# Regression Analysis of Dependent Binary Data for Estimating Disease Etiology from Case-Control Studies

("Small-Area Estimation" of Disease Etiology)

#### 7henke Wu

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- Motivation for this talk: PEFs may vary by season, a child's

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- Continuous covariates: the first  $p_1$  and  $q_1$  elements of  $X_i$  and  $W_i$ , respectively.

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- Model : J = 7: noisy presence/absence of 2 bacteria and 5 viruses in the nose
  - Causes: seven single-pathogen causes plus an "Not Specified" (NoS) cause; So L = J + 1
  - X<sub>i</sub>: enrollment date, age (< or > 1 year), disease severity for cases (severe or very severe), HIV status (+/-)
  - **W**<sub>i</sub>: **X**<sub>i</sub> minus "disease severity".

## PERCH Data: Sparsely-Populated Strata©

Table: The observed count (frequency) of cases and controls by age, disease severity and HIV status (1: yes; 0: no). The marginal fractions among cases and controls for each covariate are shown at the bottom. Regression results will be shown for the first two strata.

$age \geq 1$	very severe (VS)	HIV positive	# cases (%)	# controls (%)
	(case-only)		total: 524 (100)	total: 964 (100)
0	0	0	208 (39.7)	545 (56.5)
1	0	0	72 (13.7)	278 (28.8)
0	1	0	116 (22.1)	0
1	1	0	33 (6.3)	0
0	0	1	37 (7.1)	85 (8.8)
1	0	1	24 (4.5)	51 (5.3)
0	1	1	25 (4.8)	0
1	1	1	3 (0.6)	0
case: 25.2%	34.5%	17.0%		
control: 34.3%	-	14.1%		

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# Current Methods (per covariate stratum)

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- Fitted in a Bayesian framework; can also estimate the posterior probability of the cause of disease for an **individual** case given her measurements.
- Related to restricted latent class models (RLCM, Xu, 2017, AOS); Major differences: multiple sets of responses probabilities ("nested"); use control data ("partially-latent")

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- Gap 1a Unstable PEF estimates due to sparsely-populated strata.
- Gap 1b Informative TPR priors are often elicited for a case population and rarely for each stratum; Reusing independent prior distributions of the TPRs across all the strata will lead to overly-optimistic posterior uncertainty in  $\pi^*$ , hampering policy decisions.



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- (a) incorporates controls to estimate the PEFs  $(\pi)$ ,
- (b) specifies parsimonious functional dependence of  $\pi$  upon covariates such as additivity, and
- (c) correctly assesses the posterior uncertainty of the PEF functions and the overall PEFs  $\pi^*$  by applying the TPR priors just once.

## Okay, what is npLCM?

# Quick Technical Review: Nested Partially Latent Class Models (npLCM)

For simplicity, we assume "single-pathogen causes"

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$$L = L_1 \cdot L_0 = \left\{ \prod_{e \not e \ M'} \sum_{i=1}^{L} \pi_{\ell} \cdot \textbf{\textit{P}}_{1\ell}(\textbf{\textit{M}}_i; \Theta, \Psi, \eta) \right\} \times \prod_{i': Y_{i'} = 0} \textbf{\textit{P}}_{0}(\textbf{\textit{M}}_{i'}; \Psi, \nu)$$

$$\text{Zhenke Wu}(\textit{zhenkewu@umic}(e \not e \ M') = 1 \quad \ell = 2019 \ \text{ICSA China Conference, Tianjin}$$

Setting  $\eta_1=1$  and  $\nu_1=1$ 

Control model for multivariate binary data  $\{M_i : where Y_i = 0\}$ :

- 1.  $P_0(\mathbf{m}) = \prod_{j=1}^J \{\psi_j\}^{m_j} \{1 \psi_j\}^{1 m_j} = \Pi(\mathbf{m}; \psi)$ 
  - 1a.  $\Pi(\mathbf{m}; \mathbf{s}) = \prod_{j=1}^{J} \{s_j\}^{m_{ij}} \{1 s_j\}^{1 m_{ij}}$  is the probability mass function for a product Bernoulli distribution given the success probabilities  $\mathbf{s} = (s_1, \dots, s_J)^{\top}$ ,  $0 \le s_j \le 1$
  - 1b. Parameters  $\psi = (\psi_1, \dots, \psi_J)^{\top}$  represent the positive rates absent disease, referred to as "false positive rates" (FPRs).

Local Independence:  $M_{ij} \perp M_{ij'} \mid I = 0$ 

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- 2a-2b: Local Independence (LI):  $M_{ij} \perp M_{ij'} \mid I = \ell \neq 0$
- 2a-2b. Non-interference: disease-causing pathogen(s) are more frequently detected among cases than controls  $(\theta_\ell > \psi_\ell)$  and the non-causative pathogens are observed with the same rates

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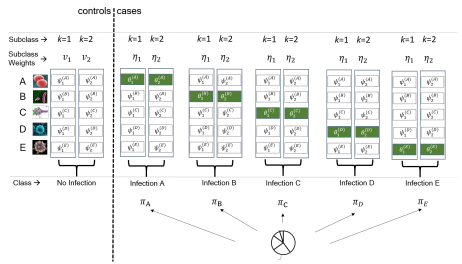
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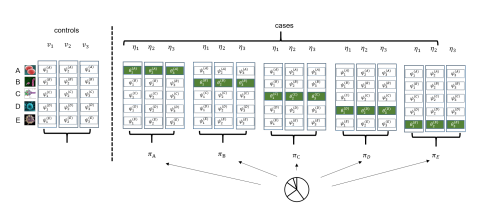
## Nested Partially-Latent Class Models (npLCM; Wu and Zeger, 2016)

Example: 5 Pathogens, 2 Subclasses; BrS Data Only



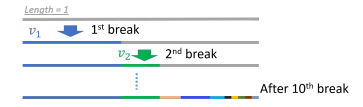
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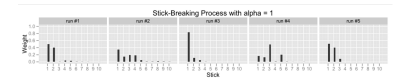
Example: 5 Pathogens, 3 Subclasses; BrS Data Only



## Encourage Few Subclasses: Stick-Breaking Prior

 $V_j \sim \text{Beta}(1, \alpha)$ ; Example: K = 10,  $\alpha = 1$ 





• On average, the first several segments receive most weights

## npLCM: Likelihood and Prior

### BrS Data Only

Likelihood

$$\begin{split} \textbf{\textit{P}}_{0}(\textbf{\textit{M}}_{i} = \textbf{\textit{m}}) &= \sum_{k=1}^{K} \nu_{k} \prod_{j=1}^{J} \left\{ \psi_{k}^{(j)} \right\}^{m_{j}} \left\{ 1 - \psi_{k}^{(j)} \right\}^{1 - m_{j}}, \\ \textbf{\textit{P}}_{1}(\textbf{\textit{M}}_{i} = \textbf{\textit{m}}) &= \sum_{j=1}^{J} \pi_{j} \sum_{k=1}^{K} \left[ \eta_{k} \left\{ \theta_{k}^{(j)} \right\}^{m_{j}} \left\{ 1 - \theta_{k}^{(j)} \right\}^{1 - m_{j}} \prod_{\ell \neq j} \left\{ \psi_{k}^{(j)} \right\}^{m_{\ell}} \left\{ 1 - \psi_{k}^{(j)} \right\}^{1 - m_{\ell}} \right], \end{split}$$

Prior:

Now, how to incorporate covariates, to which quantities?

Regression Extension for  $P_0$  and  $P_1$ : letting  $\pi_\ell$ ,  $\nu_k$ ,  $\eta_k$  depend on covariates

## Roadmap

Let three sets of parameters in an npLCM (pg.17) depend on the observed covariates

- 1x. Etiology regression function among cases,  $\{\pi_{\ell}(\mathbf{x}), \ell \neq 0\}$ , which is of primary scientific interest
- 2x. Conditional probability of measurements **m** given covariates **w** in controls:  $P_0(\boldsymbol{m}; \boldsymbol{w}) = [\boldsymbol{M} = \boldsymbol{m} \mid \boldsymbol{W} = \boldsymbol{w}, I = 0],$
- 3x. 2x above, but in the case class  $\ell$ :  $P_{1\ell}(m; w) = [M = m \mid W = w, I = \ell], \ \ell = 1, \dots, L$
- note Keep the specifications for the TPRs and FPRs  $(\Theta, \Psi)$  as in the original npLCM.

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- 5a. Use B-spline basis expansion to approximate  $f_{\ell j}^{\pi}(\cdot)$  and use P-spline for estimating smooth functions.
- 5b.  $\widetilde{\mathbf{x}}$  is the subvector of the predictors  $\mathbf{x}$ ;  $\mathbf{\Gamma}_{\ell}^{\pi} = (\boldsymbol{\beta}_{\ell i}^{\pi}, \boldsymbol{\gamma}_{\ell}^{\pi})$ .

# $P_0$ : Multivariate binary regression for controls

### Desirable properties

## Model Specification:

- Model space large enough for complex conditional dependence of M given covariates W
- Upward compatibility, or reproducibility (invariant parameter) interpretation with increasing dimensions or complex patterns of missing responses)

#### Estimation:

- Consistency and Efficiency
- Adaptivity: regularization to adapt to the difficulty of the problem, e.g., model residual dependence  $[M \mid W, I = 0]$  only if necessary; model the effect of covariates only if necessary

# Let $P_0$ depend on $W_i$

### Regression model for controls

• The pmf for controls' measurements:

Pr(
$$\mathbf{M}_i = \mathbf{m} \mid \mathbf{W}_i, I_i = 0$$
) =  $\sum_{k=1}^K \nu_k(\mathbf{W}_i) \Pi(\mathbf{m}; \Psi_k)$ ,  $\Psi_k = (\psi_k^{(1)}, \dots, \psi_k^{(J)})'$ 

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sample subclass indicator:  $Z_i \mid \mathbf{W}_i \sim \mathsf{Categorical}_{\kappa}(\nu(\mathbf{W}_i))$ 

 $M_{ii} \mid Z_i = k \sim \mathsf{Bernoulli}(\psi_k^{(j)}),$ generate measurements:

independently for i = 1, ..., J.

Regression model for controls Stick-breaking parametrization of weight functions  $\nu_k(\mathbf{W}_i) = P(Z_i = k \mid \mathbf{W}_i)$  by

$$\underbrace{h_k(\mathbf{W}_i; \mathbf{\Gamma}_k^{\nu})}_{\substack{\text{stick } k}} = \begin{cases} g(\alpha_{ik}^{\nu}) \prod_{s < k} \left\{1 - g(\alpha_{is}^{\nu})\right\}, & \text{if } k < K, \\ \prod_{s < k} \left\{1 - g(\alpha_{is}^{\nu})\right\}, & \text{if } k = K, \end{cases}$$

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We specify  $\alpha_{i\nu}^{\nu}$  via additive models at  $g^{-1}$  scale  $(g(\cdot) = 1/(1 + \exp\{-(\cdot)\}))$ :

$$\alpha_{ik}^{\nu} = \mu_{k0} + \sum_{i=1}^{q_1} f_{kj}(\mathbf{W}_{ij}; \boldsymbol{\beta}_{kj}^{\nu}) + \widetilde{\mathbf{W}}_{i}^{\top} \boldsymbol{\gamma}_{k}^{\nu}, \ k = 1, \dots, K-1.$$

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Expand the smooth functions by B-spline bases with coefficients  $\beta_{ki}^{\nu}$ ;  $\widetilde{\boldsymbol{w}}$  is a subvector of covariates  $\boldsymbol{w}$ 

## Adaptivity Considerations©

#### Proposed Model

- Prevent overfitting when the regression is easy, and improve interpretability
- We a priori place substantial probabilities on models with the following two features:
  - a) Few subclasses with effective weights (in the sense that  $\nu_k(\cdot)$  is bounded away from 0 and 1): a novel additive half-Cauchy prior for  $\mu_{k0}$ .
  - b) Smooth weight regression curves  $\nu_k(\cdot)$ : by Bayesian Penalized-Splines (P-Splines) combined with mixture priors on spline coefficients to sensitively distinguish constant  $\alpha_{\nu}^{\nu}(\cdot)$  from flexible smooth curves

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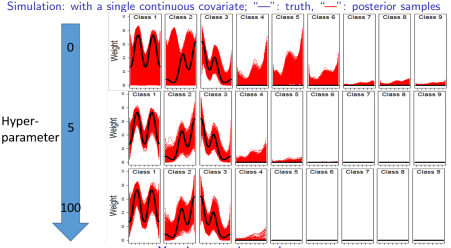
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- Encourages using a small number of effective classes (< K) to approximate the observed  $2^J$  probability contingency table in finite samples

## Inference of $\nu_k(x)$ at three hyperparameter values $s_k$



X-axis: covariate values

Y-axis: weight; 0 to 1.

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- we use  $\mu_{k0}$  from the controls (why?)

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- $\nu_{ik} = h_k(\mathbf{W}_i; \mathbf{\Gamma}_k^{\nu})$ : The S????-B???? parameterization
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Cases likelihood with covariates:

$$L_1^{\text{reg}} = \prod_{i:Y_i=1} \left\{ \sum_{\ell=1}^{L} \left[ \underbrace{\pi_{\ell}(\boldsymbol{X}_i; \boldsymbol{\Gamma}_{\ell}^{\pi})}_{\boldsymbol{PEF} \ \ell} \sum_{k=1}^{K} \left\{ \eta_{ik} \cdot \Pi(\boldsymbol{M}_i; \boldsymbol{p}_{k\ell}) \right\} \right] \right\} \quad (2)$$

- $\nu_{ik} = h_k(\mathbf{W}_i; \mathbf{\Gamma}_k^{\nu})$ : The S????-B???? parameterization
- $\eta_{ik} = h_k(\mathbf{W}_i; \mathbf{\Gamma}_k^{\eta})$

The joint likelihood for the regression model can be written as:  $L^{\text{reg}} = L_1^{\text{reg}} \times L_0^{\text{reg}}$ .

## Prior Specifications

#### Unknown parameters:

- etiology regression coefficients  $(\{\Gamma_{\ell}^{\pi}\})$ ,
- subclass mixing weight parameters for cases  $(\{\Gamma_{\nu}^{\eta}\})$  and controls  $(\{\Gamma_{\nu}^{\nu}\})$ ,
- true and false positive rates  $(\Theta = \{\theta_{\nu}^{(j)}\}, \Psi = \{\psi_{\nu}^{(j)}\}).$

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To avoid potential overfitting, we a priori introduce:

- (a) few non-trivial subclasses via novel additive half-Cauchy prior for the intercepts  $\{\mu_{k0}\}$
- (b) for continuous variable: smooth regression curves  $\pi_{\ell}(\cdot)$ ,  $\nu_k(\cdot)$  and  $\eta_k(\cdot)$  by Bayesian Penalized-splines (Lang. 2004) combined with shrinkage priors on spline coefficients (Ni et.al, 2015) (to encourage towards constant values)

Use Markov chain Monte Carlo (MCMC) algorithm to approximate joint posterior distribution

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 Posterior inference is flexible and can be obtained from any functions of model parameters and individual latent variables

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 Posterior inference is flexible and can be obtained from any functions of model parameters and individual latent variables

Fit npLCMs (w/ or w/out covariates using R package baker (https://github.com/zhenkewu/baker)

- calls Bayesian model fitting software JAGS 4.2.0 (Plummer et al., 2003) from within R
- provides functions to visualize the posterior distributions of the unknowns
- also performs posterior predictive model checking

#### Simulation Results

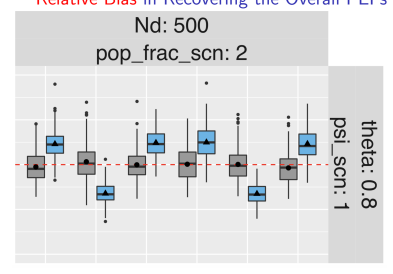
#### Simulation Results

• Simulation I: demonstrate flexible statistical inferences about the PEF functions  $\{\pi_{\ell}(\cdot)\}$  (not shown in this talk; check paper)

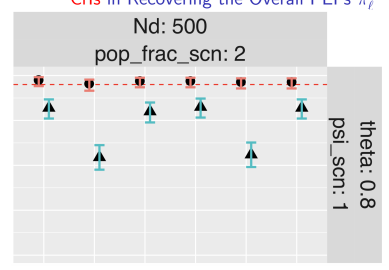
#### Simulation Results

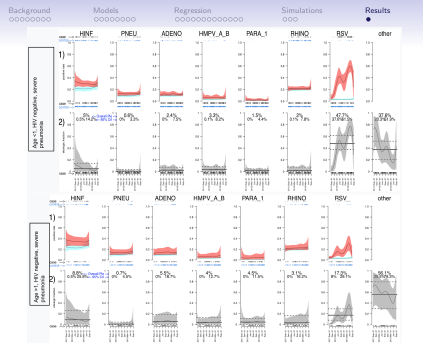
- Simulation I: demonstrate flexible statistical inferences about the PEF functions  $\{\pi_{\ell}(\cdot)\}$  (not shown in this talk; check paper)
- Simulation II: determining overall PEF  $\pi_{\ell}^*$  (empirical average) to quantify disease burdens in a population (potential policy interest)

# Simulation II: Regression Model Reduces the Percent Relative Bias in Recovering the Overall PEFs $\pi_{\ell}^*$



# Simulation II: Regression Model Produces More Valid 95% Crls in Recovering the Overall PEFs $\pi_{\ell}^*$





Context: Modern large-scale etiology studies generate complex measurements of unobserved causes of disease, and have raised the analytic needs of estimating cause-specific case fractions, or "Population Etiologic Fractions" (PEFs)

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Context: Modern large-scale etiology studies generate complex measurements of unobserved causes of disease, and have raised the analytic needs of estimating cause-specific case fractions, or "Population Etiologic Fractions" (PEFs)

Gap: Despite recent methodological advances, the need of describing the relationship between covariates and PEFs, remains unmet

Contribution: A general etiology regression framework building on npLCM that is broadly applicable to case-control studies A general framework for a class of statistical problems that can be formulated as estimating covariate-dependent class-mixing weights.

Three features of our approach:

Discussion

# Main Points Once Again

#### Three features of our approach:

• 1) allows analysts to specify a model that links important covariates to PEFs ©

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#### Three features of our approach:

- 1) allows analysts to specify a model that links important covariates to PEFs ©
- 2) produces covariate-dependent reference distribution for controls, which is critical for assigning cause-specific probabilities to a given case ☺
  - because we can compare control measurements to case measurements with similar covariate values
- 3) TPR priors are only used once; avoids overly-optimistic

### Thank You!

#### Collaborators

Laura Hammitt

Scott Zeger
Student Katherine O'Brien
Irena Chen Maria Deloria-Knoll

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Some References (More at: zhenkewu.com)

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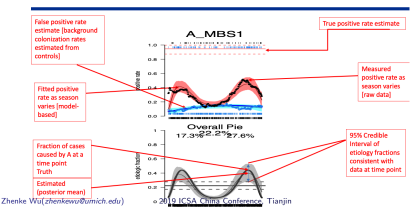
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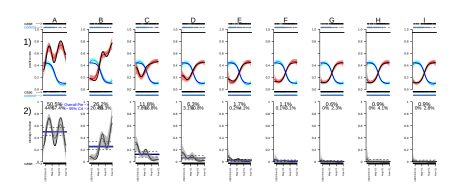
#### Simulation I Results

•  $N_d = 500$  cases and  $N_u = 500$  controls for each of two levels of S (discrete covariate); Uniformly sample the subjects' enrollment dates over a period of 300 days.

### **Etiology Regression Curves: Seasonality**

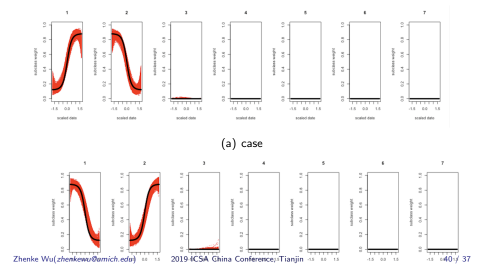


# Simulation I: Recovery of Truth $\pi_\ell^0(t,S=s)$



# Simulation I: Recovery of $\nu_k(t)$ and $\eta_k(t)$

True  $K^0 = 2$ ; Model fitted using a working number K = 7



# Appendix: Simulation II Setup

- npLCM regression analysis with  $K^* = 3$ , R = 200 replication data sets simulated under 48 different scenarios
- L = J = 3, 6, 9 causes, under single-pathogen-cause assumption, BrS measurements made on  $N_d$  cases and  $N_u$  controls for each level of X where  $N_d = N_u = 250$  or 500.
- $\phi_{\ell}(X) = \beta_{0\ell} + \beta_{1\ell} \, \mathbb{I}\{X=2\}$  take two sets of values to reflect PEF variability across X: i)  $\beta_0^i = (0,0,0,0,0,0)$ ,  $\beta_1^i = (-1.5,0,-1.5,-1.5,0,-1.5)$ ; ii)  $\beta_0^{ii} = (1,0,1,1,0,1)$  and  $\beta_1^{ii} = (-1.5,1,-1.5,-1.5,1,-1.5)$
- TPRs  $\theta_k^{(j)} = 0.95$  or 0.8 and FPRs  $(\psi_1^{(j)}, \psi_2^{(j)}) \in \{(0.5, 0.05), (0.5, 0.15)\}$ , for  $j = 1, \dots, J$ .
- $\nu_k(W) = \eta_k(W) = logit^{-1} (\gamma_{k0} + \gamma_{k1} \mathbb{I}\{W = 2\})$  where  $(\gamma_{10}, \gamma_{11}) = (-0.5, 1.5)$  and  $(\gamma_{20}, \gamma_{21}) = (1, -1.5)$ .

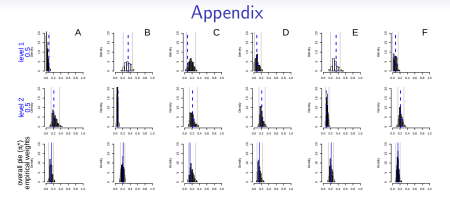


Figure: Posterior distributions of the stratum-specific (Row 1 and 2) and the overall (Bottom Row) PEFs based on a simulation with a two-level discrete covariate and L=J=6 causes. The vertical gray lines indicate the 2.5% and 97.5% posterior quantiles, respectively; The truths are indicated by vertical blue dashed lines. Row 1-2) PEFs by stratum (level = 1,2) and cause (A-F); Bottom)  $\pi_\ell^*$ : overall population etiologic fraction for cause A-F (empirical average of the two PEFs above).

## **Appendix**

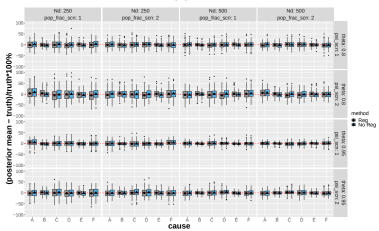
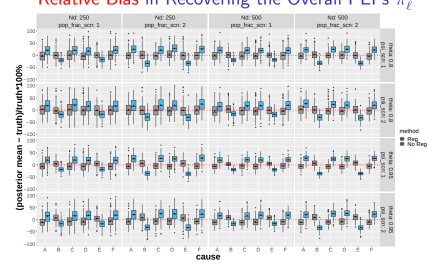
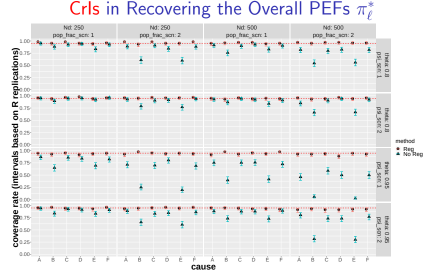


Figure: NPLCM analyses with or without regression perform similarly in terms of percent relative bias (top) and empirical coverage rates (bottom) over R=100 replications in simulations where the case and control subclass weights *do not* vary by covariates. Each panel corresponds to one of 16 combinations of true  $\frac{1}{2} \frac{1}{2} \frac{1}{2}$ 

# Simulation II: Regression Model Reduces the Percent Relative Bias in Recovering the Overall PEFs $\pi_{\ell}^*$



# Simulation II: Regression Model Produces More Valid 95%



## **Appendix**

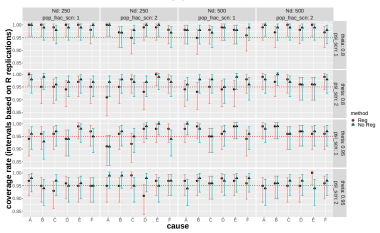


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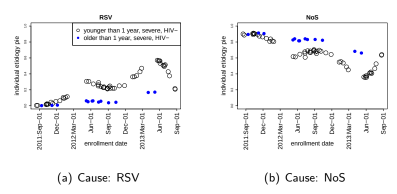


Figure: Individual etiology fraction estimates for RSV (left) and NoS (right) differ by age and season among HIV negative and severe pneumonia cases for whom the seven pathogens were *all tested negative* in the nasopharyngeal specimens.