Comparison of Random Forest and Parametric Imputation Models for Imputing Missing Data Using MICE: A CALIBER Study

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Multivariate imputation by chained equations (MICE) is commonly used for imputing missing data in epidemiologic research. The “true” imputation model may contain nonlinearities which are not included in default imputation models. Random forest imputation is a machine learning technique which can accommodate nonlinearities and interactions and does not require a particular regression model to be specified. We compared parametric MICE with a random forest-based MICE algorithm in 2 simulation studies. The first study used 1,000 random samples of 2,000 persons drawn from the 10,128 stable angina patients in the CALIBER database (Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; 2001–2010) with complete data on all covariates. Variables were artificially made “missing at random,” and the bias and efficiency of parameter estimates obtained using different imputation methods were compared. Both MICE methods produced unbiased estimates of (log) hazard ratios, but random forest was more efficient and produced narrower confidence intervals. The second study used simulated data in which the partially observed variable depended on the fully observed variables in a nonlinear way. Parameter estimates were less biased using random forest MICE, and confidence interval coverage was better. This suggests that random forest imputation may be useful for imputing complex epidemiologic data sets in which some patients have missing data.

angina, stable; imputation; missing data; missingness at random; regression trees; simulation; survival

Abbreviations: CALIBER, Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; MAR, missing at random; MICE, multivariate imputation by chained equations.
not rely on distributional assumptions and can accommodate nonlinear relations and interactions. On simulated data sets with interactions between variables, imputation of missing data using MICE with regression trees resulted in less biased parameter estimates than MICE with linear regression (7). However, regression trees may “overfit,” following the pattern of noise too closely and producing a complex model with poor predictive power in new data sets.

Random forest uses bootstrap aggregation of multiple regression trees to reduce the risk of overfitting, and it combines the predictions from many trees to produce more accurate predictions (8, 9). Random forest is widely used in genetic epidemiology (10) and has also been used for modeling survival (11, 12) and predicting response to cancer chemotherapy (13). We propose that random forest may be useful in multiple imputation of epidemiologic data sets, particularly if there are large numbers of clinical variables per participant, as may increasingly be the case (e.g., genomic or proteomic studies).

Stekhoven et al. (14) developed a random forest-based algorithm for missing data imputation called missForest. This algorithm aims to predict individual missing values accurately rather than take random draws from a distribution, so the imputed values may lead to biased parameter estimates in statistical models. Apart from a comparison between random forest and polynormous regression for imputing tumor stage using MICE (15), we are not aware of other published evaluations of multiple imputation using random forest.

In this paper, we compare a standard implementation of MICE with imputation using missForest, and we propose a new version of MICE which imputes each variable using random forest. We compare these methods in a realistically complex survival analysis based on patients with stable angina in the CALIBER (Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records) database (16) and in a simulation study with interactions.

**METHODS**

### Imputation of missing data using MICE, where each variable is imputed using random forest

Within the MICE framework, missing values of continuous variables are conventionally imputed by fitting a linear regression model for the observed values, predicting the conditional mean for each missing value, and randomly imputing a value from a normal distribution centered on this conditional mean. Our new method is derived from the “mice.impute.norm.boot” function in the “mice” package in R (4), in which linear regression is applied to a bootstrap sample of records with observed values of the variable to be imputed. The purpose of the bootstrap is to accommodate sampling variation in estimating population regression parameters, which is part of ensuring that imputations are “proper” (3). The random forest algorithm itself involves another level of bootstrap sampling. Records with missing values in the dependent variable are imputed by random draws from independent normal distributions centered on conditional means predicted using random forest. We used the “out-of-bag” mean square error as the estimator of residual variance (which we assumed to be normally distributed). Random forest fits each tree to a different bootstrap sample of the data and aggregates the results; the out-of-bag error is the mean of squared differences between each observed value and the prediction based on trees for which that observation is not included in the bootstrap sample.

For binary or unordered categorical variables, we used random forest to fit individual regression trees to a bootstrap sample of the data and imputed each missing value as the prediction of a randomly chosen tree. This is equivalent to choosing between 0 and 1 with probability according to the mean random forest prediction. Our random forest imputation functions are available from the Comprehensive R Archive Network (17).

### Simulation study based on CALIBER data

CALIBER is a database of linked routine collected electronic health records from England (16), comprising data from primary care (Clinical Practice Research Datalink (18), hospital admissions (19), the national registry of acute coronary syndromes (20), and the national death registry. The cohort consisted of patients who received a diagnosis of stable angina while registered at a general practice contributing to the Clinical Practice Research Datalink. Blood pressure, smoking status, and measurements of blood biomarkers were taken from routine clinical records before the diagnosis of stable angina, and patients were followed up for the composite endpoint of death or nonfatal myocardial infarction (see Web Appendix 1, available at http://aje.oxfordjournals.org/). The CALIBER record-linkage study has received ethical approval, and this study was approved by the Clinical Practice Research Datalink Independent Scientific Advisory Committee.

**Analysis of interest.** We investigated missing data in the context of a hypothetical analysis of associations, suggested in previous studies (21-23), between 3 commonly measured hematoletic parameters (hemoglobin concentration, lymphocyte count, and neutrophil count) and prognosis among patients with stable angina in CALIBER. The substantive analysis was a multivariable Cox model with the composite endpoint of death or nonfatal myocardial infarction, and with predictor variables specified a priori. The fully observed predictor variables were: age, age squared, sex, previous myocardial infarction, diabetes mellitus, previous stroke, peripheral arterial disease, and heart failure. Smoking status was a partially observed 3-category variable (never, former, or current smoker), and we included the following partially observed continuous variables: systolic blood pressure (mm Hg), log neutrophil count (10⁹ cells/L), log lymphocyte count (10⁹ cells/L), high-density lipoprotein cholesterol level (mmol/L), hemoglobin concentration (g/dL), and log serum creatinine concentration (µmol/L). For each of these variables, we took the mean of any observed values in the 2 years prior to the start of follow-up. We used the Efron approximation (24) for ties. We did not investigate alternative models and ignored clustering by general practice.

**Generation of sample data sets for simulation study.** For the simulation study, we created data sets with missing data...
for which we knew the “true” values, with a missingness pattern similar to that observed in the actual data set but which was missing at random, such that the MAR assumption underlying most multiple imputation approaches was satisfied (Figure 1). We denoted the entire cohort of 52,576 stable angina patients data set “A.” Patients with no missing values for any of the variables in the survival model were denoted data set “B” (13,308 patients).

We used logistic regression based on completely observed variables to investigate factors associated with a patient’s having a complete record (i.e., whether a patient was in data set B). We included the first value after cohort entry of partially observed continuous variables as auxiliary variables to help predict missing covariates in imputation models. The subset of patients with complete recording of all analysis and auxiliary variables was denoted data set “C” (10,128 patients), and they formed the basis for the resampling study. We artificially made some values of predictor variables missing in one thousand 2,000-patient random samples (with replacement) from data set C. We carried out simulations with

Figure 1. Generation of data sets with artificial missingness from a population of patients with stable angina in the CALIBER database, 2001–2010. Data sets D1, D2, …, D1,000 are samples of 2,000 patients with replacement from data set C. CALIBER, Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; MAR, missing at random; MCAR, missing completely at random; MICE, multivariate imputation by chained equations.
one of 2 missingness mechanisms: 1) MAR in a pattern similar to that of data set A or 2) an artificial pattern of missingness completely at random in which only categorical variables were missing (see Web Appendix 1 for more details).

**Multiple imputation of test data sets.** Imputation models included all of the variables in the substantive Cox model, event status, marginal Nelson-Aalen cumulative hazard (25), and the following auxiliary variables: type of endpoint, whether the practice was receiving electronic laboratory results, and the earliest recorded value after the index date for blood pressure and the 5 blood biomarkers. We imputed continuous variables using MICE with normal-based linear regression, predictive mean matching with 3 nearest neighbors, and our new random forest method. We imputed categorical variables using either MICE with logistic or polytomous regression or MICE with random forest (choice of 10 or 100 trees). We also investigated random forest with a single tree to determine whether bootstrap aggregation had an advantage over a single regression tree.

We generated 10 MICE imputations, each drawn from a separate chain with a different random seed, with 10 cycles of imputation before drawing the imputed data set. We assessed chain mixing by reviewing plots of chain mean values and standard deviations. For two of the methods (random forest with 10 trees and parametric MICE), we also calculated results using 100 imputations.

In addition to MICE, we also evaluated missForest (14), which uses random forest in an iterative way to complete a data set with missing values, where imputed values are equal to the random forest predictions rather than being randomly sampled from a conditional distribution. We generated multiple imputed data sets by running missForest using different random seeds, which leads to different random forest models being generated.

Regardless of the imputation method, all data sets were analyzed using the same multivariable semiparametric Cox model as described above. For each set of imputed data sets, the log hazard ratios from the Cox model were combined using Rubin’s rules (26), which assumes that imputed values were drawn from the appropriate Bayesian posterior. We carried out analyses using R 2.12.1 (27), with the software packages mice 2.12 (4), missForest 1.3 (28), survival 2.36-2 (29), and randomForest 4.6-6 (30). Random numbers were generated using the Mersenne Twister (31).

**Comparison of results obtained by different methods.** We considered the Cox proportional hazards model fitted to the entire data set (data set C) the “true” result for the assessment of bias and confidence interval coverage of hazard ratios. We compared the widths of 95% confidence intervals between 2 methods using paired-sample t tests. We compared coverage of 95% confidence intervals for each coefficient separately using McNemar’s test, defining discordant pairs as data sets in which the 95% confidence interval included the “true” value for one method but not the other. We compared the efficiency of the estimators by calculating their empirical standard deviations. We calculated the between-imputation variance of the estimated log hazard ratios, defined as the mean (across simulations) of the variance of the log hazard ratio estimates from the 10 imputations per data set.

**Simulation study with interactions**

We also created simulated data sets to compare the performance of methods when there were nonlinearities in the association between predictor variables. We generated 2 independent random normal variables with mean 0 and variance 1, $x_1$ and $x_2$, and a third variable $x_3$ equal to 0.5($x_1 + x_2 - x_1x_2$) + $e$, where $e$ was distributed normally with mean 0 and variance 1. Survival times were generated according to an exponential distribution with log hazard 0.5($x_1 + x_2 + x_3$). This meant that there were no interactions in the substantive model, but the default parametric imputation model for $x_3$ (which would not include any interactions) would be incorrect. Observation times were generated according to a uniform distribution in the range from 0 to the 50th percentile of survival times. If the observation time was less than the survival time, the patient was considered censored (event indicator 0, and the patient’s follow-up ended on his or her censoring date); otherwise the event indicator was 1, with follow-up ending on the date of the event.

Variable $x_3$ was made 20% MAR according to a logistic model based on $x_1$ and $x_2$, the marginal Nelson-Aalen cumulative hazard and the event indicator.

We analyzed 1,000 simulated data sets with 2,000 patients each, imputing missing data using random forest and parametric MICE (without interactions), comparing the results as above (Web Appendices 2 and 3).

**RESULTS**

**Simulation study based on CALIBER data**

The prevalence of missingness among partially observed variables ranged from 1.5% for smoking to 56.7% for lymphocyte counts and 56.8% for neutrophil counts (Web Table 1). Patients with missing data were more likely to experience the primary endpoint of death or nonfatal myocardial infarction (age- and sex-adjusted hazard ratio = 1.19, 95% confidence interval: 1.13, 1.25) (Figure 2). Patients with missing data also tended to have longer follow-up because they entered the cohort earlier (a median index date of March 1, 2002, vs. December 6, 2005; $P<0.0001$ by Wilcoxon rank-sum test). The logistic regression model showed that patients with diabetestes, peripheral arterial disease, and previous stroke were more likely to have complete records (Table 1). Web Table 2 shows that coefficients from Cox models for patients with complete data (data set C) were similar to the average results from full data analysis of the subsamples (as we would expect in the absence of small-sample bias), so we compared imputation estimates with those from data set C.

We found very little difference between results obtained using 10 MICE imputations and those obtained using 100 MICE imputations (Web Tables 3 and 4), so all results are based on 10 imputations unless stated otherwise. **Bias.** Estimates from complete-record analysis were biased for some parameters under MAR (missingness mechanism 1); this may be expected because we had introduced missingness dependent on the outcome (Table 2, Web Table 3, Web Figure 1). For example, the geometric mean hazard ratio per doubling of lymphocyte count was
0.738 from complete-record analysis but 0.799 from full-data analysis. There was no material bias with parametric MICE (mean hazard ratio = 0.806) or our random forest MICE method with 10 trees (“MICE RF 10”; mean hazard ratio = 0.807). The random forest MICE estimate for smoking (categorical) was biased towards the null (Table 3, Web Figure 2), but there was no material bias in other parameters estimated by random forest or parametric MICE (Table 2, Web Tables 3 and 4, Web Figure 3). However, imputation using single-tree random forest MICE (“MICE Tree”) or missForest produced materially biased estimates for all continuous variables missing at random (Figure 3, Table 2, Web Figure 1, Web Table 3).

Efficiency. All of the imputation methods tested produced more efficient parameter estimates than complete-record analysis. MICE with random forest produced slightly more efficient estimates than parametric MICE, and the average between-imputation variance was also lower (Tables 2 and 3, Web Tables 3–6).

Confidence intervals. Parametric MICE yielded confidence intervals with approximately 93%–95% coverage. The mean widths of confidence intervals were lower using random forest MICE than using parametric MICE ($P < 0.001$ for each comparison), but coverage was either equal or greater using random forest MICE (Tables 2 and 3, Web Tables 3–6).

For categorical variables, missForest produced imputed values which were more likely to be equal to the “true” (observed) value than the MICE methods, but confidence intervals were too small with below nominal coverage, and between-imputation variance was very small. There was no difference in bias, precision, or coverage between normal-based MICE and predictive mean matching (Web Table 5). Random forest MICE with 100 trees for continuous variables produced estimates with slightly narrower confidence intervals than random forest MICE with 10 trees (Web Table 5), but with greater bias, worse coverage of 95% confidence intervals, and 10 times the computational cost. For categorical variables, random forest MICE with 10 trees and random forest MICE with 100 trees produced almost identical results.

Simulation study with interactions

The coefficient estimate for the partially observed variable ($x_3$) was 10% biased using parametric MICE, 2.6% biased using random forest with 100 trees, and only 1.0% biased using random forest with 10 trees ($P < 0.001$ for 2-way comparisons). The bias in the $x_3$ coefficient varied with the number of trees, with 10 or 20 trees giving minimal bias (Table 4). Random forest MICE produced narrower 95% confidence intervals for the $x_3$ coefficient than parametric MICE ($P < 0.001$), and coverage was only 80% using parametric MICE as compared with 95% using random forest MICE with 5–100 trees. Further details are given in Web Appendix 2.

DISCUSSION

Summary of main findings

In this resampling study of methods for handling missing data, parametric and random forest MICE produced estimates with no material bias for a Cox model on data with artificially introduced MAR missingness. Random forest-based MICE produced more efficient estimates and narrower confidence intervals than parametric MICE, yet in some cases coverage probability was greater than 95%, suggesting that some
confidence intervals may be conservative. A possible explanation for the efficiency gain with random forest MICE is that it was able to make better use of the available information by accommodating nonlinearities among the predictors. In simulations with an interaction among the predictor variables but not in the substantive model, random forest MICE was less biased than parametric MICE, which omitted the interaction. Using missForest for multiple imputation resulted in very biased estimates and poor coverage of confidence intervals.

Overall, our results suggest that random forest imputation may be useful for imputing complex epidemiologic data sets in which some patients have missing data.

**Imputation methods for MICE**

It is important that imputation models be correctly specified for analyses to yield unbiased estimates, and random forest may help avoid the bias that can occur with parametric MICE if the latter’s imputation models are misspecified. In our main study, standard parametric MICE performed well, suggesting that the true imputation models did not contain significant nonlinearities or interactions, and hence random forest did not confer an advantage from the perspective of bias. However, the simulated data sets had interactions which were not included in the parametric MICE imputation models, and in this setting random forest MICE outperformed parametric MICE (Web Appendix 2). The default settings for MICE do not include interactions between the variables, and it is routine practice to include only those interactions that are in the substantive model, rather than actively search for all possible interactions and nonlinearities. This shows the importance of checking to be sure that the imputation models are reasonably well specified. Random forest reduces the need to investigate associations between

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**Table 2.** Comparisons Between Methods of Handling Missing Data in 1,000 Samples With Continuous Variables Missing at Random in a Pattern Similar to That of the Original Data Set (Missingness Mechanism 1), CALIBER Database, 2001–2010

| Variable and Method | Bias* of Log HR | z Score for Bias | SD of Estimated Log HR | Mean Length of 95% CI | Coverage of 95% CI, % | Between-Imputation Variance |
|---------------------|-----------------|-----------------|------------------------|-----------------------|----------------------|-----------------------------|
| Neutrophils (10⁹ cells/L), per doubling |                |                 |                        |                       |                      |                             |
| Full data           | 0.002           | 0.43            | 0.158                  | 0.564                 | 92.2                 |                             |
| Complete record     | −0.045          | −2.67           | 0.533                  | 1.677                 | 90.1                 |                             |
| MICE normal         | −0.038          | −5.15           | 0.232                  | 0.883                 | 93.4                 | 0.0243                      |
| MICE PMM            | −0.042          | −5.68           | 0.230                  | 0.889                 | 93.4                 | 0.0245                      |
| missForest          | −0.266          | 27.72           | 0.303                  | 0.781                 | 63.2                 | 0.0014                      |
| MICE RF 10 trees    | −0.024          | −4.55           | 0.165                  | 0.798                 | 97.9                 | 0.0143                      |
| Lymphocytes (10⁹ cells/L), per doubling |                |                 |                        |                       |                      |                             |
| Full data           | −0.007          | −1.23           | 0.155                  | 0.526                 | 91.6                 |                             |
| Complete record     | −0.087          | −5.87           | 0.464                  | 1.544                 | 89.8                 |                             |
| MICE normal         | 0.001           | 0.13            | 0.202                  | 0.759                 | 93.2                 | 0.0157                      |
| MICE PMM            | 0.006           | 0.99            | 0.205                  | 0.768                 | 92.4                 | 0.0162                      |
| missForest          | −0.190          | −22.21          | 0.270                  | 0.724                 | 72.5                 | 0.0011                      |
| MICE RF 10 trees    | 0.003           | 0.56            | 0.156                  | 0.727                 | 97.8                 | 0.0109                      |
| Hemoglobin, per g/dL |                |                 |                        |                       |                      |                             |
| Full data           | −0.004          | −1.99           | 0.057                  | 0.202                 | 91.6                 |                             |
| Complete record     | −0.022          | −3.91           | 0.180                  | 0.593                 | 90.8                 |                             |
| MICE normal         | −0.007          | −2.73           | 0.076                  | 0.279                 | 92.6                 | 0.0019                      |
| MICE PMM            | −0.004          | −1.47           | 0.077                  | 0.279                 | 92.7                 | 0.0019                      |
| missForest          | −0.056          | −19.96          | 0.089                  | 0.255                 | 77.3                 | 0.0001                      |
| MICE RF 10 trees    | −0.010          | −5.61           | 0.059                  | 0.261                 | 97.2                 | 0.0012                      |

Abbreviations: CALIBER, Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; CI, confidence interval; HR, hazard ratio; MICE, multivariate imputation by chained equations; PMM, predictive mean matching; RF 10 trees, random forest with 10 trees; SD, standard deviation.

* Bias was measured relative to estimates from analysis of the full data set (data set C) (Web Table 2).

b The z score is defined as the mean bias of the estimate divided by the empirical standard error from simulations, and it should lie approximately within the interval (−2, +2).

c Results for complete records were based on the 986 samples for which it was possible to estimate hazard ratios for all parameters.
predictor variables, because it should automatically accommodate nonlinearities and interactions. Imputation models should also be compatible with the substantive model (32), and random forest obviates the need to specify how the outcome should be conditioned on in the imputation models for covariates.

When using random forest for prediction, a larger number of trees is preferred in order to obtain precise predictions (30).

### Table 3. Comparisons Between Methods of Handling Missing Data in 1,000 Samples With Categorical Variables Missing Completely at Random (Missingness Mechanism 2), CALIBER Database, 2001–2010

| Variable and Method          | Bias$^a$ of Log HR | z Score for Bias$^b$ | SD of Estimated Log HR | Mean Length of 95% CI | Coverage of 95% CI, % | % Falsely Classified$^c$ |
|-----------------------------|--------------------|----------------------|------------------------|-----------------------|-----------------------|--------------------------|
| **Previous myocardial infarction** |                    |                      |                        |                       |                       |                          |
| Full data                   | 0.006              | 1.22                 | 0.154                  | 0.587                 | 94.2                  | 0                        |
| MICE logistic               | −0.013             | −2.46                | 0.168                  | 0.682                 | 95.5                  | 29.6                     |
| missForest                  | 0.002              | 0.27                 | 0.179                  | 0.625                 | 91.8                  | 17.3                     |
| MICE RF 10 trees            | −0.020             | −4.21                | 0.149                  | 0.662                 | 97.3                  | 28.5                     |
| **Diabetes mellitus**       |                    |                      |                        |                       |                       |                          |
| Full data                   | 0.010              | 2.30                 | 0.156                  | 0.592                 | 93.7                  | 0                        |
| MICE logistic               | 0.016              | 3.21                 | 0.171                  | 0.685                 | 95.7                  | 32.0                     |
| missForest                  | 0.014              | 2.73                 | 0.182                  | 0.627                 | 90.8                  | 19.7                     |
| MICE RF 10 trees            | −0.021             | −4.25                | 0.149                  | 0.668                 | 97.5                  | 30.7                     |
| **Previous stroke**         |                    |                      |                        |                       |                       |                          |
| Full data                   | 0.005              | 0.86                 | 0.198                  | 0.707                 | 94.0                  | 0                        |
| MICE logistic               | −0.005             | −0.58                | 0.207                  | 0.828                 | 95.5                  | 17.9                     |
| missForest                  | 0.004              | 0.65                 | 0.211                  | 0.763                 | 92.9                  | 8.4                      |
| MICE RF 10 trees            | −0.011             | −1.79                | 0.183                  | 0.808                 | 97.9                  | 16.7                     |
| **Peripheral arterial disease** |                  |                      |                        |                       |                       |                          |
| Full data                   | 0.016              | 2.59                 | 0.199                  | 0.730                 | 93.6                  | 0                        |
| MICE logistic               | −0.002             | −0.21                | 0.218                  | 0.858                 | 94.8                  | 15.5                     |
| missForest                  | 0.028              | 4.18                 | 0.223                  | 0.788                 | 91.9                  | 7.0                      |
| MICE RF 10 trees            | 0.005              | 0.94                 | 0.192                  | 0.834                 | 97.1                  | 14.5                     |
| **Heart failure**           |                    |                      |                        |                       |                       |                          |
| Full data                   | 0.015              | 2.47                 | 0.191                  | 0.653                 | 91.7                  | 0                        |
| MICE logistic               | 0.015              | 2.22                 | 0.207                  | 0.759                 | 93.8                  | 14.6                     |
| missForest                  | 0.001              | 0.08                 | 0.216                  | 0.696                 | 89.4                  | 7.2                      |
| MICE RF 10 trees            | −0.034             | −5.78                | 0.190                  | 0.746                 | 95.5                  | 13.7                     |
| **Smoking status: current vs. never** |            |                      |                        |                       |                       |                          |
| Full data                   | 0.019              | 2.62                 | 0.264                  | 0.969                 | 93.9                  | 0                        |
| MICE logistic               | 0.023              | 2.65                 | 0.292                  | 1.092                 | 94.0                  | 52.4                     |
| missForest                  | −0.036             | −3.56                | 0.308                  | 1.062                 | 91.5                  | 35.0                     |
| MICE RF 10 trees            | −0.098             | −12.92               | 0.237                  | 1.072                 | 95.5                  | 50.0                     |
| **Smoking status: former vs. never** |      |                      |                        |                       |                       |                          |
| Full data                   | 0.011              | 1.66                 | 0.247                  | 0.908                 | 93.6                  | 0                        |
| MICE logistic               | −0.008             | −0.82                | 0.266                  | 1.022                 | 94.1                  | 52.4                     |
| missForest                  | 0.045              | 5.34                 | 0.270                  | 0.980                 | 93.2                  | 35.0                     |
| MICE RF 10 trees            | −0.060             | −8.81                | 0.212                  | 1.000                 | 97.1                  | 50.0                     |

Abbreviations: CALIBER, Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; CI, confidence interval; HR, hazard ratio; MICE, multivariate imputation by chained equations; RF 10 trees, random forest with 10 trees; SD, standard deviation.

$^a$ Bias was measured relative to estimates from analysis of the full data set (data set C) (Web Table 2).

$^b$ The $z$ score is defined as the mean bias of the estimate divided by the empirical standard error from simulations, and it should lie approximately within the interval $(-2, +2)$.

$^c$ Percentage of imputed values that were different from the “true” (observed) missing value.
However, when imputing continuous variables using random forest MICE, bias seemed to tend towards a nonzero limit as the number of trees increased, with 10 or 20 trees giving minimal bias (Table 4). It is possible that the relationship between the number of trees and bias may have the same functional form but with a different direction of bias, asymptotic limit, and optimal number of trees, depending on the data. This phenomenon warrants further investigation.

A disadvantage of random forest is that the “models” are complex and not easily interpretable, although arguably this is not a shortcoming for the purpose of imputation. Another disadvantage is that random forest can be biased in

![Figure 3. Bias in estimates of log hazard ratios for partially observed variables with data missing at random (missingness mechanism 1) in 1,000 samples of patients with stable angina in the CALIBER database, 2001–2010. A) log neutrophil count (10⁹ cells/L); B) log lymphocyte count (10⁹ cells/L); C) hemoglobin concentration (g/dL). The solid horizontal line is the “true” log hazard ratio from the full data set (data set C); the dashed lines show ±1 empirical standard error. The boxes span the interquartile range (25th–75th percentiles), and the whiskers extend to the most extreme data point, which is no more than 1.5 times the interquartile range from the box. Circles represent outliers. The light gray boxes show results from simulations with 50% complete records, and the dark gray boxes show results from simulations with 25% complete records. CALIBER, Cardiovascular Disease Research using Linked Bespoke Studies and Electronic Records; MICE, multivariate imputation by chained equations; PMM, predictive mean matching; RF, random forest.](image)
some situations, due to random forest predictions of continuous variables at the extremes of their range being biased towards less extreme values (33). This is because a random forest prediction effectively consists of a weighted average of observed values of the variable being predicted; unlike model-based prediction, it is unable to extrapolate beyond observed values. In a simulation study, we found that random forest imputation led to bias when the distribution of missing values was very different from that for observed values (17), although in such situations any kind of imputation may produce poor results. However, we did not find such a bias in our CALIBER study because missing and observed values had similar distributions. Another limitation of our random forest MICE method is the assumption that the residuals from the random forest regression are normally distributed with constant variance.

On these 2,000-patient data sets, computation time was 3 times as long for random forest MICE with 10 trees as for parametric MICE (137 seconds per data set vs. 48 seconds per data set on a computer with an Intel Xeon 3.47-GHz processor (Intel Corporation, Santa Clara, California)), but on a 10,000-patient data set, random forest took 6.5 times as long. However, random forest may yield a saving in analyst time because there is theoretically less need for transformation of fully observed variables or investigation of nonlinearities and interactions. It is also possible to include a large number of related predictor variables in random forest models without encountering problems due to collinearity.

We included missForest and rfImpute in our study as examples of algorithms for completing single data sets (14). They replace missing values with predicted values rather than draw from a distribution, such that the imputed values do not have the correct joint distribution, leading to biased parameter estimates. Better predictions do not mean better coverage of confidence intervals; it is important that imputation methods incorporate the correct amount of variation in order to produce unbiased estimates with correct coverage of confidence intervals (34).

There was no difference in the results between linear regression and predictive mean matching. This was probably because the partially observed continuous variables in our data were approximately normally distributed; predictive mean matching may be preferred for variables that are not (conditionally) normally distributed (35).

**Limitations**

Although this study had strengths (it was based on real data, and the analysis was realistically complex), it also had important limitations. The most important limitation in producing general recommendations is that it was based on a single analysis of a single study, so results should be generalized to other data sets with caution.

A limitation of our resampling methodology was that in order to avoid excessive computing time we used only 10 imputations for most of the comparisons, leading to noisy estimates of between-imputation variability. To save time, we also restricted the number of cycles of MICE to 10, and although we evaluated plots of chain means and standard deviations between cycles for a few runs, this is a crude way of assessing chain convergence; it is possible that the chains may not have converged by the end of every run.

We ignored practice-level clustering at the imputation and analysis stages, for simplicity. If patients from the same practice are more similar than patients from different practices, the variance of parameter estimates might be underestimated, and parameter estimates may also be biased. This could be properly accounted for by using hierarchical models for analysis and imputation (36).

**Recommendations for further development**

We consider random forest multiple imputation to be promising, but it should be tested on a larger range of data sets and in simulations to explore whether it gives unbiased estimates where there are nontrivial nonlinearities or
interactions in imputation models, such that a standard parametric MICE imputation which ignores them gives biased results. Random forest tuning parameters (such as the number of trees and number of nodes) should be further investigated.

Conclusions

MICE is one of the recommended methods for multiple imputation in electronic health-record data, and we have shown that standard parametric MICE and our new random forest MICE method work reasonably well under artificially introduced missingness at random in a realistically complex data set. Random forest imputation should be further investigated in situations where MICE with default parametric imputation models produces biased results.

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