TUTORIAL IN BIOSTATISTICS





STRATOS guidance document on measurement error and misclassification of variables in observational epidemiology: Part 2—More complex methods of adjustment and advanced topics

We continue our review of issues related to measurement error and misclassification in epidemiology. We further describe methods of adjusting for biased estimation caused by measurement error in continuous covariates, covering likelihood methods, Bayesian methods, moment reconstruction, moment-adjusted imputation, and multiple imputation. We then describe which methods can also be used with misclassification of categorical covariates. Methods of adjusting estimation of distributions of continuous variables for measurement error are then reviewed. Illustrative examples are provided throughout these sections. We provide lists of available software for implementing these methods and also provide the code for implementing our examples in the Supporting Information. Next, we present several advanced topics, including data subject to both classical and Berkson error, modeling continuous exposures with measurement error, and categorical exposures with misclassification in the same model, variable selection when some of the variables are measured with error, adjusting analyses or design for error in an outcome variable, and categorizing continuous variables measured with error. Finally, we provide some advice for the often met situations where variables are known to be measured with substantial error, but there is only an external reference standard or partial (or no) information about the type or magnitude of the error.

KEYWORDS

 $Bayesian\ methods,\ bias\ analysis,\ distribution\ estimates,\ likelihood\ methods,\ moment\ reconstruction,\ multiple\ imputation$

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1 | INTRODUCTION

In the first part of this article we presented the basic concepts underlying the effects of measurement error and misclassification of variables, described validation and other types of studies that provide information regarding the statistical properties of the error involved in measurement, discussed study design and impact of measurement error on sample size, and presented some methods of adjusting inference for measurement error in simple but commonly occurring situations in epidemiology. In this second part, we present some more complex methods of adjusting estimates or inference for measurement error and misclassification (Section 2), discuss methods to estimate a distribution of an outcome subject to error (Section 3), review the software available for performing such analyses (Section 4), and describe some recent developments regarding more advanced problems (Section 5). The methods described in both Parts 1 and 2 of our tutorial are based on knowledge of the type and magnitude of the measurement error. In Section 6, we provide advice on how to deal with the all-too-common situations in which such information is imperfect or not available for study participants.

2 | ANALYSIS OF STUDIES WHERE ONE OR MORE OF THE MAJOR COVARIATES IS MEASURED WITH ERROR—MORE COMPLEX METHODS OF ADJUSTMENT

In Section 6 of Part 1, we described two methods of adjusting estimates of association between exposure and outcome when a continuous exposure is measured with error—regression calibration (RC) and simulation-extrapolation (SIMEX). Each of these methods is conceptually simple. For RC, the exposure measured with error is replaced by a predicted value of the true exposure and the main analysis proceeds as usual, albeit with adjustment for the SEs of the estimated association parameters. With SIMEX, one repeatedly introduces more measurement error to approximate a curve for the relationship between the measurement error variance and the regression coefficient in order to estimate the value of that parameter in the absence of measurement error. Sections 2.1-2.4 deal with some more complex but general methods for continuous variables that have measurement error. Some of these methods, such as the Bayesian approach or multiple imputation (MI), can also handle covariate misclassification. Section 2.5 discusses approaches for categorical variables subject to misclassification.

2.1 | Likelihood methods

Likelihood methods are pervasive in statistics. This section considers maximum likelihood estimation in measurement error problems. However, likelihood is also a building block for Bayesian inference, which will be discussed in Section 2.2. In the measurement error literature, discussion of maximum likelihood methods is given in the books by Carroll et al, ¹ Buonaccorsi, ² and Yi. ³

Figure 1 illustrates the steps in obtaining the likelihood function in order to carry out measurement error adjustment and perform the likelihood analysis. For non-Berkson error (ie, classical or linear measurement error model), these steps are as follows:

Step 1: Perform a likelihood analysis. One must specify a parametric model for every component of the data. Any likelihood analysis begins with the model one would use if X were observable. We denote the likelihood of this model as $f_{Y|X,Z}(Y|X,Z,\beta)$, where β denotes the parameters of the model. For example, in logistic regression, with $H(s) = \exp(s)/\{1 + \exp(s)\}$, the likelihood function is: $\{H(\beta_0 + X^T\beta_X + Z^T\beta_Z)\}^Y\{1 - H(\beta_0 + X^T\beta_X + Z^T\beta_Z)\}^{1-Y}$.

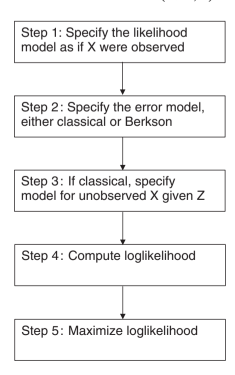
Step 2: Choose the error model. This could be a classical error model, a linear measurement error model, a Berkson model, and so on. Presuming non-Berkson error, the likelihood of the model for X^* given (X, Y, Z) can be denoted by $f_{X^*|X,Y,Z}(X^*|X,Y,Z,\alpha)$, where α denotes the parameters of the model. In the case of nondifferential classical measurement error, for example, if the measurement error is normally distributed with constant variance σ_U^2 , then $f_{X^*|X,Y,Z}(X^*|X,Y,Z,\sigma_U^2) = (2\pi\sigma_U^2)^{-\frac{1}{2}} \exp\{-(X^*-X)^2/(2\sigma_U^2)\}$. Note that this is a slightly stronger version of nondifferential error, which in general only requires X^* to be conditionally independent of Y given X. Here, X^* is conditionally independent of Y and Z given X.

Step 3: If one has a classical or linear measurement error model, specify a distribution for the unobserved X given the observable covariates Z, which we call $f_{X|Z}(X|Z,\gamma)$. The need to estimate the distribution of the unobserved X given Z is described in detail in chap. 8 of Carroll et al.¹ For example, one might assume that X is normally distributed with mean $\gamma_0 + Z^T \gamma_Z$ and variance σ_X^2 . In this example, $f_{X|Z}(X|Z,\gamma) = (2\pi\sigma_X^2)^{-\frac{1}{2}} \exp\{-(X-\gamma_0-Z^T\gamma_Z)^2/(2\sigma_X^2)\}$.

Step 4: Form the likelihood. When X is not observed and is continuous, the likelihood function of the observed (Y, X^*) given Z is $\int f_{Y|X,Z}(Y|X,Z,\beta)f_{X^*|X,Z}(X^*|X,Z,\alpha)f_{X|Z}(X|Z,\gamma)dX$. If X is discrete, the likelihood is $\sum f_{Y|X,Z}(Y|X,Z,\beta)f_{X^*|X,Z}(X^*|X,Z,\alpha)f_{X|Z}(X|Z,\gamma)$.

Step 5: Find the values of the parameters (β, α, γ) that maximize the likelihood.

As a brief aside, note that Steps 2 through 4 are specific to non-Berkson error and the analogous procedures for Berkson error would be rather different. Typically, a nondifferentiality assumption would be needed, so that the Step 1 specification is in fact for $(Y|X, X^*, Z)$. Then the other required specification is the Berkson model for $(X|X^*, Z)$, and the product of the two specified densities describes $(Y, X|X^*, Z)$. This is then integrated to yield a likelihood function based on $(Y|X^*, Z)$.



Steps 4 and 5 (or their counterpart in the case of Berkson error) involve the sometimes hard work of computing and maximizing the likelihood function to obtain parameter estimates. Because *X* is latent, that is, unobservable, these steps can be difficult or time-consuming, because one must integrate out the possibly high dimensional latent variable. Below, we provide a few details about computing and maximizing the likelihood function.

The overall likelihood based on a sample of n individuals is the product of each individual's likelihood function. Typically, one maximizes the logarithm of the overall likelihood in the unknown parameters. There are two ways to maximize the likelihood function. The most direct is to compute the likelihood function itself, and then use numerical optimization techniques. The second general approach is to view the problem as a missing-data problem, and then use missing-data techniques; see for example Little and Rubin, 4 Tanner, 5 and Geyer and Thompson. 6

Computing the likelihoods analytically is usually easier if X is discrete, as the conditional likelihoods are simply sums of terms. For likelihoods in which X is continuous, standard numerical methods for integration, such as Gaussian quadrature, can be applied. When sufficient computing resources are available, the likelihood can be computed using Monte Carlo techniques.

There are many computer routines for minimizing functions. Since we want to maximize the log likelihood, it is typical to multiply the log likelihood by -1 and then minimize it: the inverse of the Hessian matrix in such a computation serves as an estimate of the joint covariance matrix of all the parameters. See Section 4.4 for further comments on software for performing likelihood-based analyses.

The above description covers cases where X is not observed. In cases where X is observed for a subset of individuals in an internal validation study, the likelihood of the observed (Y, X, X^*) conditional on Z must be computed for those individuals separately from the remainder of the participants and then the two sets of likelihoods combined. Similarly, if the internal validation study involves measurement of, not X, but an unbiased measurement X^{**} of X, then an additional measurement error model must be specified for X^{**} and the likelihood of the observed (Y, X^*, X^{**}) , conditional on Z, computed separately for the individuals having measurements of X^{**} .

To illustrate the likelihood approach, we use an example already introduced in Part 1 of this article (Section 6). To recap briefly, the Observing Protein and Energy Intake (OPEN) study⁷ was a dietary intake validation study using unbiased reference measurements, conducted in 483 adult volunteers. Participants reported on their dietary intake using a food frequency questionnaire (FFQ), provided two 24-hour urine samples for measuring potassium intake, and provided samples for measuring total energy intake through a technique known as doubly labeled water. The target dietary measure is considered to be average daily potassium density intake, that is, the ratio of potassium intake to total energy intake. The questionnaire responses are considered to have linear measurement error and the urinary biomarker data are considered to have classical measurement error (since they measure only a single day's intake). The issue to be addressed is the association of log potassium density intake with a person's body mass index (BMI). The dataset is referenced as "Selected OPEN data" (2018).⁸

In this analysis, although each participant provided urine samples to measure potassium intake, we assume, as in Part 1, that these were available in only the first 250 participants, so that the "reference" measure is available in only a subsample of the 483 participants. The analysis was performed using the CALIS procedure in SAS. The models required in Steps 1-3, namely the BMI outcome model (model for Y), the FFQ log potassium density intake measurement error model (model for X*) and the log potassium density intake exposure model (model for X) are specified in the upper part of Table 1. Note that in this example X itself is not observed, even in the validation subset, and instead the biomarker log potassium density X^{**} , an unbiased measure of X, is observed in a subset. The measurement error model for X^{**} must also be included (see Table 1). Estimates of the regression coefficients in the outcome model are presented in the middle part of Table 1. These results are later compared with those of Bayesian methods (Section 2.2), moment reconstruction (MR) (Section 2.3), MI (Section 2.4), and RC (Part 1, Table 2)—see the discussions at the end of Sections 2.2, 2.3, and 2.4.

2.2 | Bayesian methods

Arguably there are advantages and disadvantages to taking a Bayesian standpoint when addressing measurement error problems. Perhaps the biggest advantage of a Bayesian approach to measurement error correction is an inherent logical and conceptual simplicity. After specifying appropriate submodels and prior distributions for the unknown parameters therein, the remaining steps are then a matter of computation. A joint posterior distribution of the unknown parameters ensues, and all inferences stem from this in a logical manner. However, this does require committing to specific prior distributions for all unknown parameters, and not all users will be comfortable with this.

TABLE 1 Analyses of the association of log potassium density intake with body mass index, using maximum likelihood estimation and Bayesian methods: data from the 483 participants in the OPEN study

Notation				
X = true log potassium densit	$y; Z_1 = \text{Sex}; Z_2 = \text{Age}; X^* =$	= FFQ log potassi	ium density; X^{**} = bion	narker log potassium density; $Y = BMI$
Models				
Outcome model	$Y X,Z_1,Z_2 \sim N(\beta_0 + \beta_X)$	$X + \beta_1 Z_1 + \beta_2 Z_2,$	${\sigma_{arepsilon}}^2)$	
Measurement error model 1	$X^* X,Z_1,Z_2 \sim N(\alpha_0 + \alpha_X)$	$X + \alpha_1 Z_1 + \alpha_2 Z_2,$	(σ_U^2)	
Measurement error model 2	$X^{**} X \sim N(X,\sigma_{X^{**}}^2)$			
Exposure model	$X Z_1,Z_2 \sim N(\gamma_0 + \gamma_1 Z_1 + \gamma_2 Z_1)$	$+\gamma_2 Z_2, \sigma_X^2)$		
Maximum likelihood results fo	r outcome model			
Coefficient	Estimated coefficient	SE	95% CI	P-value
Log potassium density, β_X	-7.19	1.25	-9.64, -4.74	<.001
Sex (F v M), β_1	-0.03	0.51	-1.03, 0.97	.95
Age (years), β_2	0.09	0.03	0.03, 0.15	.004
Bayesian results for outcome m	odel			
Coefficient	Estimated coefficient	Posterior SD	95% credible limits	Posterior probability of being >0
Log potassium density, β_X	-6.08	1.43	-9.38, -3.78	< 0.005
Sex (F v M), β_1	-0.30	0.51	-1.30, 0.71	0.27
Age (years), β_2	0.08	0.03	0.02, 0.15	>0.995

As soon as it was recognized that Markov Chain Monte Carlo (MCMC) computational techniques greatly expand the "domain of applicability" for Bayesian methods,⁹ applications to measurement error adjustment quickly followed.¹⁰⁻¹² An overview of the Bayesian approach to adjusting for measurement error is provided by Gustafson.¹³ Bartlett and Keogh¹⁴ make specific comparisons between Bayesian adjustments for measurement error and RC (Part 1, Section 6.1), maximum likelihood (Part 2, Section 2.1), and MI (Part 2, Section 2.4).

To give a specific example, suppose one has a parametric exposure model $f_{X|Z}(X|Z,\gamma)$ for (X|Z) parameterized by γ , a parametric outcome model $f_{Y|X,Z}(Y|X,Z,\beta)$ for (Y|X,Z) parameterized by β , and a parametric measurement error model $f_{X^*|X,Y,Z}(X^*|X,Y,Z,\alpha)$ for $(X^*|Y,X,Z)$ parameterized by α . Note that these are the same specifications as required for a likelihood analysis, as described in Section 2.1. Furthermore, assume there is a validation subsample, such that additionally the actual exposure X is observed for the first n of the N study subjects. Then the joint posterior density of all parameters (in this instance $[\alpha, \beta, \gamma]$) and latent variables (in this instance $X_{[n+1]:N}$) can be expressed as

$$f_{\text{post}}(\boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\gamma}, X_{(n+1):N} \mid X_{1:n}, (X^*, Y, Z)_{1:N}) \propto f_{X|Z}(X_{1:N} \mid Z_{1:N}, \boldsymbol{\gamma}) \times f_{Y|X,Z}(Y_{1:N} \mid (X, Z)_{1:N}, \boldsymbol{\beta}) \times f_{X^*|Y,X,Z}(X^*_{1:N} \mid (Y, X, Z)_{1:N}, \boldsymbol{\alpha}) \times f_{\text{prior}}(\boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\gamma}),$$

where the four terms on the right-hand side are, in order, the exposure model density, the outcome model density, the measurement error model density, and the prior density of all the parameters. MCMC methods can be applied to draw simulated samples from this joint posterior density, hence the drawn (α, β, γ) values (upon ignoring the drawn $X_{(n+1):N}$ values) are representative of the posterior distribution of the unknown parameters given the observed data. Bayesian point and interval estimates are thereby computed as appropriate summaries of this MCMC output. Note that this approach frees the user from having to explicitly join together two likelihood functions, one for the unvalidated observations and another for the validated observations as described in Section 2.1. Arguably this is a simplifying feature of proceeding in a Bayesian fashion.

As was seen to be the case with likelihood methods, using the Bayesian paradigm to "glue together" three submodels for exposure, outcome, and measurement has the appealing feature that uncertainty propagates across these submodels in a manner that is both principled and automatic. For instance, reported uncertainties (say posterior SDs or credible intervals directly computed from the MCMC output) about estimated outcome model parameters fully acknowledge the uncertainty about measurement error model parameters and exposure model parameters. So the data analyst is less

burdened by issues of whether uncertainties are correctly propagated than is the case for, say, RC (see Part 1, Section 6.1) or SIMEX (see Part 1, Section 6.2) approaches.

Against this coherence and logical simplicity, there are some challenges. General-purpose MCMC software for Bayesian analysis is available, including packages such as WinBUGS, JAGS, and STAN. However, some very specialized submodel specifications may not be supported by some packages. There is also the more pervasive issue that MCMC works better for some models and datasets than others, such that there is a need to examine the sampled draws to rule out problems with convergence and/or mixing of the MCMC algorithm. (MCMC methods draw realizations of a Markov chain specially constructed to have the posterior distribution of parameters and latent variables as its stationary distribution, relying on the fact that a Markov chain converges to its stationary distribution under weak assumptions.) Bayesian computing is not yet at the level of an "automated black box." In addition, it can be more challenging to relax modeling assumptions when working under the Bayesian paradigm. That is, going from parametric to semiparametric or nonparametric analysis becomes quite intricate, even though there has been much research on Bayesian nonparametric methods over the last decade. Articles by Sarkar et al^{15,16} and Sinha and Wang¹⁷ are recent examples that bring nonparametric Bayes technology into measurement error adjustment problems. Finally, as with all Bayesian analyses, some see the requisite specification of a prior distribution of the unknown parameters as a blessing, while others perceive it as a curse.

Much of the measurement error literature presumes a "hard" source of information about the measurement error magnitude, via observed replicates of a measurement X^* that has classical measurement error, or data from a validation subsample. Implicit here is the notion that if the amount of data increases in the right way, then the values of *all* the parameters, including those describing the measurement error process, would be revealed, that is, estimated consistently. However, Bayesian methods also offer the alternative possibility of using "soft" information. For instance, in the absence of replicates or a validation study, subject-area experts could assert a range of plausible measurement error magnitudes; a prior distribution that puts the vast majority of its mass on this plausible range could then be chosen. Of course, this sort of uncertainty would not diminish as more data are collected; so one must bear in mind that the final answer incorporates the usual statistical uncertainty arising because the sample size is finite, as well as the uncertainty in the experts' opinions about the measurement error magnitude. Related, there is no "free lunch." If one places a very diffuse (or even "improper") prior distribution on the measurement error magnitude, a correspondingly diffuse (or even "improper") posterior distribution will result. No useful measurement error correction can arise without either data or expert opinion to inform the magnitude of the measurement error.

We illustrate the Bayesian approach through the same example presented for the maximum likelihood analysis in Section 2.1. The analysis was performed in RJAGS. The exposure model, outcome model and measurement error models are the same models for the likelihood approach in Section 2.1 and are specified in the upper part of Table 1. Prior distributions with minimal information were adopted for the parameters of the exposure model, outcome model, and measurement error models; all regression coefficients were given normal priors (with mean zero and variance 1000) and precision (reciprocal of variance) parameters were given gamma priors (with shape and rate both set to 0.01). Estimates of the regression coefficients in the outcome model are presented at the bottom of Table 1. The results are rather similar to those of maximum likelihood presented in the middle of Table 1, and we will see they are also similar to those of the MR and MI approaches to be discussed in the next section. The posterior SD for the target parameter (describing the relationship between potassium intake and BMI) (1.43) is comparable to the SE obtained for the maximum likelihood estimate (1.25).

2.3 | MR and moment-adjusted imputation

MR and moment-adjusted imputation (MAI) are methods for handling covariate measurement error in which the goal is to construct a quantity $X_{\rm M}(X^*,Y)$ that has the same distribution as X, and such that $(X_{\rm M},Y)$ has the same joint distribution as (X,Y). If covariates Z are also to be included in the regression of Y on X, then the above distributions are conditional on Z. The quantity $X_{\rm M}(X^*,Y)$ is generally constructed by estimating moments of the joint distribution of (X,Y) from validation data, and then is substituted for X into the desired outcome regression model to produce an estimate β_X . SEs that account for the extra variability in the resulting estimate of β_X , which comes from the uncertainty in the parameter estimates used to construct $X_{\rm M}(X^*,Y)$, are necessary and can be obtained using the bootstrap. The bootstrap sample in this case is stratified on membership in the validation subset.

In MR,¹⁸ $X_{\rm M}(X^*,Y,Z)$ is constructed by matching only the first two moments of the joint distribution for (X,Y). In the case of classical measurement error this is achieved by defining $X_{\rm M}(X^*,Y,Z)$ as:

$$X_{\rm M}(X^*,Y,Z) = E(X^*|Y,Z) + G\{X^* - E(X^*|Y,Z)\},\tag{1}$$

where $G = \text{var}(X \mid Y, Z)^{1/2}\{\text{var}(X^* \mid Y, Z)\}^{-1/2}$. This expression can be extended to the linear measurement error model by replacing the first $E(X^* \mid Y, Z)$ on the right hand side with $E(X \mid Y, Z)$, keeping G as before. When the measurement error parameters are assumed known and the error is nondifferential, MR is equivalent to RC in linear regression, and is therefore consistent. When the measurement error is nondifferential and the error model parameters are estimated in an ancillary study, MR is not equivalent to RC, but both are consistent. MR is also consistent for logistic regression with normally distributed covariates, unlike RC which is only approximately consistent (Carroll et al^{1[p. 91]}). Under conditions of differential measurement error, MR is still consistent as the necessary moments are estimated conditional on Y, and in this it can prove advantageous compared to RC, which is biased.

MAI is an extension of MR, in which the moments of $(X_M(X^*, Y, Z), Y \mid Z)$ match more than the first two moments of $(X, Y \mid Z)$. Thomas et al²⁰ recommended matching the first four moments and observed that when X is normal, its performance is similar to RC in linear and nonlinear regression models. However, MAI has been shown superior to RC for logistic regression where the distribution of X is far from normal. MR and MAI may also be used for several covariates measured with error. 18,21

We illustrate MR with an example from the OPEN study, similar to the one in the previous two sections. We consider the regression of BMI on log sodium intake, while controlling for age and sex. We choose this example as there is evidence in the data that the measurement error in self-reported FFQ sodium intake is differential with respect to the BMI outcome variable, so that RC is an inappropriate method of adjustment. The estimated regression coefficients of the unadjusted model, using FFQ-reported sodium intake, are presented in the second column of part A of Table 2. The estimated coefficient for log sodium intake is 1.13, and its z-value 1.97. However, the z-value (unlike for a single covariate with nondifferential error) is invalid because the error is differential (see Part 1, Table 1). Parts B1 and B2 of Table 2 show the results of the models for E(X|Y,Z) and $E(X^*|Y,Z)$, which are both needed for the construction of $X_M(X^*,Y,Z) = E(X|Y,Z) + G\{X^* - E(X^*|Y,Z)\}$. The third column of Part A of Table 2 shows the MR adjusted estimated coefficients for log sodium intake, age, and sex. The SEs are obtained by bootstrap. One can see that the estimated coefficient for log sodium intake is 12.21, about 10 times larger than the unadjusted estimate, with a z-value of 4.53. This means that a 30% increase in sodium intake (change in log sodium intake of 0.26) is associated with an increase of 3.2 BMI units (95% CI: 1.8-4.5), rather than 0.3 units obtained from the unadjusted model. (Note that this very large effect cannot be directly causal since sodium is a micronutrient, supplying no calories. However, it indicates that high sodium intake is associated with higher BMI, probably because sodium intake is strongly correlated with energy intake.) There is also a notable change in the coefficient for sex from a nonsignificant negative association (z = -0.46) in the unadjusted analysis to a significant positive association (z = 3.16), that is, higher BMI in women than men for a given age and sodium intake (but see also the result for MI, given in Section 2.4).

MR can also be used as an alternative to RC when measurement error is nondifferential. In that case, it can be more efficient or less efficient than RC depending on the type of data at hand. If we apply MR to the same problem as used in Part 1, Section 6.1.1, namely the regression of BMI on log potassium density, we obtain a remarkably improved result over that from RC, in terms of the variance of the adjusted estimate. One can directly compare the estimates for RC presented in Part 1, Table 2 with those obtained for MR here in the third column of Table 3. The estimated regression coefficient for log potassium density is -8.13 (compared to -3.76 for RC) with a bootstrap SE of 1.77 (compared to 2.49 for RC). For the full result of the final model, see the third column of Table 3. In this case, MR is more efficient than RC and reveals a significant negative association of potassium density with BMI. The circumstances that cause the greater precision of MR over RC are (i) the quite strong association between outcome and exposure and (ii) the availability of the biomarker in more than 50% of the participants. In many studies, the association between the outcome and exposure variable is much weaker (eg, in studies of disease incidence) and the validation data are available in a much smaller proportion of participants, causing RC to be more efficient than MR (see Freedman et al¹⁹). We will further consider how MR compares to MI for this example in the next section.

2.4 | Multiple imputation

When there is an internal validation subset, in which (X, X^*, Z, Y) are all observed, then the case of measurement error is really just a problem of missing data.²² If individuals in the validation sample are a random sample of the main study population then the unobserved X is missing completely at random; if the sample is dependent by design on covariates, then X is missing at random (Little and Rubin, $^{4[\text{chap.1}]}$). In either case, the distribution of $X \mid X^*, Z, Y$

TABLE 2 Analyses of the association of log sodium intake with body mass index, using analyses unadjusted for covariate measurement error, moment reconstruction, and multiple imputation

Part A: Outcome regression parameter estimates			
Variable	Unadjusted analysis β (SE)	Moment reconstruction β (SE)	Multiple imputation β (SE)
Log sodium intake	1.13 (0.57)	12.21 (2.70 ^a)	11.02 (2.65 ^b)
Sex (F vs M)	-0.23 (0.51)	3.16 (1.00 ^a)	2.75 (1.45 ^b)
Age (years)	0.037 (0.029)	0.052 (0.035 ^a)	0.051 (0.070 ^b)
Part B: Regression models needed for Moment Recon	struction		
B1: Model of biomarker log sodium intake on BMI, sex, and age ^c	Estimated coefficient	SE	z-Value
Intercept	8.03	0.19	43.4
BMI (kg/m²)	0.031	0.004	7.81
Sex (F vs M)	-0.29	0.04	-7.00
Age (years)	-0.0027	0.0025	-1.11
Residual variance	0.0965		
B2: Model of FFQ log sodium intake on BMI, sex and age	Estimated coefficient	SE	z-Value
Intercept	8.43	0.170	49.53
BMI (kg/m²)	0.0071	0.0036	1.97
Sex (F vs M)	-0.28	0.04	-7.27
Age (years)	-0.0062	0.0023	-2.68
Residual variance	0.1736		
Part C: Regression models needed for multiple imput	tation		
Model of biomarker log sodium intake on FFQ log sodium, BMI, sex, and age ^{c, d}	Estimated coefficient	SE	z-Value
Intercept	7.54	0.45	16.93
FFQ log sodium intake	0.059	0.049	1.22
BMI (kg/m^2)	0.030	0.004	7.47
Sex (F vs M)	-0.27	0.04	-6.22
Age (years)	-0.0022	0.0025	-0.91
Residual variance	0.0963		

Note: Data from the 483 participants in the OPEN study.

is the same for those in the validation sample as for those who are not; thus, the values of X can be imputed from a model for $X \mid X^*$, Z, Y. MI, in which the unobserved values X are imputed m times, allows estimation of the coefficients in the outcome model and their SEs (Little and Rubin, $^{4[\operatorname{chap}.10]}$). Under a correctly specified model for $X \mid X^*$, Z, Y, MI will produce consistent estimates for β_X and consistent SEs. Like MR and MAI, and unlike RC, MI can handle differential measurement error, since Y is used for imputing the unknown X. The same procedure can be used if the error is assumed to be nondifferential, and a more efficient version of MI may also be constructed under this assumption. Freedman et al on that in circumstances where RC outperformed default MI, the "nondifferential" MI method performed similarly to RC. Here, we focus entirely on the default version, which accommodates differential error.

^aFrom a bootstrap sample of 5000.

^bFrom 500 multiple imputations.

^cBased on a random subsample of 250 participants.

^dUsed to impute biomarker log sodium intake.

TABLE 3 Analyses of the association of log potassium density intake with body mass index, using analyses unadjusted for measurement error, moment reconstruction, and multiple imputation

Variable	Unadjusted ANALYSIS β (SE)	Moment reconstruction β (SE)	Multiple imputation β (SE)
log potassium density	-1.69 (0.93)	-8.13 (1.77 ^a)	-7.28 (2.03 ^b)
Sex (F vs M)	-0.38 (0.49)	0.04 (0.53 ^a)	0.05 (0.73 ^b)
Age (years)	0.039 (0.29)	0.101 (0.035 ^a)	0.094 (0.048 ^b)

Note: Data from the 483 participants in the OPEN study.

As with any setting, the success of MI relies on having sufficient data to build a reliable imputation model and on correct specification of that model. For this reason, MI is generally not recommended when only an external validation study is available, and coincident measures of Y are not available. 19,23

It is also possible to use MI when there is an internal validation subset in which, instead of X, a measure of X that has classical measurement error is obtained, as well as X^* , Y and Z (a calibration study—see Part 1, Section 4.2). However, in this case, implementation of the method is a little more involved than usual MI. Details are provided in sect. A2 of appendix A in Freedman et al.¹⁹ This method is the one used in the examples that follow. Note, Keogh and White²⁴ described an MI approach for use in the setting of a replicates study, assuming availability of repeated measures of the error-prone covariate in some individuals, and assuming classical error. More recently, another approach for the setting of a validation or replicates study has been described²⁵ based on a modification of the substantive model compatible imputation approach for missing data described by Bartlett et al,²⁶ and accompanying software is available in R.²⁷

To illustrate MI, we use the same two examples as given for MR in Section 2.3. First, we consider the regression of BMI on log sodium intake, while controlling for age and sex. The results are presented in Table 2. Part C of the table shows the model that is used as a basis for imputing the unknown values of true log sodium intake. Note that this is based on a regression of the biomarker log sodium intake on the FFQ log sodium intake, BMI, age, and sex. One can see that the main variables influencing the imputation are BMI and sex. Note the strong effect of BMI in the imputation model does not imply differential error. The fourth column of Part A shows the estimated model of BMI on log sodium intake, age, and sex, based on 500 MIs. Generally, using a relatively large number of MIs is recommended, since one is imputing 100% of the values for the unknown true log sodium intake, although no formal recommendations have been established on the number of imputations required in this context. The results are similar, although not identical, to those obtained from MR (third column, part A of Table 2). The coefficient for log sodium intake is again about 10 times the unadjusted estimate and is highly statistically significant. The coefficient for sex is large and positive but, unlike for MR, does not attain conventional statistical significance. Discrepancies between the results of the two methods are not very common, but a careful analyst would perform both methods, where possible, to check on the stability of results. What is clear from the results of both methods is that the association of BMI with sodium intake appears far stronger than that indicated in the unadjusted analysis.

As mentioned with MR, one may use MI also in cases of nondifferential measurement error. We applied MI to the example of Part 1, Section 6.1.1, which was the analysis of the association of BMI with log potassium density. The results are shown in the final column of Table 3. The results are in accord with those of MR, indicating a strong negative association of BMI with potassium density intake. The same remarks made in Section 2.3, about the relative efficiency of MR compared to RC, apply also to MI. The results of MR and MI presented in Table 3, showing a strong negative association between BMI and log potassium density, are rather similar to those for the likelihood method presented in Table 1. However, the SE (1.25) of the regression coefficient for log potassium density intake is considerably smaller for the likelihood method than for the MR and MI analyses (1.77 and 2.03, respectively). A possible explanation for this is that the maximum likelihood analysis included all the data, whereas the MR and MI analyses, for simplicity, omitted 13 participants who provided one urine sample only (instead of two) for the measurement of potassium density. When the datasets are identical one would expect MI and maximum likelihood to yield very similar estimates and SEs.

The performance of MI has been compared with other methods, including RC and MR in settings of linear and logistic regression^{19,23,28} and Cox regression.^{22,29} These authors found that the optimal method depends on the size of the validation subset and degree of measurement error. Shepherd et al²⁸ noted MI worked well in the setting of correlated covariate

^aFrom a bootstrap sample of 5000.

^bUsing 500 multiple imputations, where the imputation model for the biomarker log-potassium intake was based on a random subsample of 250 participants and included the self-reported FFQ log-potassium density, sex, age (years), and BMI.

and outcome measurement error in the linear model. With censored survival data, implementing MI can be especially challenging; Bang et al²⁹ recommended implementing multiple methods to compare sensitivity of results to assumptions since in reality one rarely knows the true model for the error structure. In the case of a linear outcome and linear nondifferential measurement error model, a method of moments (MOM) approach can also be applied.² In this case we expect the performance of MOM to be similar to that of RC, as seen by Shaw et al.³⁰

2.5 | Analysis of studies where one or more categorical covariates are subject to misclassification

Section 2 of this article and Section 6 of Part 1 have addressed various methods that can be applied when a continuous covariate *X* is measured with error. The "methods menu" one can choose from when faced with a categorical covariate subject to misclassification is similar, but not identical.

Likelihood (Section 2.1) and Bayesian methods (Section 2.2) transfer directly and simply from the continuous covariate case to the discrete case. In fact, these methods are arguably more attractive in the discrete case, since concern about possible model misspecification (for the distribution of the unobservable X) is typically reduced. If X is binary and there are no precisely measured covariates, then X must follow a Bernoulli distribution, so that there is no concern about misspecification. If there are precisely measured covariates Z, then a model for $X \mid Z$ is required, and misspecification could arise. For instance, a logistic regression relationship between X and Z might be posited, and might be wrong. But more fundamental concerns about the *shape* of the X distribution do not apply when X is categorical. So likelihood and Bayesian methods, as per Sections 2.1 and 2.2, can be applied as described. The variable type for X is not particularly consequential for these methods.

In some simple situations, notably when Y and Z (if applicable) are also categorical, closed-form estimation of parameters is sometimes possible. This literature dates back to at least Barron³¹ who proposed the closed-form "matrix method" applicable when there are main study data in the form of a 2 by 2 table for binary (X^*, Y) and validation data in the form of a 2 by 2 by 2 table for binary (X, X^*, Y) . Subsequently, Marshall³² proposed an alternative closed-form estimator, known as the "inverse matrix method," by framing the classification in terms of predictive values rather than specificity and sensitivity. Later work, notably that of Morrissey and Spiegelman, ³³ Lyles³⁴ and Greenland³⁵ served to both (i), quantify the efficiency of these closed-form estimators relative to iteratively computed maximum-likelihood estimators, and (ii), understand nuances of how the various methods work under differential and nondifferential misclassification assumptions.

In more involved contexts, the expectation-maximization (EM) algorithm is quite straightforwardly applied to compute maximum likelihood estimates of parameters in the $(Y \mid X, Z)$ model.³⁶ Also, just as a categorical X variable is particularly amenable to the EM algorithm for likelihood estimation, it is also particularly amenable to MCMC methods (and Gibbs sampling specifically) for computing Bayesian estimates. See Joseph et al,³⁷ Gustafson et al,³⁸ Johnson et al,³⁹ and Prescott and Garthwaite⁴⁰ for examples.

Both RC and MR are less obvious strategies to pursue explicitly when X is categorical. However, some of the estimators discussed in Morrissey and Spiegelman,³³ Lyles,³⁴ and Greenland³⁵ indeed end up having a RC spirit. That is, they can be viewed as replacing X with an estimate of $E(X \mid X^*, Z)$. Also, MI can certainly be applied to problems involving a categorical X. In fact, this approach will be rather similar to a Bayesian analysis using MCMC computation.

The SIMEX method, described in Part 1, Section 6.2, has been extended to handle a categorical X variable that is subject to misclassification, using a method termed MC-SIMEX. Suppose we have a regression model with a discrete covariate X which is subject to misclassification.⁴¹ The misclassification process is described by the matrix Π , which is defined by its components

$$\pi_{ij} = \Pr(X^* = i | X = j), i = 1, \dots, r; \quad j = 1, \dots, r.$$
 (2)

 Π is a $r \times r$ matrix, where r is the number of possible outcomes for X. MC-SIMEX employs the function ($s \ge 0$) defined by:

$$\beta_{X^*}(s) = \beta_{X^*}(\Pi^s),\tag{3}$$

where $\beta_{X^*}(\Pi^s)$ denotes the value of the coefficient β^* when X^* is subject to misclassification by Π^s , defined as $E\Lambda^sE^{-1}$, with Λ being the diagonal matrix of eigenvalues and E the corresponding matrix of eigenvectors. For integer values of s,

TABLE 4 Data from a study of risk factors for sudden infant death syndrome (Kraus et al⁴³): Y = sudden infant death syndrome (case = 1, control = 0); X = antibiotic use during pregnancy according to medical record (yes = 1, no = 0); $X^* =$ antibiotic use during pregnancy according to mother's report (yes = 1, no = 0)

	X = 0	X = 1	X unobserved	Total
$Y=0, X^*=0$	168	16	479	663
$Y=0, X^*=1$	12	21	101	134
$Y=1, X^*=0$	143	17	442	602
$Y=1, X^*=1$	22	29	122	173

 $\Pi^{1+s} = \Pi^s * \Pi$, where * denotes matrix multiplication, and for s = 0, $\Pi^0 = I_{r \times r}$. The central idea of the MC-SIMEX method is to add extra misclassification to X^* . Namely, if X^* has misclassification probabilities Π in relation to variable X, and $X^*(s)$ is related to X^* by the misclassification matrix Π^s , then $X^*(s)$ is related to X by the misclassification matrix Π^{1+s} , when these two misclassification mechanisms are independent. Thus, the SIMEX algorithm can be applied to misclassification in the same manner as the original SIMEX. For details, including the variance estimation, see for example, Küchenhoff et al. 42

As an example of adjusting for misclassification in a binary explanatory variable X, we consider the study reported by Kraus et al⁴³ on risk factors for sudden infant death syndrome (SIDS). Here X is defined as an indicator of maternal use of antibiotics during pregnancy, as ascertained from medical record review, while X^* indicates the mother's self-report of antibiotic use on a questionnaire. The study employed a case-control design (though the same analysis would apply for a cross-sectional or prospective study), recruiting 797 controls (Y = 0) and 775 cases (Y = 1) of SIDS. Since medical record review was only conducted for a subset of 217 of the controls and 211 of the cases, we are presented with a misclassified data problem with an internal validation study. The data are presented in Table 4. If we ignore the X measurements available for the validated study subjects and simply focus on the (X^* , Y) association, we estimate a log odds ratio (OR) of 0.35 (corresponding to OR = 1.42), with a SE of 0.13, indicating a positive association.

A number of authors have illustrated misclassification adjustment methods using data from this study, including Greenland⁴⁴ and Chu et al, ⁴⁵ who contrast multiple methods, including maximum likelihood methods, Bayes methods, and SIMEX. Suspecting the possibility of differential misreporting, they applied a likelihood ratio test for the null hypothesis of conditional independence of X^* and Y given X to the data on validated subjects, obtaining some evidence against the null (P = 0.096). Thus, we focus on an adjustment method that allows for differential misclassification. While Section 2.1 alludes to general implementation challenges for likelihood methods arising because X is latent, obtaining maximum likelihood estimates in the present context is actually quite straightforward. As detailed by Lyles,³⁴ we can reparameterize the problem in terms of the (X^*,Y) and $(X|X^*,Y)$ distributions, rather than the (X,Y) and $(X^*|X,Y)$ distributions, with all the study units contributing to estimation of (X^*,Y) but only the validated study units contributing to the estimation of $(X|X^*,Y)$. Lyles³⁴ shows that one can then work back to obtain closed-form estimates and SEs of parameters in the original parameterization. Applying this method to the data at hand results in an estimated (X,Y) log odds-ratio of 0.19 (corresponding to OR = 1.21), with a SE of 0.22. Note that this adjustment for misclassification pushes the point estimate toward the null, as can arise when misclassification is differential. For comparison, if we presume nondifferential misclassification, then the maximum likelihood estimate of the (X,Y) log odds-ratio, as determined by numerical maximization, is 0.40 (corresponding to OR = 1.49), with a SE of 0.19. As must arise when nondifferential misclassification is presumed, relative to the naïve estimate, the adjustment moves the estimate away from the null, and increases the corresponding measure of uncertainty. Generally, this example is another where the medical conclusions to be drawn from the analysis are changed substantially by taking reporting error into account.

3 | ANALYSIS METHODS FOR ESTIMATING DISTRIBUTIONS

In Part 1, Section 3.4, we briefly discussed the impact of measurement error on estimating the distribution of a random variable Y. In this section, we consider methods for estimating the distribution of Y using error-prone observations Y^* . In contrast to prior sections, here the error-prone variable measures the outcome of interest (Y) rather than a covariate (X). We focus on the case where Y is continuous. For example, it might be of interest to know selected percentiles of the distribution of Y, or related quantities such as the interquartile range. Alternatively, one might wish to know what proportion of the distribution falls above or below specific cut-points. In Section 3.4 of Part 1, we briefly considered how such distributional quantities can be biased if estimated by an error-prone Y^* . Most of the methods presented here focus

on the case where Y^* follows the classical measurement error (model (1) in Section 2.1 of Part 1). Development of these methods has been most pronounced in the field of nutrition, where the desired random variable Y is the long-term average daily consumption ("usual intake") of a food or nutrient, and Y^* is typically the reported intake from a 24-hour dietary recall, which queries everything eaten or drunk during the previous 24 hours. Under this scenario, a substantial amount of the measurement error U is assumed to be due to day-to-day variation in diet that makes a single day an imprecise proxy for a long-term average. Other sources of systematic error are routinely assumed to be zero in these methods, although there are exceptions.

3.1 | A simple case

Consider the simplest case of model (1) (Part I), where $E(Y) = \mu_Y$, E(U) = 0, $var(Y) = \sigma_Y^2$ and $var(U) = \sigma_U^2$. Then $E(Y^*) = \mu_Y$ and $var(Y^*) = \sigma_Y^2 + \sigma_U^2$. It follows that the distribution of.

$$\widetilde{Y} = E(Y^*) + \sqrt{\frac{\sigma_Y^2}{\sigma_Y^2 + \sigma_U^2}} (Y^* - E(Y^*)),$$
(4)

has the same first two moments (μ_Y, σ_Y^2) as the distribution of Y. This approach, applied to interval data, was used by the National Research Council (NRC),⁴⁹ where replicated observations Y^* were available to permit separate estimation of the required variance components in Equation (4). In general, for this classical measurement error setting, one can estimate var(U) by the within-person variance when there are replicate measurements; var(Y) can then be estimated by subtracting the estimate of var(U) from $var(Y^*)$. If V and V are normally distributed, then V^* is as well, and matching the first two moments of V is equivalent to fully characterizing the distribution. Thus, the empirical distribution of V may be used as an estimate of the distribution of V. This approach of constructing a set of representative observations to be used as a basis for an empirical distribution estimator can be extended to the more complex cases discussed below.

3.2 Use of normality transformations

The NRC report⁴⁹ highlighted the fact that 24-hour recall data (Y^*) tend to be skewed, suggesting that the normality assumption is not tenable in the original scale. Therefore, transformations are routinely applied to observed data as a first step in distribution estimation. This is a complication that requires a careful choice of assumption about how Y and Y^* are related. In the NRC analysis, formula (4) was applied to log transformed data, and each estimated percentile of the distribution of \widetilde{Y} was exponentiated to obtain the corresponding percentile in the original scale. This approach is consistent with the model.

$$g(Y^*) = g(Y) + U, (5)$$

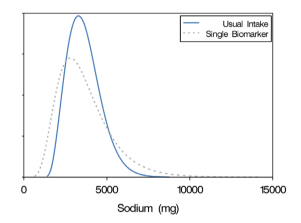
where $g(\cdot)$ is an invertible transformation. That is, Y^* is unbiased for Y on the transformed scale (and therefore biased for Y on the original scale). The transformation is also presumed to result in well-behaved (e.g., normally distributed) errors U.

We illustrate this approach with an example taken from data obtained in the OPEN study (for a short description of the study, see the example given in Section 2.1). Here we consider estimating the distribution of usual sodium intake in a population typical of those participating in the study. Besides the self-reported intakes (which have some bias), two measurements of 24-hour urinary sodium were available, a biomarker for sodium intake that is thought to be unbiased, but subject to random day-to-day variation and random assay error. Due to this random variation, these measurements (our Y^*) have error when the target is to measure usual (ie, long-term average) intake, Y. In addition, sodium intakes tend to have a skew distribution that is approximately log-normal in shape. We therefore assume model (5) where g is the logarithmic function. The value of the shrinkage factor $\sqrt{\frac{\sigma_Y^2}{\sigma_Y^2 + \sigma_U^2}}$ shown in Equation (4) (but applied on the logarithmic scale) was 0.72, indicating a relatively large day-to-day variation in sodium intake. Table 5 shows the percentiles of the distribution estimated from a single biomarker measurement Y^* assuming it has no random error (the incorrect assumption) vs that based on model (5). The latter is calculated from applying the NRC method to the first measurement of $\log(Y^*)$,

TABLE 5 Estimated percentiles of the distribution of usual sodium intake (mg/day) among a population typical of the participants in the OPEN study, using a single measurement of urinary sodium (unadjusted method) vs the NRC method applied to the log value of this measurement

Percentile	Unadjusted method (mg/day)	NRC method (mg/day)
5	1810	2233
10	2150	2530
25	2879	3126
50	3948	3928
75	5322	4876
90	6649	5729
95	7686	6363

FIGURE 2 Smoothed estimated densities of usual sodium intake per day among the OPEN study participants based on (i) a single 24-hour urinary sodium determination (single biomarker) and (ii) the NRC method of adjusting for the measurement error in a single determination (usual intake) [Colour figure can be viewed at wileyonlinelibrary.com]



estimating σ_Y^2 and σ_U^2 from the repeat measurement, and followed by back-transformation. Figure 2 presents the density functions after smoothing. Both Table 5 and Figure 2 show the substantial shrinkage of the distribution obtained when using the adjustment for measurement error.

Later authors, beginning with Nusser et al⁵⁰ assumed the model:

$$g(Y^*) = \mu + U,\tag{6}$$

where μ is the individual's long-term average of the transformed Y^* , and

$$Y = E_U(Y^*|\mu) = E_U(g^{-1}(\mu + U)|\mu),$$

which assumes that Y^* is unbiased for Y on the original scale. Because $g(\cdot)$ is typically nonlinear, estimating the distribution of Y now requires integration over the distribution of U. It is often impossible to decide from the available data which model, (5) or (6), is the more appropriate.

3.3 | Model-assisted vs model-based approaches

The NRC method uses the empirical distribution of \widetilde{Y} as the basis for estimating the distribution of Y. However, the routine use of normality transformations in later approaches permits the use of exact percentiles from a normal distribution as the basis for estimation. In small samples, the empirical distribution of \widetilde{Y} can be quite granular, leading to a granular approximation of the distribution of Y. Using normal distribution percentiles allows smooth estimated distributions of Y, but at the expense of relying on the normality assumption (after appropriate transformation). The NRC approach and extensions such as the Multiple Source Method method have been characterized as "model-assisted," in contrast to the other, "model-based" approaches. 55

3.4 | Extensions for inclusion of covariates, semicontinuous data, and multivariate estimation

A more general version of model (6) is given by the mixed-effects model.

$$g(Y^*) = \mu(Z) + r + U, \tag{7}$$

where $\mu(Z)$ is a function of observed covariates Z of an individual, r is a random effect that is constant across replicate observations on the same subject, and U is random within-subject error. Then,

$$Y = E(Y^* \mid r, Z) = E(g^{-1}(\mu(Z) + r + U) \mid r, Z).$$
(8)

Under (7-8), the variation in the distribution of Y comes in part from the variation explained by covariates Z and in part from residual deviations r. Model (7) can be fit using standard software for (nonlinear) mixed models. The software typically requires assuming normality of the random effects, which favors the use of model-based approaches. To maintain the proper distribution of covariates, a Monte Carlo simulation approach is often used to generate a dataset for calculating the empirical distribution mentioned earlier. In this approach, predictions of $\mu(Z)$ from sampled individuals are added to multiple randomly generated r values before integrating over the (presumed normal) errors U.

Many dietary components are "episodically consumed," where the single-day reported intakes (Y^*) can be zero, even if long-term intake (Y) is positive. The observed data Y^* can therefore have a large proportion of zeros, as well as a skewed distribution of positive values. In these situations, simple transformations applied to Y^* will not even approximate a normal distribution. Models for such semicontinuous data are motivated by writing the average value of Y^* as a conditional expectation:

$$E(Y^*) = E(Y^*|Y^* > 0) \Pr(Y^* > 0) + E(Y^*|Y^* = 0) \Pr(Y^* = 0) = E(Y^*|Y^* > 0) \Pr(Y^* > 0).$$
 (9)

This formulation expresses the average daily consumption as the product of the average consumption on consumption days $E(Y^* \mid Y^* > 0)$ and the probability of consuming on a specific day $Pr(Y^* > 0)$. This approach led several authors 51,53,54,56,57 to consider models that used binary indicators of zero vs nonzero consumption Y^* to inform estimation of the probability part of the model and used the transformed nonzero values of Y^* to inform estimation of the amount part of the model. These methods were further extended (Freedman et al 58 and Zhang et al 59) to allow flexible joint modeling of multiple components, which permits analysis of ratios and high-dimensional indices. A detailed description of these extended models is beyond the scope of this work.

3.5 Nonparametric estimation of distribution functions

There is a very large literature on nonparametric estimation of distribution functions. The articles concentrate on estimating the density function, and this is often called *density deconvolution*. This literature has two major themes. The first uses kernel density functions, while the second usually uses infinite mixtures of normal and/or Bsplines. For the first, see Carroll and Hall, ⁶⁰ Stefanski and Carroll, ⁶¹ and Fan, ⁶² while for the case of heteroscedastic measurement error, see Delaigle and Meister. ⁶³ For the second, see Staudenmayer et al ⁶⁴ and Sarkar et al. ¹⁵ For multivariate density deconvolution, see Masry ⁶⁵ and Sarkar et al. ⁶⁶ The articles by Sarkar et al ^{15,66} are very general, allowing heteroscedastic measurement error with unknown distributions for that measurement error, as well as of course unknown distributions for the latent variable. There is substantial software available for these estimation methods. References to online sources of the available code, which cover methods for kernel-based deconvolution and Bayesian semiparametric density deconvolution, are provided in Table 6. Nonparametric maximum likelihood is another approach considered by several authors. ^{67,68}

4 | SOFTWARE FOR ANALYSIS

One of the main barriers in the past to the use of the analysis methods described in Sections 2 and 3 was the lack of specific software for implementing them. The situation is now gradually improving. Here, we describe software programs,

TABLE 6 Software for estimating distributions (see notes in Section 4.3)

Package/procedure	Website location or information	Notes	References
SAS code for National Research Council method	https://www.nal.usda.gov/sites/default/files/fnic_uploads// dietary_planning_full_report.pdf Pages 207-208	The National Research Council method was devised for estimating distributions of usual dietary intake from reports of a single day's intake. SAS code for implementing the National Research Council method can be found in the source given in the References column.	National Research Council ⁴⁹
PC-SIDE (Iowa State University)	http://www.side.stat.iastate.edu/pc-side.php	Created with the same intention as the National Research Council method. Implements the method of Nusser et al which employs a nonparametric transformation of the data to normality, deconvolutes, and back-transforms to the original scale	Nusser et al ⁵⁰
MCRA 8.2	https://mcra.rivm.nl/	Implements the Monte-Carlo Risk Assessment (MCRA) method devised by researchers at Wageningen University and Research, Netherlands, This method was created for assessing risk from chemicals in the diet and includes the facility of creating a distribution of X from X^* .	van der Voet et al ¹⁵¹
NCI Method SAS macros	https://epi.grants.cancer.gov/diet/usualintakes/macros.html	Created for analyzing dietary data. For X^* measured in all individuals that, after suitable transformation, has classical measurement error. A substantial subsample should have at least one repeat value of X^* . X^* may be univariate, bivariate, or multivariate, and may have excess zeros. Allows inclusion of covariates in the modeling.	Tooze et al ⁵¹ ; Freedman et al ⁵⁸ ; Zhang et al ⁵⁹
SPADE	http://rivm.nl/en/Topics/S/SPADE/Access_to_SPADE	The Statistical Program to Assess Dictary Exposure (SPADE) is written in R and made available by the Netherlands National Institute for Public Health and Environment. X^* is univariate and may have excess zeros.	Dekkers et al ⁵³
MSM	https://msm.dife.de/	The Multiple Source Method (MSM) was developed by the German Institute of Human Nutrition for estimating distributions of usual dietary intake. Currently, there is a warning on the website regarding use of the program with covariates.	Haubrock et al ⁵⁴
Matlab and R code	https://researchers.ms.unimelb.edu.au/~aurored/links.html#Code	https://researchers.ms.unimelb.edu.au/~aurored/links.html#Code Nonparametric kernel density methods; including those in Delaigle and Meister 2008; Stefanski and Carroll 1990 and several other methods	Delaigle and Meister ⁶³ ; Stefanski and Carroll ⁶¹
Rcode	https://abhrastat.github.io/software.html	Bayesian semiparametric density deconvolution methods, including those in Sarkar et al 2014; Sarkar et al 2018, and other methods	Sakar et al ^{15,66}
Package <i>decon</i> in R language	https://cran.r-project.org/web/packages/decon/decon.pdf	Written by Wang XF and Wang B, this package computes the deconvolution kernel density estimator and its bandwidths from <i>X*</i> that has normal or Laplace homoscedastic errors or normal heteroscedastic errors with known variances. There the quality of its "tuning parameter" for (bandwidth) estimation has potential problems (Delaigle, 2014).	Wang and Wang ¹⁵² ; Delaigle ¹⁵³

macros, or packages that are now available for performing some of the methods described in those sections. Note that software for performing RC and SIMEX was described in Part 1, Section 7. We also provide the code that conducted our analyses of the OPEN data at https://github.com/PamelaShaw/STRATOS-TG4-Guidance-Paper.

4.1 | Software for Bayesian methods

Over the last several decades, the most common software used for applied Bayesian work has been *BUGS*. ⁶⁹⁻⁷¹ Fitting a Bayesian model to data using *BUGS* involves specifying model and prior distributions within the *BUGS* language, and then using both a *BUGS* interface and a *BUGS* engine to get the work done. In particular, the engine carries out MCMC sampling of the posterior distribution of parameters and latent variables given observed data. The interface serves to deliver the model and prior specifications and the data to the engine, and to then process the Monte Carlo output from the engine into inferential quantities. While the *BUGS* language is unique, a common point of confusion is that there are multiple possible engines and interfaces. Commonly used engine/interface combinations include *WinBUGS/R2WinBUGS*⁷² and *JAGS/rjags*. ⁷³ A worked example of implementation using R and JAGS is provided in supplementary materials to the article of Bartlett and Keogh 2018. ⁷⁴

Generally, and for measurement error modeling specifically, using *BUGS* is intermediate between a completely "do-it-yourself" workflow and a fully automated macro. The user need not have in-depth knowledge of MCMC algorithms, that is, the user is not required to select, code, and tune a particular algorithm. However, the user must express the chosen model and prior distributions in the *BUGS* language. On balance this seems a plus, as these specifications are then much more customizable than would be the case with a fully automated macro having a "hard-wired" model specification. Three specific examples of measurement error models expressed in the *BUGS* language appear in sect. 9.3 of Lunn et al.⁷¹ Many problems could be approached by extending one of these examples.

Regardless of how the workflow is implemented, and as alluded to in Section 2.2, using MCMC to compute point and interval parameter estimates requires somewhat more scrutiny and oversight than with other statistical methods. Issues of sampler "burn-in" and "mixing" arise, so that some level of human judgment is needed to assess whether the amount of Monte Carlo sampling utilized is indeed sufficient to numerically approximate posterior quantities well. This process is streamlined, but not automated, with the *BUGS* interfaces mentioned above. Simple summaries and diagnostic plots are readily provided to attest to the trustworthiness of the computational output.

Recently, the *Stan* probabilistic programming language has been developed, along with an *R* interface.⁷⁵ Compared to *BUGS* engines, *Stan* makes use of rather different MCMC algorithms, with excellent performance reported in many contexts. Chapter 11 of the *Stan* Reference Manual⁷⁶ illustrates the coding of a measurement error model in Stan.

One package that is specifically written for Bayesian analysis of measurement error problems in R is BayesME, available at http://www.stat.tamu.edu/~carroll/matlab_programs/software.php. This package is based on Sarkar et al^{15,16} and deals with nonparametric density and regression estimation when the measurement error is heteroscedastic, unknown, and may depend on X.

4.2 | Software for MR, MAI, and MI

When a validation substudy is available (Section 4.2), that is, in which the true X is observed, MI may be implemented using standard MI packages. Available packages include mice, and smcfcs in R, and smcfcs in R, and smcfcs in Stata, and PROC MI in SAS. The smcfcs package in R has been extended to accommodate measurement error correction in the settings of a validation substudy or a replicates substudy, and allows measurement error and missing data to be addressed simultaneously. 27

No packages are available for the implementation of MR or MAI. Thus, MR and MAI require a program to construct the "predicted" values of *X*. However, from thereon those predicted values may be used in standard regression programs to yield the measurement error adjusted estimates of the regression coefficients. Valid SEs of these estimates may then be obtained by bootstrap methods.

4.3 | Software for estimating distributions

As mentioned in Section 3, several packages or macros are available for estimating the distribution of a variable Y, using measurements Y^* that have classical measurement error. Most, but not all, of these packages have been developed for nutritional data but may be used for other types of data. All packages require that all individuals have at least one measurement of Y^* and that a substantial number have one or more repeat measurements. The user will note that some of these programs deal not only with continuous Y^* variables, but also with semicontinuous Y^* that have a positive probability of a zero value. Nevertheless, most of them assume that Y is continuous even when Y^* is semicontinuous, and they yield an estimated continuous distribution. SEs are typically obtained via replication methods such as the bootstrap (or under specific survey designs, balanced repeated replication). The packages and macros are summarized in Table 6. The resources listed in Table 6 include software for methods relying on a variety of necessary assumptions, including the semiparametric and nonparametric methods described in Section 3.5.

4.4 | Software for other methods

Section 2.1 includes some comments on what is needed to program likelihood methods. These methods are often implemented though custom-built programs. Rabe-Hesketh et al⁷⁹ describe how to conduct maximum likelihood estimation in *Stata* when *X* is normally distributed. In many problems of measurement error, the *SAS* procedures *MIXED*, *NLMIXED*, and *CALIS* can be used, and were used for the NCI Method *SAS* macros (see Table 6). The *SAS* manual has a nice introduction illustrating how *CALIS* can be used for measurement error modeling (see: https://support.sas.com/documentation/cdl/en/statug/63033/HTML/default/viewer.htm#statug_calis_sect001.htm), and the procedure can be used for nonlinear modeling as well.

The R software package SIMEX, introduced in Part 1 for error in a continuous exposure, also includes the *mcsimex* function, which implements the MC-SIMEX adaption for misclassified categorical exposures described in Section 2.5. 80.81 There is also a website that has *R* and *Matlab* programs (http://www.stat.tamu.edu/~carroll/matlab_programs/software. php), to deal with measurement error that is a mixture of classical error and Berkson error, as often occurs in radiation research and in other fields (see Section 5.1 for this topic).

5 | SPECIAL TOPICS

In Part 1 and in previous sections of this second part we have provided information regarding the effects of measurement error and misclassification on estimates obtained from some standard analyses, how to adjust for these effects and the software available to implement such adjustments. However, there is much that can be added to this basic information. In this section, we present a few selected more advanced topics. The first two topics concern situations where the data are affected by a mixture of types of errors. The third topic involves model building and variable selection in the presence of measurement error, the fourth topic involves the design and analysis of studies whose main outcome variable is measured with error, and the fifth topic involves categorization of continuous exposures that are measured with error.

5.1 Analysis of data subject to both Berkson and classical measurement error

We focus on epidemiologic projects involving an exposure that is measured by two or more methods, some of which involve Berkson errors and some classical-type measurement errors, which are then combined into a single measure. This occurs for example in radon studies, ⁸² and in radiation studies, such as at Hiroshima, ⁸³ the Nevada Test Site Thyroid Disease Study, ^{84,85} the Hanford Thyroid Disease Study, ⁸⁶⁻⁸⁸ and studies of the Chernobyl nuclear accident. ⁸⁹⁻⁹¹ There is a similar literature in occupational epidemiology, where direct measurements of exposure are taken on individuals, but other measurements of the same exposure are "grouped," for example, the time spent in the location of the specific exposure (eg, in a uranium mine), with an overall estimate of exposure at that location being derived from the combined information. In environmental epidemiology, exposure to pollutants might be based partly on a spatial model of pollution in that region and the amounts of time spent by the individual in different locations within the region, and partly on some direct measurements taken from an individual.

In all these types of problems, there is then a *calculated dose* X^* , a true dose X that is unobserved, and exactly measured covariates Z that are included in the outcome model and that are potentially related to X. We assume that, other than X, the covariates in the outcome model are measured exactly.

Analysis of such data is based on the statistical idea of linking the direct and indirect measurements via some version of a latent variable. Specifically, a *latent intermediate variable*, *L* links the true and calculated doses through a model such as

$$X = L + U_{\text{Berk}};$$

$$X^* = L + U_{\text{Clas}};$$

$$L = f(Z, \theta) + \varepsilon.$$

L can be hard to interpret; in some settings, it might be useful to think of L as an average underlying dose for a given set of covariates. Here, $U_{\rm Berk}$ is the component of Berkson error with mean zero and variance $\sigma_{\rm Berk}^2$, $U_{\rm Clas}$ is classical error with mean zero and variance $\sigma_{\rm Clas}^2$, $f(Z,\theta)$ describes the relationship of L to the covariates Z, and ε is the remaining variability not explained by the covariates or the measurement errors, with mean zero and variance σ_{ε}^2 . In the Nevada Test Site example, X= the true radiation dose, $X^*=$ the derived radiation dose, which relied on participant characteristics such as age, sex, and self-reported milk consumption; and we assume X and X^* are related via a latent variable L, where L may be a function of other precisely measured covariates Z determining exposure such as age, sex, distance from test site, and so on. If $U_{\rm Berk}$ has zero variance, then the above model is a purely classical measurement error model; if $U_{\rm Clas}$ has zero variance then the above model is a purely Berkson model. Carroll et al $^{1({\rm pp193-196})}$ provide more details on the analysis of such joint models, giving examples of the use of RC (see also Reeves et al 82 and Mallick et al 84) and maximum likelihood. In practice, knowledge of the sizes of the measurement error variances $\sigma_{\rm Berk}^2$ and $\sigma_{\rm Clas}^2$ is critical to analysis. This can be particularly difficult for Berkson errors (see Part 1, Section 4.2). In case of such difficulty, sensitivity analyses can be conducted.

The impact of a mixture of Berkson and classical errors depends critically on the ratio of σ_{Berk}^2 to σ_{Clas}^2 . When this ratio is very large and Berkson error dominates, the impact is close to that expected from purely Berkson error. When the ratio is small, the impact is close to that expected from classical error; and when the ratio is near one, and the Berkson error is nondifferential, then the impact is an average of the impacts of both—in other words estimated regression coefficients are attenuated, although to a lesser degree than with purely classical error, and loss of power is similar to that found with either classical error or Berkson error.

The literature given above includes a host of variations on the model given above, often specific to the application. For example, Li et al⁸⁵ consider the same model as above, but give reasons to allow the Berkson errors to be correlated among groups of individuals. For the Chernobyl accident, Masiuk et al⁹¹ argue that a better model has classical additive heteroscedastic measurement errors as well as Berkson multiplicative measurement errors.

5.2 | Analysis of exposure variables subject to both measurement error and misclassification

In previous sections, we have described methods for dealing with two distinct cases: (i) a continuous variable measured with error and (ii) a categorical variable subject to misclassification. What can be done, though, if data have a combination of both?

The only work we are aware of that combines issues of measurement error and misclassification in a single analysis is that of Spiegelman et al,⁹² White et al,⁹³ and Yi et al.⁹⁴ Each of these articles considers a main study/validation study design, where the validation study includes both the true and the error-prone observations of the variables subject to measurement error and misclassification, while the much larger main study has only the error-prone versions. In addition, White et al⁹³ considers a main study/replicates study design.

White et al⁹³ proposed a RC approach for a continuous outcome when there is both a continuous covariate and binary covariate subject to measurement error, and discuss the necessary supportive data for an identifiable model depending on whether a validation study or replicate data are available. Spiegelman et al⁹² consider a binary outcome, and use logistic regression with maximum likelihood to obtain estimates and inference. Yi et al⁹⁴ consider methods applicable for all generalized linear models, with a binary covariate subject to misclassification and a continuous covariate

measured with error. They discuss methods based on (i) full maximum likelihood, as in Spiegelman et al;⁹² (ii) an estimating function method based on ideas of semiparametric methods such as in Tsiatis and Ma⁹⁵ and Ma and Tsiatis;⁹⁶ (iii) an augmented RC method; and (iv) an augmented SIMEX method. Methods (ii)-(iv) aim at providing robustness to distributional assumptions about the error-prone continuous variable.

Of methods (ii)-(iv), the augmented SIMEX method is easiest to describe. We denote by $X_{\rm cont}$ and $X_{\rm cat}$ the continuous and categorical predictors subject to measurement error and misclassification, respectively. Their mismeasured versions are $X_{\rm cont}^*$ and $X_{\rm cat}^*$. The procedure is based on the idea that if $X_{\rm cont}$ were observed, then one has a simple misclassification problem, with $X_{\rm cat}$ misclassified, which can be solved by misclassification methods described in Section 2.5, for example by positing a model for the misclassification distribution of $X_{\rm cat}^*$ given $Z, X_{\rm cont}$, and $X_{\rm cat}$. Then one applies ordinary SIMEX to the method that would have been used if $X_{\rm cat}$ had been observed.

The augmented SIMEX and RC methods have the advantage that they are easily implemented using regular software for SIMEX or RC, once one has computer code also for solving a misclassification problem.

5.3 | Variable selection when some covariates are measured with error

In many biomedical settings, one seeks to develop a parsimonious regression model from a set of candidate predictors. We saw in Part 1, Section 3, that when there is at least one covariate in a multivariable regression model that is measured with error, the estimated coefficients for that X and any other covariates can be subject to bias because of the underlying correlation structures. Furthermore, as discussed in Part 1, Sections 3.1.3 and 3.2.3, when there are multiple such error-prone or misclassified covariates in a regression model, the direction of the bias can be in either direction (towards or away from the null). One way of viewing the cause of this bias in general linear regression is that measurement error induces bias in the least squares estimating equation so that its expectation at the true parameter vector β is no longer zero. Thus, many of the statistics used in variable selection procedures, such as the deviance or P-values associated with regression coefficients, will also be biased. Consequently, measurement error in one or more covariates puts any model selection procedure at added risk of selecting an incorrect set of variables and estimating regression coefficients with bias. This is not a concern for prognostic modeling, where only the risk prediction is of interest; however, it is a concern when one wishes to interpret the model coefficients or infer biological importance of the variables selected. Zhang et al⁹⁷ also remark that even when risk prediction is the sole interest, problems in model selection can occur when the measurement error structure in the data used to develop the prediction model is different from that in the data used for prediction.

There are many modeling procedures used to perform variable selection. Some methods have been developed to accommodate measurement error in variable selection and we highlight a few here. These methods have focused on penalized regression approaches, which use a penalty function to effectively control the model dimension. Whereas conventional stepwise procedures have been shown to be subject to instability and overfitting, 98-101 penalized regression procedures are becoming increasingly popular because of their better operating characteristics, particularly when there is a large number of candidate predictors relative to the sample size. 99,101,102 Penalized regression methods typically add a penalty to the usual parameter estimating equation (eg, the score), which then addresses both dimension reduction and parameter estimation in a single step.

For linear and partially linear regression models, Liang and Li¹⁰³ develop a corrected score type approach in which a term, proportional to β^{T} var(U) β , which offsets the bias caused by classical measurement error in the covariate vector X^* , is subtracted from the estimating equation to then achieve consistent estimation. Here var(U) is the measurement error covariance matrix. To this end the authors propose minimizing the following adjusted least squares expression:

$$\frac{1}{2} \sum_{i=1}^{n} \{ Y_i - X_i^{*T} \beta - \nu(Z_i) \}^2 - \frac{n}{2} \beta^T \text{var}(U) \beta,$$
 (10)

where $v(Z_i)$ is a general function of a precisely observed covariate Z, which is estimated with local splines. This adjusted least squares expression is incorporated into a general penalized regression framework and the authors discuss choices of the penalty, such as the L_0 , L_1 , or the smoothly clipped absolute deviation penalty (SCAD), that provide a variable selection framework. The authors show that under certain conditions, asymptotically, this procedure can perform as well as if the true model were known. The method assumes that var(U) is either known or can be estimated from repeat measurements of X^* . These authors also develop a similar penalized quantile regression procedure, building on the work of He and Liang¹⁰⁴ who had developed a quantile regression to handle covariate measurement error.

Ma and Li¹⁰⁵ develop a more general penalized estimating variable selection approach that can be applied to both parametric and semiparametric measurement error models. Their method is applicable to any consistent estimating equation, including generalized linear models, and can be applied to a large class of regression models.

Currently, we are not aware of any available software to implement the methods described above; however, software for penalized regression methods is now widely available and could be used as a base for building the required software for selection of error-prone variables.

In this section, we have highlighted only a few approaches in detail. Other approaches to address variable selection in high dimensional data include Datta and Zou,¹⁰⁶ Loh and Wainwright,¹⁰⁷ Sorensen et al,¹⁰⁸ Yang and Xia,¹⁰⁹ Tian and Xue,¹¹⁰ and Wang et al.¹¹¹ Zhang et al⁹⁷ discuss generally model selection in the setting linear regression models with measurement error.

5.4 Design and analysis when the outcome variable is measured with error

The effects of measurement error in outcome variables were outlined in Part 1, Section 3.3. Here, we discuss implications for study design and analysis and summarize methods for correcting for the effects of measurement error and misclassification of the type that produces biased estimates.

5.4.1 | Classical error in a continuous outcome

Classical measurement error in a continuous outcome variable in a linear regression does not result in biased estimates of regression coefficients (Part 1, Section 3.3.1). Therefore, a standard linear regression analysis can be used without any alterations. However, classical error in a continuous outcome results in lower precision of estimated regression coefficients and this should be accounted for in the study design. Consider the linear regression model $Y = \beta_0 + \beta_X X + \varepsilon$, where Y denotes the error-free outcome, and the corresponding linear regression model $Y^* = \beta_0^* + \beta_X^* X + \varepsilon^*$ using the error-prone outcome Y^* . The variance of the estimate is $\text{var}(\hat{\beta}_X^*) = \text{var}(\varepsilon^*)(X^TX)^{-1}$ and the relationship between variances of estimates from models using Y and Y^* is $\frac{\text{var}(\hat{\beta}_X^*)}{\text{var}(\hat{\beta}_X)} = \frac{\text{var}(\varepsilon^*)}{\text{var}(\varepsilon)}$. Let n_Y denotes the sample size required to achieve a desired SE for $\hat{\beta}_X$. To achieve the same SE for $\hat{\beta}_X^*$ therefore requires a sample size of $n_Y \frac{\text{var}(\varepsilon^*)}{\text{var}(\varepsilon)}$, where $\text{var}(\varepsilon^*) > \text{var}(\varepsilon)$.

5.4.2 | Systematic error in a continuous outcome

When the outcome variable has linear measurement error (see model (2) in Part 1, Section 2.1), that is $Y^* = \alpha_0 + \alpha_Y Y + U$, where U has mean zero and is independent of Y, the unadjusted regression coefficients estimated using $Y^* = \beta_0^* + \beta_X^* X + \varepsilon^*$ will be biased. Consistent estimates could be obtained by using $\frac{Y^* - \alpha_0}{\alpha_Y}$ in place of Y^* . This of course requires values for α_0 and α_Y , which may be known from previous studies or may need to be estimated. Buonaccorsi^{112,113} and Buonaccorsi and Tosteson¹¹⁴ devised methods for obtaining unbiased intervention effect estimates in this setting, when there is available either a validation study or replicates of an unbiased measure (eg, a biomarker) in a substudy. These methods were summarized by Carroll et al. ^{1(sect.15)} First, consider the setting with a validation study available, in which Y (as well as X) is observed. An estimate of β_X can be obtained from the data in the validation subset, and we denote this by $\widehat{\beta}_X^{(1)}$. This estimate is consistent but clearly inefficient because it is based on only a subset of the data. A second estimate, $\widehat{\beta}_X^{(2)}$, can be obtained from a regression of Y^* on Y. The variance-covariance matrix for $\widehat{\beta}_X^{(1)}$ and $\widehat{\beta}_X^{(2)}$, denoted Σ , can be obtained using a stacked estimating approach (Carroll et al ^{1[appendix A]} and Keogh et al ¹¹⁵). Note that the variance for $\widehat{\beta}_X^{(1)}$ and $\widehat{\beta}_X^{(2)}$ will have increased uncertainty from the added variability in U, as well as from the uncertainty in the estimated parameters $\widehat{\alpha}_0$ and $\widehat{\alpha}_Y$. Alternatively, bootstrapping can be used. An efficient estimator of β_X is then given by the "best weighted combination."

$$\widehat{\beta}_X^C = (J^T \Sigma^{-1} J)^{-1} J^T \Sigma^{-1} (\widehat{\beta}_X^{(1)T}, \widehat{\beta}_X^{(2)T})^T$$

where J = (I, I) and I is the identity matrix with the same number of rows as there are elements of β_X . The above result is general and extends to regressions with multiple covariates having a vector of regression coefficients β_X . In the simple setting of a single covariate X, the efficient estimator is

$$\widehat{\beta}_{X}^{C} = \frac{\widehat{\beta}_{X}^{(1)} \sigma_{(2)}^{2} + \widehat{\beta}_{X}^{(2)} \sigma_{(1)}^{2} - (\widehat{\beta}_{X}^{(1)} + \widehat{\beta}_{X}^{(2)}) \sigma_{(12)}}{\sigma_{(1)}^{2} + \sigma_{(2)}^{2} - 2\sigma_{(12)}},$$

where $\Sigma = \begin{pmatrix} \sigma_{(1)}^2 & \sigma_{(12)} \\ \sigma_{(12)} & \sigma_{(2)}^2 \end{pmatrix}$ is the variance-covariance matrix. This approach, based on a weighted combination of estimates, extends to the setting in which a replicates study is available instead of a validation study (see Part 1, Section 4.2). Suppose that two repeats of an unbiased measurement (eg, biomarkers) $Y_1^{**} = Y + U_1$ and $Y_2^{**} = Y + U_2$, with U_1 and U_2 independent, are available for a subset of individuals. A consistent estimate of β_X , again denoted $\widehat{\beta}_X^{(1)}$, can be estimated in the replicates substudy from a regression of $(Y_1^{**} + Y_2^{**})/2$ on X. The parameters α_0 and α_Y can be estimated in the replicates substudy using an MOM approach (Carroll et al^{1[sect.15]}).

If the values of parameters α_0 and α_Y are not known from a previous study, the need to estimate them should be accommodated at the design stage by incorporating plans and resources to conduct a validation or replicates substudy. Further research is needed to establish methods for optimal design of such studies, including the incorporation of information on the relative cost of the systematic-error-prone and biomarker measures.

5.4.3 Differential error in a continuous outcome

In some studies, the outcome measure is prone to differential error. This can arise in intervention studies with a self-reported outcome when participants are aware of their intervention group. We consider a measurement error model of the form $Y^* = \alpha_{0X} + \alpha_{YX}Y + U$ for two groups X = 0 and 1. This is a generalization of the linear measurement error model considered above. Differential error gives rise to biased estimates of the intervention effect; additional information is needed to estimate the form of the differential error so as to obtain consistent estimates of the intervention effect. Keogh et al¹¹⁵ described methods for analysis in the setting of dietary intervention trials in which the main differential-error-prone outcome measure is from a self-report and unbiased biomarkers are available in a replicates substudy, based on the Buonaccorsi¹¹² approach outlined above. The Buonaccorsi method extends directly to the differential error setting, with $\hat{\beta}_X^{(2)}$ being based on a regression of $\frac{Y^* - \hat{\alpha}_{0X}}{\hat{\alpha}_{YX}}$ on X. In particular, Keogh et al¹¹⁵ investigated the contribution of $\hat{\beta}_X^{(2)}$ to the estimator $\hat{\beta}_X^{(2)}$ (the weighted combination of $\hat{\beta}_X^{(1)}$ and $\hat{\beta}_X^{(2)}$) under different assumptions. It was shown theoretically that in the case of nondifferential error the combined estimator will be more precise than $\hat{\beta}_X^{(1)}$, while in the case of differential error nearly all the information about the intervention effect comes from the validation or replicates study and that $\hat{\beta}_X^{(2)}$ adds little in large samples. However, via simulation studies Keogh et al¹¹⁵ found that in finite samples, it is advantageous to use the self-report data in addition to the replicate biomarkers to estimate the intervention effect when the reliability of self-report measurements is comparable to that of the biomarker.

5.4.4 | Misclassification of a binary outcome

Section 2.5 noted that "matrix methods" can be used for handling a misclassified binary exposure when the outcome is also binary. Matrix methods can also be applied directly when instead it is the outcome that is misclassified, but they work only in very simple settings. Binary outcomes are more typically analyzed using logistic regression, and methods for correcting the impact of outcome misclassification in logistic regression analysis have also been devised.

For a study of *n* individuals, the full likelihood can be written:

$$L = \prod_{i=1}^{n} \Pr(Y^* = y_i^* | X = x_i) = \prod_{i=1}^{n} \sum_{y=0}^{1} \Pr(Y = y | X = x_i) \Pr(Y^* = y_i^* | Y = y, X = x_i), \tag{11}$$

where the logistic model of interest is logit($Pr(Y = 1 | X = x_i)$) = $\beta_0 + \beta_X x_i$. The misclassification probabilities in the second term of the likelihood, $Pr(Y^* = y_i^* | Y = y, X = x_i)$, can be expressed in terms of $Pr(Y = y | X = x_i)$ and the sensitivity

(Sn) and specificity (Sp) of Y^* . Magder and Hughes¹¹⁶ described estimation of β_0 and β_X using an EM algorithm. They described approaches for when the sensitivity and specificity are known, when they can be estimated from a previous validation study, and when they need to be estimated from the data. In the last situation, there are issues of identifiability if the model is saturated, and smoothing assumptions concerning the relation between covariates and outcome are needed to proceed. The authors caution against using this approach unless the smoothing assumptions are strongly believed. The probabilities of Y = y given Y^* and X are estimated in the E-step, and the logistic regression parameters β_0 and β_X are estimated in the M-step. Neuhaus, ¹¹⁷ Lyles and Lin, ¹¹⁸ and Lyles et al ¹¹⁹ instead used direct maximum likelihood estimation based on (11). They used the result that for the case of nondifferential misclassification, the likelihood can be expressed in terms of Sn and Sp, as follows:

$$L = \prod_{i=1}^{n} \left\{ (1 - \operatorname{Sp}) \Pr(Y = 0 | X = x_i) + \operatorname{Sn} \Pr(Y = 1 | X = x_i) \right\}^{y_i^*} \times \left\{ \operatorname{Sp} \Pr(Y = 0 | X = x_i) + (1 - \operatorname{Sn}) \Pr(Y = 1 | X = x_i) \right\}^{1 - y_i^*}$$
(12)

They considered sensitivity analyses assuming particular values for Sn and Sp, and making use of internal or external validation data. SAS code was provided.

A number of other authors have also considered sensitivity analyses for investigating the impact of outcome misclassification, incorporating uncertainty in the specified values for sensitivities and specificities. Fox et al¹²⁰ described probabilistic sensitivity analyses that involve simulating the data that would have been observed if there were no misclassification, given sensitivities and specificities. They focused on misclassified exposures but noted that the methods could also be applied for a misclassified outcome. Lyles and Lin¹¹⁸ described a "predictive value weighting" for handling misclassified exposures, which can also be applied for misclassified outcomes, and in the more complex scenario of misclassification in both outcome and exposures. This has been implemented in the *pvw* module in *Stata*. ¹²¹

Edwards et al 122 applied MI to handle misclassified outcome data when there is an internal validation study. A Bayesian approach can also be taken by assigning priors to the sensitivity and specificity. Some special considerations are needed when Y represents case or control status in a case-control study. 116,119,123

5.5 Misclassification due to categorizing continuous exposures measured with error

We have presented in Sections 2.2 and 3.2 of Part 1, and in Section 2.5 of this second part, problems arising from and methods for dealing with misclassified categorical variables. In this section, we discuss the special case where the categorical variable has been formed by categorizing an observed continuous variable. Despite the resulting loss of information, in epidemiologic analyses, continuous exposure variables are often categorized using either prespecified cut-points or estimated quantiles of the variable's distribution. Flegal et al¹²⁴ published a key result showing that dichotomization of a continuous exposure that is subject to nondifferential measurement error leads to a binary exposure that has differential misclassification. Later work of Brenner and Blettner¹²⁵ and Delpizzo and Borghes¹²⁶ also stress this point. Although differential measurement error and misclassification may, in general, lead to bias in the estimated regression coefficient in any direction (see Part 1, Section 3), the simulations of Flegal et al¹²⁴ demonstrated relative risk (RR) estimates that were attenuated. These simulations were based on a univariate linear logistic regression with a continuous exposure X prone to classical measurement error and dichotomization using a prespecified cut-point.

Considering the same assumptions regarding the exposure X, Gustafson and Le¹²⁷ extended the results of Flegal et al. First, they provided analytic expressions for linear regression in addition to numerical results for linear logistic regression. Second, they considered the effect of changing the prespecified cut-point c. Third, they considered the inclusion of a second precisely measured continuous covariate Z in the regression and the effect of correlation ρ between X and Z. Finally, they considered situations where the true regression of the outcome Y was linear in Z and a weighted average of the continuous and dichotomized X, that is,

$$E(Y \mid X, Z) = \beta_0 + \beta_1 \{ (1 - \omega)X + \omega I(X > c) \} + \beta_2 Z.$$
(13)

This form of regression allowed a more general investigation of the effects of covariate dichotomization, that is, it considers the situation where the truth lies somewhere "in between" the extremes of dichotomization leading to a completely right model specification vs a completely wrong model specification. Their results demonstrated that, when the true regression contains a linear exposure on the continuous scale (ω < 1), it can be beneficial to dichotomize imprecise

continuous exposures, as this can reduce bias from the analysis on the continuous scale. For example, in the case of the linear regression (with $\omega = 0$),

$$E(Y \mid X, Z) = \beta_0 + \beta_1 X + \beta_2 Z, \tag{14}$$

where $X \sim N(0, 1)$, $X^* \sim N(0, 1 + \text{var}(U))$, $Z \sim N(0, 1)$, and $\rho = \text{cor}(X, Z)$, the multiplicative biases in the estimated regression coefficients arising in the analysis with continuous X^* and categorized $W^* = I(X^* > c)$ are given by, respectively,

$$\frac{\beta_{1X^*}}{\beta_1} = \frac{1}{1 + \text{var}(U)/(1 - \rho^2)} \text{ and } \frac{\beta_{1W^*}}{\beta_{1W}} = \theta \left\{ \frac{R(c) - \rho^2 \phi(c)}{R(\theta c) - \rho^2 \theta \phi(\theta c)} \right\}, \tag{15}$$

where $\theta = \frac{1}{\sqrt{1+var(U)}}$, $R(c) = \frac{\Phi(c)\{1-\Phi(c)\}}{\phi(c)}$ and $\phi(.)$, $\Phi(.)$ are the probability density and cumulative probability functions of the standard normal distribution. Some values of var(U), ρ and c lead to $\frac{\beta_1 W^*}{\beta_1 W} > \frac{\beta_1 X^*}{\beta_1}$. Gustafson and Le¹²⁷ point out that no general statements can be made about how the bias due to dichotomization depends on the choice of threshold or the strength of correlation between predictors. The authors produce several graphs of attenuations comparing continuous and dichotomized observed main exposure for $0 < var(U)^{1/2} < 1.2$, c = 1, 2, 3, and $\rho = 0, 0.3, 0.6, 0.9$. In all those cases, attenuation was greater in the continuous case, that is, $\frac{\beta_1 W^*}{\beta_1 W} > \frac{\beta_1 W^*}{\beta_1 W$

Recently Keogh et al¹²⁸ investigated several methods of adjusting for misclassification due to dichotomizing a continuous error-prone predictor. In contrast to previous work, a measured continuous exposure was specified to follow a linear measurement error model, thereby allowing for systematic error. In addition to methods using estimated misclassification probabilities, the authors considered applying two RC-based methods, MI and MR to the continuous exposure followed by dichotomization, and also a SIMEX method. Simulation studies were used to compare the methods when either the true exposure or reference measurements with classical error were available in a validation subsample. The underlying relationship between the continuous exposure and the outcome in the simulations was a univariate linear logistic regression, and dichotomization was based on a predefined cutpoint. In that study, RC and SIMEX methods failed to correct adequately for bias because both methods assume nondifferential error (the failure of RC was also confirmed by Dalen et al¹²⁹). However, MI and MR performed well. Methods using estimated misclassification probabilities also performed well, provided differential misclassification was assumed (see also Dalen et al¹³⁰). It is important to note that the latter methods are restricted to estimating ORs, while MI and MR could, in principle, be used with different regression models, with quantile-based categorization, and could also accommodate covariate adjustment. Extending MI and MR to those cases as well as to the case of regression of the outcome on a nonlinear function of the exposure remains an important area for further research.

6 | EXTERNAL, IMPERFECT OR MISSING REFERENCE INSTRUMENTS

While Part 1 and earlier sections of this article have made it clear that both theory and software are available for handling errors in measurement or classification of variables, ultimately their use hinges on the knowledge and data available regarding the measurement error or misclassification model that relates the imperfect observed exposure X^* to the true value X. Unfortunately, all too often, information about the error model is incomplete or missing. In this section, we describe how one might approach the problem of measurement error or misclassification when there is lack of knowledge about such a model.

The ideal setting for applying methods to address measurement error or misclassification of variables is one in which an internal validation study has been done, which would directly relate the imperfect observed exposure or outcome to its true value in the population of interest. We consider analysis options when such a validation study is not available.

In this case, there may be data from another cohort, namely an external validation study, or there may be an imperfect reference instrument that can provide partial information about the nature of the measurement error. We also discuss approaches for settings where there is little to no information available regarding the error-prone data.

6.1 Using an external validation study

Internal validation studies are the ideal because their data can be used directly with all the methods of measurement error analysis described previously. In their absence, external studies can be used to specify the measurement error model (or its associated RC model) in the external data and to estimate its parameters. However, the use of this external study data in the analysis of the primary study is then reliant on the assumption of transportability for the model of $X^*|X,Z$ (see also Part 1, Section 4.2). Transportability means that the specified model relating the error prone X^* with the true data (X,Z) holds with the same parameter values in both studies, and that the relevant parameter estimates and their SEs obtained in the external study can be used without bias in analysis of the primary study. See, for example, Guo et al¹³¹ who provide a MI method that addresses covariate measurement error in a regression model using information from an external validation study that provided information regarding only the covariate error, without measurement of the outcome or other study variables. Further discussion of this article appears in Liao et al.¹³² Buonaccorsi² also provides some discussion of error correction methods that rely on external data. In this case, also assuming nondifferential error, one can use the external data to inform the RC approach, as described in Part 1, Section 6.1; however, the calculation of the SEs would need to be different, particularly if the original external data were not available.

The assumption of transportability may be reasonable if external independent data come from a similar population with measurements obtained with the same or a very similar instrument. Consider, for example, a nutritional study with dietary intake measured by a FFQ. If the external validation study provides independent data from the same population with the same FFQ plus a reference instrument, the error model that is estimated in the validation study could be assumed transportable to the primary study and used for adjusting its results for measurement error. If, however, the same FFQ is used in a somewhat different population, or a different version of FFQ is used in that population, the transportability assumption may not be fully justified.

A different example of possible problems with transportability includes the situation when the distribution of X^* given (X, Z), the measurement error model, is the same in both primary and external studies, but the distributions of true exposure X given Z are different. Since by Bayes' theorem the regression of X given (X^*, Z) depends on both of these distributions, the RC model is not transportable. Carroll et al^{1(sect.2.2.5)} give an example of this phenomenon related to blood pressure measurement.

Since adopting the correct error model is critical for an appropriate adjustment for measurement error, whenever there is doubt about the transportability of an external validation study, it is advisable to conduct a sensitivity analysis by considering some possible variation in the relevant error parameters and their effect on the results of the primary study. Thus, while external validation studies can undoubtedly provide worthwhile information about the measurement error, they often do not entirely exempt the investigator from conducting a sensitivity analysis. However, in comparison to the situations in Sections 6.2 and 6.3 that follow, where there is less information regarding the measurement error, in the case of external validation studies, the sensitivity analysis could involve a more restricted range of parameter values.

6.2 Methods that use an imperfect reference instrument

Often, exposures in epidemiological studies are known to be measured with substantial error, but the corresponding measurement error model is not known due to absence of appropriate reference instruments. Typical examples include most dietary exposures (see Part 1, Section 4.3.1) and characteristics of physical activity such as measures of moderate to vigorous activity (see Part 1, Section 4.3.2). In some such cases, an instrument less biased than the main study instrument, but nevertheless biased, is used as the reference instrument (calibration study with imperfect reference instrument). Examples in nutrition and physical activity studies are given in Part 1, Sections 4.3.1 and 4.3.2.

As mentioned later in Section 6.3, in the absence of knowledge of the measurement error model, a bias or sensitivity analysis is recommended using a plausible set of parameters (or their distribution) for the model. When the measurement error model is estimated using an imperfect reference, sensitivity analysis is also recommended. Although the error model parameters are imperfectly estimated, they may nevertheless be used together with supplementary information to choose

the range of parameters for the sensitivity analysis. Thiébaut et al¹³³ provided a good example of this approach. They reported the estimated RR for breast cancer associated with a 2-fold increase in fat density (percent of total energy provided by fat). In the Results section, they first report the unadjusted RR estimate of 1.15 (95% CI 1.05-1.26) based on FFQ data. They then report the RR adjusted for measurement error based on a 24-hour recall (imperfect) reference validation study: 1.32 (95% CI 1.11-1.58). Finally, in the Discussion section, they use data from the OPEN validation study⁷ to adjust for the bias that may have occurred due to use of an imperfect reference instrument for fat density. The adjustment for bias is calculated by comparing attenuation factors based on the imperfect reference (24-hour recall) with that based on the perfect reference (recovery biomarkers) for protein density that has a recovery biomarker, and transporting the ratios of these estimates to the case of fat density, which does not have a recovery biomarker. This procedure, which was justified by the substantial correlation between protein and fat intake, gave an estimated RR of 1.46 (no confidence limits for this estimate were provided). Although they did not perform a formal sensitivity analysis in the last analysis, it is clear that their approach was moving in that direction.

This example includes the elements of how a sensitivity analysis may be constructed from knowledge of measurement error in exposures similar to that being considered. For exposure X (eg, fat intake), the measured exposure X^* (using instrument I_Q , for example, an FFQ) is compared to an imperfect reference instrument X_{lmp}^{**} (using instrument I_R , for example, a 24-hour recall) to obtain an estimated measurement error model M^* . For some similar exposure X_1 (eg, protein intake), information is available on the measured exposure X_1^* (using instrument I_Q , the FFQ), its imperfect reference instrument $X_{1,lmp}^{**}$ (using instrument I_R , the 24-hour recall), and also an unbiased reference measurement X_1^{**} (eg, 24-hour urinary nitrogen excretion). The availability of both the imperfect reference measurement and an unbiased reference measurement for X_1 enables one to learn about the relationship between the true measurement error model M_1 estimated using measurement X_1^{**} and the model X_1^{**} estimated using the imperfect reference measure $X_{1,lmp}^{**}$. This information about X_1^{**} is then applied to the estimated measurement error model X_1^{**} for the exposure of real interest X_1^{**} , to yield the desired range of parameters for the true measurement error model X_1^{**} for the exposure X_1^{**} .

In this approach, the choice of the "similar" exposure X_1 will, of course, depend on the context. For dietary intakes, it will be the intake of another dietary component, one that has an unbiased reference measurement; for physical activity measures such as moderate or vigorous activity measured by a physical activity diary and compared to an accelerometer reference, it could be total energy expenditure that can be measured unbiasedly by doubly labeled water.

A different approach to dealing with studies using imperfect reference instruments starts with the question of whether using an imperfect reference instrument to adjust for measurement error is preferable to making no adjustment whatsoever. In other words, if one uses an imperfect reference to estimate the measurement error model and then uses this model to adjust risk parameter estimates in the health outcome model, would these adjusted estimates, even if biased, still have less bias than unadjusted ones? If that could be demonstrated, it could motivate the use of these imperfectly adjusted estimates in preference to the unadjusted ones. This approach is less demanding than conducting a sensitivity analysis, since it involves applying the measurement error adjustment for just one measurement error model, but it is also less complete.

The issue has been studied in nutritional epidemiology. Freedman et al¹³⁴ published the results of analyzing such a question using data from the OPEN study, and more recently updated their results using data from the five validation studies included in the Validation Studies Pooling Project.¹³⁵ They concluded that, on average, 24-hour recall-based calibration of a FFQ reduced, but did not eliminate, the bias in the risk estimates in multivariate risk models that included energy, and protein, potassium and sodium intake densities, in comparison with unadjusted estimates. Although those results, as well as similar results using linear measurement error models in the sensitivity analysis conducted by Buonaccorsi et al,¹³⁶ indicate that using a 24-hour recall as a reference instrument to adjust for measurement error would improve the analysis of studies in nutritional epidemiology, there remain some doubts. The improvement has been demonstrated in only a handful of nutrients (those which have unbiased biomarkers), and may not transfer to all other dietary components, especially episodically consumed dietary components, for which the measurement error model is highly nonlinear.¹³⁷

A general limitation of this approach is that even if the resulting estimates are less biased than unadjusted estimates, they are nevertheless biased. Therefore, presenting them as the best estimates available does not reveal the full extent of the underlying uncertainty, and is a less complete approach than conducting a sensitivity analysis.

6.3 | Approaches when there is no reference instrument

If we have no knowledge about the measurement error model, then we have to make assumptions about it. Note that ignoring measurement error is one (incorrect) assumption, which is akin to assuming there was no error in measurement.

In this section, we outline alternatives to this naïve approach. Investigations can still be undertaken to understand the potential impact of measurement error or misclassification on study results. This approach involves three steps. First, a measurement error model is posited. Second, study results are produced that are corrected for measurement error, under the assumed measurement error model. This may involve direct reanalysis of the data, or post hoc adjustment of estimated outcome model parameters. Finally, assumptions about the parameters in the assumed error model are typically varied in a sensitivity analysis to examine the robustness of study results to a range of assumptions about the measurement error.

Bias analysis, sometimes also referred to as uncertainty analysis or probabilistic sensitivity analysis, follows the above general approach for quantifying the potential effects of measurement error. This method focuses on sources of systematic and random error. Bias analyses have several goals: (i) to estimate the direction and magnitude of the bias in study results induced by the errors in the data, (ii) to make explicit the sources of suspected errors and the degree of uncertainty that they introduce into study results, and (iii) to efficiently guide future research by elucidating what associations are sensitive to the underlying amount of measurement error or misclassification and would best benefit from future replicates or validation studies to estimate that error. ¹³⁸

Methods for bias analyses are well established in the statistical and epidemiologic literature and apply to estimating the impact of sources of bias that go beyond just measurement error, such as unmeasured confounding or nonignorable dropout (Lash et al, ¹³⁸ Greenland et al, ¹³⁹ Greenland, ¹⁴⁰ Fox and Lash, ¹⁴¹ Fox, ¹⁴² MacLehose et al, ¹⁴³ Lash and Ahern ¹⁴⁴). A fundamental step of bias analysis is to thoroughly review the study's subject selection and retention, methods of data collection, and other opportunities for confounding, selection bias, and measurement error. 138 Once those potential sources of bias have been identified, mathematical models are developed for the relationship between the underlying true data with biases removed and the study data. For this endeavor, distributions rather than a single set of parameter values are used to generate a sensitivity analysis for the results of the bias analysis. In the absence of any validation data or other studies to inform the selection of parameters, educated guesses can be used to posit such relationships. 142 In this last step, one option is to assign a prior distribution from which to draw the necessary error parameters, which allows for a Bayesian analysis that naturally integrates the uncertainties coming from the submodels for exposure, outcome, and measurement. Choice of this prior in the absence of validation studies could similarly be informed by expert option, as discussed in Section 2.2. Lash et al¹³⁸ provide a review of best practices for bias analysis. One challenge to this approach is its reliance on proper specification of the mathematical form of the measurement error, such as additive or multiplicative. This choice is likely best informed by validation data but could also be made part of the sensitivity analysis in the absence of such data.

A practical example of bias analysis can be seen in the study by Jurek et al, ¹⁴⁵ who sought to quantify the impact that exposure misclassification may have had on a study by Ross et al ¹⁴⁶ reporting on the effect of maternal supplement use on the risk of leukemia in children with Down syndrome. Because of a lack of an internal validation study, Jurek et al ¹⁴⁵ developed their misclassification models and parameter distributions from a mixture of expert opinion, a literature review of validation studies of similar exposure instruments, and limitations set by the data themselves. Using several error model scenarios, including both differential and nondifferential misclassification and a formula to adjust the estimated OR for the underlying exposure misclassification, they conducted a sensitivity analysis for the induced bias. Their bias analyses revealed that data that were corrected for the reporting bias in supplement use generally yielded a stronger protective effect than the naïve analysis that ignored the misclassification. The uncertainty was increased in all scenarios.

He et al¹⁴⁷ provided an alternative approach for examining the potential effects of measurement error. These authors considered an accelerated failure time model for mortality in the Bussleton Health Study cohort that included two error-prone covariates, serum cholesterol and systolic blood pressure (SBP), as well as other assumed precise covariates. The errors in cholesterol and blood pressure were assumed to be independent and to follow the classical measurement error. Lacking a replicates substudy, the authors considered several possibilities for the size of the underlying measurement error variance and applied a SIMEX approach to re-estimate the regression parameters for each value of the assumed measurement error variance. With this exercise, the authors were able to conclude that even under small to moderate classical measurement error, the factors determining mortality remained the same, with the most uncertainty about the magnitude of the effect of SBP. Such analyses motivate future replicates studies to gather multiple measures of SBP in similar settings to better understand the magnitude of the measurement error variance and the relationship of SBP to mortality. The decreased sensitivity of results to the measurement error in cholesterol suggest that for this exposure such studies may be of secondary importance to that for SBP.

7 | CONCLUSION

In this two-part tutorial, we have presented basic information needed to understand the impact of measurement error and misclassification on results of epidemiological research studies, and methods available to adjust for such error. In Part 2, we have also presented some more advanced methods to address covariate and outcome measurement error, but our review is not exhaustive. Some notable methods not considered here include conditional score and corrected score methods to address covariate error. For density estimation, there is also only minimal detail regarding deconvolution kernel estimators and other nonparametric density estimation methods. The reader is referred to some recent textbooks (Carroll et al, Buonaccorsi, Yi, Gustafson, 13) for introductions to these and other methods not considered.

Our impression is that the problem of measurement error and misclassification is being seriously neglected in the design of many epidemiologic studies and in the presentation of their results. Has, 148, 149 Barriers to satisfactory handling of such problems include lack of validation studies required to quantify the amount and type of error, lack of appreciation and understanding of the effects of such error, and lack of knowledge of the methods and software required to adjust for these effects. Publication of this article is part of a wider effort by our STRATOS Topic Group to bring these problems to the attention of the biostatistical and epidemiologic communities; and on a broader perspective, our work on publishing this guidance article is part of the general aim of STRATOS to strengthen the analytic thinking underlying observational studies. How

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DATA AVAILABILITY STATEMENT

The OPEN Study data that illustrate the methods presented in this paper are available upon request to RFAB@mail.nih.gov. The request should specify the dataset used in analyses presented in the papers by Keogh et al (2020) and Shaw et al (2020). More information about these data can be obtained at https://epi.grants.cancer.gov/past-initiatives/open/.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of this article.

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