# Leveraging the causal effects of stochastic interventions to evaluate vaccine efficacy in two-phase trials

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SER: "Methods for the thorny challenges of real studies"



#### The burden of HIV-1

- The HIV-1 epidemic the facts:
  - now in its fourth decade,
  - 2.5 million new infections occurring annually worldwide,
  - new infections outpace patients starting antiretroviral therapy.
- Most efficacious preventive vaccine: 31% reduction rate.
- Question: To what extent can HIV-1 vaccines be improved by modulating immunogenic CD4+/CD8+ response profiles?

#### HVTN 505 trial examined new antibody boost vaccines

- HIV Vaccine Trials Network's (HVTN) 505 vaccine efficacy;
   randomized controlled trial, n = 2504 (Hammer et al. 2013).
- Question: How would HIV-1 infection risk in week 28 have changed had immunogenic response (due to vaccine) differed?
- Immunogenic response profiles only available for second-stage sample of n=189 (Janes et al. 2017) due to cost limitations.
- Two-phased sampling mechanism: 100% inclusion rate if HIV-1 positive in week 28; based on matching otherwise.

#### Two-phase sampling censors the complete data structure

- Complete (<u>unobserved</u>) data  $X = (W, A, Y) \sim P_0^X \in \mathcal{M}^X$ , as per the full HVTN 505 trial cohort (Hammer et al. 2013):
  - W (baseline covariates): sex, age, BMI, behavioral HIV risk,
  - A (exposure): immune response profile for CD4+ and CD8+,
  - Y (outcome of interest): HIV-1 infection status at week 28.
- Observed data  $O = (C, CX) = (W, C, CA, Y); C \in \{0, 1\}$  is an indicator for inclusion in the second-stage sample.

#### **NPSEM** with static interventions

 Use a nonparametric structural equation model (NPSEM) to describe the generation of X (Pearl 2009), specifically

$$W = f_W(U_W); A = f_A(W, U_A); Y = f_Y(A, W, U_Y)$$

- Implies a model for the distribution of counterfactual random variables generated by interventions on the process.
- A static intervention replaces f<sub>A</sub> with a specific value a in its conditional support A | W.
- This requires specifying a particular value of the exposure under which to evaluate the outcome a priori.

#### **NPSEM** with stochastic interventions

- Stochastic interventions modify the value A would naturally assume by drawing from a modified exposure distribution.
- Consider the post-intervention value  $A^* \sim G^*(\cdot \mid W)$ ; static interventions are a special case (degenerate distribution).
- Such an intervention generates a counterfactual random variable  $Y_{G^*} := f_Y(A^*, W, U_Y)$ , with distribution  $P_0^{\delta}$ , .
- We aim to estimate  $\psi_{0,\delta} := \mathbb{E}_{P_0^\delta}\{Y_{G^\star}\}$ , the counterfactual mean under the post-intervention exposure distribution  $G^\star$ .

#### Stochastic interventions for the causal effects of shifts

Díaz and van der Laan (2012; 2018)'s stochastic interventions

$$d(a, w) = \begin{cases} a + \delta, & a + \delta < u(w) & \text{(if plausible)} \\ a, & a + \delta \ge u(w) & \text{(otherwise)} \end{cases}$$

- Our estimand is  $\psi_{0,d} := \mathbb{E}_{P_0^d} \{ Y_{d(A,W)} \}$ , mean of  $Y_{d(A,W)}$ .
- Statistical target parameter is  $\Psi(P_0^X) = \mathbb{E}_{P_0^X} \overline{Q}(d(A, W), W)$ , counterfactual mean of the *shifted* outcome mechanism.
- For HVTN 505,  $\psi_{0,d}$  is the counterfactual risk of HIV-1 infection, had the observed value of the immune response been altered under the rule d(A, W) defining  $G^*(\cdot \mid W)$ .

#### One-step and targeted minimum loss estimators

One-step estimator relies on a bias correction:

$$\Psi_n^+ = \frac{1}{n} \sum_{i=1}^n \overline{Q}_n(d(A_i, W_i), W_i) + D^*(P_0^X)(X_i).$$

■ TML estimator constructs  $\overline{Q}_n^{\star}$  via a (logistic) tilting model

$$\Psi_n^{\star} := \Psi(P_n^{\star}) = \frac{1}{n} \sum_{i=1}^n \overline{Q}_n^{\star}(d(A_i, W_i), W_i).$$

In this case, the efficient influence function (EIF) is

$$D^*(P_0^X)(x) = H(a,w)(y-\overline{Q}(a,w)) + \overline{Q}(d(a,w),w) - \Psi(P_0^X),$$
 where the auxiliary ("clever") covariate is

$$H(a, w) = \mathbb{I}(a + \delta < u(w)) \frac{g_0(a - \delta \mid w)}{g_0(a \mid w)} + \mathbb{I}(a + \delta \geq u(w)).$$

# Augmented estimators for two-phase sampling designs

- Rose and van der Laan (2011) introduce the IPCW-TMLE, to be used when observed data is subject to two-phase sampling.
- Initial proposal: correct for two-phase sampling by using a loss function with inverse probability of censoring weights:

$$\mathcal{L}(P_0^X)(O) = \frac{C}{\pi_0(Y, W)} \mathcal{L}^F(P_0^X)(X)$$

- When the sampling mechanism  $\pi_0(Y, W)$  can be estimated by a parametric form, this procedure yields an efficient estimator.
- However, when machine learning is used, this is insufficient.

#### Efficient estimation under two-phase sampling

- When the sampling mechanism is not known by design, it is best to employ a flexible (ML) estimator of  $\pi_0(Y, W)$ .
- Then, the IPCW augmentation must be applied to the EIF:

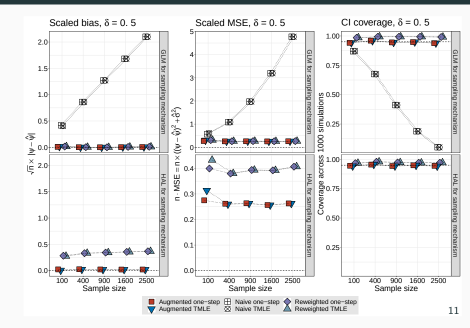
$$D(P_0^X)(o) = \frac{c}{\pi_0(y, w)} D^F(P_0^X)(x) - \left(1 - \frac{c}{\pi_0(y, w)}\right) \cdot \mathbb{E}(D^F(P_0^X)(x) \mid C = 1, Y = y, W = w),$$

- Expresses observed data EIF  $D^F(P_0^X)(o)$  in terms of full data EIF  $D^F(P_0^X)(x)$ ; inclusion of second term induces efficiency.
- The expectation of the full data EIF  $D^F(P_0^X)(x)$ , taken only over units selected by the sampling mechanism (i.e., C=1).

#### **Emergent property: multiple robustness**

- A unique multiple robustness property combinations of  $(g_0, \overline{Q}_0) \times (\pi_0, \mathbb{E}(D^F(P_0^X)(x) \mid C = 1, Y, W)).$
- We now have a semiparametric-efficient and robust procedure for assessing the effect of the intervention  $d(a, w) = a + \delta$ .
- Due to the construction of the IPCW-TMLE, the resultant estimator is robust and efficient under two-phase sampling.
- This allows us to assess how posited shifts in the assayed immune responses would have affected HIV-1 infection risk.

### Finding the "best" efficient estimator: relative performance



# Helping to fight the HIV-1 epidemic

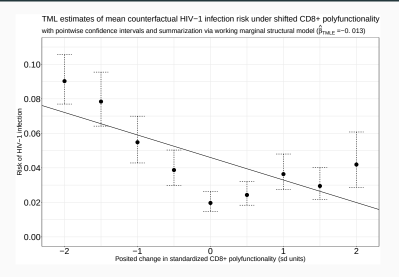


Figure 2: Analysis of HIV-1 risk as a function of CD8+ immunogenicity, using R package txshift (https://github.com/nhejazi/txshift.)

#### Big picture takeaways

- Vaccine efficacy evaluation helps to develop enhanced vaccines better informed by biological properties of the target disease.
- HIV-1 vaccines modulate immunogenic response profiles as part of their mechanism for lowering HIV-1 infection risk.
- Stochastic interventions constitute a flexible framework for considering realistic treatment/intervention policies.
- Large-scale (vaccine) trials often use two-phase designs need to (carefully!) accommodate for sampling complications.
- We've developed robust, open source statistical software for assessing stochastic interventions in observational studies.

#### Thank you

For more, check out https://arxiv.org/abs/TODO

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Appendix

# From the causal to the statistical target parameter

#### **Assumption 1:** *Consistency*

$$Y_i^{d(a_i,w_i)} = Y_i$$
 in the event  $A_i = d(a_i, w_i)$ , for  $i = 1, \ldots, n$ 

#### **Assumption 2: SUTVA**

 $Y_i^{d(a_i,w_i)}$  does not depend on  $d(a_j,w_j)$  for  $i=1,\ldots,n$  and  $j\neq i$ , or lack of interference (Rubin 1978; 1980)

#### Assumption 3: Strong ignorability

$$A_i \perp \!\!\! \perp Y_i^{d(a_i,w_i)} \mid w_i$$
, for  $i=1,\ldots,n$ 

### From the causal to the statistical target parameter

# Assumption 4: Positivity (or overlap)

 $a_i \in \mathcal{A} \implies d(a_i, w_i) \in \mathcal{A} \text{ for all } w \in \mathcal{W}, \text{ where } \mathcal{A} \text{ denotes}$  the support of A conditional on  $W = w_i$  for all i = 1, ... n

- This positivity assumption is not quite the same as that required for categorical interventions.
- In particular, we do not require that the intervention density place mass across all strata defined by W.
- Rather, we merely require the post-intervention quantity be seen in the observed data for given  $a_i \in A$  and  $w_i \in W$ .

# Literature: Díaz and van der Laan (2012)

- Proposal: Evaluate outcome under an altered intervention distribution e.g.,  $P_{\delta}(g_0)(A = a \mid W) = g_0(a \delta(W) \mid W)$ .
- Identification conditions for a statistical parameter of the counterfactual outcome  $\psi_{0,d}$  under such an intervention.
- Show that the causal quantity of interest  $\mathbb{E}_0\{Y_{d(A,W)}\}$  is identified by a functional of the distribution of X:

$$\psi_{0,d} = \int_{\mathcal{W}} \int_{\mathcal{A}} \mathbb{E}_{P_0^{\mathsf{X}}} \{ Y \mid A = d(a, w), W = w \} \cdot$$

$$q_{0,A}^{\mathsf{X}} (a \mid W = w) \cdot q_{0,W}^{\mathsf{X}} (w) d\mu(a) d\nu(w)$$

 Provides a derivation based on the efficient influence function (EIF) with respect to the nonparametric model M.

### Literature: Haneuse and Rotnitzky (2013)

- Proposal: Characterization of stochastic interventions as modified treatment policies (MTPs).
- Assumption of piecewise smooth invertibility allows for the intervention distribution of any MTP to be recovered:

$$g_{0,\delta}(a \mid w) = \sum_{j=1}^{J(w)} I_{\delta,j}\{h_j(a, w), w\}g_0\{h_j(a, w) \mid w\}h_j'(a, w)$$

- Such intervention policies account for the natural value of the intervention A directly yet are interpretable as the imposition of an altered intervention mechanism.
- Identification conditions for assessing the parameter of interest under such interventions appear technically complex (at first).

### Literature: Young et al. (2014)

- Establishes equivalence between g-formula when proposed intervention depends on natural value and when it does not.
- This equivalence leads to a sufficient positivity condition for estimating the counterfactual mean under MTPs via the same statistical functional studied in Díaz and van der Laan (2012).
- Extends earlier identification results, providing a way to use the same statistical functional to assess  $\mathbb{E}Y_{d(A,W)}$  or  $\mathbb{E}Y_{d(W)}$ .
- The authors also consider limits on implementing shifts d(A, W), and address working in a longitudinal setting.

# Literature: Díaz and van der Laan (2018)

- Builds on the original proposal, accommodating MTP-type shifts d(A, W) proposed after their earlier work.
- To protect against positivity violations, considers a specific shifting mechanism:

$$d(a, w) = \begin{cases} a + \delta, & a + \delta < u(w) \\ a, & \text{otherwise} \end{cases}$$

- Proposes an improved "1-TMLE" algorithm, with a single auxiliary covariate for constructing the TML estimator.
- Our (first) contribution: implementation of this algorithm.

### Nonparametric conditional density estimation

- To compute the auxiliary covariate H(a, w), we need to estimate conditional densities  $g(A \mid W)$  and  $g(A \delta \mid W)$ .
- There is a rich literature on density estimation, we follow the approach proposed in Díaz and van der Laan (2011).
- To build a conditional density estimator, consider

$$g_{n,\alpha}(a \mid W) = \frac{\mathbb{P}(A \in [\alpha_{t-1}, \alpha_t) \mid W)}{\alpha_t - \alpha_{t-1}},$$

for  $\alpha_{t-1} \leq a < \alpha_t$ .

- This is a classification problem, where we estimate the probability that a value of A falls in a bin  $[\alpha_{t-1}, \alpha_t)$ .
- The choice of the tuning parameter *t* corresponds roughly to the choice of bandwidth in classical kernel density estimation.

#### Nonparametric conditional density estimation

- Díaz and van der Laan (2011) propose a re-formulation of this classification approach as a set of hazard regressions.
- To effectively employ this proposed re-formulation, consider

$$\mathbb{P}(A \in [\alpha_{t-1}, \alpha_t) \mid W) = \mathbb{P}(A \in [\alpha_{t-1}, \alpha_t) \mid A \ge \alpha_{t-1}, W) \times \Pi_{j=1}^{t-1} \{1 - \mathbb{P}(A \in [\alpha_{j-1}, \alpha_j) \mid A \ge \alpha_{j-1}, W)\}$$

- The likelihood of this model may be expressed to correspond to the likelihood of a binary variable in a data set expressed via a long-form repeated measures structure.
- Specifically, the observation of  $X_i$  is repeated as many times as intervals  $[\alpha_{t-1}, \alpha_t)$  are before the interval to which  $A_i$  belongs, and the binary variables indicating  $A_i \in [\alpha_{t-1}, \alpha_t)$  are recorded.

# Density estimation with the Super Learner algorithm

- To estimate  $g(A \mid W)$  and  $g(A \delta \mid W)$ , use a pooled hazard regression, spanning the support of A.
- We rely on the Super Learner algorithm of van der Laan et al. (2007) to build an ensemble learner that optimally weights each of the proposed regressions, using cross-validation (CV).
- The Super Learner algorithm uses V-fold CV to train each proposed regression model, weighting each by the inverse of its average risk across all V holdout sets.
- By using a library of regression estimators, we invoke the result of van der Laan et al. (2004), who prove this likelihood-based cross-validated estimator to be asymptotically optimal.

### Key properties of TML estimators

Asymptotic linearity:

$$\Psi(P_n^*) - \Psi(P_0^X) = \frac{1}{n} \sum_{i=1}^n D(P_0^X)(X_i) + o_P\left(\frac{1}{\sqrt{n}}\right)$$

Gaussian limiting distribution:

$$\sqrt{\textit{n}}(\Psi(\textit{P}^{\star}_\textit{n}) - \Psi(\textit{P}^{X}_\textit{0})) \rightarrow \textit{N}(0, \textit{Var}(\textit{D}(\textit{P}^{X}_\textit{0})(\textit{X})))$$

Statistical inference:

Wald-type confidence interval : 
$$\Psi(P_n^\star) \pm z_{1-\frac{\alpha}{2}} \cdot \frac{\sigma_n}{\sqrt{n}}$$
,

where  $\sigma_n^2$  is computed directly via  $\sigma_n^2 = \frac{1}{n} \sum_{i=1}^n D^2(\cdot)(X_i)$ .

#### Algorithm for TML estimation

- 1. Construct initial estimators  $g_n$  of  $g_0(A, W)$  and  $Q_n$  of  $\overline{Q}_0(A, W)$ , perhaps using data-adaptive regression techniques.
- 2. For each observation *i*, compute an estimate  $H_n(a_i, w_i)$  of the auxiliary covariate  $H(a_i, w_i)$ .
- 3. Estimate the parameter  $\epsilon$  in the logistic regression model

$$\operatorname{logit} \overline{Q}_{\epsilon,n}(a,w) = \operatorname{logit} \overline{Q}_n(a,w) + \epsilon H_n(a,w),$$

or an alternative regression model incorporating weights.

4. Compute TML estimator  $\Psi_n$  of the target parameter, defining update  $\overline{Q}_n^*$  of the initial estimate  $\overline{Q}_{n,\epsilon_n}$ :

$$\Psi_n = \Psi(P_n^*) = \frac{1}{n} \sum_{i=1}^n \overline{Q}_n^*(d(A_i, W_i), W_i).$$

# Algorithm for IPCW-TML estimation

- 1. Using all observed units (X), estimate sampling mechanism  $\pi(Y, W)$ , perhaps using data-adaptive regression methods.
- 2. Using only observed units in the second-stage sample C=1, construct initial estimators  $g_n(A, W)$  and  $\overline{Q}_n(A, W)$ , weighting by the sampling mechanism estimate  $\pi_n(Y, W)$ .
- 3. With the approach described for the full data case, compute  $H_n(a_i, w_i)$ , and fluctuate submodel via logistic regression.
- 4. Compute IPCW-TML estimator  $\Psi_n$  of the target parameter, by solving the IPCW-augmented EIF estimating equation.
- 5. Iteratively update estimated sampling weights  $\pi_n(Y, W)$  and IPCW-augmented EIF, updating TML estimate in each iteration, until  $\frac{1}{n} \sum_{i=1}^{n} \mathsf{EIF}_i < \frac{1}{n}$ .

# A linear modeling perspective

- Briefly consider a simple data structure: X = (Y, A); we seek to model the outcome Y as a function of A.
- To posit a linear model, consider  $Y_i = \beta_0 + \beta_1 A_i + \epsilon_i$ , with error  $\epsilon_i \sim N(0,1)$ .
- Letting  $\delta$  be a change in A,  $Y_{A+\delta} Y_A$  may be expressed

$$\mathbb{E}Y_{A+\delta} - \mathbb{E}Y_A = [\beta_0 + \beta_1(\mathbb{E}A + \delta)] - [\beta_0 + \beta_1(\mathbb{E}A)]$$
$$= \beta_0 - \beta_0 + \beta_1\mathbb{E}A - \beta_1\mathbb{E}A + \beta_1\delta$$
$$= \beta_1\delta$$

• Thus, a *unit shift* in A (i.e.,  $\delta=1$ ) may be seen as inducing a change in the difference in outcomes of magnitude  $\beta_1$ .

#### A causal inference perspective

- Consider a data structure:  $(Y_a, a \in A)$ .
- To posit a linear model, let  $Y_a = \beta_0 + \beta_1 a + \epsilon_a$  for  $a \in \mathcal{A}$ , with error  $\epsilon_a \sim N(0, \sigma_a^2) \ \forall a \in \mathcal{A}$ .
- For the counterfactual outcomes  $(Y_{a'+\delta}, Y_{a'})$ , their difference,  $Y_{a'+\delta} Y_{a'}$ , for some  $a' \in \mathcal{A}$ , may be expressed

$$\mathbb{E}Y_{a'+\delta} - \mathbb{E}Y_{a'} = [\beta_0 + \beta_1(a'+\delta) + \mathbb{E}\epsilon_{a'+\delta}] - [\beta_0 + \beta_1a' + \mathbb{E}\epsilon_{a'}]$$
$$= \beta_1\delta$$

■ Thus, a *unit shift* for  $a' \in A$  (i.e.,  $\delta = 1$ ) may be seen as inducing a change in the difference in the counterfactual outcomes of magnitude  $\beta_1$ .

#### Slope in a semiparametric model

• Consider the stochastic intervention  $g^*(\cdot \mid W)$ :

$$\mathbb{E}Y_{g^*} = \int_W \int_a \mathbb{E}(Y \mid A = a, W) g(a - \delta \mid W) \cdot da \cdot dP_0(W)$$

$$= \int_W \int_z \mathbb{E}(Y \mid A = z + \delta, W) g(z \mid W) \cdot dz \cdot dP_0(W),$$
defining the change of variable  $z = a - \delta$ .

• For a semiparametric model,  $\mathbb{E}(Y \mid A = z, W) = \beta z + \theta(W)$ :

$$\mathbb{E}Y_{g^*} - \mathbb{E}Y = \int_{W} \int_{z} \left[ \mathbb{E}(Y \mid A = z + \delta, W) - \mathbb{E}(Y \mid A = z, W) \right]$$
$$g(z \mid W) \cdot dz \cdot dP_{0}(W)$$
$$= \left[ \beta(z + \delta) + \theta(W) \right] - \left[ \beta z + \theta(W) \right]$$
$$= \beta \delta$$

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