

**DECART Summer School 2018:** 

Causal Inference Module

**Special Topics** 

### Topic 1: Doubly Robust Estimator



#### **Problem Setup**

• Consider inverse propensity weighted (IPW) estimator for the mean of  $Y_i(1)$  in the study population  $\mu_t = E[Y(1)]$  as discussed in the earlier session, we have

$$\hat{\mu}_t = n^{-1} \sum_{i=1}^n \frac{A_i Y_i}{\hat{e}(X_i)}$$

where  $\hat{e}(X_i)$  is the propensity score estimated using logistic regression, we will write this as  $\hat{e}(X_i) = e(X_i; \hat{\beta})$  to reflect the fact that this is a parametric model.

- Why does this work?
  - By the law of large numbers, this should estimate the mean of a term in the sum with  $\hat{\beta}$  replaced by the quantity it estimates.



#### **Consistency of IPW Estimator**

• If  $e(X; \beta) = e(X)$ , the true propensity score

$$E\left[\frac{AY}{e(X)}\right] = E\left[\frac{AY(1)}{e(X)}\right] = E\left[E\left\{\frac{AY(1)}{e(X)}|Y(1),X\right\}\right]$$
$$= E\left\{\frac{Y(1)}{e(X)}E(A|Y(1),X)\right\} = E\left\{\frac{Y(1)}{e(X)}E(A|X)\right\}$$
$$= E\left\{\frac{Y(1)}{e(X)}e(X)\right\} = E(Y(1))$$



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It is worth noting:

- i) The consistency depends on the fact the model used to estimate e(X) is correctly specified.
- ii) The estimator only makes use of outcome data with A=1, ignores the information from subjects with A=0



### Improve efficiency through data augmentation -- AIPW Estimator

Modified estimator (Augmented Inverse Propensity Weighted estimator):

$$\hat{\mu}_{t} = n^{-1} \sum_{i=1}^{n} \left[ \frac{A_{i}}{e(X_{i}; \hat{\beta})} Y_{i} - \frac{\{A_{i} - e(X_{i}; \hat{\beta})\}}{e(X_{i}; \hat{\beta})} m_{t}(X_{i}; \hat{\alpha}) \right]$$

>  $e(X; \beta)$  is a postulated model for the true propensity score e(X) = E(A|X) (fitted by <u>logistic regression</u>)

>  $m_t(X; \alpha)$  is postulated model for the true regression E(Y|A=1,X) (fitted by <u>least square</u>)

By the law of large numbers, this should estimate the mean of a term in the sum with  $\hat{\beta}$  and  $\hat{\alpha}$  replaced by the quantity they estimate.



#### **Double Robustness**

$$\begin{split} E\left[\frac{A}{e(X;\beta)}Y - \frac{\{A - e(X;\beta)\}}{e(X;\beta)}m_t(X;\alpha)\right] \\ &= E\left[\frac{A}{e(X;\beta)}Y(1) - \frac{\{A - e(X;\beta)\}}{e(X;\beta)}m_t(X;\alpha)\right] \\ &= E\left[Y(1) + \frac{\{A - e(X;\beta)\}}{e(X;\beta)}\{Y(1) - m_t(X;\alpha)\}\right] \\ &= E[Y(1)] + E\left[\frac{\{A - e(X;\beta)\}}{e(X;\beta)}\{Y(1) - m_t(X;\alpha)\}\right] \\ \text{the second term} = &E\left\{Y(1) - m_t(X;\alpha)\}E\left[\frac{\{A - e(X;\beta)\}}{e(X;\beta)}|Y(1),X\right]\right\} \\ &= &E\left[\frac{\{A - e(X;\beta)\}}{e(X;\beta)}E[\{Y(1) - m_t(X;\alpha)\}|A,X]\right] \end{split}$$

- > If propensity model is correctly specified,  $e(X; \beta) = E(A|X)$ , then the second term = 0
- > If outcome model is correctly specified,  $m_t(X; \alpha) = E(Y(1)|X)$ , then the second term = 0

#### **Double Robustness**

- When either one model is correctly specified, we obtain an unbiased estimator.
- When both models are correctly specified, the resulting estimator is not only unbiased but also more efficient. (incorporate more information)
- Offers protection against mismodeling.



# Topic 2: Time Varying Confounding and Marginal Structural Model



#### **Marginal Structural Models**

 If the treatment can be quantified on at least an interval scale, we may consider models of the form:

(MSM1) 
$$E[Y(a)] = \beta_0 + \beta_1 a$$
, or  
(MSM2)  $E[Y(a)] = \beta_0 + \beta_1 a + \beta_2 a^2$ , or  
(MSM3)  $E[Y(a)] = f(a)$  for some functional form  $f(\cdot)$ 

- Under (MSM1), ATE contrasting treatment 1 to treatment 0 is  $\beta_1$  (essentially what we did in ipw example)
- Under (MSM3), ATE = f(1) f(0)
- These are called marginal structural models.
- <u>Structural</u> because the models are based on the counterfactual outcomes *Y*(*a*)
- Marginal because the models are based on the marginal distributions of each Y(a)



### Marginal Structural Models with Effect Modification

- *V* = baseline factors
- Models of the form E[Y(a)|V] = f(a,V) can be used to model modification of the causal effect of A by the factors in V
- For example,

$$E[(Y(a))|V] = \beta_0 + \beta_1 a + \beta_2 V + \beta_3 a \times V$$
 Causal effect is then  $\beta_1 + \beta_3 V$ 

Estimate model parameters by fitting regression model

$$E(Y|V,A) = \beta_0 + \beta_1 A + \beta_2 V + \beta_3 A \times V$$

using weighted regression with weights  $W^A$  or  $SW^A$ 

• Consider stabilized weights as  $SW^A(V) = f(A|V)/f(A|L,V)$ 

#### **Multiple Levels of Treatment**

- Assume treatment A has k levels, a = 1, 2, ..., k
- Could use <u>multinomial logistic regression</u> to estimate f(A|L) = Pr(A|L) for each A = a.
- Then define inverse probability of treatment weights for treatment A as:

$$W^A = 1/f(A|L)$$

- Stabilized weights:  $SW^A = f(A)/f(A|L)$
- Then use weighted regression to estimate parameters of *a* marginal structural model for the effect of the treatment;
- e.g.  $E[Y(a)] = \beta_0 + \beta_1 a$ , estimate  $\beta_0$  and  $\beta_1$  based on weighted regression of Y on A using weights  $W^A$  or  $SW^A$



#### **Evaluating the Causal Effect**

- For linear MSMs, this is equivalent to a 2-step procedure where we first obtain  $\widehat{E}(Y(a))$  as  $\frac{\sum_{i} 1_{[Ai=a]} Y_{i}}{\sum_{i} W_{i}^{a}}$  for each a, and then regress the  $\widehat{E}(Y(a))$  on a.
- Problem: Some treatment levels may be much more common than others, but the IPW weights give equal overall weight to each value of A in the regression
- Solution is to use stabilized weights:  $SW^A = f(A)/f(A|X)$
- The stabilized weights give more weight to treatment values a which are more common in the dataset



#### **Continuous Treatment**

• When the treatment is continuous (e.g. dosage):

$$Pr(A = a | X) = 0$$
 for all A and X.

cannot use standard propensity weighting approach.

- stabilized weights are OK:  $SW^A = f(A)/f(A|X)$  where f(A) and f(A|X) now represents the density of A and the conditional density of A given X.
- To estimate f(A|X), one regresses A on X, heavily dependent on the assumed conditional distribution of the error term, some choices in literature:
  - i) normal distribution
  - ii) truncated normal distribution
  - iii) t distribution
  - vi) quantile binning

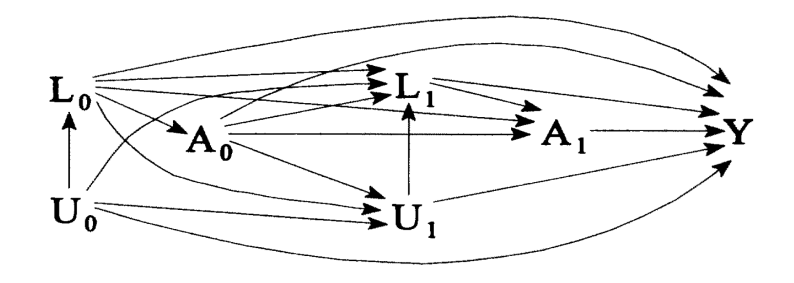


#### **Time-Varying Treatment: Notation**

- Consider a study with K followup visits, indexed by k = 0, 1, ..., K
- $A_k$ : treatment at kth visit
- L<sub>k</sub>: covariates measured at kth visit
- We assume the outcome Y is evaluated at visit K+1
- $\bar{A}_k = (A_0, A_1, ..., A_k)$ : the treatment history through the kth visit
- $\overline{L}_k = (L_1, L_2, ..., L_k)$ : the covariate history through the kth visit
- $Y(\bar{a}_K) = Y(a_1, a_2, ..., a_K)$ : the counterfactual outcome under the treatment history  $a_1, a_2, ..., a_K$ .



## Time Varying Treatment Framework with No Unmeasured Confounders



NUCA for this case:  $Y(\bar{a}_K) \perp A_k \mid \bar{L}_k, \bar{A}_{k-1}$ 

### Marginal Structural Models for Time Dependent Treatments

• For continuous Y:  $E(Y(\bar{a}_K)) = f(\bar{a}_K, V)$  If  $a_k = 0$  or 1, a simple MSM is

$$E(Y(\bar{a}_K)) = \beta_0 + \beta_1 cum(\bar{a}_K)$$
, where  $cum(\bar{a}_K) = \sum_k a_k$ .

When effect modification is of interest:

$$E(Y(\bar{a}_K)) = \beta_0 + \beta_1 cum(\bar{a}_K) + \beta_2 V + \beta_3 V \times cum(\bar{a}_K)$$

• For dichotomous Y,  $logit(Pr(Y(\bar{a}_K) = 1)) = f(\bar{a}_K, V)$ 



#### Weight for Longitudinal MSMs:

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

$$sw = \prod_{k=0}^{K} \frac{Pr(A_k | \overline{A}_{k-1}, V)}{Pr(A_k | \overline{A}_{k-1}, \overline{L}_k)}$$

- where  $\bar{A}_{-1}$  is defined to be 0, and V includes a set of baseline covariates including modifiers of the treatment effect, subset of baseline  $L_0$ .
- Use the pooled logistic regression to estimate these probabilities
- May also calculate stabilized censoring weight following similar procedure to account for drop-out, final weight is then the product of the treatment weight and the censoring weight.



## Why traditional regression methods fail in the time-varying case

- For the case of time-varying treatment, the confounders would also be time-varying. There may be treatment-confounder feedback.
- If time-varying treatments and confounders, and confounders are affected by prior treatment
- > Adjusting for confounder at time t masks (partially?) the effect of treatment prior to time t.
- > IP weighting controls confounding because they can handle treatment-confounder feedback



# Topic 3: Revisit - Yule-Simpson's Paradox



Table 1: Yule-Simpson's Paradox

Population			
	Survive	Die	Survive Rate
Treatment	20	20	50%
Control	16	24	40%
Male			
	Survive	Die	Survive Rate
Treatment	18	12	60%
Control	7	3	70%
Female			
	Survive	Die	Survive Rate
Treatment	2	8	20%
Control	9	21	30%

Example from Pearl 2000



#### Revisit - Yule-Simpson's Paradox

Notation:

Treatment Assignment T: 0 - control, 1 -treat

Outcome Y: 0 - die, 1 - survive

Covariate X: 0 – female, 1 – male.

The unadjusted treatment effect (ATE) is

$$\widehat{ATE}_{unadj} = \widehat{P}(Y = 1|T = 1) - \widehat{P}(Y = 1|T = 0) = 0.50 - 0.40 = +0.10$$

• The IPW (adjusted) estimator is

$$\widehat{ATE}_{adj} = \frac{\frac{1}{\widehat{P}(T=1|X=0)} \times 2 + \frac{1}{\widehat{P}(T=1|X=1)} \times 18}{80} - \frac{\frac{1}{\widehat{P}(T=0|X=0)} \times 9 + \frac{1}{\widehat{P}(T=0|X=1)} \times 7}{80}$$
$$= \frac{\frac{2}{10/40} + \frac{18}{30/40}}{80} - \frac{\frac{9}{30/40} + \frac{7}{10/40}}{80} = (0.40 - 0.50) = -0.10$$

Two estimates in opposite directions, whom should we trust?



No easy answers from "association" perspective

We have to think of "causality"

This is a good place where we can make use of the theories and tools we learnt from Causal Diagrams

Thoughts?

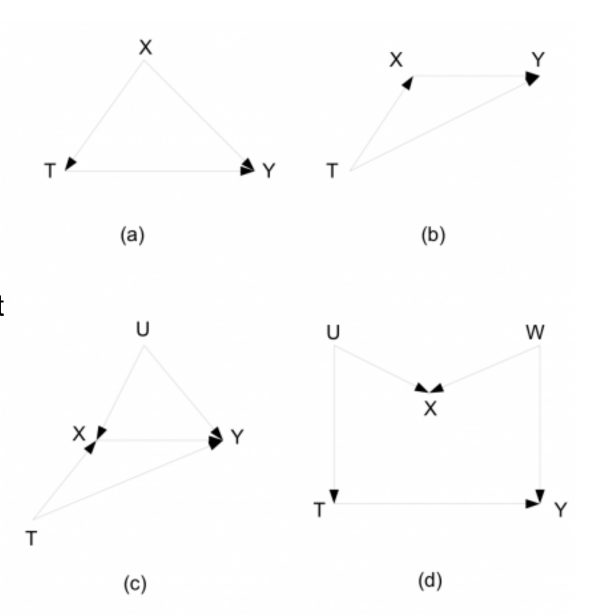


Figure 4 Simpson's paradox: possible DAGs



No easy answers from "association" perspective

We have to think of "causality"

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Think about back-door criterion, for structure (a), you may want to use the adjusted estimator while in structures (b) and (c), you may want the unadjusted estimator.

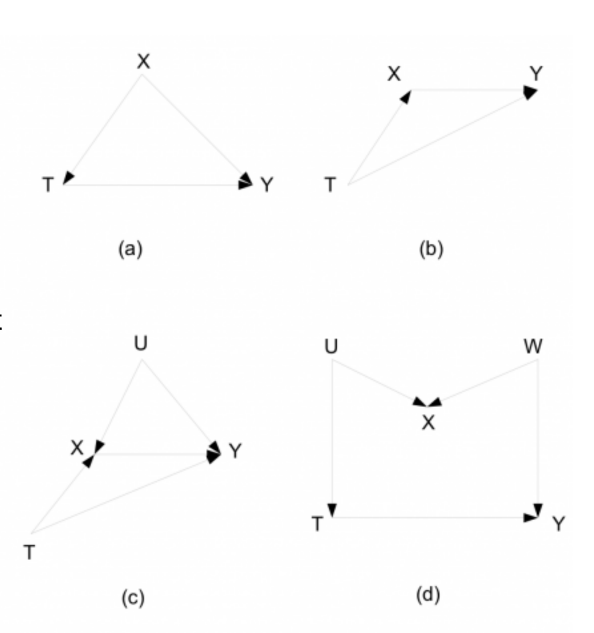
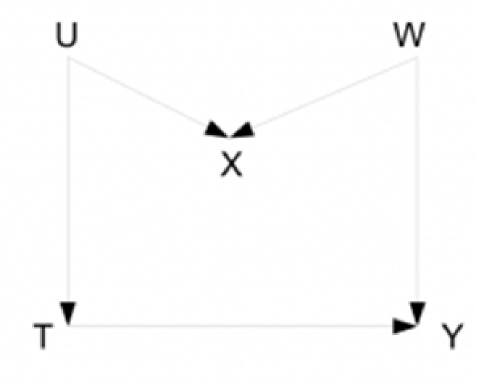


Figure 4 Simpson's paradox: possible DAGs



#### Bias Introduced under M-structure

- X is a pretreatment variable;
- There is a V-Structure (collider {U,X,W}), controlling X actually opens up back-door path T to Y (U and W are not independent any more!)
- Should we rely on the unadjusted estimator for causality?
- Some empirical studies suggest that the cost for not adjusting for the confounding (X) may be dominating in a lot of cases.



(d)

