# Sensitivity to Unobserved Confounding in Studies with Factor-structured Outcomes

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#### Causal Inference From Observational Data

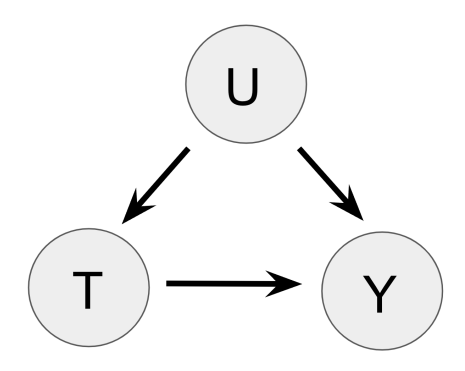
- ullet Consider a treatment T and outcome Y
- Interested in the population average treatment effect (PATE) of T on Y:

$$E[Y|do(T=t)] - E[Y|do(T=t')]$$

In general, the PATE is not the same as

$$E[Y|T=t]-E[Y|T=t']$$

#### Confounders



Need to control for U to consistently estimate the causal effect

## **Confounding bias**

- ullet Observed data regression of T on Y fails because the distribution of U varies in the two treatment arms
- We try to condition on as many observed confounders as possible to mitigate potential confounding bias
- Commonly assumed that there are "no unobserved confounders" (NUC) but this is unverifiable
- Sensitivity analysis is a tool for assessing the impacts of violations of this assumption

### A Motivating Example

**HEALTH** > NUTRITION & DIET

## 7 Science-Backed Health Benefits of Drinking Red Wine

Yep, moderate red wine consumption is healthy—and here's the proof.

By Ashley Zlatopolsky Updated on November 5, 2022

Fact checked by Emily Peterson

## A Motivating Example

The New York Times

## Even a Little Alcohol Can Harm Your Health

Recent research makes it clear that any amount of drinking can be detrimental. Here's why you may want to cut down on your consumption beyond Dry January.

#### The Effects of Light Alcohol Consumption

- Observational data from the National Health and Nutrition Examination Study (NHANES) on alcohol consumption.
- Light alcohol consumption is positively correlated with blood levels of HDL ("good cholesterol")
- Define "light alcohol consumption" as 1-2 alcoholic beverages per day
- Non-drinkers: self-reported drinking of one drink a week or less
- Control for age, gender and indicator for educational attainment

#### HDL and alcohol consumption

```
1 summary(lm(Y[, "HDL"] ~ drinking + X))
Call:
lm(formula = Y[, "HDL"] ~ drinking + X)
Residuals:
   Min 10 Median 30
                              Max
-5.0855 -0.6127 -0.0512 0.6389 4.2383
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
                  0.091105 2.476 0.013412 *
(Intercept) 0.225550
drinking 0.597399 0.091917 6.499 1.11e-10 ***
    0.006409 0.001452 4.415 1.09e-05 ***
Xage
0.689557 0.049426 13.951 < 2e-16 ***
Xeduc 0.194338 0.051161 3.799 0.000152 ***
```

What must be true for this correlation to be non-causal?

#### Blood mercury and alcohol consumption

```
1 summary(lm(Y[, "Methylmercury"] ~ drinking + X))
Call:
lm(formula = Y[, "Methylmercury"] ~ drinking + X)
Residuals:
   Min 10 Median 30
                               Max
-2.3570 -0.7363 -0.0728 0.6242 4.1127
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.442044 0.096385 4.586 4.91e-06 ***
drinking 0.364096 0.097244 3.744 0.000188 ***
Xage
    0.008186 0.001536 5.330 1.14e-07 ***
-0.062664 0.052290 -1.198 0.230966
Xeduc 0.269815 0.054126 4.985 6.95e-07 ***
```

But... no plausible causal mechanism in this case

#### **Residual Correlation**

```
hdl_fit <- lm(Y[, "HDL"] ~ drinking + X)
   mercury fit <- lm(Y[, "Methylmercury"] ~ drinking + X)</pre>
 4 cor.test(hdl fit$residuals, mercury fit$residuals)
   Pearson's product-moment correlation
data: hdl fit$residuals and mercury fit$residuals
t = 3.7569, df = 1437, p-value = 0.0001789
alternative hypothesis: true correlation is not equal to 0
95 percent confidence interval:
 0.04718758 0.14953581
sample estimates:
      cor
0.0986225
```

Residual correlation might be indicative of confounding bias

#### **Sensitivity Analysis**

- NUC unlikely to hold exactly. What then?
- Calibrate assumptions about confounding to explore range of causal effects that are plausible
- Robustness: quantify how "strong" confounding has to be to nullify causal effect estimates
- Well established methods for single outcome analyses

#### Multi-outcome Sensitivity Analysis

- If we measure multiple outcomes, is there prior knowledge that we can leverage to strengthen causal conclusions?
- What might residual correlation in multi-outcome models mean for potential for confounding?
- How do results change when we assume a priori that certain outcomes cannot be affected by treatments?
  - Null control outcomes (e.g. alcohol consumption should not increase mercury levels)

#### **Standard Assumptions**



Assumption (Latent Ignorability)

U and X block all backdoor paths between T and Y (Pearl 2009)



Assumption (Latent positivity)

f(T = t | U = u, X = x) > 0 for all u and x



**Assumption (SUTVA)** 

There are no hidden versions of the treatment and there is no interference between units

## Single-outcome Sensitivity Analysis

#### Result (Cinelli and Hazlett 2020)

Assume the outcome is linear in the treatment and confounders (no interactions). Then the squared omitted variable bias for the PATE is

$$ext{Bias}_{t_1,t_2}^2 \, \leq \, rac{(t_1-t_2)^2}{\sigma_{t|x}^2} \Bigg(rac{R_{T\sim U|X}^2}{1-R_{T\sim U|X}^2}\Bigg) \, R_{Y\sim U|T,X}^2 \, .$$

ullet  $R^2_{T\sim U|X}$ : partial fraction of treatment variance explained by confounders (given observed covariates)

ullet  $R^2_{Y\sim U|T,X}$ : partial fraction of outcome variance explained by confounders (given observed covariates and treatment)

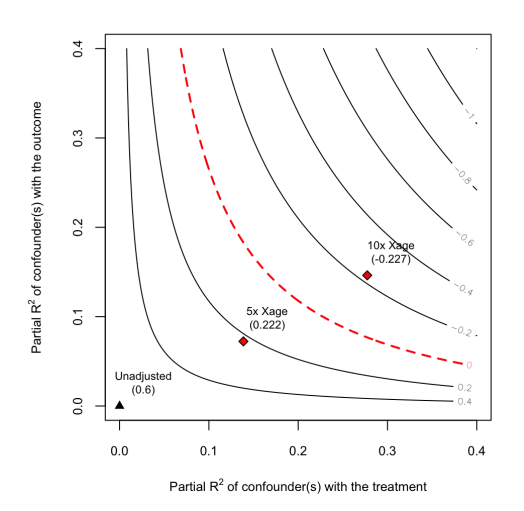
#### Robustness

- $\bullet$  How big do  $R^2_{T\sim U|X}$  and  $R^2_{Y\sim U|T,X}$  need to be to nullify the effect?
- ullet  $RV^1$  smallest value of  $R^2_{T\sim U|X}=R^2_{Y\sim U|T,X}$  needed to nullify effect (Cinelli and Hazlett 2020)
- ullet XRV smallest value of  $R^2_{T\sim U|X}$  assuming  $R^2_{Y\sim U|T,X}=1$  needed to nullify effect (Cinelli and Hazlett 2022)

### **Calibrating Sensitivity Parameters**

- ullet What values of  $R^2_{Y \sim U|T,X}$  and  $R^2_{T \sim U|X}$  might be reasonable?
- Can use observed covariates to generate benchmark values:
  - lacksquare Compute  $R^2_{T\sim X_j|X_{-j}}$  for all covariate  $X_j$
  - lacksquare Compute  $R^2_{Y\sim X_i|X_{-i},T}$  for all covariate  $X_j$
- Use domain knowledge to reason about most important confounders

#### Sensitivity of HDL Cholesterol Effect



From the sensemakr documentation (Cinelli, Ferwerda, and Hazlett 2020)

## Models with factor-structured residuals

Assume the **observed data** mean and covariance can be expressed as follows:

$$E[Y \mid T=t, X=x] = \check{g}(t,x) \ Cov(Y \mid T=t, X=x) = \Gamma\Gamma' + \Lambda,$$

ullet  $\Gamma$  are factor loading matrices,  $\Lambda$  is diagonal

#### A Structural Equation Model

- ullet U (m-vector) and X are possible causes for T (scalar) and Y (q-vector)
- ullet X are observed but U are not.

$$egin{aligned} U &= \epsilon_U \ T &= f_\epsilon(X,U) \ Y &= g(T,X) + \Gamma \Sigma_{u|t,x}^{-1/2} U + \epsilon_y, \end{aligned}$$

ullet This SEM is compatible the factor structured residuals,  $Cov(Y|T,X) = \Gamma\Gamma' + \Lambda$ 

#### A Structural Equation Model

$$egin{aligned} U &= \epsilon_U \ T &= f_\epsilon(X,U) \ Y &= g(T,X) + \Gamma \Sigma_{u|t,x}^{-1/2} U + \epsilon_y \end{aligned}$$

- ullet Confounding bias is  $\Gamma \Sigma_{u|t,x}^{-1/2} \mu_{u|t,x}$
- $\mu_{u|t,x}$  and  $\Sigma_{u|t,x}$  are the conditional mean and covariance of the unmeasured confounders
  - User specified sensitivity parameters

#### **A Sensitivity Specification**

• Interpretable specification for  $\mu_{u|t,x}$  and  $\Sigma_{u|t,x}$  parameterized by a single m-vector,  $\rho$ :

$$\mu_{u|t,x} = rac{
ho}{\sigma_{t|x}^2}ig(t-\mu_{t|x}ig)\,,$$

$$\Sigma_{u|t,x} = I_m - rac{
ho 
ho'}{\sigma_{t|x}^2},$$

- ullet ho is the partial correlation vector between T and U
- Define  $0 \le R^2_{T \sim U|X} := \frac{||\rho||_2^2}{\sigma_{t|x}^2} < 1$  to be the squared norm of the partial correlation between T and U given X

#### Multi-Outcome Assumptions

Assumption (Homoscedasticity)

Cov(Y|T=t,X=x) is invariant to t and x. Implies factor loadings,  $\Gamma$ , are invariant to t and x

Assumption (Factor confounding)

The factor loadings,  $\Gamma$ , are identifiable (up to rotation) and reflect all potential confounders. (Anderson and Rubin 1956)

To identify factor loadings,  $\Gamma$ ,  $(q-m)^2-q-m\geq 0$  and each confounder must influence at least three outcomes

#### **Bounding the Omitted Variable Bias**

#### Theorem (Bounding the bias for outcome $Y_j$ )

Given the structural equation model, sensitivity specification and given assumptions, the squared omitted variable bias for the PATE of outcome  $Y_j$  is bounded by

$$ext{Bias}_{j}^{2} \, \leq \, rac{(t_{1} - t_{2})^{2}}{\sigma_{t|x}^{2}} \Bigg(rac{R_{T \sim U|X}^{2}}{1 - R_{T \sim U|X}^{2}}\Bigg) \parallel \Gamma_{j} \parallel_{2}^{2}.$$

- ullet The bound on the bias for outcome j is proportional to the norm of the factor loadings for that outcome
- ullet A single sensitivity parameter,  $R^2_{T\sim U|X}$ , shared across all outcomes

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- ullet Assume we have null control outcomes,  ${\mathcal C}$
- $\check{\tau}$  are the vector of PATEs under NUC
- ullet  $\Gamma_{\mathcal{C}}$  are the factor loadings for the null control outcomes,  $\mathcal{C}$
- Need at least  $R^2_{T\sim U|X} \geq R^2_{min}$  of the treatment variance to be due to confounding to nullify the null controls

#### Theorem (Bias with Null Control Outcomes)

Assume the previous structural equation model and sensitivity specification. Then the squared omitted variable bias for the PATE of outcome  $Y_j$  is bounded by

$$ext{Bias}_j \in \left[\Gamma_j \Gamma_\mathcal{C}^\dagger \check{ au}_\mathcal{C} \pm \parallel \Gamma_j P_{\Gamma_\mathcal{C}}^\perp \parallel_2 \sqrt{rac{1}{\sigma_{t|x}^2} igg(rac{R_{T \sim U|X}^2}{1 - R_{T \sim U|X}^2} - rac{R_{min}^2}{1 - R_{min}^2}igg)}
ight],$$

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ight],$$

## Robustness under Factor Confounding

- $RV_j^\Gamma$  smallest value of  $R^2_{T\sim U|X}$  needed to nullify the effect for outcome j under factor confounding
- ullet  $RV_j^\Gamma$  can be smaller or larger than  $RV^1$
- ullet  $RV_i^\Gamma \geq XRV$  by definition
- $RV_{j,NC}^{\Gamma}$  smallest value of  $R_{T\sim U|X}^2$  needed to nullify the effect for outcome j and the assumed null controls

### **Simulation Study**

Gaussian data generating process

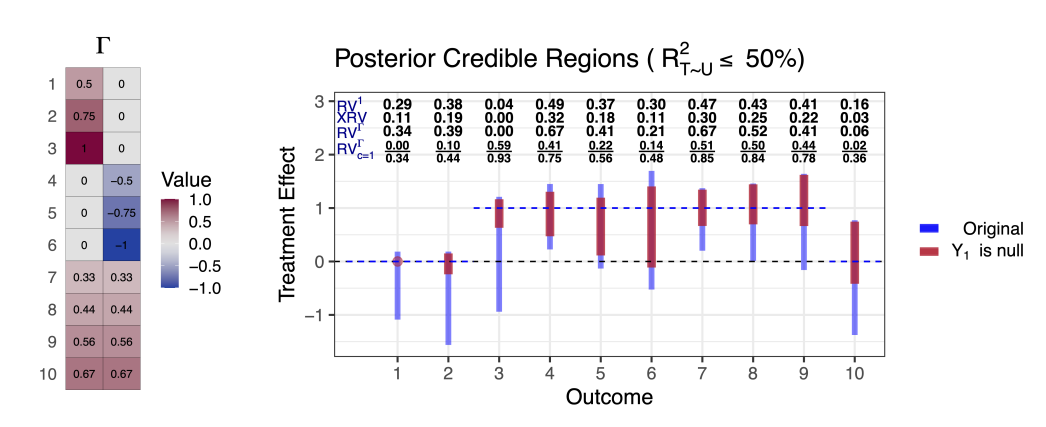
$$T = eta'U + \epsilon_T \ Y_j = au_j T + \Gamma' \Sigma_{u|t}^{-1/2} U + \epsilon_y$$

- ullet  $R^2_{T\sim U|X}=0.5$  from m=2 unmeasured confounders
- ullet  $au_j=0$  for  $Y_1,Y_2$  and  $Y_{10}$
- $au_j = 1$  for all outher outcomes

### **Simulation Study**

- Fit a Bayesian linear regression on the 10 outcomes given then treatment
- Assume a residual covariance with a rank-two factor structure
- ullet Plot ignorance regions assuming  $R^2_{T\sim U} \leq 0.5$
- ullet Plot ignorance regions assuming  $R^2_{T\sim U} \leq 0.5$  and  $Y_1$  is null

## **Simulation Study**



#### The effects of light drinking

- Measure ten different outcomes from blood samples:
  - natural: HDL, LDL, triglycerides, potassium, iron, sodium, glucose
  - environmental toxicants: mercury, lead, cadmium.
- Measured confounders: age, gender and indicator for highest educational attainment
- Residual correlation in the outcomes might be indicative of additional confounding bias

## Aside: reparametrizing $R^2_{T\sim U|X}$

- ullet  $R^2_{T\sim U|X}$  is unnatural for binary treatments
- ullet  $\Lambda$ -parameterization  $\leftrightarrow$   $R^2_{T\sim U|X}$ -parameterization

Fix a  $\Lambda_{\alpha}$  such that

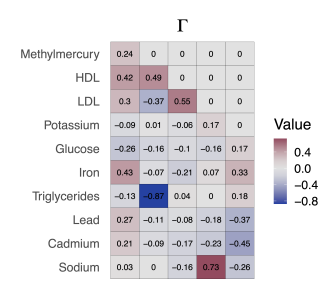
$$Pr\left(\Lambda_lpha^{-1} \leq rac{e_0(X,U)/(1-e_0(X,U))}{e(X)/(1-e(X))} \leq \Lambda_lpha
ight) = 1-lpha$$

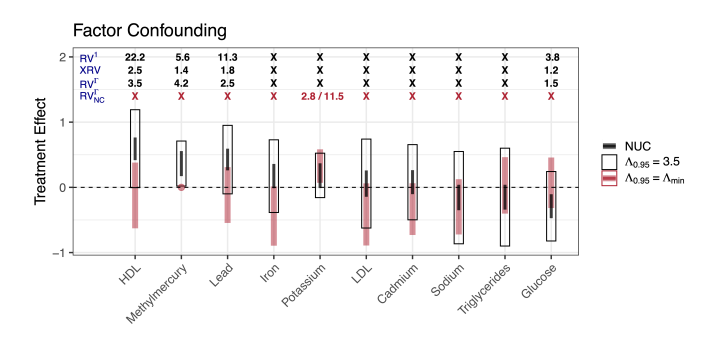
• Related to the marginal sensitivity model (Tan 2006)

#### **Benchmark Values**

- Use age, gender and an indicator of educational attainment to benchmark
- $\frac{1}{3.5} \leq \mathrm{Odds}(X)/\mathrm{Odds}(X_{-age}) \leq 3.5$  for 95% of observed values
- $\bullet$  For gender and education indicators the odds change was between  $\frac{1}{1.5}$  and 1.5
- Assume light drinking has no effect on methylmercury levels

#### Results: NHANES alcohol study





### **Takeaways**

- Prior knowledge unique to the multi-outcome setting can help inform assumptions about confounding
- Sharper sensitivity analysis, when assumptions hold
- Negative control assumptions can potentially provide strong evidence for or against robustness

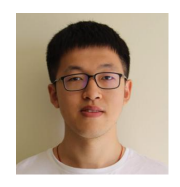
#### **Future directions**

- Identification with multiple treatments multiple outcomes
  - Collaboration on effects of pollutants on multiple heath outcomes
- Sensitivity analysis for more general models / forms of dependence.

#### References

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#### Thanks!







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Preprint: Sensitivity to
Unobserved Confounding in
Studies with Factorstructured Outcomes
https://arxiv.org/abs/2208.06552