

On the role of marginal confounder prevalence – implications for the high-dimensional propensity score algorithm

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ABSTRACT

Purpose The high-dimensional propensity score algorithm attempts to improve control of confounding in typical treatment effect studies in pharmacoepidemiology and is increasingly being used for the analysis of large administrative databases. Within this multi-step variable selection algorithm, the marginal prevalence of non-zero covariate values is considered to be an indicator for a count variable's potential confounding impact. We investigate the role of the marginal prevalence of confounder variables on potentially caused bias magnitudes when estimating risk ratios in point exposure studies with binary outcomes.

Methods We apply the law of total probability in conjunction with an established bias formula to derive and illustrate relative bias boundaries with respect to marginal confounder prevalence.

Results We show that maximum possible bias magnitudes can occur at any marginal prevalence level of a binary confounder variable. In particular, we demonstrate that, in case of rare or very common exposures, low and high prevalent confounder variables can still have large confounding impact on estimated risk ratios.

Conclusions Covariate pre-selection by prevalence may lead to sub-optimal confounder sampling within the high-dimensional propensity score algorithm. While we believe that the high-dimensional propensity score has important benefits in large-scale pharmacoepidemiologic studies, we recommend omitting the prevalence-based empirical identification of candidate covariates. Copyright © 2015 John Wiley & Sons, Ltd.

KEY WORDS—confounder prevalence; high-dimensional propensity score; risk ratio; bias formula; pharmacoepidemiology

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INTRODUCTION

The high-dimensional propensity score (hd-PS) algorithm introduced and established by Schneeweiss and colleagues^{1–6} attempts to reduce residual confounding in typical treatment effect studies in pharmacoepidemiology and is increasingly being used by large, collaborative drug safety networks such as the US Mini-Sentinel⁷ and the Canadian Network for Observational Drug Effect Studies.⁸

It is assumed that adjustment for large numbers of covariates, as commonly available in administrative databases, may serve as sufficient proxy for unobserved

confounders, which, if not considered, would yield biased estimates of treatment effects. In general, an ideal hd-PS algorithm will either select all relevant confounders in a given data set or, because of the latent correlation structure of all potential confounder variables, select enough confounder-associated variables so that unobserved or non-selected confounders are sufficiently mirrored in the final propensity score model. In order to achieve confounder-adjusted treatment or exposure effect estimates, the fitted propensity score usually serves as a balancing score in a multivariable outcome model, as matching criterion or is used to define inverse probability weights for the estimation of marginal treatment or exposure effects.

The hd-PS variable selection algorithm considers the marginal prevalence of non-zero values of a covariate (either binary or count variable) as an indicator for its potential confounding magnitude. Accordingly,

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candidate covariates to be included in the final propensity score model are empirically identified. The idea of assessing the potential confounding impact of extraneous variables is not entirely new and has received broad attention in the epidemiological literature.^{9–14} However, so far, the proposed formulas and illustrations focus mainly on the conditional prevalence of confounder variables, that is, dissimilarities in confounder prevalence between the exposure groups. In this article, we investigate the theoretical justification for assessing potential bias impact based on the marginal prevalence of confounder variables as implied by the hd-PS algorithm.

The article is structured as follows: Section 2 provides a brief review of the high-dimensional propensity score covariate selection procedure. Section 3 gives formal insights in confounding mechanisms and the role of marginal confounder prevalence. Section 4 provides a comprehensible data example and illustrates the formally determined boundaries for bias magnitudes in dependency on marginal confounder prevalence and confounder-outcome association. Section 5 closes with discussion and conclusion.

THE HIGH-DIMENSIONAL PROPENSITY SCORE COVARIATE SELECTION PROCEDURE

The hd-PS confounder selection algorithm is essentially comprised of three steps: First, among all count variables (codes from the claims database), candidate empirical covariates are selected according to their marginal prevalence (proportion of subjects with a count value of at least one). Here, variables with marginal prevalence values closer to 0.5 are given higher priority. Second, the selected count variables are recoded into three binary variables indicating (i) if a subject has at least one count; (ii) if a subject has a count value greater or equal to the median of all non-zero counts in this variable; and (iii) if a subject has a count value greater or equal to the 75th percentile of all non-zero counts. Third, after dichotomization, ranking and selection of potential confounder variables according to a multiplicative bias term that reflects the relative magnitude of bias potentially caused by each of the created dummy variables. This calculation is performed using a formula based on an earlier work of Bross⁹:

$$ARR_{ED} = RR_{ED} \cdot \frac{P_{C1} (RR_{CD} - 1) + 1}{P_{C0} (RR_{CD} - 1) + 1} \quad (1)$$

Here, ARR_{ED} refers to the apparent (unconditional) relative risk on a binary outcome variable D associated

with a binary point-exposure or treatment E . The relative risk between exposed and unexposed individuals conditional on the binary confounding variable C is given by RR_{ED} . Furthermore, RR_{CD} depicts the relative risk between subjects with and without the confounder attribute, and $P_{C1} = P(C=1|E=1)$ and $P_{C0} = P(C=1|E=0)$ are the prevalence of the confounder in exposed and unexposed individuals, respectively.

CONFOUNDING AND THE ROLE OF MARGINAL CONFOUNDER PREVALENCE

The maximum possible magnitude of the multiplicative bias term of a confounder variable is a clearly defined function of $P(C)$, $P(E)$, and RR_{CD} . According to the law of total probability, the marginal prevalence of C is given by the weighted sum of the respective conditional prevalence values:

$$P(C) = P(C = 1|E = 1) \cdot P(E = 1) + P(C = 1|E = 0) \cdot P(E = 0) \quad (2)$$

As indicated by Equation (1), and as universally known as the key point in confounding mechanism, the magnitude of confounder-induced bias is strictly increasing with increased imbalance of the confounder distribution in the exposure groups. Accordingly, at fixed values of $P(E)$ and RR_{CD} , the highest magnitude of bias potentially caused by a confounder variable appears in the most extreme cases of pure imbalance either if $P(C=1|E=1)=1$ and $P(C=1|E=0)=0$ or if $P(C=1|E=1)=0$ and $P(C=1|E=0)=1$. In these cases, the effect of the confounder variable cannot be differentiated from the effect of the exposure so that RR_{CD} immediately bonds with RR_{ED} and the apparent relative risk becomes simply the product of these two effect measures. Therefore, the maximum reachable magnitude of the multiplicative bias term corresponds to the value of RR_{CD} or $1/RR_{CD}$ in the worst two possible confounder distribution scenarios among the exposure groups. Equation (2) illustrates the crucial implications $P(C)=P(E)$ or $P(C)=1-P(E)$ induced by such extreme confounding situations. However, because neither $P(C)$ nor $P(E)$ are restricted to a value of 0.5 and the bias magnitude increases with decreasing confounder balance among the exposure groups, the rationale to pre-select potential confounders according to marginal prevalence values close to 0.5 may lead to an inappropriate choice of propensity score variables.

Table 1. Hypothetical data example demonstrating relevant confounding magnitude in estimating an exposure-outcome association in presence of a low prevalent binary confounder

I				II				III				IV			
Total sample				Total sample				Confounder = "No"				Confounder = "Yes"			
Outcome	No	Exposure		Confounder	No	Exposure		Outcome	No	Exposure		Outcome	No	Exposure	
		No	Yes			No	Yes			No	Yes			No	Yes
	Yes	25	488		22	22	948		11	11	474		14	14	14
		12	475		15	15	15		11	11	474		1	1	1
		37	963		37	37	963		22	22	948		15	15	15

EXAMPLE DATA AND ILLUSTRATION

The following example provides a simple hypothetical data scenario in which the marginal prevalence of a confounder variable is low but the magnitude of bias caused is still substantial.

Panel I of Table 1 displays the unconditional exposure-outcome association indicating an about 1.5-fold higher risk for exposed individuals. Exposure (E) is high prevalent (96%) in the study population. Panel II shows for the same study sample the distribution of a low prevalent (3%) binary confounding variable (C). A strong imbalance of the confounder between the exposure groups is present, as $P(C=1|E=1)=0.02$ and $P(C=1|E=0)=0.41$. Panels III and IV indicate, in contrast to Panel I, no exposure-outcome association conditional on the confounder variable. Thus, despite the given low marginal confounder prevalence, the

confounder-associated multiplicative bias still yields a magnitude of 1.5 in this example scenario. This is because the confounder distribution is strongly imbalanced between the exposure groups, a fact which cannot be deduced from the marginal confounder prevalence.

Figure 1 provides a comprehensive illustration of the described multivariable impact of $P(C)$, $P(E)$, and RR_{CD} on the confounding magnitude potentially caused by C. We simply plotted the multiplicative bias term from Equation (1) for all possible values resulting from a grid $\{P_{C0}, P_{C1}\} \in \{(0, 0.1, \dots, 0.9, 1) \times (0, 0.1, \dots, 0.9, 1)\}$ conditional on the respective marginal confounder prevalence $P(C)$. We considered nine exemplary configurations by setting $RR_{CD} = \{1, 1.5, 2\}$ and $P(E) = \{0.1, 0.5, 0.75\}$. In the resulting graphs, we added horizontal dashed lines on angle of the respective value of RR_{CD} and $1/RR_{CD}$ as well as vertical dashed lines for values of $P(C)$ and $1 - P(C)$. As can

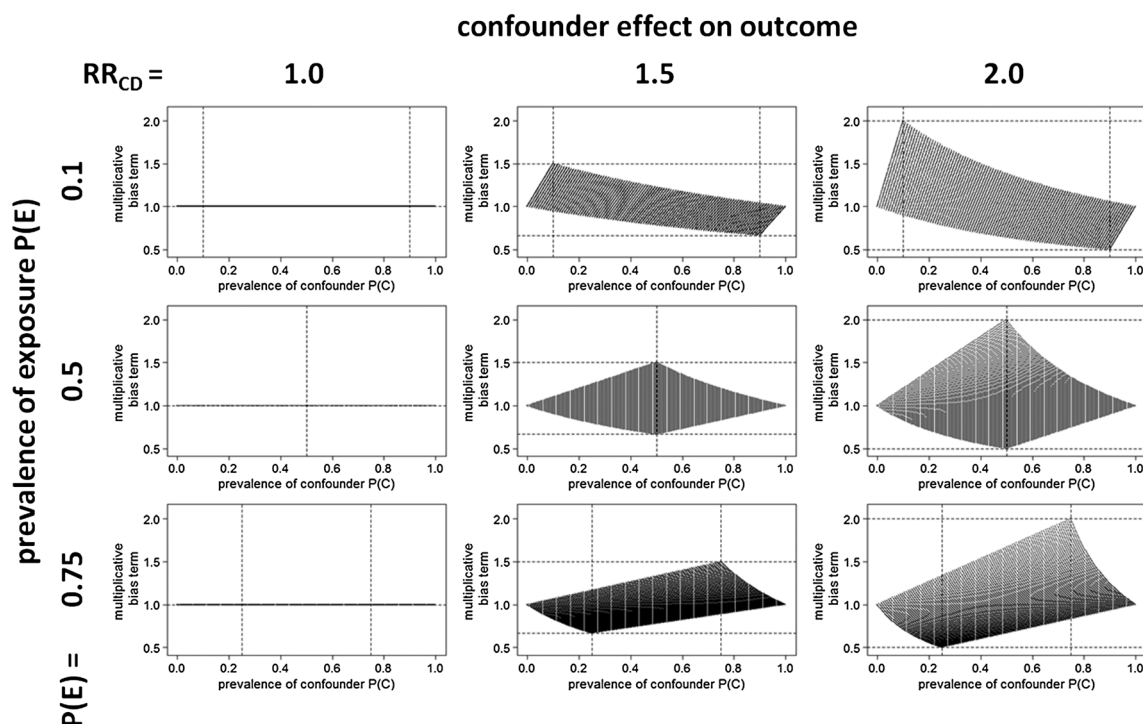


Figure 1. Multiplicative bias term depending on $P(C)$, RR_{CD} and $P(E)$

be seen from the figure, the maximum multiplicative bias magnitude for a given scenario corresponds to the value of RR_{CD} (or $1/RR_{CD}$) and is achieved at the respective value of $P(C)=P(E)$ (or $1-P(E)$). Therefore, only in the case of $P(E)=0.5$, a pre-selection of confounders according to their marginal prevalence value would be appropriate. However, because the multiplicative bias term already reflects the impact of the marginal prevalence on the confounding magnitude, such pre-selection would be superfluous.

DISCUSSION AND CONCLUSION

We formally explained and illustrated the definite role of the marginal prevalence of an uncontrolled binary variable on its confounding impact when estimating risk ratios in point-exposure studies with a binary outcome. We showed that low prevalent confounder variables can become highly influential in scenarios where the prevalence of at least one exposure category is low.

Propensity score methods such as the hd-PS are commonly not used in situations where the exposure prevalence is low. However, in the analysis of large pharmacoepidemiological data sets, low relative frequencies of exposed individuals often translate to sufficient absolute numbers that allow for reliable effect estimation using propensity score methods.

In light of the presented results, while we believe that the high-dimensional propensity score has important benefits in large-scale pharmacoepidemiologic studies in administrative data, we recommend the deletion of the prevalence-targeted pre-selection step within the hd-PS confounder selection procedure.

CONFLICT OF INTERESTS

The authors declare that there is no conflict of interest in the context of the submitted work.

KEY POINTS

- The marginal distribution of a potentially confounding variable is not an indicator for the magnitude of bias in estimating relative risks that failure to adjust for this variable may cause.
- Semi-automated confounder selection procedures such as the high-dimensional propensity score algorithm should not consider the marginal prevalence of a candidate confounding covariate as a selection criterion.
- Established bias formulas for relative risk estimates sufficiently approximate potential bias magnitudes induced by a single independent confounding variable.

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