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An Introduction to Sensitivity Analysis for Unobserved Confounding in Non-Experimental Prevention Research

Weiwei Liu,

NORC at the University of Chicago

S. Janet Kuramoto, and American Psychiatric Institute for Research and Education

Elizabeth A. Stuart

Department of Mental Health and Department of Biostatistics, Bloomberg School of Public Health, Johns Hopkins University

Abstract

Despite that randomization is the gold standard for estimating causal relationships, many questions in prevention science are left to be answered through non-experimental studies often because randomization is either infeasible or unethical. While methods such as propensity score matching can adjust for observed confounding, unobserved confounding is the Achilles heel of most nonexperimental studies. This paper describes and illustrates seven sensitivity analysis techniques that assess the sensitivity of study results to an unobserved confounder. These methods were categorized into two groups to reflect differences in their conceptualization of sensitivity analysis, as well as their targets of interest. As a motivating example we examine the sensitivity of the association between maternal suicide and offspring's risk for suicide attempt hospitalization. While inferences differed slightly depending on the type of sensitivity analysis conducted, overall the association between maternal suicide and offspring's hospitalization for suicide attempt was found to be relatively robust to an unobserved confounder. The ease of implementation and the insight these analyses provide underscores sensitivity analysis techniques as an important tool for non-experimental studies. The implementation of sensitivity analysis can help increase confidence in results from non-experimental studies and better inform prevention researchers and policymakers regarding potential intervention targets.

Keywords

sensitivity analysis; causal inference; unobserved confounding; suicide prevention

This paper describes and illustrates a set of tools, broadly known as "sensitivity analysis," to help understand the robustness of non-experimental findings to a potential unobserved confounder. The goal of this paper is to discuss the relevance of common sensitivity analysis methods to prevention research and to provide an easy-to-understand guide for interested prevention researchers. We do not intend to discuss the technical nuances of sensitivity analysis or to provide a comprehensive listing of all of the methods used to assess sensitivity. However, interested readers can refer to works cited throughout the paper for details and description of additional methods.

Correspondence to: Weiwei Liu.

Many studies in prevention research aim to investigate causal effects—either the effects of early risk factors or experiences on later outcomes, such as the effects of adolescent drug use on outcomes such as unemployment and drug use during adulthood (Stuart & Green, 2008), or the effects of particular interventions, such as the Good Behavior Game (Kellam, Brown, Poduska, Ialongo, Wang, Toyinbo et al., 2008). Understanding such causal effects is crucial for determining what risk or protective factors should be targeted to improve outcomes or whether a preventive intervention is effective. In some situations researchers can randomize individuals to receive the intervention or the control condition, a method considered the gold standard for estimating causal effects (Flay, Biglan, Boruch, Castro, Gottfredson, Kellam et al., 2005). However, randomization is often infeasible or unethical, sometimes due to the nature of the factor under study (such as drug abuse or childhood maltreatment). In those cases questions of causal inference are left to be answered using non-experimental methods.

The Achilles heel of non-experimental studies is that the exposed and unexposed (or treatment and control) groups may not be comparable, a phenomenon formally known as confounding (Rothman, Greenland, & Lash, 2008). While observed confounding can be addressed with methods such as propensity score matching (Stuart, 2010), methods to assess the consequences of unobserved confounding are less readily available, and researchers tend to shy away from the issue. Being able to determine the robustness of study findings to potential unobserved confounding is crucial for theory testing and development. In addition, the knowledge of to what extent the conclusions drawn from those studies are robust to potential unobserved confounding can help with policy decision making.

The origin of sensitivity analysis to unobserved confounding has been attributed to a study by Cornfield, Haenszel, Hammon, Lilienfeld, Shimkin, and Wynder (1959), which quantified the role of unobserved confounding in the observed relationship between smoking and lung cancer (Cornfield et al., 1959). Specifically, Cornfield et al. (1959) showed that an unobserved confounder, such as a genetic factor, would need to lead to a nine-fold increase in the odds of smoking in order to explain away the association between smoking and lung cancer, and it was asserted that such a strong unobserved confounder was very unlikely to exist. Since then, more recent examples demonstrate the use of sensitivity analysis to assess unobserved confounding in various fields, such as sociology, criminology, psychology, and prevention science (e.g., DiPrete & Gangl, 2004; Harding, 2003; Haviland, Nagin, & Rosenbaum, 2007; Kitahata, Gange, Abraham, Merriman, Saag, Justice et al., 2009; Liu, in press), though such examples are still relatively rare. For example, Haviland et al. (2007) found that the estimated effect of gang involvement on subsequent violence was robust to a fairly weak unobserved confounder but may be sensitive to an unobserved confounder moderately associated with gang involvement and subsequent violence. Harding (2003) found that the estimated effect of neighborhood choice on high school dropout was robust to an unobserved confounder if high and low poverty neighborhoods were compared. However, the effect was sensitive to an unobserved confounder if high and moderate poverty neighborhoods were compared or if moderate and low poverty neighborhoods were compared. As a final example, Liu (in press) found that an observed relationship between high school dropout and subsequent adult criminal offending estimated using propensity score matching remained significant even after taking into account an unobserved confounder strongly associated with both high school dropout and criminal offending.

Conceptually, sensitivity analysis can be understood from both a statistical and an epidemiological perspectives (Luiz & Cabral, 2010). Although the two perspectives differ in their approaches to conceptualize sensitivity analysis, they both assess how strong the effects of the unobserved covariate on the exposure and/or the outcome would have to be to change the study inference. From a statistical perspective, Rosenbaum (2002, 2010) emphasizes the difference between randomized trials and non-experimental studies; that is,

in a non-experimental study, exposed and unexposed groups may differ on an unobserved characteristic even after matching on observed characteristics. In other words, individuals with the same observed covariates may have different probabilities of being exposed if they have different unobserved covariates. A sensitivity parameter is used to quantify the difference in the odds of exposure for two individuals with the same observed covariates (or the same propensity score) but diverge on unobserved covariates. The goal is to determine the smallest value of this parameter that will change the *p*-value of the "true" outcome-exposure association to a non-significant level.

Sensitivity analysis can also be understood from an epidemiological perspective (e.g., Harding, 2003), as a tool to assess the extent to which a significant association found between observed variables could be due to unobserved confounding. Sensitivity parameters are used to quantify the strengths of the associations between a hypothetical unobserved confounder and the exposure and outcome. The goal is to arrive at a "true" association between the exposure and the outcome, adjusting for the hypothetical unobserved confounder with various values of the sensitivity parameters.

We selected seven methods to present in this paper. In selecting these methods, we attempted to keep a balance between introducing a broad range of methods that accommodate different interests and data availability and selecting the methods that are relatively straightforward to understand and easy to apply to a number of different settings. For example, the selected methods capture the above-mentioned two different perspectives to conceptualize sensitivity analysis. Additionally, these seven methods have different targets of interest. Specifically, the first group, i.e. Rosenbaum's three approaches (Gastwirth, Krieger, & Rosenbaum, 1998) focus on the statistical significance of the "true" outcome-exposure association, while the second group, i.e. Greenland's (1996), Harding's (2003), Lin et al.'s (1998) and VanderWeele and Arah's (VanderWeele & Arah 2011, Arah, Chiba, & Greenland 2008) approaches obtain the point estimate of the "true" outcomeexposure association with a 95% confidence interval. Other differences between these approaches (such as study design and outcome distribution) are described later in the manuscript and summarized in Table 1. Importantly, the seven selected methods are all relatively straightforward to understand and can be computed by hand or with standard statistical software.

Motivating Study

As a demonstration, we apply the seven sensitivity analysis methods to investigate potential unobserved confounding in a recent study by Kuramoto, Stuart, Runeson, Lichtenstein, Langstrom, and Wilcox (2010), which examined the association between maternal suicide and their offspring's hospitalization for suicide attempt. Parental suicide has been examined as a risk factor for adolescent's suicide attempt (Wilcox, Kuramoto, Lichtenstein, Langstrom, Brent, & Runeson, 2010). However, the association between parental suicide and offspring's risk has been somewhat equivocal, with most of the research in this area coming from short-term prospective or cross-sectional studies of referred samples. Additionally, most previous studies compared offspring of suicide decedents with offspring of living parents, which cannot clarify our understanding on the impact of parental suicide over and beyond the impact of sudden parental death. Understanding the causal mechanisms, and not just associations, is particularly crucial to suicide prevention to identify populations in which prevention efforts can be targeted. For example, if parental suicide was found to indeed cause offspring's suicide attempt, then prevention efforts should target individuals whose parents died from suicide. If it was found that such a relationship can be easily explained away by some other factors, it is important to search for those factors that may better explain hospitalization for suicide attempt. However, we cannot

randomly assign parental suicide, leaving the investigation of such a relationship relying solely on non-experimental studies. With such a goal, Kuramoto et al. (2010) compared 5,600 offspring who lost a mother to suicide before age 18 (exposed group) with 2,872 offspring who lost a mother to an accident (unexposed group). This comparison group allowed a clearer distinction of risk, over and beyond the stress and disruption associated with sudden parental death. Propensity score matching was used to make the exposed and unexposed groups as similar as possible on observed characteristics such as deceased parent and surviving parent's psychiatric hospitalization. One to one nearest neighbor propensity score matching with replacement (Stuart 2010) was used. The authors concluded that maternal suicide was associated with a 1.86-fold increased risk (95% CI=(1.49, 2.32))¹ of their offspring being hospitalized for suicide attempt, as compared to matched offspring who lost their mother to an accident, a result in congruent with the increasing body of literature on the impact of parental suicide on offspring's risk for suicidal behavior (Wilcox et al., 2010; Niederkrotenthaler, Floderus, Alexanderson, Rasmussen, & Mittendorfer-Rutz., 2012).

While the exposed and unexposed groups were matched on a large number of characteristics, potential unobserved confounders remain a concern. For example, the observed association may be partly explained by genetic predisposition to suicidal behavior, which has been suggested to be associated with both suicide and offspring's suicide attempt (Brent & Mann, 2005; Lieb, Bronisch, Hofler, Schreier, & Wittchen, 2005). In order to accommodate some methods (e.g., primal sensitivity analysis) that are most easily applied to a 1:1 matched setting, and ease the comparison of these methods with other methods, the 1:1 matching with replacement was modified to resemble a 1:1 match without replacement. This resulted in 5,600 matched pairs, with a total sample size of 11,200. In these pairs, 233 offspring of suicide decedents and 128 offspring of accident decedents were hospitalized for suicide attempt. This modification did not change the inferences about the association between maternal suicide and offspring's suicide attempt.

Setting, Assumptions and Notation

In order to demonstrate and motivate the use of sensitivity analysis, we focus on a relatively simple setting with a binary exposure, a binary outcome, and a binary unobserved confounder. The focus on binary exposures and outcomes is common in the causal inference literature (Stuart, 2010), helps simplify the sensitivity analysis techniques, and is a common assumption in that setting (e.g., VanderWeele & Arah 2011; Harding 2003). The binary unobserved confounder can also be thought of as a combination of a number of unobserved confounders (Lin, Psaty, & Kronmal, 1998). Some of the methods discussed in this paper can be generalized to accommodate continuous normally distributed unobserved confounders, although the computation is more involved. The basic ideas remain the same except that when the confounder is continuous, the relationships between the unobserved confounder and the exposure and the outcome are expressed as mean differences rather than OR's (a brief discussion is provided later in this paper). In addition, some of these methods can be generalized to accommodate continuous or censored outcomes (e.g. Lin et al.'s approach; see Table 1).

Two common assumptions are made (though not necessarily required by every method, as discussed later in the paper) to apply the seven methods to the motivating example. First, we assume that the relationships between the unobserved confounder and the exposure and the outcome do not vary as a function of the observed covariates. As VanderWeele and Arah (2011) discussed when presenting their more general approach, it becomes virtually

¹The original study estimated hazard ratio of 1.80 with a 95% confidence interval of 1.19, 2.74.

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impossible to allow the specified parameters to differ across levels of the observed covariates when multiple observed covariates are involved. Second, we assume no three way interaction between the exposure, the outcome, and the unobserved confounder, an assumption made by most studies using sensitivity analysis (e.g., Harding, 2003). While of course these two assumptions are not always met in reality, demonstrating methods to accommodate violation of these assumptions is beyond the scope of this paper. A brief discussion is provided later in this paper; see VanderWeele and Arah (2011) and Lin et al. (1998) for more detailed discussion of those approaches.

The following notation will be used throughout the paper:

- x = binary treatment status/exposure
- y = binary outcome
- u = unobserved binary confounder
- c = observed confounders
- p(x) = prevalence of the exposure

p(u) = prevalence of the unobserved confounder

p(u|x=1) = prevalence of the unobserved confounder among the exposed group

p(u|x=0) = prevalence of the unobserved confounder among the unexposed group

ORyu = odds ratio of the relationship between the outcome and unobserved confounder

ORxu = odds ratio of the relationship between the exposure and unobserved confounder

 $ORyx \bullet c$ = observed odds ratio of the relationship between the outcome and the exposure from the data, adjusted for *c* (but not for *u*)

 $ORyx \cdot cu =$ true (bias-free/bias-adjusted) odds ratio of the relationship between the outcome and the exposure, adjusted for both *c* and *u*

Generally when performing sensitivity analyses, researchers should specify a range of parameter values that are suggested by the literature or based on the relationships between observed confounders and the exposure and outcome (e.g., Harding 2003) to examine the sensitivity of study inferences under different specifications. This approach is particularly useful when some parameters require outside knowledge that is not easily obtained. However, for demonstration purposes and to make the results from different methods comparable, we specify a single set of parameters to be used in the motivating example. Since the relationship between the hypothetical unobserved confounder (*u*; such as genetic predisposition) and the exposure (ORxu) and the outcome (ORyu) (net of all the covariates matched on) were not readily available in the literature, they were obtained by examining the relationships of the observed confounders (c) available in the motivating study with the exposure (having a mother die of suicide; ORxc) and the outcome (offspring's hospitalization for suicide attempt; ORvc). The ORs between the observed confounders and the exposure ranged from 0.98 to 7.39, with the strongest factor being the deceased parent's history of psychiatric hospitalization (ORxc=7.39). The ORs between the observed confounder and the outcome (y) ranged from 0.95 to 1.84, with the strongest factor being the psychiatric hospitalization of the surviving parent prior to the death of the parent (ORyc=1.84). To err on the conservative side, we fixed the values of ORxu and ORyu at these two highest values of the observed OR's (i.e., ORxu=7.39; ORyu=1.84). Some approaches require the specification of p(u|x=1) and p(u|x=0) instead of ORxu. The prevalence of an unobserved confounder such as genetic predisposition to suicidal behavior in the general population is not available; hence, we specified a range of p(u|x=0) from 1%

to 25% with the fixed *ORxu*=7.39 to obtain varying p(u|x=1). In the remainder of the paper, we introduce each of the seven methods and apply these methods to the motivating example. Details of the computations, sample R code, and links to relevant Excel spreadsheets and web equation solvers can be found in the eAppendix.

Rosenbaum's Approaches

Rosenbaum's approaches, in general, are interested in finding the thresholds of the association(s) between the unobserved confounder and the exposure (ORxu) and/or between the unobserved confounder and the outcome (ORyu) that would render the test statistics of the study inference (ORyx•cu) insignificant. This method is most frequently used when the observed confounders have been dealt with using matching methods (such as propensity score matching) that form matched pairs of exposed and unexposed individuals who are similar on the observed covariates. These approaches are further broken down into primal, dual and simultaneous analysis (Gastwirth et al. 1998), which differ in their specified parameters. Primal sensitivity analysis varies the association between the unobserved confounder and the exposure (ORxu, with an upper bound denoted), while setting ORyu at infinity. In contrast, dual sensitivity analysis varies the association between the unobserved confounder and the outcome (ORvu, with an upper bound denoted), while setting ORxu at infinity. Simultaneous sensitivity analysis varies both ORxu and ORyu. The primal and dual sensitivity analyses are, therefore, special cases of the simultaneous sensitivity analysis. We focus our further discussion on primal and simultaneous sensitivity analysis, as the steps involved in dual sensitivity analysis are similar to primal sensitivity analysis, except that the parameter that is varied is ORyu instead of ORxu.

Primal Sensitivity Analysis

ORxu estimates are bounded by : 1/ORxu, where 1^2 . The upper (p+) and lower (p-) bounds on the probability of being exposed, accounting for u, can then be calculated. In particular, p+ is of most interest and can be expressed as /1+.

A modified McNemar's exact test (McNemar, 1947) is then used to examine the association between x and y, accounting for u (i.e., by computing an upper-bound p-value using p+ instead of the observed probability of exposure of 0.5 in the matched pairs). In Equation 1, T as the total number of discordant pairs (those where the outcomes differ within the pair), and a as the number of discordant pairs, in which the exposed had an outcome and the unexposed did not. This is repeated with different values of to find the value of at which the upper-bound p-value becomes non-significant (e.g., p>0.05). A higher value of required to render the upper-bound p-value non-significant is preferred, as it indicates that $ORyx \cdot cu$ is more robust to unobserved bias, i.e., a stronger association between the unobserved confounder and the exposure is necessary for the $ORyx \cdot cu$ to become nonsignificant.

upperbound
$$P-value = \sum_{a}^{T} {T \choose a} (p^{+})^{a} (1-p^{+})^{T-a}$$
 (Eq. 1)

The primal sensitivity analysis for a 1:1 matched design with a binary outcome can be implemented by hand, using the 'rbounds' package in R (Keele, 2010) or using an available Excel spreadsheet (Love 2008; see the eAppendix for details). These tools can also be used for continuous outcomes; for example, by using a function written for Stata (Gangl, 2004).

 $^{^{2}}$ Note that this is not a loss of generality; if the unobserved confounder is negatively associated with exposure status, we could simply redefine the unobserved confounder to meet this scenario.

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Simultaneous Sensitivity Analysis

The simultaneous sensitivity analysis allows researchers to vary not only ORxu (with an upper bound) but also ORyu (with an upper bound). The goal is to find the combinations of and at which $ORyx \cdot cu$ becomes statistically non-significant. The steps are similar to primal sensitivity analysis. One first specifies values for and , which can be used to calculate the upper and lower bounds of the probability of being exposed given the unobserved confounder (p+ and p-). In particular, p+ can be calculated using Equation 2, where p() = /(1+) and p() = /(1+). We then use p+ and the numbers of discordant pairs to calculate the upper bound p-value of $ORyx \cdot cu$ using McNemar's exact test.

$$p^+ = p(\pi) \times p(\theta) + (1 - p(\pi)) \times (1 - p(\theta))$$
 (Eq. 2)

A combination of values of and for which the test-statistic becomes non-significant is a point at which the result is sensitive to an unobserved confounder. Although there is no specific package available, this analysis can be hand computed or easily programmed using Excel or R for a 1-1 matched pair design (see the eAppendix for details).

Application of Rosenbaum's Approaches

Primal sensitivity analysis

Using the counts of discordant pairs, i.e., a=226 and T=347, and different values of four methods (hand computation, 'rbounds' package in R, computation using R codes and Love's spreadsheet) were used for the analysis. The results suggest that when 1.55, the association between maternal death by suicide and offspring's hospitalization for suicide attempt would no longer be significant (with a p-value of .054).

Simultaneous sensitivity analysis

Using the counts of discordant pairs, as well as the specified parameters (=7.39, =1.84), we computed p^+ and the upper-bound p-value by hand and by using R codes. Results suggest moderate sensitivity of the study inference to an unobserved confounder, as when is 7.39 and is 1.84, *ORyx•cu* is no longer significant (with a p-value of .08).

Methods to Obtain ORyx•cu with Confidence Interval

While Rosenbaum's approaches were primarily concerned with finding the point at which effects became non-significant, the following methods quantify the unobserved confounder under certain specifications and then arrive at an estimate of the target of interest, $ORyx \cdot cu$ (i.e., the true relationship between x and y), and an associated confidence interval, adjusting for the unobserved confounder. Two groups of methods are presented. The first group (Greenland's and Harding's approaches) utilizes the association between x, y and u to create the actual data as if u was observed, which is then used to estimate the $ORyx \cdot cu$. The second group (Lin et al.'s and VanderWeele and Arah's approaches) utilizes the association between x, y, and u to compute an adjustment or bias factor that is then used to obtain the $ORyx \cdot cu$.

Greenland's and Harding's approaches

Both approaches break down the observed $2x^2$ table of x and y into eight combinations of x, y and u, imagining the data that would be observed if u was observed. Table 2a presents the

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observed 2x2 cross-tabulation of x and y. Table 2b presents the 2x2 cross-tabulation of x and y stratified by u, which we would see if u was observed. The goal is to estimate the cell counts a-h by specifying aspects of u so that Table 2b can be re-created.

Greenland's and Harding's approaches slightly differ in the necessary parameter inputs and in the manner in which these *a*-*h* cell counts are obtained. In his paper, Harding (2003) set *u* to be evenly distributed in the population (i.e., p(u)=0.5), but this can be easily modified. Harding's approach requires the specification of *ORyu* and *ORxu*. Given those values and the observed data ((a+e), (e+f), (c+g), (d+h)), these eight cell counts (*a*-*h*) can be solved using Equation 3.

$$af/be=ORyu$$
 $ch/dg=ORyu$ $ag/ce=ORxu$ $bh/fd=ORxu$
 $a+b+c+d=e+f+g+h$ (Eq. 3)

Instead of specifying *ORyu* and *ORxu* Greenland (1996) specifies the prevalence of the unobserved confounder in the unexposed individuals (p(u|x=0)) and the exposed individuals (p(u|x=1)).

Greenland's approach then finds the cell counts *e*-*h* using the values specified by the user (p(u|x=1), p(u|x=0) and ORyu) through Equation 4.

$$\begin{array}{ll} g=p(u|x=1)\times(c+g) & e=p(u|x=0)\times(a+e) \\ h=\frac{ORyu\times(d+h)\times g}{ORyu\times g+(c+g)-g} & f=\frac{ORyu\times(b+f)\times e}{ORyu\times e+(a+e)-e} \end{array} \tag{Eq. 4}$$

Instead of specifying p(u|x=1) and p(u|x=0), an alternative approach is to specify *ORxu* and p(u|x=0) and then calculate the implied p(u|x=1) using Equation 5.

$$p(u|x=1) = \frac{ORxu \times p(u|x=0)}{1 - p(u|x=0) + ORxu \times p(u|x=0)} \quad \text{(Eq. 5)}$$

Once we know four of the eight cell counts, the values of a-d can be easily obtained from the observed data and the values of e-h using simple algebra.

For both Harding's and Greenland's approaches, *ORyx•cu* is then estimated by using these cell counts as frequency weights to re-create a dataset that contains information on the unobserved confounder. From this re-created data, a weighted logistic regression can be performed to obtain *ORyx•cu*, including its confidence interval (Harding, 2003). It is also possible to supplement the original data by explicitly creating the number of observations that reflect each cell count and then performing regular logistic regression.

Lin et al.'s and VanderWeele and Arah's approaches

Although both Lin et al.'s (1998) and VanderWeele and Arah's (VanderWeele and Arah 2011, Arah et al. 2008) approaches also aim to arrive at the *ORyx•cu* with a confidence interval, they use a slightly different approach than the Greenland and Harding approaches discussed above. It is important to note that Lin et al.'s and VanderWeele and Arah's approaches both allow relaxing the no-three-way-interaction assumption. In addition, VanderWeele and Arah's approach allows the relationships between the unobserved confounder and the exposure and outcome to vary as a function of observed covariates. To ease the comparison between these methods to other methods presented above, we maintain these assumptions in the motivating example.

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In Lin et al.'s approach, the information about the relationship between *y* and *u* is first summarized by an equation that estimates an adjustment factor (*AF*) derived from the following parameters specified by the researcher: p(u|x=1), p(u|x=0), OR(yu|x=1) and OR(yu|x=0) using Equation 6.

$$AF = \frac{OR(yu|x=1) \times p(u|x=1) + (1-p(u|x=1))}{OR(yu|x=0) \times p(u|x=0) + (1-p(u|x=0))} \quad (\text{Eq. 6})$$

When assuming no three way interaction, i.e. OR(yu|x=1) = OR(yu|x=0), the equation is simplified to Equation 7.

$$AF = \frac{ORyu \times P(u|x=1) + (1-p(u|x=1))}{ORyu + P(u|x=0) + (1-p(u|x=0))} \quad \text{(Eq. 7)}$$

ORyx•cu can then be calculated by dividing *ORyx•c* by *AF*. The confidence interval for *ORyx•cu* can be obtained using the same *AF* and dividing the upper and lower bounds of the confidence interval for *ORyx•c* by the *AF*.

VanderWeele and Arah (VanderWeele and Arah 2011; Arah et al. 2008) proposed a more general framework to assess sensitivity to an unobserved confounder, which can accommodate continuous or categorical outcomes and exposures and observed and unobserved confounders, while also relaxing the two assumptions discussed in the beginning of this paper. Briefly, they suggest that the bias for any outcome distribution can be estimated as long as one can specify two conditions for each level of the observed confounders, c. 1) the relationship between u and y across different levels of x (u denotes a chosen reference value for u), e.g., $\{E(y|x=x1, u=u, c) - E(y|x=x1, u=u, c)\}$ and $\{E(y|x=x1, u=u, c)\}$ $x=x^2$, u=u, c) – $E(y|x=x^2, u=u, c)$, and 2) the comparison between the prevalence of uwhen x is at different levels with the overall prevalence of u set, e.g., p(u|x=x1, c)-p(u|c) and p(u|x=x2, c)-p(u|c). Interested readers can refer to VanderWeele and Arah (2011) for formulas to apply this method. While attractive in its generality, VanderWeele and Arah (2011) also discuss the challenges in using such a general setting as it requires a large number of parameters to be specified. They recommended simplifying the approach for specific settings. Applying this general method to our specific setting (i.e. x, y, and u are all binary) and assumptions (i.e., 1) the relationships between y and u and between x and u are the same across different levels of c; 2) no three way interaction between x, y, and u), we can estimate the bias in the ORyx•c using the following simple Equation 8, which is essentially equivalent to Equation 7 in Lin et al.'s approach:

$$bias = \frac{1 + (ORyu - 1)p(u|x=1)}{1 + (ORyu - 1)p(u|x=0)}$$
 (Eq. 8)

Similar to Lin et al.'s approach, we can then divide the *ORyx•c* by this bias term to obtain *ORyx•cu*, as well as its confidence intervals.

Application of Methods to Obtain ORyx•cu with Confidence Interval

As a reminder, to ease the comparison between methods, we fixed the values of ORxu and ORyu at the two highest observed values (ORxu=7.39; ORyu=1.84). For methods that require the prevalence of the unobserved confounder, we specified a range of p(u|x=0) from 1% to 25% and then obtained p(u|x=1) using p(u|x=0) and ORxu=7.39 (see the eAppendix for detailed computation and sample R code). Results are summarized in Table 3 for the

range of p(u|x=0) specified. Similar results were observed across different methods, with $ORyx \cdot cu$ estimates decreasing with increasing p(u|x=0). However, the confidence interval never included one, suggesting that the study inference is not sensitive to an unobserved binary confounder that is associated with seven-fold increased odds of having a mother die from suicide as compared to accident and approximately two-fold increased odds of the offspring being hospitalized for suicide attempt.

Comparisons and Synthesis of Methods

The seven approaches described differed in their targets of interest, implementable study designs, as well as the necessary specifications of parameters. Rosenbaum's approaches focus on obtaining the value (upper bound of ORxu) and/or (upper bound of ORyu) at which *ORyx*•*cu* becomes non-significant. Given that it uses information on the actual number of pairs in the study, this method reflects the uncertainty of the analysis associated with sample size. The results of sensitivity analyses may then change as sample size changes: the values of and tend to be slightly larger as sample size increases. In other words, the study conclusion tested may appear to be more robust when the sample size is large. The other class of methods we described, which we labeled the $ORyx \cdot cu$ with confidence interval approaches, obtains both the point estimate of the treatment effect and its confidence interval. This group of methods does not use the actual sample from the original study, thus sensitivity results do not change as a function of the sample size. Additionally, these seven methods also differ in the study designs in which they can be implemented. The software to implement Rosenbaum's approach can generally only be applied to 1:1 matching designs. Compared to Rosenbaum's approaches, the ORyx•cu with confidence interval approaches are more flexible in that they accommodate any study design.

Each of these techniques requires the specification of a different set of parameters related to the unobserved confounder. The differences in the parameters specified across methods suggest that researchers should choose a method based on the parameters that they feel comfortable specifying, or on which there is existing literature. The advantage of Rosenbaum's approach is that users are able to directly specify the values of both *ORyu* and *ORxu*, which may be more readily obtained from the literature. However, unlike the other approaches descried in the paper, Rosenbaum's approach does not allow users to conduct sensitivity analysis on results from published studies, since it requires the number of discordant pairs, which is not always readily available. Greenland's, Lin et al.'s and VanderWeele and Arah's approach does not require the specification of *p*(*u*/*x*=*1*).While Harding's approach does not require the specification of these parameters, the computation is more involved.

As a motivating example, we applied these seven methods to assess the sensitivity of an observed relationship between maternal suicide and offspring's hospitalization for suicide attempt to an unobserved binary confounder, such as genetic predisposition to suicidal behavior. Rosenbaum's primal approach suggests that the study inference will be no longer significant when the upper bound of *ORxu* is greater than 1.55. While this would imply sensitivity to unobserved confounding, it is important to keep in mind that this method assumes a situation in which the unobserved confounder perfectly predicts the outcome of interest, e.g., genetic predisposition to suicidal behavior perfectly predicts offspring's hospitalization for suicide attempt. As a result, this method may overstate the study sensitivity.

The simultaneous approach yielded an inference closer to the approaches that estimate *ORyx•cu* with its confidence interval, which suggests that the association between maternal

suicide and offspring's hospitalization for suicide attempt is relatively robust to an unobserved confounder. However, the simultaneous approach still suggested that the Kuramoto et al. (2010) study is somewhat sensitive to an unobserved confounder. This evidence of this heightened sensitivity may relate to the fact that Rosenbaum's approaches depend on the sample size in the original study and, in particular, the number of discordant pairs, which is relatively small in the Kuramoto et al. study since the outcome is rare.

Other Sensitivity Analysis Methods

Although we focused on a few particular sensitivity analysis techniques because of their relative ease of implementation and fairly intuitive explanations, other sensitivity analysis methods are available. For example, Schneeweiss (2006) described methods that quantify the unobserved confounder under certain specifications and then arrive at an estimate of ORyx•cu, but do not provide an estimate of the confidence interval. Harding (2009) used a sensitivity analysis based on omitted variable bias calculations for ordinary least squares regression (an approach relatively common in economics; Harding, 2009). Ridgeway (2006) and McCaffrey, Ridgeway, and Morral (2004) discussed a method that can be applied to studies that utilize propensity score weights and is available in the 'twang' package for R (Ridgeway, 2006). Methods are also available that yield nonparametric bounds on the treatment effects, without characterizing the relationship between the unobserved confounder with the exposure and the outcome (Manski, Sandefur, McLanahan, & Power, 1992). A method using Monte Carlo methods and Bayesian analysis techniques to allow sensitivity parameters to come from specified distributions is described in McCandless, Gustafson, & Levy (2007) and Steenland & Greenland (2004), with R and WinBUGS code provided. One of the advantages of simulations methods is that the confidence intervals obtained may be more accurate than those provided by the simpler approaches. See Arah et al. (2008) for a practical example of sensitivity analysis using Monte Carlo simulation.

Discussion

While observed confounding in non-experimental studies can be addressed by methods such as propensity score matching, researchers have not had good tools to handle potential unobserved confounding. This paper discussed different methods of sensitivity analysis that quantify the sensitivity of study results to unobserved confounding. When applying these methods to Kuramoto et al. (2010), they yielded a fairly consistent result suggesting that the relationship between maternal suicide and offspring's hospitalization for suicide attempt is relatively robust to an unobserved confounder strongly associated with maternal suicide and moderately associated with offspring's hospitalization for suicide attempt. The sensitivity analysis gives us the confidence to conclude that the observed association between parent's suicide and off spring's suicide attempt is likely to be causal. Our study conclusion suggests that prevention strategies need to particularly focus on individuals whose parents died from suicide, for example by ensuring access to counseling services. Future studies should replicate the study findings to establish true causality, and of course most studies, this one included, also have other limitations, such as measurement error and other threats to validity. In addition, future studies should further investigate the mechanism of such a relationship, such as behavioral changes that might mediate the relationship between parent's suicide and offspring's hospitalization for suicide attempt.

Most of the methods described in this paper can be implemented relatively easily with available software packages or by hand. The preferred sensitivity analysis for a particular study may be driven by the target of interest, the study design, the details of a potential unobserved confounder that can be easily specified using external sources, and its ease of implementation. However, as the methods discussed rely on slightly different sets of

assumptions and come from different perspectives, it may also be helpful to explore several sensitivity analysis tools. In addition, it is important to note that the second set of methods ($ORyx \cdot cu$ with confidence interval) do not require the original data and can be implemented using just the results in a published study. This allows researchers to conduct sensitivity analyses for published results, enabling them to determine how much confidence should be placed in those results.

Although more general and more complex approaches are available, the availability of these simple approaches leaves researchers little excuse for not performing sensitivity analyses when conducting non-experimental studies. Although we are not aware of their use in that context, these methods may also be useful for non-experimental comparisons conducted within the context of randomized trials, such as to handle non-compliance (e.g., Jo and Stuart, 2009). We hope that this paper will raise awareness and the use of these important methods. By giving researchers insight into how sensitive studies may be to unobserved confounding, these methods can serve a critical role in non-randomized studies aiming to establish causal relationships. Importantly, these methods can inform prevention researchers and policy makers when drawing conclusions from non-experimental studies.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Summary of Sen	sitivity A	nalysis								
	Rosenbaum				ORys	*cu with Confidence Interval				_
	Primal		Dual	Simultaneous	Gree	ıland	Harding	Lin	VanderWeele & Arah	_
Target of interest	Non-signific	ance of test statistic			ORyx with c	•cu :onfidence interval				
Study Design	1-1 matched	pairs			Any s	tudy design				-
Outcome Distribution	Any				Binar	, ,	Binary	Binary, continuous, or censored	Any	
Parameters obtained from the data	•	Number of discordant pair	SI			• 2x2 joint cells from x and y	• 2x2 joint cells from x and y	• ORyx+c	• ORyx*c	
Parameters set by the method	•	ORyu= infinity	ORxu= infinity	• None		• None	• None	• None	• None	
Parameters explicitly set	•	ORxu	• ORyu	 ORyu 		• ORyu	ORyu	OR(yu x=1)	• ORyu	-
by the user				• ORxu		 ORxu 	• ORxu	• $OR(yu x=0)$	• $p(u x=1)$	
						• p(u x=0)		• p(u x=1)	• p(u x=0)	_
								• p(u x=0)		_
Pros		Reflects uncertainty given	1 sample size			Can vary both ORyu and ORy	xu			
	•	Easy to implement				 Yields confidence interval 				
	•	Does not require the speci-	ification of the prevalence of u			 Can accommodate any study 	design			
						Do not need the raw data to c	onduct sensitivity analysis			
	.	Gives a conservative resul	It	Can vary both and OR x1	ORyu	 Easier to implement (e.g. command to Hardino's 	Does not require the specification of the mevalence of u	Easy to implement	Easy to implement	-
						method)		Can accommodate more general settings	Can accommodate more general settings	
								Allows three way interaction between x, y and u	Can accommodate the violation of assumptions	
Cons		Confidence interval for the	e estimate not easily available			 Need to specify p(u x=1) 	More involved computation	Need to specify p(u x=1)	Need to specify p(u x=1)	
		Implementation limited to Requires the number of dis	o matching design scordant nairs, which may not always be readily	available		or preta-uj				
										_
Software for implementat ion	•••	rbounds in Stata or R Love's Excel spreadsheet	Hand computation (or with standard statistical software)	Hand comput with standard software)	ttion (or statistical	 Hand computation(or with standard statistical software) Regression analysis 	System equation solver (or with standard statistical software) Regression analysis	Hand computation (or with standard statistical software)	Hand computation (or with standard statistical software)	

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Table 1

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Table 2a

Observed Data: 2x2 Table for x and y

	y=0	y=1
x=0	a + e	b + f
x=1	c + g	d + h

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Table 2b

Underlying True Data: 2x2 Table for x and y Controlling for u

		y=0	y=1
u =0	x=0	а	b
	x=1	с	d
u =1	x=0	е	f
	x=1	g	h

Table 3

Estimated ORyx•cu (95%CI) Using Sensitivity Analysis Methods that Obtain ORyx•c and Confidence Interval^a

p(u x=0)	10/2	207	1002	150%	2006	750/2
Approach	T /0	0	10.01	0/ CT	0/07	0/ 57
Greenland	1.77 (1.42,2.22)	1.57 (1.24,1.98)	1.46(1.15, 1.86)	1.42 (1.12,1.82)	1.41 (1.10,1.80)	1.40 (1.10,1.79)
$\operatorname{Harding}^{b}$	1.77 (1.42, 2.22)	1.57 (1.24, 1.98)	1.46(1.15,1.86)	1.42 (1.11,1.82)	$1.40\ (1.10, 1.80)$	1.41 (1.10,1.80)
Lin/VanderWeele and Arah	1.77 (1.42, 2.21)	1.57 (1.26, 1.96)	1.46 (1.17, 1.82)	1.42 (1.14,1.77)	1.41 (1.13,1.75)	1.41 (1.13,1.76)

^aORyu=1.84, ORxu=7.39, ORyx•c=1.86

b p(u) was obtained from the specified p(u|x=0), p(x)=50% and ORxu=7.39.