 2: estimate regression parameters for pseudo-population

**effect modification** variables  $V$  can be included (e.g.

$\beta_0 + \beta_1 a + \beta_2 V a + \beta_3 V$ ; technically not marginal anymore),

$$SW^A(V) = \frac{f[A|V]}{f[A|L]} \text{ more efficient than } SW^A$$

**Censoring** measuring joint effect of  $A$  and  $C$

$$E[Y^{a,c=0}] \text{ is of interest}$$

$$\text{standardization } E[Y|A=a] = \int E[Y|L=l, A=a, C=0] dF_L[l]$$

$$\text{IP weights } W^{A,C} = W^A \times W^C \quad (\text{uses } n) \quad \text{or} \\ SW^{A,C} = SW^A \times SW^C \quad (\text{uses } n^{c=0})$$

**g-estimation** can only adjust for confounding, not selection bias  
 $\rightarrow$  use IP weights

**G-Estimation** (additive) structural nested models

$$\text{logit } \Pr[A=1|H(\psi^\dagger), L] = \alpha_0 + \alpha_1 H(\psi^\dagger) + \alpha_2 L$$

$$H(\psi^\dagger) = Y - \psi^\dagger A$$

find  $\psi^\dagger$  which renders  $\alpha_1 = 0$ ; 95 %-CI: all  $\psi^\dagger$  for which  $p > 0.05$   
closed-form solution for linear models

**derivation:**  $H(\psi^\dagger) = Y^{a=0}$

$$\text{logit } \Pr[A=1|Y^{a=0}, L] = \alpha_0 + \alpha_1 Y^{a=0} + \alpha_2 L$$

$Y^{a=0}$  unknown, but because of exchangeability  $\alpha_1$  should be zero

$$Y^{a=0} = Y^a - \psi_1 a$$

equivalent to  $Y^{a=0} = Y^{a=1} - \psi_1$ , but using no counterfactuals

**structural nested mean model**

$$\text{additive: } E[Y^a - Y^{a=0}|A=a, L] = \beta_1 a (+\beta_2 a L)$$

$$\text{multiplicative: } \log\left(\frac{E[Y^a|A=a, L]}{E[Y^{a=0}|A=a, L]}\right) = \beta_1 a (+\beta_2 a L)$$

multiplicative is preferred if  $Y$  always positive, but does not  
extend to longitudinal case

semi-parametric: agnostic about  $\beta_0$  and effect of  $L \rightarrow$  robust  $\uparrow$

**no time-varying:** no nesting; model equals marginal structural  
models with missing  $\beta_0, \beta_3$  (unspecified “no treatment”)

**sensitivity analysis:** unmeasured confounding ( $\alpha_1 \neq 0$ ) can be  
examined: do procedure for different values of  $\alpha_1 \rightarrow$  plot  $\alpha_1$  vs.  
 $\psi^\dagger \rightarrow$  how sensitive is estimate to unmeasured confounding?

**effect modification:** add  $V$  in both g-estimation equations

**doubly robust estimators** exist

**IP Weighting** inverse probability of treatment (g-formula)

$$E[Y^a] = E\left[\frac{I(A=a)Y}{f[A|L]}\right]; W^A = \frac{1}{f[A|L]}; SW^A = \frac{f(A)}{f[A|L]}$$

unknowns can be estimated non-parametrically or modeled

**pseudo-population:** everyone is treated & untreated ( $L \nrightarrow A$ )

**FRCISTG** (*fully randomized causally interpreted structured graph*): probability tree for  $L \rightarrow A \rightarrow Y$ , can be used to  
calculate/visualize simulation of values for  $A$

**for discrete  $A, L$**   $f[a|l] = \Pr[A=a, L=l]$

**estimators:** Horvitz-Thompson; Hajek (modified version)

**stabilized weights  $SW^A$**  should have an average of 1 (check!)

$\rightarrow$  pseudo-population same size  $\rightarrow$  CI width  $\downarrow$

**Standardization and IP Weighting** are equivalent,

**but** if modeled, different “no misspecification” assumptions:

standardization: outcome model

IP weighting: treatment model

**doubly robust estimators:** reduce model misspecification bias,

consistent if either model is correct; **e.g.:**

$$1. \text{ fit outcome regression with variable } R = \begin{cases} +W^A & \text{if } A=1 \\ -W^A & \text{if } A=0 \end{cases}$$

2. standardize by averaging

## 2.2.1 Time-varying A

**IP Weighting**

$$W^{\bar{A}} = \prod_{k=0}^K \frac{1}{f(A_k|\bar{A}_{k-1}, \bar{L}_k)}$$

$$SW^{\bar{A}} = \prod_{k=0}^K \frac{f(A_k|\bar{A}_{k-1}, \bar{L}_k)}{f(A_k|\bar{A}_{k-1}, \bar{L}_k)}$$

**Doubly Robust Estimator** sequential estimation

1. estimate  $\hat{f}(A_m | \bar{A}_{m-1}, \bar{L}_m)$  (e.g. logistic model), use it to calculate at each time  $m$ :  $\widehat{W}^{\bar{A}_m} = \prod_{k=0}^m \frac{1}{\hat{f}(A_k | \bar{A}_{k-1}, \bar{L}_k)}$  and modified IP weights at  $m$ :  $\widehat{W}^{\bar{A}_{m-1}, a_m} = \frac{\widehat{W}^{\bar{A}_{m-1}}}{\hat{f}(a_m | \bar{A}_{m-1}, \bar{L}_m)}$
2. with  $\widehat{T}_{K+1} := Y$ , recursively for  $m = K, K-1, \dots, 0$ :
  - (a) fit outcome regression on  $\widehat{T}_{m+1}$  with variable  $\widehat{W}^{\bar{A}_m}$
  - (b) calculate  $\widehat{T}_m$  using the outcome model with  $\widehat{W}^{\bar{A}_{m-1}, a_m}$
3. calculate standardized mean outcome  $\widehat{E}[Y^{\bar{a}}] = E[\widehat{T}_0]$

**valid**, if treatment or outcome model correct, or treatment correct until  $k$  and outcome otherwise ( $k+1$  robustness)

**G-Estimation** nested equations: for each time  $k$

**structural nested mean models** separate effect of each  $a_k$

$$E[Y^{\bar{a}_{k-1}, a_k, \bar{a}_{k+1}} - Y^{\bar{a}_{k-1}, \bar{a}_{k+1}} | \bar{L}^{\bar{a}_{k-1}} = \bar{l}_k, \bar{A}_{k-1} = \bar{a}_{k-1}] = a_k \gamma_k(\bar{a}_{k-1}, \bar{l}_k, \beta)$$

## 2.3 Advanced Methods

**TMLE** targeted minimum loss-based estimation

$$O = (W, A, Y) \sim P_0$$

target  $\Psi(P_0) = \Psi(\bar{Q}_0, Q_{W,0}) = \psi_0$ ,

often:  $E_{W,0}[E_0(Y|A=1, W) - E_0(Y|A=0, W)]$

**first step**: outcome model  $\bar{Q}_n^0(A, W)$  estimating  $\bar{Q}_0$  (part of  $P_0$ )

- super learning is often used here, but leads to a biased estimate
- not all of  $f(Y|A, W)$  needs to be estimated, just the relevant portion, *typically average outcome*  $E_0(Y|A, W) \rightarrow$  efficiency  $\uparrow$

**second step**: update  $\bar{Q}_n^0(A, W)$  to  $\bar{Q}_n^1(A, W)$  using treatment model  $g_n$  estimating  $g_0 = P_0(A|W)$

1. model  $g_n$ , super learning is a popular choice here, too
2. calculate  $n$  clever covariates:  $H_n^*(A, W) = \begin{cases} \frac{1}{g_n(1|W)} & \text{if } A_i=1 \\ \frac{1}{g_n(0|W)} & \text{if } A_i=0 \end{cases}$
3. update  $\bar{Q}_n^0$ , by estimating  $\epsilon_n$  with offset logistic regression:  $\text{logit} \bar{Q}_n^1(A, W) = \text{logit} \bar{Q}_n^0(A, W) + \epsilon_n H_n^*(A, W)$  (converges after first update), then calculate counterfactuals

- goal: bias reduction, get optimal bias-variance trade-off
- removes all asymptotic bias, if consistent estimator is used here

**third step**: use empirical distribution for  $Q_{W,0}$  in a substitution estimator, e.g.:  $\psi_n^{TMLE} = \frac{1}{n} \sum_{i=1}^n [\bar{Q}_n^1(1, W_i) - \bar{Q}_n^1(0, W_i)]$

**advantages**: doubly robust (consistent if either outcome or treatment model is correctly specified), asymptotically efficient (if both are correct), substitution estimator (more robust to outliers and sparsity)

**closed form inference based on the influence curve**:

$$IC_n^*(O_i) = \overbrace{\left[ \frac{1(A_i=1)}{g_n(1, W_i)} - \frac{1(A_i=0)}{g_n(0, W_i)} \right] [Y - \bar{Q}_n^1(A_i, W_i)]}^a + \overbrace{\bar{Q}_n^1(1, W_i) - \bar{Q}_n^1(0, W_i) - \psi_{TMLE, n}}^b$$

TMLE sets the mean of the IC,  $\overline{IC}_n$ , to zero ( $b$  has already mean zero, see third step, the first part of  $a$  is the clever covariate)

sample variance is then:  $S^2(IC_n) = \frac{1}{n} \sum_{i=1}^n (IC_n(O_i) - \overline{IC}_n)^2$

standard error of estimator:  $\sigma_n = \sqrt{\frac{S^2(IC_n)}{n}}$

95% CI:  $\psi_{TMLE, n} \pm z_{0.975} \frac{\sigma_n}{\sqrt{n}}$ ; p-value:  $2 \left[ 1 - \Phi \left( \left| \frac{\psi_{TMLE, n}}{\sigma_n / \sqrt{n}} \right| \right) \right]$

**calculations**

$$H_k(\psi^\dagger) = Y - \sum_{j=k}^K A_j \gamma_j(\bar{A}_{j-1}, \bar{L}_j, \psi^\dagger)$$

function  $\gamma_j$  can be, e.g. constant ( $\psi_1$ ), time-varying only ( $\psi_1 + \psi_2 k$ ), or dependent on treatment/covariate history

$$\text{logit Pr}[A_k = 1 | H_k(\psi^\dagger), \bar{L}_k, \bar{A}_{k-1}] =$$

$$\alpha_0 + \alpha_1 H_k(\psi^\dagger) + \alpha_2 w_k(\bar{L}_k, \bar{A}_{k-1})$$

find  $\alpha_1$  that is closest to zero

a closed form estimator exists for the linear case

**Censoring**  $\bar{C}$ : monotonic type of missing data

**standardization**:

$$\int f(y | \bar{a}, \bar{c}=\bar{0}, \bar{l}) \prod_{k=0}^K dF(l_k | \bar{a}_{k-1}, c_{k-1}=0, \bar{l}_{k-1})$$

**IP weighting**:

$$SW^{\bar{C}} = \prod_{k=1}^{K+1} \frac{1 \cdot \Pr(C_k = 0 | \bar{A}_{k-1}, C_{k-1} = 0)}{\Pr(C_k = 0 | \bar{A}_{k-1}, C_{k-1} = 0, \bar{L}_k)}$$

$$\mathcal{L}(O) = \overbrace{\Pr(Y|A, W)}^{Q_Y} \overbrace{\Pr(A|W)}^g \overbrace{\Pr(W)}^{Q_W}$$

$H(A, W)$  depends on target parameter and loss function but is a function of the propensity score update initial fit

$$\bar{Q}_n^* = \bar{Q}_n^0 + \epsilon H(A, W)$$

valid inference, good finite sample performance,

$H(A, W)$  comes from the influence curve, targeting ensures mean of efficient influence curve  $D^*(P)$  is zero

TMLE solves  $P_n D^*(P_n^*) = 0$

TMLE is a substitution estimator

$\psi_n^{TMLE} = \frac{1}{2} \sum_{i=1}^n \bar{Q}_n^*(1, W_i) - \frac{1}{2} \sum_{i=1}^n \bar{Q}_n^*(0, W_i)$  therefore mean of  $b$  is zero

targeting step makes sure  $a$  also has mean zero

MLE solves  $\sum_{i=1}^n H(A_i, W_i) [Y_i - \bar{Q}_n^*(A_i, W_i)] = 0$  where  
 $\bar{Q}_n^*(A_i, W_i) = \hat{e}H(A, W) + \bar{Q}_n^0$  therefore obvious choice:

$$\left| \begin{array}{l} H(A, W) = \frac{A}{g(1, W)} - \frac{1-A}{g(0, W)} \\ \text{influence curve based inference: asymptotic linearity} \\ \sqrt{n} (\psi_n^{TMLE} - \psi_0) \xrightarrow{D} N(0, \sigma^2) \end{array} \right.$$



### 3 Longitudinal Data

**Time-Varying Treatments** compare 2 treatments

treatment history up to  $k$ :  $\bar{A}_k = (A_0, A_1, \dots, A_k)$

shorthand: always treated  $\bar{A} = \bar{1}$ , never treated  $\bar{A} = (\bar{0})$

**static strategy:**  $g = [g_0(\bar{a}_{-1}), \dots, g_K(\bar{a}_{K-1})]$

**dynamic strategy:**  $g = [g_0(\bar{l}_0), \dots, g_K(\bar{l}_K)]$

**stochastic strategy:** non-deterministic  $g$

optimal strategy is where  $E[Y^g]$  is maximized (if high is good)

**Sequential Identifiability** sequential versions of

**exchangability:**  $Y^g \perp\!\!\!\perp A_k | \bar{A}_{k-1} \quad \forall g, k = 0, 1, \dots, K$

*conditional exchangeability:*

$$(Y^g, L_{k+1}^g) \perp\!\!\!\perp A_k | \bar{A}_{k-1} = g(\bar{L}_k), \bar{L}^k \quad \forall g, k = 0, 1, \dots, K$$

**positivity:**  $f_{\bar{A}_{k-1}, \bar{L}_k}(\bar{a}_{k-1}, \bar{l}_k) \neq 0 \Rightarrow$

$$f_{A_k | \bar{A}_{k-1}, \bar{L}_k}(a_k | \bar{a}_{k-1}, \bar{l}_k) > 0 \quad \forall (\bar{a}_{k-1}, \bar{l}_k)$$

**consistency:**

$$Y^{\bar{a}} = Y^{\bar{a}^*} \quad \text{if } \bar{a} = \bar{a}^*; \quad Y^{\bar{a}} = Y \quad \text{if } \bar{A} = \bar{a};$$

$$\bar{L}_k^{\bar{a}} = \bar{L}_k^{\bar{a}^*} \quad \text{if } \bar{a}_{k-1} = \bar{a}_{k-1}^*; \quad \bar{L}_k^{\bar{a}} = \bar{L}_k \quad \text{if } \bar{A}_{k-1} = \bar{a}_{k-1}$$

**generalized backdoor criterion** (static strategy): all backdoors into  $A_k$  (except through future treatments) are blocked  $\forall k$

**static sequential exchangeability for  $Y^{\bar{a}}$**

$$Y^{\bar{a}} \perp\!\!\!\perp A_k | \bar{A}_{k-1}, \bar{L}_k \quad \text{for } k = 0, 1, \dots, K$$

use SWIGs to visually check d-separation

**time-varying confounding**  $E[Y^{\bar{a}} | L_0] \neq E[Y | A = \bar{a}, L_0]$

**Treatment-Confounder Feedback**  $A_0 \rightarrow L_1 \rightarrow A_1$ :

an unmeasured  $U$  influencing  $L_1$  and  $Y$  turns  $L_1$  into a collider;

traditional adjustment (e.g. stratification) biased: use g-methods

**g-null test** sequential exchangeability & sharp null true  $\Rightarrow$

$Y^g = Y \quad \forall g \Rightarrow Y \perp\!\!\!\perp A_0 | L_0$  &  $Y \perp\!\!\!\perp A_1 | A_0, L_0, L_1$ ; therefore:

if last two independences don't hold, one assumption is violated

**g-null theorem:**  $E[Y^g] = E[Y]$ , if the two independences hold ( $\Rightarrow$  sharp null: only if strong faithfulness (no effect cancelling))

# References

*If no citation is given, the information is taken from the book (Hernán and Robins, 2020)*

Hernán, M. A. and Robins, J. M. (2020). *Causal inference: what if*. Boca Raton: Chapman & Hall/CRC.

Petersen, M. L. and van der Laan, M. J. (2014). Causal models and learning from data: integrating causal modeling and statistical estimation. *Epidemiology (Cambridge, Mass.)*, 25(3):418–426.

Schomaker, M., Luque-Fernandez, M. A., Leroy, V., and Davies, M.-A. (2019). Using longitudinal targeted maximum likelihood estimation in complex settings with dynamic interventions. *Statistics in medicine*, 38(24):4888–4911. ISBN: 0277-6715 Publisher: Wiley Online Library.

Van der Laan, M. J., Polley, E. C., and Hubbard, A. E. (2007). Super learner. *Statistical applications in genetics and molecular biology*, 6(1). Article 24.

Van der Laan, M. J., Rose, S., et al. (2011). *Targeted learning: causal inference for observational and experimental data*, volume 4. Springer.

Young, J. G., Cain, L. E., Robins, J. M., O'Reilly, E. J., and Hernán, M. A. (2011). Comparative effectiveness of dynamic treatment regimes: an application of the parametric g-formula. *Statistics in biosciences*, 3:119–143.