Accounting for Complex Sample Designs in Multiple Imputation Using the Finite
Population Bayesian Bootstrap

By

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DEDICATION

To my mom and dad
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ABSTRACT

Multiple imputation (MI) is a well-established method to handle item-nonresponse in sample surveys. Survey data obtained from complex sampling designs often involve features that include unequal probability of selection, clustering and stratification. Because sample design features are frequently related to survey outcomes of interest, the theory of MI requires including them in the imputation model to reduce the risks of model misspecification and hence to avoid biased inference. However, in practice multiply-imputed datasets from complex sample designs are typically imputed under simple random sampling assumptions and then analyzed using methods that account for the design features. Less commonly-used alternatives such as including case weights and/or dummy variables for strata and clusters as predictors typically require interaction terms for more complex estimators such as regression coefficients, and can be vulnerable to model misspecification and difficult to implement.

We develop a simple two-step MI framework that accounts for complex sample designs using a weighted finite population Bayesian bootstrap (FPBB) method to generate draws from the posterior predictive distribution of the population. Imputations may then be performed assuming IID data. We propose different variations of the weighted FPBB for different sampling designs, and evaluate these methods using three studies. Simulation results show that the proposed methods have good frequentist properties and are robust to model misspecification compared to alternative approaches. We apply the proposed method to accommodate missing data in the Behavioral Risk Factor Surveillance System, the National Automotive Sampling System and the National...
Health and Nutrition Examination Survey III when estimating means, quantiles and a variety of model parameters.
CHAPTER 1
INTRODUCTION

1.1 Objectives

Probability sample surveys where each member of the population has a known, non-zero probability of being selected form the backbone of empirical research in social science and public health. To assure broad representativeness, many sample survey designs use unequal probabilities of selection, selection of subjects in stages (introducing clustering) and stratification (For theoretical accounts of sampling methods, see Cochran, 1977 and Särndal et al., 1992). Accordingly, analysis methods for survey data need to take into account these complex sample design features.

It is notable that even the most well-designed sample surveys are imperfect in various ways. Missing data presents a particular challenge. Unit nonresponse occurs when sampled individuals fail to participate in the survey at all. Item nonresponse occurs when sampled individuals do not respond to certain questions. This is common in large scale surveys that include an extensive collection of questions. One principled approach to handle item nonresponse is multiple imputation (MI) (Rubin 1987). The key to success with MI lies in specifying an imputation model that reasonably describes the conditional distribution of the missing data given the observed data. Since complex sample design features frequently are related to survey variables, it is important to include them in the imputation model to reduce the risks of model misspecification and hence to avoid biased inference (Reiter, Raghunathan & Kinney, 2006). This thesis concerns methods that use
MI to deal with item nonresponse while accounting for complex sample designs. It has two objectives:

(1) To illustrate the bias that can arise using existing MI methods to account for complex sample designs when the imputation model is misspecified;

(2) To propose and evaluate a modified MI framework that accounts for complex sample designs with simpler modeling and with no resort to design-based estimators.

We explore the impact of the interrelationship among the data, the sampling mechanism and the missingness mechanism on the performances of the existing and the proposed MI methods. We use both simulated data and real survey data. The following assumptions are made throughout the thesis:

(1) There are no unit nonresponse problems;

(2) The data are missing at random (MAR).

Section 1.2 reviews approaches to survey sampling inference in the complete data context, with a focus on Bayesian approach. Section 1.3 reviews the standard multiple imputation (MI) as a well-established method to handle item nonresponse that has a Bayesian justification. The importance of accounting for complex sample designs in MI is discussed from a theoretical perspective. Section 1.4 and 1.5 point out the limitations of current techniques, and introduce the proposed method that builds upon the method reviewed in section 1.2. Finally, section 1.6 outlines the research question each chapter addresses.

1.2 Bayesian approach to survey sampling inference in the complete data context
Consider a finite population of size $N$ with two types of measurement on the population units: $\{Y_i, Z_i\}$ for $i = 1,...,N$, where $Y$ represents a single survey outcome of interest, and $Z$ represents the design variable used for sample selection. We consider three types of $Z$ in this thesis: (1) size measure for probability-proportional-to-size (PPS) sampling, (2) cluster indicators for two-stage cluster sampling, and (3) stratum indicators for stratified sample design. Typically $Z$ is known for all units of the population to the sampler, but not necessarily to the data analyst. Let $I_i = \{I_1, ..., I_N\}$ denote the vector of sample indicator variables, such that $I_i = 1$ if unit $i$ is sampled and 0 otherwise. We use subscripts $s$ and $ns$ to denote the selected sample of size $n$ and the nonsampled part of the population of size $N-n$, respectively. Thus both $Y$ and $Z$ divide into two parts: $\{(Y_s, y_{ns}), (Z_s, z_{ns})\}$. Let $w_i = \{w_s, i = 1,...,n.\}$ denote the sampling weights, e.g. $w_i = 1/\pi_i = \sum_{i=1}^{N} Z_i / nZ_i$ for a single-stage PPS sampling design. Figure 1.1 illustrates the data from a sample survey in the absence of missing data.

![Figure 1.1 The data from a sample survey in the absence of missing data](image-url)
In survey sampling, the objective is usually to learn about some population quantity denoted by $Q(Y)$, e.g. population mean/quantile/total, etc., through relating $Y_s$ to $Y_{ns}$ in some fashion. There are two general inferential frameworks to accomplish this: 1) the *design-based* approach (Hansen, Hurwitz & Madow, 1953; Cochran, 1977) treats the survey outcome $Y$ as a fixed quantity, and imposes a random distribution on the sample inclusion indicator $I$. The statistical distribution of an estimator for $Q(Y)$ is thus induced by the sampling design. While the design-based framework brings in an objectivity element by minimizing the use of modeling assumptions, this objectivity is lost in the presence of nonsampling errors like nonresponse; 2) the *model-based* approaches, which include the frequentist modeling (Royall, 1970; Valliant et al., 2000) and the Bayesian modeling (Ericson, 1969; Basu, 1971). These regard the survey outcome $Y$ as a random variable as well as $I$, and assume a model to predict $Y_{ns}$ from $Y_s$. Both variants assume that $Y$ comes from some parametric family of distributions indexed by the parameter $\theta$.

While the frequentist modeling treats $\theta$ as fixed, and bases inferences on repeated sampling from the model, the Bayesian modeling specifies a prior distribution on $\theta$ in addition to $Y$. The posterior distribution of $\theta$ given the observed sample $p(\theta | Y_s)$, and hence the posterior predictive distribution of the nonsampled population values given the sampled data $p(Y_{ns} | Y_s) = \int p(Y_{ns} | \theta, Y_s)p(\theta | Y_s)d\theta$, serves as the basis for inference about $Q(Y)$.

The Bayesian paradigm provides the most satisfying inferential approach to survey inference when it is done right. That is by incorporating complex sample design features to avoid sensitivity to model misspecification, using noninformative priors to
avoid subjectivity, and being frequentist calibrated, i.e. having good repeated sampling properties (Little, 2004; Little & Zheng, 2007). Bayesian finite population inference is thus proposed as a means to harmonize design- and model-based approaches for sample survey inference (Little, 2006, 2011; Gelman, 2007), and is exemplified by a variety of work in the complete data context. When design variables or the selection probabilities are known for all units in the population, Zheng and Little (2003, 2005) and Chen, Elliott and Little (2010) propose robust Bayesian predictive inference that improves efficiency over design-based estimators, using penalized splines under fairly weak model assumptions. In situations where the sizes of the non-sampled units are unavailable, Little and Zheng (2007) and Sangeneh, Keener and Little (2011) consider a two-step procedure to assure ignorable sampling (Sugden and Smith, 1984) with a PPS sampling design. In the first step, the nonsampled sizes \(Z_{ns}\) are predicted by a modified Bayesian Bootstrap (BB) procedure that adjusts for unequal probability sample selection \(w_s\), i.e. 
\[ p(Z_{ns} | Z_s, w_s) \]; the nonsampled survey outcomes \(Y_s\) are then predicted using a penalized spline model relating \(Y\) with \(Z\), i.e. 
\[ p(Y_{ns} | Y_s, Z) . \]

The modified BB considered by Little and Zheng (2007) is a noninformative Bayesian method closely related to an offshoot of the Bayesian approach to surveys, known as the “Pseudo-Bayesian approach” (Ghosh & Meeden, 1997; Cohen, 1997; Dong, Elliott & Raghunathan, 2014). The modified BB is of particular interest in this thesis. Details of how our proposed methodology in this thesis builds upon and compares with these methods will be discussed in section 1.5 and in later chapters.

1.3 Standard multiple imputation as a calibrated Bayesian method to deal with item nonresponse
Consider the same finite population as in section 1.2, and assume item nonresponse occurs on the outcome variable $Y$ and a covariate $X$ is completely observed. Let $R_i = \{R_i, \ldots, R_N\}$ denote the response indicator variable for $Y$, such that $R_i = 1$ if unit $i$ provides a value for $Y$ and 0 otherwise. Write $R = (R_s, R_{ns})$, where $R_s$ is observable from the sample and $R_{ns}$ is unobservable for the nonsampled population. We use subscripts $obs$ and $mis$ to denote the responding and the non-responding units, respectively. Thus $Y_s$ further breaks down to $Y_{s,obs}$ and $Y_{s,mis}$, and $Y = (Y_{s,obs}, Y_{nobs})$ if we recombine $Y_{s,mis}$ with $Y_{ns}$ as unobserved data $Y_{nobs}$. The observed data may be thought as the outcome of two random processes: sampling and responding. We illustrate in Figure 1.2 the data from a sample survey with item nonresponse occurring on the survey outcome. Three types of population quantities are of interest in the thesis: the mean/proportion of $Y$, the regression coefficients of $Y$ on $X$, and the quantiles of a continuous outcome $Y$.

![Figure 1.2](image)

The Bayesian modeling paradigm deals with nonresponse naturally, since unknowns about the finite population given the observed data can be generated from a
predictive distribution. We thus consider multiple imputation (MI) to deal with item nonresponse which has a Bayesian justification. The basic idea of MI is to replace each missing value with a set of plausible values which can then be combined in a simple way for inference using complete-data analysis techniques. The fundamental conceptualization of MI is Bayesian, where the posterior distribution of the model parameters $\theta$ of interest is obtained by averaging the completed data posterior of $\theta$ over the posterior predictive distribution of the missing data (Rubin, 1987, Result 3.1):

$$p(\theta|Y_{s,obs},X_s,Z,R,I) = \int p(\theta|Y_s,X_s,Z,R,I) p(Y_{s,mis}|Y_{s,obs},X_s,Z,R,I) dY_{s,mis} \quad [1.1]$$

Assuming ignorable sampling $p(I|Y,R,Z) = p(I|R,Y,Z)$ and $p(R|Y,Z) = p(R|Y_{obs},Z)$, [1.1] becomes

$$p(\theta|Y_{s,obs},X_s,Z) = \int p(\theta|Y_s,X_s,Z) p(Y_{s,mis}|Y_{s,obs},X_s,Z) dY_{s,mis} ,$$

allowing the sampling and response mechanism to be ignored in the modeling.

This integration is typically accomplished using Markov Chain Monte Carlo data augmentation (Tanner & Wong, 1987). Let $t$ index the iteration. Draws from the posterior predictive distribution of the missing data $p(Y^{(t)}_{s,mis}|Y_{s,obs},X_s,Z)$ are obtained by iterating between draws of the model parameters conditional on the “filled in data” $p(\theta^{(t)}|Y_{s,obs},Y^{(t-1)}_{s,mis},X_s,Z)$ and imputations of the missing data conditional on the observed data and draw of the model parameter $p(Y^{(t)}_{s,mis}|Y_{s,obs},\theta^{(t)},X_s,Z)$ . Rubin (1987) develops simple combining rules to obtain estimates of posterior means and variances using only a finite number ($M$) of independent draws $\{Y^{(1)}_{s,mis},...,Y^{(M)}_{s,mis}\}$ of the imputed data together with $Y_{s,obs}$, i.e. multiple “completed” datasets $Y^{(l)}_{comp} = \{Y_{s,obs},Y^{(l)}_{s,mis}\}$, $l=1,...,M$. He shows that inferences obtained using these combining rules have good frequentist properties for relatively small $M$ (5-20):
Posterior Mean: \( \hat{Q} = M^{-1} \sum_{l=1}^{M} Q(y^{(l)}_{\text{comp}}) \)  

Posterior Variance: \( V = U + (1 + M^{-1})B \),

where \( Q(y^{(l)}_{\text{comp}}) \) is the point estimate obtained from the \( l^{th} \) completed dataset \( y^{(l)}_{\text{comp}} \),

\[
U = M^{-1} \sum_{l=1}^{M} \text{vár}(Q(y^{(l)}_{\text{comp}})) \]

is the within imputation variance calculated as the average of variance estimates based on the \( M \) completed datasets, \( B = (M - 1)^{-1} \sum_{l=1}^{M} (\hat{Q} - Q(y^{(l)}_{\text{comp}}))^2 \) is the between imputation variance.

Typically this is a combined design and model set up, where an imputation model is used to predict missing data in the sample. In other words, prediction of \( p(Y_{s,\text{mis}}|Y_{s,\text{obs}}) \) is model-based, and design-weighted estimators (i.e. \( Q(y^{(l)}_{\text{comp}}) \) and \( \text{vár}(Q(y^{(l)}_{\text{comp}})) \)) are used to estimate the finite population quantities of interest once \( Y_{s,\text{mis}} \) are filled in by model predictions, i.e. predictions of \( p(Y_{n}\|Y_{s}) \) is design-based (Reiter et al., 2006; Yuan & Little 2007a).

Rubin (1987) combines nonsampled and missing data into a single framework:

\[
p(Y_{\text{obs}}\|Y_{s,\text{obs}}, X, Z, R, I; \theta, \varphi, \phi) \propto \int p(Y, X, Z; \theta) p(R|Y, X, Z; \phi) p(I|Y, X, Z; \varphi) dR_{\text{ns}} \quad [1.3]
\]

where \( \theta, \varphi \) and \( \phi \) denote the parameter that indexes the distribution of \( Y, I \) and \( R \), respectively, and they are assumed a priori independent. He also gives conditions for proper imputation that ensure randomization validity for a calibrated Bayes in the sense of Little (2006, 2011). Rubin’s conditions include: (i) point estimation is approximately unbiased for the scientific estimand of interest \( Q(Y) \), and (ii) actual interval coverage equals the nominal interval coverage, over repeated imputation and sampling processes.
These in turn require *ignorability* assumptions of two random mechanisms— the specified sampling mechanism \( I \) and the posited missing data mechanism \( R \). The *ignorability conditions* (Rubin, 1976; Little, 1982) are:

- **Ignorable sampling:**
  \[
  p(I \mid R_{s}, Y_{s}, X_{s}, Z_{s}; \phi) = p(I \mid R_{s}, Y_{s, obs}, X_{s}, Z_{s}; \phi) \\
  \iff p(Y_{nobs} \mid Y_{s, obs}, X_{s}, Z_{s}, R_{s}, I; \theta, \phi, \phi) = p(Y_{nobs} \mid Y_{s, obs}, X_{s}, Z_{s}; \theta, \phi)
  \]

- **Ignorable missing data:**
  \[
  p(R_{s} \mid Y_{s}, X_{s}, Z_{s}; \phi) = p(R_{s} \mid Y_{s, obs}, X_{s}, Z_{s}; \phi) \\
  \iff p(Y_{nobs} \mid Y_{s, obs}, X_{s}, Z_{s}, R_{s}; \theta, \phi) = p(Y_{nobs} \mid Y_{s, obs}, X_{s}, Z_{s}; \theta)
  \]

Thus explicit modeling for the sample indicator \( I \) and the response indicator \( R_{s} \) is not necessary in equation [1.3].

Rubin (1976) and Little and Rubin (2002) formalized the concept of *missing data mechanism*. By their definition, ignorable missing data always implies missing at random (MAR). That is, given the observed data \((Y_{s, obs}, X, Z)\), the missingness mechanism does not depend on the unobserved data \((Y_{nobs})\). MAR is a common assumption in practice and is typically assumed for implementing MI. Other missing data mechanisms include missing completely at random (MCAR), i.e. \( p(R_{s} \mid Y_{s}, X, Z; \phi) = p(R_{s} \mid \phi) \), and not missing at random (NMAR), i.e. the missingness also depends on the unobserved data \((Y_{nobs})\). We focus on MAR in this thesis. We do not consider MCAR which is often too ideal for real world sample surveys, or NMAR, which requires special statistical techniques (e.g. selection and pattern mixture models) beyond the scope of this thesis.

To make the MAR assumption plausible, we need a sufficiently rich imputation model that ideally includes in the covariate space \( \{X, Z\} \) all variables that are related to
the sample selection and the missingness, and are potentially predictive of the outcome variable of interest \((Y)\). For a complex sample survey, this implies that features such as stratification and clustering as well as unequal inclusion probabilities need to be built into the imputation model via design variables (Rubin, 1996).

1.4 Fully parametric techniques to account for complex sample designs in MI

Despite expert recommendations, imputers seldom account for sample designs when using available software packages to construct imputation models. They rely instead, on use of design-based estimators at the analysis stage to account for design effects. This can lead to biased point estimation and below-nominal confidence interval coverage (Reiter et al., 2006). More complex methods utilize design variables \((Z)\) as covariates in models. In the setting where the design issue of interest lies in the probability of selection, Elliott and Little (2000) and Elliott (2007, 2008, 2009) account for unequal probabilities of inclusion by considering weight strata and treat the stratum means as random effects (which they term “weight smoothing models”). Their ideas to shrink the mean across weight strata are recently considered for accommodating survey weights in MI using random-effects imputation models (Carpenter et al., 2012). With stratified multistage sampling, Reiter et al. (2006) and Schenker et al. (2006) consider dummies for fixed stratum effects and fixed or random cluster effects in the imputation model. We call these MI techniques “fully parametric MI”.

In practice, however, as more covariates and design variables are included in the model, the inferential conclusions become more valid conditionally but possibly more sensitive to model misspecification (Gelman et al., 2004). This leads to “uncongenial”
imputation (Meng, 1994)—since oftentimes, correct parametric or nonparametric approximations of the functional forms of design variables as well as their interactions with other covariates are needed. Pfeffermann (2011) points out that, modeling the relationship between the design variables and the covariates in order to integrate out the effect of the former can be complicated. In particular, uncongeniality concerns regarding the incorporation of survey weights in the imputation model for domain estimation (Kim et al., 2006), overestimation of MI variance under the fixed cluster effects model (Andridge, 2011), and convergence issues about random cluster effects models with high dimensional data and/or hierarchical data structure (Yucel & Raghunathan 2006; Zhao & Yucel, 2009) all limit the development of adequate software packages to apply these MI techniques.

The importance of accounting for sample designs in multiple imputation, and the limitation of fully parametric MI techniques to do this, creates an awkward dissonance between the theory and practice of multiple imputation. This also provides a good motivation for an alternative methodology that ideally satisfies the following properties:

- Consistent with the Bayesian derivation of MI
- Easier implementation and less expensive computation
- Less prone to model misspecification
- Do not require design-based estimators for complex population quantities of interest under complex sampling designs (e.g. domain estimation, quantile estimation, and the ratio of two medians for which there seems no readily available frequentist interval)
1.5 Proposed methodology to account for complex sample designs in MI

This thesis develops a modified MI framework (termed “two-step MI”) to accommodate complex sample designs, which avoids the complication of modeling design variables \((Z)\) and obviates the need for design-based analysis.

In the first step, we regard the nonsampled part of the target population as missing by design, and create synthetic populations that contain item-level missing data. Thus we obtain \(p\left(Y_{ns}, X_{ns}, R_{ns}, w_{ns} \mid Y_s, X_s, R_s, w_s\right)\), where \(Y_{ns} = (Y_{ns,obs}, Y_{ns,miss})\), \(w_s\) is the design weight for sampled units. In the second step, we multiply impute the missing values in the synthetic populations using standard parametric MI techniques assuming that the data are independent and identically distributed. Thus we obtain \(p(Y_{mis} \mid Y_{obs}, X)\), where \(Y_{mis} = (Y_{s,mis}, Y_{ns,mis})\), \(Y_{obs} = (Y_{s,obs}, Y_{ns,obs})\). This is equivalent to a situation where a census is conducted but not all respondents answer all the survey questions, requiring multiple imputation to be performed on the entire target population.

Note that the proposed method follows a “synthesize/reverse design-then-impute” procedure, different from the procedure of Reiter (2004) who used the parametric MI to simultaneously impute missing data and generate synthetic data in a “two-step” fashion for disclosure risk limitation purpose. Reiter follows a “impute-then-synthesize” procedure in which modeling design variables in the imputation model remains an issue.

Note also that although the proposed MI framework requires one step to deal with nonsampled data, it is an intermediate step toward our ultimate goal of treating item-level missing data.

To realize the first step in the proposed MI procedure, we consider a ‘Pseudo-
Bayesian approach’, i.e. the Polya posterior of Ghosh & Meeden (1997), which is equivalent to the finite population Bayesian bootstrap of Lo (1988). The approach ensures objectivity relative to a standard Bayesian approach by assuming a noninformative prior that is dominated by a nonparametric likelihood function. Because the sampling designs considered in this thesis all involve unequal selection probabilities, we consider the weighted version of the finite population Bayesian bootstrap (‘weighted FPBB’) (Cohen, 1997; Little & Zheng, 2007; Dong et al., 2014). We also develop several adapted versions of the weighted FPBB pertinent to accommodating different sample design features.

Though built upon the modified BB in Little and Zheng (2007), we have a different implementation of it because we have a different objective in the missing data context. We create the posterior joint distribution of the nonsampled population 

\((Y_{ns}, X_{ns}, R_{ns})\) based on the weighted FPBB, whereas they ‘impute’ the unknown design variable \(Z_{ns}\) only. Our purpose is to reverse design effects while retaining population-level multivariate relationships among variables in the process. As stated previously, \(Z_{ns}\) is not imputed in our case, since we focus on model predictions of item-level missing data, and do not consider further capitalizing on \(Z\) to model the relationships of the design variable \((Z)\) and the outcome \((Y)\), as they do in their second step. Further, our implementation assumes no auxiliary information is available for the distribution of \(Z\) (e.g. the population mean of \(Z\)), and therefore do not adjust the weighted FPBB as they do to create synthetic populations that satisfy such a restriction.

1.6 Outline of chapters

Chapter 2 proposes a two-step MI framework to account for sampling weights.
We combine the method of Ghosh and Meeden (1997), who propose a Polya posterior to generate a noninformative Bayesian approach to finite population sampling, with that of Cohen (1997), who proposes a method to generate draws from Polya posteriors using data obtained from weighted sample designs in a non-clustered setting. We then modify the standard MI combining rules for inference that follow immediately from the rules developed in Raghunathan et al. (2003) for combining synthetic datasets. While the literature recognizes the importance of using weights in the analysis of complex sample survey data, methods incorporating weights in conjunction with multiple imputation to adjust for item nonresponse are underdeveloped. Imputation models that simply include weights or weight-related design variables as covariates can quickly become very complicated. They can be susceptible to misspecification if the inclusion probabilities are related to survey variables \( Y \) and/or the missing data mechanism in a nonlinear fashion.

The new procedure allows the sampling mechanism \( (I) \) and missing data mechanism \( (R) \) to be simultaneously disentangled, so that imputation can be performed assuming IID. As a result, data users need only to apply simple unweighted estimation methods to the imputed population datasets.

Our simulation assuming a PPS sampling design shows that the proposed method has good frequentist properties under MAR, which contrasts with standard approaches that are prone to model misspecification. We also apply the proposed method to estimation of means and linear regression, log-linear regression, and general location models using data from the BRFSS.

Chapter 3 extends the proposed MI framework to accommodate clustering effects as well as sample weight effects in two-stage unbalanced cluster samples. We extend the
approach of Chapter 2, this time combining the method of Meeden (1999), who proposes ‘a two-stage Polya posterior’ approach to a simple balanced two-stage cluster sample with equal selection probability at both cluster and element levels, with that of Cohen (1997). Because this approach requires that both first- and second-stage weights be known, we also propose an alternative that uses a standard Bayesian bootstrap at the resampling of the clusters. We show that this method also produces draws from the posterior predictive distribution of the population that incorporate both clustering and weighting components of the sample design while requiring only the final weights that are usually supplied in public databases. Their performances are evaluated under different population models and different degrees of clustering effects. Small and large sample behaviors of alternative methods are also investigated. While the framework of fully parametric MI does not seem to provide a direct and robust technique to deal with sample weights in hierarchical models, the proposed method turns out to be an easy alternative that recovers most of the information in the data generating mechanisms. We apply the different MI techniques to the analysis of passenger vehicle injury data from the National Automotive Sampling System – Crashworthiness Data System (NASS-CDS) survey, in particular, estimates of mean “delta-v” (instantaneous deceleration velocity) and its associated injury.

Chapter 4 develops a general purpose MI approach to account for various sample design features in a highly stratified multistage sample. Our specific focus is on evaluating the performance of the methods in comparison with existing MI techniques, with respect to several frequently encountered yet not previously addressed issues in the statistical analysis of missing data. These include: (i) accommodating stratification and
multi-stage sampling in the imputation process; (ii) the employment of logistic imputation models for estimating probabilities of rare events; and (iii) the estimation of population quantiles with multiply imputed data. In the simulation study, the proposed procedures demonstrate fairly good coverage properties for nonsmooth statistics over the entire support of the distribution for a continuous variable of interest, and yield quite stable parameter estimates in the case of sparse data. We argue that the proposed methods offer a computationally feasible solution to problems that are not well handled by current MI techniques. This method is applied to accommodate missing body mass index (BMI) data in the analysis of BMI percentiles using NHANES III data.

Chapter 5 summarizes findings in the thesis and points out both the advantages and limitations of the proposed methodology. We also point to several directions of future research, including the applications of the proposed methodology to domain/small area estimation, adaptations of it to unit nonresponse and to propagation of uncertainty in unit nonresponse adjustments, and possible extensions of it to incorporate auxiliary information.
CHAPTER 2
A TWO-STEP SEMIPARAMETRIC METHOD TO ACCOMMODATE
SAMPLING WEIGHTS IN MULTIPLE IMPUTATION

2.1 Introduction

Both item nonresponse and sampling weights are typical features of survey data obtained from complex sample designs. Item nonresponse occurs when some respondents do not answer all the items in a survey questionnaire. Both “don’t know” and refusal answers are considered as item nonresponse. Sampling weights arise as a correction factor to compensate for over- or under-representation of units in the target population due to unequal selection probabilities. Examples of unequal probability sampling designs include: i) disproportionate stratified sampling, where sampling units in each stratum have differential selection probabilities, and ii) probability proportional to size (PPS) sampling, in which the probability of selection for a sampling unit is proportional to a positive size measure (Z) known for all population units. For example, the Behavior Risk Factor Surveillance System (BRFSS) has both a substantial amount of missing data on income measures as well as survey weights that adjust for oversampling of adults in smaller sized households and for selection bias by poststratifying and raking to known control totals for basic demographics.

When the amount of item-level missing values is nontrivial and the data are not missing completely at random (MCAR), typical solutions for missing data like the
complete case analysis often lead to reduction of statistical power, and problematic inferences of the target population. Multiple imputation (MI) (Rubin, 1987, 1996) is a more principled method for addressing item-level missing data, which has a Bayesian conceptualization. The basic idea is to fill in missing data with \( M \) sets of plausible values. These values are obtained as repeated draws from the posterior predictive distribution of the missing components of the sample \( Y_{s,mis} \) given its observed components \( Y_{s,obs} \), i.e.

\[
p(Y_{s,mis} | Y_{s,obs}),
\]

where \( p(\cdot) \) denotes the probability density function. (Typically \( M \) is small, e.g. \( M=3\sim5 \), but larger \( M (10\sim20) \) may be needed to obtain stable estimates of variance when the fraction of missing information is large.) The production of multiple “completed” datasets \( \{(Y_{s,obs}, Y_{s,mis}^{(1)}), \ldots, (Y_{s,obs}, Y_{s,mis}^{(M)})\} \) is typically done by an “imputer” who has access to the data to develop reasonable models for generating the predictive distribution of \( Y_{s,mis} \), allowing the “analyst” to then analyze each of the \( M \) imputed datasets, and combine the point and variance estimates using the combining rules developed by Rubin (1987). Examples of this approach include imputation for blood alcohol concentration in the Fatal Accident Reporting System (FARS) (Heitjan & Little, 1991) and income imputation in the National Health Interview Survey (NHIS) (Schenker et al., 2006).

The implementation of multiple imputation typically assumes missing at random (MAR), that is, given the observed data, the reason for the missing data does not depend on the unobserved data. To make MAR plausible, it is important to let the imputation model condition on all variables (including sample design variables) that are either predictive of the outcome \( (Y) \) or the missing data mechanism \( (R) \). In settings where the observed data are obtained using unequal probability sampling design, however, data are
typically imputed using models where variables are assumed to be independent and identically distributed (IID). They are then analyzed using a design-weighted approach that accounts for the unequal selection probability. This can lead to biased point estimation and below-nominal confidence interval coverage.

A simple and seemingly straightforward way to incorporate sampling weights in MI is using fully parametric techniques. One option is to require the imputer’s model to be conditioned on a few key design variables that determine the individuals’ probabilities of inclusion, such as measure of size and stratification variables (e.g. demographics, socioeconomic status, as well as geographical characteristics, etc.). Another option is to summarize the design information by using weights as a covariate in the imputation, perhaps after log transformation or categorization in “weight strata” and modeling them as dummy indicators. However, the modeling task may be complicated by attempts to include all interactions of weights (or weight-related design variables) with other covariates in the model, particularly the interaction of the weights with domain indicators (Meng 1994; Kim et al. 2006; Seaman et al. 2006). Moreover, this approach typically requires the functional form of the interaction to be modeled correctly, perhaps using a spline or other non-parametric form to be robust against model misspecification (Elliott & Little 2000; Zheng & Little 2005; Breidt, Claeskens, & Opsomer, 2005). It can be a challenging task to come up with a robust imputation model that is attentive to sampling weights and sufficiently captures all relevant aspects of the distribution of \( Y \) of interest.

This chapter develops a modified MI framework to account for sampling weights from single-stage sampling designs. The primary goals are:

1) to propose a two-step MI procedure. In the first step, nonparametric models
are used to generate the posterior predictive distribution of the population that includes the item-level missing data. In the second step, parametric models assuming IID are used to impute the missing values in the created populations. We consider utilizing the weighted finite population Bayesian bootstrap (weighted FPBB) to account for sampling weights in the first step;

2) to illustrate the impact, during the imputation process, of ignoring sampling weights or unequal probabilities of selection, on the bias properties of both the MI point estimator and the MI variance estimator; and

3) to compare the performances of the proposed two-step MI and the fully parametric MI in terms of robustness to different degrees of model misspecification. The comparison will be made under different scenarios, defined by the associations of the design variable \((Z)\) used for determining the selection probabilities with both the outcome variable \((Y)\) where missing data occur, and the latent variable \((T)\) which defines the response mechanism.

The rest of this chapter is organized as follows. Section 2.2 provides a detailed overview of the proposed two-step semiparametric multiple imputation procedure. (We term it “semiparametric” because the design feature, in particular the weights, are accommodated non-parametrically, whereas the actual imputation is conducted under a standard parametric model.) Section 2.3 discusses point estimation and inference using the MI datasets from the proposed procedure. Section 2.4 provides a simulation study in the context of a single-stage probability-proportional-to-size (PPS) sample design. We estimate population means and regression coefficients under settings where sampling weights are associated to differing degrees with both the outcome and the probability of
nonresponse, and where failure to account for design in the imputation procedure has differing degrees of impact. Section 2.5 applies the proposed procedure to estimate means, linear and loglinear regression models, and general location models describing marginal and joint distributions of income and health insurance accessibility. Section 2.6 discusses possible extensions to incorporate other design features beyond sampling weights, as well as extensions to deal with unit non-response weights.

2.2 A Two-Step Semiparametric MI Procedure

2.2.1 Overview and Notation

Bayesian finite population inference (Ericson 1969) has been proposed as a means to harmonize design- and model-based approaches for sample survey inference (Little 2004, 2011; Gelman 2007). Under this approach, we focus on the posterior predictive distribution of the finite population quantity of interest (e.g., population mean, population regression parameter) obtained from the posterior predictive distribution for the non-sampled elements of the population. To make matters more concrete, consider the setting where we have a scalar outcome \( Y \), sampling weight \( w \) based on a single-stage PPS sampling design, and no missing data. Our complete data consist of the vector of sampling indicators \( I \) for the population, sampled \( Y \), for which \( I = 1 \), the non-sampled \( Y \), for which \( I = 0 \), and similarly \( w \) and \( w \). Given the sampling weights, the sampling mechanism generating \( I \) is assumed to be ignorable \( ( p(I | Y, w) = p(I | w) ) \), and can be ignored in the modeling. Assuming a model for the outcome given the sampling weights \( p(Y | \theta, w) \) parameterized by \( \theta \) with prior \( p(\theta) \), the posterior predictive distribution for the non-sampled elements of the population \( Y \) is given by
\[p(Y_{ns} | Y_s, w_s) \propto \int p(Y_{ns} | Y_s, \theta, w) p(\theta | Y_s, w) p(w_{ns} | w_s) d\theta dw_{ns}\]  \[2.1\]

Previous work has tackled estimation of this predictive distribution in a variety of manners. Zheng and Little (2003, 2005) and Chen, Elliott and Little (2010) assumed that the sampling weights were known for all subjects, so that \(w_s = w\), reducing [2.1] to

\[p(Y_{ns} | Y_s, w) \propto \int p(Y_{ns} | Y_s, \theta, w) p(\theta | Y_s, w) d\theta ; \]

these authors then obtained draws from the posterior predictive distribution under fairly weak modeling assumptions (parametric regression model for \(p(Y | \theta, w)\) based on penalized splines). Little and Zheng (2007) and Zangeneh, Keener, and Little (2011) considered the situation in which weights (or equivalently the size measure \(Z\)) are observed only for the sample (as in a public use data setting), and obtained predictive draws for \(p(w_{ns} | w_s)\) under a Dirichlet model with a non-informative (Haldane) prior; the resulting predictive draw of the population of weights \(w\) was then used as covariates in Little and Zheng to obtain posterior predictive draws of \(Y_{ns}\).

Dong, Elliott, and Raghunathan (2014) consider a different factorization of [2.1]:

\[p(Y_{ns} | Y_s, w_s) \propto \int p(Y_{ns}, w_{ns} | Y_s, w_s) p(Y_s, w_s) dw_{ns}\]  \[2.2\]

The parameter \(\theta\) is dropped because the draws of \(p(Y_s, w_s)\) are made directly from the posterior of the empirical joint CDF of \(Y_s, w_s\) using a Bayesian bootstrap (BB) procedure (Rubin 1981). Draws from \(p(Y_{ns}, w_{ns} | Y_s, w_s)\) are then made using a weighted finite population Bayesian bootstrap (FPBB) procedure described in Cohen (1997).

Here we extend the approach of Dong, Elliott, and Raghunathan to accommodate missing data. We assume that, had we taken a census of the entire population, we could have observed a vector of response indicators \(R = (R_s, R_{ns})\), where \(R_s\) corresponds to the response indicators observed in the sample, and \(R_{ns}\) to the response indicators associated...
with the non-sampled elements. We then divide the sampled $Y_s = (Y_{s,obs}, Y_{s,mis})$ into the fully-observed and missing elements, corresponding to the sampled $Y$ values associated with $R_s = 1$ and $R_s = 0$ respectively, and similarly the non-sampled $Y_{ns} = (Y_{ns,obs}, Y_{ns,mis})$ into those that would have been observed had they been sampled ($R_{ns} = 1$), and those that would have had missing values ($R_{ns} = 0$). We also assume a fully-observable covariate $X = (X_s, X_{ns})$ consisting of the sampled and nonsampled elements respectively. Note that we can combine the observed from the sampled and nonsampled parts of the population to obtain the potentially “observable” $Y_{obs} = (Y_{s,obs}, Y_{ns,obs})$, and similarly $Y_{mis} = (Y_{s,mis}, Y_{ns,mis})$.

We assume ignorable missingness, so that $p(R | Y, X, w) = p(R | Y_{obs}, X, w)$, allowing $R$ to be ignored in the model along with $I$. Extending [2.2] to incorporate item-level missingness (and thus the covariate $X$ used for imputation) then yields

$$p(Y_{ns}, X_{ns} | Y_s, X_s, w_j) = p(Y_{ns,obs}, Y_{ns,mis}, X_{ns} | Y_{s,obs}, Y_{s,mis}, X_s, w_j) \propto \int p(Y_{ns,obs}, Y_{ns,mis}, X_{ns}, w_{ns} | Y_{s,obs}, Y_{s,mis}, X_s, w_j) p(Y_{s,obs}, Y_{s,mis}, X_s, w_j) dw_{ns}$$

[2.3]

To obtain the posterior predictive distribution based on fully-observed data, we need to integrate the left side of [2.3] with respect to $Y_{mis} = (Y_{s,mis}, Y_{ns,mis})$.

$$p(Y_{ns,obs}, X_{ns} | Y_s, obs, X_s, w_j) \propto \int p(Y_{ns,obs}, X_{ns} | Y_{s,obs}, X_s, w_j) p(Y_{mis} | Y_{s,obs}, X_s, w_j) dY_{mis}$$

To accomplish this, we reintroduce a parametric model for $p(Y | \theta, X, w)$, and integrate out with respect to the posterior distribution of $\theta$:

$$p(Y_{mis} | Y_{s,obs}, X_s, w_j) = \int p(Y_{mis} | Y_{s,obs}, X_s, \theta, w_j) p(\theta | Y_{s,obs}, X_s, w_j) d\theta$$.

Thus [2.3] becomes

$$p(Y_{mis} | Y_{s,obs}, X_s, w_j) = \int \int p(Y_{ns,obs}, X_{ns} | Y_{mis}, X_{obs}, X_s, w_j) p(Y_{mis} | Y_{s,obs}, X_s, \theta, w_j) p(\theta | Y_{s,obs}, X_s, w_j) d\theta dY_{mis}$$

[2.4]

$$p(Y_{ns,obs}, X_{ns} | Y_{s,obs}, X_s, w_j) \int \int p(Y_{mis} | Y_{s,obs}, X_s, \theta, w_j) p(\theta | Y_{s,obs}, X_s, w_j) d\theta dY_{mis}$$

where the last equality in [2.4] follows from the fact that the unobserved (but potentially
observable) elements of the population are generated using the finite population Bayesian bootstrap.

A key result from [2.4] is that, since the unobserved elements of the population have been generated non-parametrically, we can implement the integration in [2.4] by use of a Gibbs sampler that iterates between obtaining draws of $\theta$ conditional on the entire population, and draws of the missing data conditional on $\theta$ and the observable elements of the population

$$p(Y_{\text{mis}} | Y_{\text{obs}}, X, \theta, w) = p(Y_{\text{mis}} | Y_{\text{obs}}, X, \theta)$$
$$p(\theta | Y, X, w) = p(\theta | Y, X)$$

Since the nonparametric procedure generates draws from the joint posterior distribution of all variables for the nonsampled population, the relationships of the weights and other variables are maintained in the draws. It is thus sufficient to develop a parametric model for $Y$ that does not involve the weights: $p(Y | \theta, X, w) = p(Y | \theta, X)$. This model does need to condition on weights, however, when the weights are further associated with the missing data mechanism.

Figure 2.1 shows the creation of a single imputed synthetic population dataset under the proposed two-step MI procedure: a) shows the original sample data, b) the result from the BB-weighted FPBB procedure, and c) the result from the (model-based) imputation procedure. The shaded area represents observed data and ‘?’ represents missing data. We discuss in detail the derivation and implementation of the proposed two-step MI procedure below.
2.2.2 Step 1: Undo Sampling Weights through Synthetic Data Generation

The Pólya’s Urn Scheme

The Polya urn distribution (Feller, 1968) is defined by construction as follows: suppose we have an urn containing a finite number \( n \) of balls of different colors. A ball is randomly drawn from the urn and another ball with the same color from outside of the urn is added back to the urn along with the originally picked one. Repeat this selection process until \( m \) balls have been selected; the resulting sample is termed a ‘Pólya sample of size \( m \)’.

The Pólya Posterior

The Polya posterior (Ghosh & Meeden, 1997) derives its name from the Polya urn distribution. It is a noninformative Bayesian procedure which can be used when little or no prior information is available. One advantage of the Polya posterior is that it has a stepwise Bayes justification (Hsuan, 1979) and leads to admissible procedures.

Assume that a simple random sample of size \( n \) is drawn from a finite population.
of size $N$. Let $y = \{y_1, \ldots, y_n\}$ denote the sample, where $y$ represents the realized value of a response variable $Y$. Let $\{d_1, d_2, \ldots, d_K\}$ denote the set of $K$ distinct values in the sample and $\hat{\lambda} = \{\hat{\lambda}_1, \hat{\lambda}_2, \ldots, \hat{\lambda}_K\}$ the vector of probabilities, such that $\Pr(y_i = d_j | \hat{\lambda}) = \hat{\lambda}_j$, for $i = 1, 2, \ldots, n$, $j = 1, \ldots, K$, and $\sum_{j=1}^{K} \hat{\lambda}_j = 1$. Let $n_j$ and $n_j'$ be the number of units taking value $d_j$ in the sample and in the nonsampled part of the population, respectively, for $j = 1, 2, \ldots, K$, and $\sum_{j=1}^{K} n_j = n$, $\sum_{j=1}^{K} n_j' = N - n$. Assuming a noninformative Haldane prior of $\hat{\lambda}$: $\hat{\lambda} \sim \text{Dir}(0, \ldots, 0)$, $p(\hat{\lambda}) \propto \prod_{j=1}^{K} \hat{\lambda}_j^{-1}$, together with a multinomial distribution for the counts of sample data: $n_1, \ldots, n_K | \hat{\lambda} \sim \text{Mult}(n; \hat{\lambda})$,

$$p(n_1, \ldots, n_K | \hat{\lambda}) = \frac{n!}{\prod_{j=1}^{K} n_j!} \prod_{j=1}^{K} \hat{\lambda}_j^{n_j} \propto \prod_{j=1}^{K} \hat{\lambda}_j^{n_j}$$, yields a Dirichlet posterior distribution of $\hat{\lambda}$:

$$\hat{\lambda} | n_1, \ldots, n_K \sim \text{Dir}(n_1, \ldots, n_K), \ p(\hat{\lambda} | n_1, \ldots, n_K) \propto \prod_{j=1}^{K} \hat{\lambda}_j^{n_j - 1}$$.

The Posterior predictive distribution of counts in the nonsampled data thus follows a compound multinomial distribution $n_1', \ldots, n_K' | n_1, \ldots, n_K \sim \text{Mult}(N - n; \hat{\lambda})$:

$$p(n_1', \ldots, n_K' | n_1, \ldots, n_K) = \frac{\int_{0}^{1} \cdots \int_{0}^{1} \prod_{j=1}^{K} p(n_1', \ldots, n_K' | n_1, \ldots, n_K, \hat{\lambda}) p(n_1, \ldots, n_K | \hat{\lambda}) d\hat{\lambda}_1 \cdots d\hat{\lambda}_K}{\int_{0}^{1} \cdots \int_{0}^{1} \prod_{j=1}^{K} p(n_1, \ldots, n_K | \hat{\lambda}) d\hat{\lambda}_1 \cdots d\hat{\lambda}_K}$$

$$= \frac{\int_{0}^{1} \cdots \int_{0}^{1} \prod_{j=1}^{K} \hat{\lambda}_j^{n_j + n_j' - 1} (1 - \sum_{j=1}^{K-1} \hat{\lambda}_j)^{n_k - 1} d\hat{\lambda}_1 \cdots d\hat{\lambda}_K}{\int_{0}^{1} \cdots \int_{0}^{1} \prod_{j=1}^{K} \hat{\lambda}_j^{n_j - 1} (1 - \sum_{j=1}^{K-1} \hat{\lambda}_j)^{n_k - 1} d\hat{\lambda}_1 \cdots d\hat{\lambda}_K}$$

$$= \frac{\prod_{j=1}^{K} \Gamma(n_j + n_j') / \Gamma(n_j)}{\Gamma(N) / \Gamma(n)}$$, \hspace{1cm} [2.5]

where $\Gamma(\cdot)$ denotes the gamma function.
Ghosh and Meeden (1997) show that this posterior predictive distribution is in fact a Pólya urn distribution for the probability of seeing $n_j$ balls with color $d_j$, for $j=1,2,...,K$ in some specified order. They also discuss the close relationship between the Polya posterior and the Dirichlet process priors (Ferguson, 1973) and the Bayesian bootstrap (BB) (Rubin, 1981). The Polya posterior is operationally and inferentially equivalent to the finite population Bayesian Bootstrap (FPBB) of Lo (1988), which is just the BB adapted to finite population sampling problems.

**The weighted Polya Posterior/weighted FPBB**

Formula [2.5] can be generalized to the case when the sampled units bear different weights, i.e. when the realized sample is selected with unequal probabilities (Cohen, 1997). To be consistent with the specific PPS sample we are considering in this chapter, we adapt the notation as follows: denote the PPS sample as

$$(Y_i, X_i, w_i, R_i) = \{(Y_i, X_i, w_i, R_i), i = 1,...,n_\text{.}\},$$

where $w_i = \sum_i^N Z_i / n Z_i$ for size variable $Z_i$ and sample and population sizes $n$ and $N$, and $Y_i = Y_{i,\text{obs}}$ if $R_i = 1$ and $Y_i = Y_{i,\text{mis}}$ if $R_i = 0$. Let $\{d_1, d_2, ..., d_K\}$ denote the set of $K$ distinct vectors of $(Y_i, X_i, w_i, R_i)$ in the sample and $\zeta = \{\zeta_1, \zeta_2, ..., \zeta_K\}$ the vector of probabilities that $\Pr((Y_i, X_i, w_i, R_i) = \tilde{d}_j | \zeta) = \zeta_j$, for $i = 1,2,...,n$, $j = 1,...,K$, and $\sum_{j=1}^K \zeta_j = 1$. Let $n_j$ and $n_j'$ be the number of units taking vector $\tilde{d}_j$ in the sample and in the nonsampled part of the population, respectively, for $j = 1,2,...,K$, and $\sum_{j=1}^K n_j = n$, $\sum_{j=1}^K n_j' = N - n$. For convenience, assume that all sampled units have distinct vector of values, i.e. $K = n$. Let $w_i$ denote the sampling
weight for the $i^{th}$ unit in the sample, which is normalized to sum up to $N$, i.e. $\sum_{i=1}^{n} w_i = N$.

Then $w_i$ can be thought as the posterior expectation of the count of units in the population that have the same value as the $i^{th}$ unit in the sample. Again assuming a noninformative Haldane prior of $\zeta : \zeta \sim \text{Dir}(0,...0)$ together with multinomially distributed weighted counts in the data $p(w_1,...,w_K | \zeta) \propto \prod_{j=1}^{K} \zeta_j^{w_j}$ yields a Dirichlet posterior distribution of $\zeta : \zeta | w_1,...,w_K \sim \text{Dir}(w_1,...,w_K)$,

$$p(\zeta | w_1,...,w_K) \propto \prod_{j=1}^{K} \zeta_j^{w_j-1} \tag{2.6}$$

The posterior predictive distribution of counts in the nonsampled data then follows a compound multinomial distribution with an adjusted parameter $\zeta^* = \{\zeta_1^*,...,\zeta_K^*\}$, i.e.

$$n_1^*,...,n_K^* | w_1,...,w_K \sim \text{Mult}(N-n;\zeta^*)$$

$$p(n_1^*,...,n_K^* | w_1,...,w_K) = \frac{\int_{0}^{1} \int_{0}^{1} \prod_{j=1}^{K} \zeta_j^{w_j+n_j-1} (1-\sum_{j=1}^{K-1} \zeta_j^{w_j+n_j-1} d\zeta_1^*...d\zeta_{j-1}^* ) d\zeta_j^*...d\zeta_K^*}{\int_{0}^{1} \int_{0}^{1} \prod_{j=1}^{K} \zeta_j^{w_j+n_j-1} (1-\sum_{j=1}^{K-1} \zeta_j^{w_j+n_j-1} d\zeta_1^*...d\zeta_{j-1}^* ) d\zeta_j^*...d\zeta_K^*} \tag{2.7} = \frac{\prod_{j=1}^{K} \Gamma(w_j+n_j) / \Gamma(w_j)}{\Gamma(2N-n) / \Gamma(N)}$$

In Little and Zheng (2007), $\zeta_j^* = C \cdot \zeta_j \cdot (w_j-1)$, for $j = 1,...,K$, where $C$ is a constant that satisfies $\sum_j \zeta_j^* = 1$; In this thesis, we follow the weighted Polya urn sampling suggested by Cohen (1997) and use a formula shown in equation [2.10].

**The Adapted-weighted FPBB method**

The adapted-weighted FPBB consists of two stages. The first stage resamples the original sample using the standard Bayesian bootstrap assuming IID, and the second
stage reverses/undoes the sampling weights using the weighted FPBB. This two-stage algorithm is similar in spirit to the standard parametric Bayesian method, where the first stage is equivalent to drawing values of the parameter ($\zeta$) from its posterior distribution given the counts in sampled data ($n_1, ..., n_K$) and the second stage draws the predicted counts in the nonsampled data ($n_1', ..., n_K'$) given the drawn parameter. Note that in Little and Zheng (2007), the first stage is replaced by drawing the parameter directly from a Dirichlet posterior distribution given by [2.6]. The method is described as follows:

- **Resampling using the standard Bayesian Bootstrap (BB)**

  The standard Bayesian Bootstrap of Rubin (1981) assuming IID is used to generate $L$ replicate BB samples each of size $n$, i.e. $\{(Y_s^{(l)}, X_s^{(l)}, w_s^{(l)}, R_s^{(l)}), l = 1, ..., L\}$. This essentially generates the posterior of the empirical joint CDF (denoted by $f$) of all the variables in the population given their realized values in the sample data set. Or equivalently, the posterior distribution of the parameter vector $\zeta$ is drawn given the sample, i.e.

  $$f^{(l)}(Y, X, w, R) \mid (Y_s^{(l)}, X_s^{(l)}, w_s^{(l)}, R_s^{(l)}) \Leftrightarrow \left(\zeta^{(l)} \mid Y_s^{(l)}, X_s^{(l)}, w_s^{(l)}, R_s^{(l)}\right) \sim Dir(n_1, ..., n_K)$$

  [2.8]

  for $l = 1, ..., L$, where $\zeta^{(l)} = (\zeta_1^{(l)}, ..., \zeta_K^{(l)})$.

  This stage captures the sampling variability. The uncertainty in the posterior draws of the parameter $\zeta^{(l)}$ is reflected in the varying counts of distinct units in the original sample being selected in different replicate BB samples. Let $\alpha_s(i)$ denote the number of times unit $i$ is selected in the $l^{th}$ replicate BB sample, for $l = 1, ..., L$. We incorporate this source of uncertainty in computing “the $l^{th}$ bootstrap weight for unit $i$”,

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i.e. \( w^{(l)}_i = w_i \cdot \alpha_l(i) \), where \( w_i \) denotes the original sampling weight for unit \( i \). Note that unequal inclusion probabilities still exist in these created BB samples. The bootstrap weights are carried forward as input weights to the next stage.

- **Undo Sampling Weight using the weighted Polya posterior/weighted FPBB**

The weighted Polya posterior in equation [2.7] is used to create \( B \) synthetic populations for each of the \( L \) BB sample obtained from the previous stage, i.e.

\[
\{(Y^{(l)}, X^{(l)}, R^{(l)}), (Y^{(l)}_{ns}, X^{(l)}_{ns}, R^{(l)}_{ns})\}, \text{ for } b = 1, \ldots, B, l = 1, \ldots, L.
\]

Specifically, it draws predicted counts of the distinct nonsampled units given that of the BB sample, which simultaneously adjusts for unequal inclusion probabilities, i.e.

\[
\left( n^{(lb)}_1, \ldots, n^{(lb)}_K \mid w^{(l)}_1, \ldots, w^{(l)}_K \right) \sim \text{Mult}(N - n; \zeta^{(l)*}),
\]

for \( b = 1, \ldots, B, l = 1, \ldots, L \).

This stage captures the variability due to “imputing” the nonsampled cases within the \( l^{th} \) BB sample (under the same posterior draw of the parameter \( \zeta^{(l)*} \)). The distribution in Equation [2.7] does not lend itself to direct calculation and thus needs to be approximated using Monte Carlo simulation. Specifically, we apply a procedure suggested by Cohen (1997) to simulate the posterior predictive distribution of the counts in nonsampled part of the population through generating \( B \) synthetic populations for each of the \( L \) Bayesian bootstrap samples:

i) Take a Pólya sample of size \( N - n \), denoted by \( (Y^{(lb)}_{ns}, X^{(lb)}_{ns}, R^{(lb)}_{ns}) \) from the urn \( (Y^{(l)}, X^{(l)}, R^{(l)}) \). In this process, each \( (Y^{(l)}_i, X^{(l)}_i, w^{(l)}_i, R^{(l)}_i) \), for \( i = 1, \ldots, n \) in the urn is selected with probability
where $w_i^{(l)}$ is the bootstrap weight for the $i^{th}$ unit in the $l^{th}$ replicate BB sample, and $l_{i,k-1}$ is the number of selections of unit $i$ up to $(k-1)^{th}$ selection, setting $l_{i,0} = 0$.

ii) Form the weighted FPBB synthetic population

$$P_{(b)}^{(l)} = \{ (Y_i^{(l)}, X_i^{(l)}, R_i^{(l)}), (Y_{ns}^{(lb)}, X_{ns}^{(lb)}, R_{ns}^{(lb)}) \}$$

so that it has exact size $N$.

The heuristic interpretation of [2.10] for adjusting the selection probability based on the initial bootstrap weight is as follows. Let $k = 1, 2, ..., N-n+1$, and $i = 1, 2, ..., n$, before making any FPBB selection of nonsampled units in the population from the complex BB sample (which could also be seen as $n$ balls with distinct colors in the original urn), i.e. when $k = 1$ and $l_{i,k-1} = l_{i,0} = 0$, the probability of selecting unit $i$ with weight $w_i^{(l)}$ is

$$\zeta_i^{(l)} = \frac{w_i^{(l)} - 1 + l_{i,k-1} \times \left( \frac{N-n}{n} \right)}{N-n + (k-1) \times \left( \frac{N-n}{n} \right)}, \; k = 1, 2, ..., N-n+1. \quad [2.10]$$

where $w_i^{(l)}$ is the bootstrap weight for the $i^{th}$ unit in the $l^{th}$ replicate BB sample, and $l_{i,k-1}$ is the number of selections of unit $i$ up to $(k-1)^{th}$ selection, setting $l_{i,0} = 0$. As we proceed with the FPBB selection, we adjust this selection probability according to the number of times each unit among the $n$ sampled units was selected during the FPBB procedure, with each unit now representing $(N-n)/n$ among the $(N-n)$ units to be selected during one FPBB whenever it is selected once. After each selection, the denominator of the probability function needs to be inflated to reflect the total units being represented during the whole FPBB selection so far, while the numerator also needs to be inflated to reflect the total units represented by unit $i$ in the process. Therefore we obtain the probability form as in
formula [2.10].

The first step (i.e. the two-stage adapted-weighted FPBB algorithm) results in the following “unweighted” synthetic populations, where $L$ and $B$ are the numbers of datasets generated from first- and second-stage, respectively:

$$
P_{(b)}^{(l)} = (Y^{(lb)}_b, X^{(lb)}_b, R^{(lb)}_b) = (P_{(b)obs}^{(l)}, Y^{(lb)}_{mis}) , b = 1, ..., B, l = 1, 2, ..., L ,$$

where

$$
P_{(b)obs}^{(l)} = ((Y^{(l)}_{s,obs}, X^{(l)}_s, R^{(l)}_s), (Y^{(lb)}_{ns,obs}, X^{(lb)}_{ns}, R^{(lb)}_{ns}))$$

and $Y^{(lb)}_{mis} = (Y^{(l)}_{s,mis}, Y^{(lb)}_{ns,mis})$ consist of the observed and unobserved data in the $lb$th FPBB synthetic population dataset respectively.

### 2.2.3 Step 2: Multiply Impute Missing Data through Parametric Models

Now that we have undone the sampling design, we are ready to perform conventional MI under an IID assumption. Following the standard MI procedure or approximations such as SRMI (Raghunathan, Lepkowski, Van Hoewyk, & Solenberger, 2001), we obtain draws from the posterior predictive distribution $p(Y^{(lb)}_{mis} | P_{(b)obs}^{(l)})$.

Without the need to include weights in the imputation model due to an IID FPBB population generated from the previous step, our task can now be concentrated on correctly modeling the covariates as well as interactions among them whenever necessary.

Note that the elimination of the weights from the FPBB population does not obviate the need to account for the weights in the imputation process, if the probability of selection ($I$) and non-response ($R$) are associated with each other (i.e.,

$$p(R | Y_{obs}, X, w) \neq p(R | Y_{obs}, X))$$. This step results in $M$ imputed synthetic datasets for each of the $L \times B$ FPBB synthetic populations generated from the first step,

$$P_{(b)M}^{(l)} = (P_{(b)1}^{(l)} , P_{(b)2}^{(l)} , ..., P_{(b)M}^{(l)}),$$

for $b = 1, 2, ..., B, l = 1, 2, ..., L$.

### 2.3 Point and Variance Estimates for the Two-Step MI Procedure
In the absence of missing data, Dong, Elliott and Raghunathan (2014) showed that, conditional on $P^{\text{syn}} = \{P_{(1)}^{(1)}, ..., P_{(B)}^{(1)}, ..., P_{(1)}^{(L)}, ..., P_{(B)}^{(L)}\}$, i.e. the $L$ synthetic populations obtained after $B$ FPBB samples, the posterior predictive distribution of a scalar population statistic $Q(Y) = Q$ is given by

$$Q \mid P^{\text{syn}} \sim t_{L-1}(\bar{Q}_L, (1 + L^{-1})V_L),$$

[2.11]

where $t_{L-1}()$ denotes t distribution with $L-1$ degrees of freedom, $\bar{Q}_L = \frac{1}{L} \sum_{i} \tilde{Q}^{(i)}$ and

$$V_L = \frac{1}{L-1} \sum_{i} (\tilde{Q}^{(i)} - \bar{Q}_L)^2$$ for $\tilde{Q}^{(i)} = \lim_{b \to \infty} \frac{1}{B} \sum_{b} q^{(ib)}$. Here $q^{(ib)}$ is the estimate of $Q$ obtained from the $b^{th}$ FPBB synthetic population within the $l^{th}$ Bayesian Bootstrap sample; in practice we estimate $\tilde{Q}^{(i)}$ by $\hat{Q}^{(i)} = \frac{1}{B} \sum_{b} q^{(ib)}$. The result follows immediately from Section 4.1 of Raghunathan, Reiter, and Rubin (2003), and is based on the standard Rubin (1987) multiple imputation combining rules, treating $Y_{ns}$ as missing data and $Y_{s}$ as observed data.

The average “within” imputation variance is zero, since the entire population is being synthesized; hence the posterior variance of $Q$ is entirely a function of the between-imputation variance, and the degrees of freedom is simply given by the number of BB samples. The result assumes that $\mathbb{E}(q^{(ib)}) = Q$ – a result guaranteed by the adapted-weighted FPBB estimator – as well as a sufficiently large sample size for Bayesian asymptotics to apply.

Here we have the additional need to impute the missing data within each of the synthetic population datasets, yielding $P^{\text{imp}} = \{P_{(11)}^{(1)}, ..., P_{(1M)}^{(1)}, ..., P_{(B1)}^{(1)}, ..., P_{(BM)}^{(1)}, ..., P_{(B1)}^{(L)}, ..., P_{(BM)}^{(L)}\}$.

However, the similar result holds as in [2.11]:

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\[ Q \mid P_{\text{imp}} \sim t_{L-1}(\widehat{Q}_L, (1 + L^{-1})V_L) \]  

where as in the fully-observed sample data case \( \bar{Q}_L = \frac{1}{L} \sum_l \widehat{Q}^{(i)} \) and

\[ V_L = \frac{1}{L-1} \sum_l (\widehat{Q}^{(i)} - \bar{Q}_L)^2, \] but now \( \widehat{Q}^{(i)} = \lim_{B \to \infty, M \to \infty} \frac{1}{BM} \sum_b \sum_m q^{(bwm)} \), where \( q^{(bwm)} \) is an estimate of \( Q \) obtained from the \( m^{th} \) imputation of the \( b^{th} \) synthetic population within the \( l^{th} \) Bayesian Bootstrap sample; in practice we estimate \( \bar{Q}_L \) by \( \hat{Q}^{(i)} = \frac{1}{BM} \sum_b \sum_m q^{(bwm)} \). The result again is based on the standard multiple imputation combining rules, where now \((Y_{\text{mis}}, X_{\text{mis}}, R)\) and \(Y_{\text{mis}}\) are missing data and \((Y_{\text{obs}}, X, R)\) is observed; the generation of the synthetic population again sets the within imputation variance to 0. We now require \( E(q^{(bwm)}) = Q \), which implies that our imputation model for \( Y_{\text{mis}} \) is correctly specified (including any associations with probabilities of selection), as well as the standard sufficiently large sample size for the \( t \) approximation to be reasonable.

Note that our point estimator is the same as that derived in (Reiter, 2004), but our variance estimator differs. This is a result of the fact that Reiter must condition on a released sample from the synthetic data, whereas here we are conditioning on the entire synthetic population, substantially reducing the complexity of the analytical approximation to the posterior distribution of \( Q \).

### 2.4 Simulation Study

A simulation study is designed to investigate the inferential properties of the proposed method. The two-step MI procedure is compared with the existing fully
parametric methods under a total of 4 simulation designs defined by crossing the following two factors:

(1) Associations of the probabilities of selection with the mechanism generating the data. We call the design ‘outcome relevant’ if the probabilities of selection are correlated with the outcome variable $Y$, otherwise we term it an ‘outcome irrelevant’ design.

(2) Associations of the probabilities of selection with the mechanism generating the missing values. We use ‘MAR_X’ (weight independent missingness) and ‘MAR_X,W’ (weight dependent missingness) respectively to denote respective situations where the missing data mechanism is dependent on the fully-observed covariate $X$ only, and where it depends on probabilities of selection as well as the covariate.

For each of the four simulation designs, we analyze the data using three imputation models with differing degrees of model misspecification corresponding to different amounts of design information incorporated into the imputation model. *Model 1* ignores weights altogether in the imputation process, which is a procedure typically adopted by survey practitioners. Under Model 1, we assume that the data are resulted from simple random sampling for the purposes of imputation. *Model 2* includes log(weight) as a scalar summary of design information in the imputation model. This is a simple way to compensate for the naïve SRS assumption in Model 1 by adding the weight variable as a separate predictor in the imputation model. Model 2 is relatively easy to implement in practice, and has been adopted by the MI project on NHIS missing income data (Schenker et al., 2006). *Model 3* includes both log(weights) and its interactions with the covariate in the imputation model. All three imputation models will
be tested with both the fully parametric MI method and the proposed two-step synthetic MI procedure. Table 2.1 illustrates the setup of the simulation design where the differing imputation models are nested within the four cells defined by the two association factors.

Table 2.1 Strength of association of the sampling weight with both the missingness and the outcome

<table>
<thead>
<tr>
<th>Association with missingness (M)</th>
<th>Association with outcome variable (Y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>Irrelevant design, MAR_X</td>
</tr>
<tr>
<td></td>
<td>Relevant design, MAR_X</td>
</tr>
<tr>
<td>High</td>
<td>Irrelevant design, MAR_X,W</td>
</tr>
<tr>
<td></td>
<td>Relevant design, MAR_X,W</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Imputation Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>3</td>
</tr>
</tbody>
</table>

2.4.1 Description of the Study Design

The simulation design is described below:

Step 1. Population data generation:

The population involves three variables: the outcome variable $Y$, a covariate $X$, and a size variable $Z$ based on which probability-proportionate-to-size without replacement (PPSWOR) sampling is conducted. We consider two versions of the outcome variable $Y$, one in which the outcome is associated with the covariate, the probability of selection, and their interaction ($Y_1$), and the other in which the outcome is associated only with the covariate ($Y_2$). The joint distribution of $Z$, $X$, and $Y$ is given by:
\[
\log Z \sim N(2,1)
\]
\[
X \mid Z \sim N(0.1 \log Z, \sigma^2_x)
\]
\[
Y_i \mid X, Z \sim N(0.2 \times X + 0.6 \log Z + 0.5 \times X \times \log Z, \sigma^2_{y_i})
\]
\[
Y_2 \mid X, Z \sim N(0.2 \times X, \sigma^2_{y_2})
\]

Thus \((Y_1, X, Z)\) constitutes the “relevant design” population and \((Y_2, X, Z)\) constitutes the “irrelevant design” population. Figure 2.2 shows the scatter plots of \(Y_1\) and \(Y_2\) versus the size measure \(Z\). Both populations have size \(N=4,000\). For each population, we drew 100 independent samples of size \(n=200\) without replacement, with inclusion probability for the \(i\)th unit \(\pi_i = nZ_i / \sum_{j=1}^{N} Z_j\). We call the 100 PPSWOR samples “before deletion (BD) samples”.

![Scatter plots](image)

**Figure 2.2** Scatter plots of survey variable \(Y\) versus size variable \(Z\), under relevant and irrelevant designs, respectively.

**Step 2.** Impose missingness on the complete data under MAR mechanism:

Probit models are used as deletion functions to create missing data in the outcome
variable $Y$ for each of the 100 simulations. Both $X$ and $Z$ are assumed to be completely observed. For both populations, we define two types of missing data mechanisms by generating two different latent variables $T_1$ and $T_2$ for the deletion function, which correspond to the MAR _X condition and MAR _X,W condition respectively.

Specifically, $T_1 = -0.635 + 0.4X + e$ and $T_2 = -0.55 + 0.4X - 0.5 \log Z + 0.4X \ast \log Z + e$, where $e \sim N(0,1)$. The outcome is then missing if $T_j > 0$ (i.e., $P(M = 1|T_j) = \Phi(T_j)$, $j = 1,2.$, where $\Phi(.)$ corresponds to the standard normal CDF. This yields a moderate missingness fraction of 30% in all four designs corresponding to the four cells defined in Table 2.1.

**Step 3. Impute missing data under different imputation models:**

MI was performed separately for each of the 100 replication samples under each of the 12 simulation scenarios. All simulations were programmed using the R software (Version 2.14.2). In particular, the mice package (van Buuren & Groothuis-Oudshoorn, 2011) was used to implement the imputation, and the survey package was used for the design-based analyses under the fully parametric method (Lumley, 2004). For the proposed method, the wtpolyap function in package polyapost (Meeden & Lazar, 2012) may be used to create the weighted FPBB synthetic populations.

**Step 4. Evaluation of performance:**

We consider estimating the population mean of $Y$ (i.e. $\bar{Y}$) and the population regression coefficients of $Y$ on $X$: $Y = X^T \beta$, where $\beta = (\beta_0, \beta_1)^T$ and $\beta_0$ and $\beta_1$ are the intercept and the slope, respectively. We focus on five quantities to evaluate the performance of the two methods under comparison: bias, empirical root mean square error (RMSE), empirical interval coverage, empirical variance, and empirical estimated variance.


· **Bias.** Let \( Q = (\hat{\gamma}, \beta_0, \beta_1) \) denote the set of the three population parameters we are interested in, and \( q = (\hat{\gamma}, \hat{\beta}_0, \hat{\beta}_1) \) denote their corresponding point estimates (Pt.est). The raw bias of each of the parameter estimates \( q \) is calculated as the difference between its averaged value over the 100 simulations and the true parameter value \( Q \).

\[
\text{Bias} = E(q_r) - Q = \frac{1}{100} \sum_{r=1}^{100} q_r - Q
\]

where \( q_r \) is the point estimate associated with the \( r^{\text{th}} \) simulation.

· **Empirical Root Mean Square Error (RMSE).** As a measure of accuracy which accounts for bias and efficiency simultaneously, RMSE is a key statistic we will consider in comparing the overall performance of the methods.

\[
\text{RMSE} = \sqrt{\text{MSE}} = \sqrt{E(q_r - Q)^2} = \sqrt{\frac{1}{100} \sum_{r=1}^{100} (q_r - Q)^2}
\]

· **Empirical interval coverage (95% CI cov.).** We report the actual coverage rates of a nominal 95% interval based on the 100 simulation repetitions under each simulation scenario. Normal approximation was used for inference under the proposed method.

· **Empirical Estimated Variance & Empirical Variance.** We calculated these two quantities and compared them with each other. They should be approximately equal if the variance estimator is unbiased.

\[
\text{Est.Var} = E[v(q_r)] = \frac{1}{100} \sum_{r=1}^{100} V_{\text{ML}}(q_r), \quad \text{SE} = \frac{1}{100} \sum_{r=1}^{100} \sqrt{V_{\text{ML}}(q_r)}
\]

\[
\text{Emp.Var} = \text{Var}(q_r) = \frac{1}{100-1} \sum_{r=1}^{100} (q_r - E(q_r))^2, \quad \text{Emp.SE} = \sqrt{\text{Emp.Var}}
\]

2.4.2 **Simulation Results**

In deciding how many synthetic populations \( B \) are needed, we conducted a preliminary study based on the before deletion (BD) data. Equivalence of the BD
variance estimates to the actual sample variance estimates is desired. Simulation results are shown in Table 2.2. Note that the BD variance estimation under the synthetic method was calculated as the between synthesis variance, using the variance formula \((1 + L^{-1})V_L\) in [2.11]. We observe that as we increase \(B\), the variance estimate decreases, and stabilizes close to the actual sample variance for \(B \geq 20\). Therefore, we use \(B=20\) in the after deletion (AD) simulation.

Table 2.3 and Table 2.4 present the results from our simulation study. Each table is divided into two parts, containing the results from MAR _X condition and MAR _X,W condition respectively. Within each condition, we compare our new method with the fully parametric method. The three columns indicated by ‘X’, ‘X,W’ and ‘XW’ each correspond to the estimates under the three imputation models described in section 2.4.1.

We first examine the results in Table 2.3. When the design is relevant to the outcome variable \(Y\) yet uncorrelated with missingness (MAR _X), obvious advantages can be observed of the synthetic methods over the fully parametric method. For the fully parametric method to work properly under this design, the imputation model has to be correctly specified, otherwise all inferences based on this method are invalid---not only there is substantial bias associated with all three parameter estimates, but a corresponding disruption in coverage rates as well. This is particularly poor when the design is completely ignored in the model. For example, under the X only imputation model, the relative bias of the estimated mean and the slope is as high as \(\frac{0.260}{1.343} = 19.4\%\) and \(\frac{0.210}{1.435} = 14.6\%\), respectively, with 95% CI coverage rates as low as 79% and 67%, respectively. Including \(W\) as a main effect in the imputation model improves the
performance for estimating the mean but not the slope--the relative bias of the slope is

\[ \frac{0.137}{1.435} = 9.5\% \] with only 79\% coverage. These confirm the argument by Kim et al. (2006) and Seaman et al. (2011) that including survey weights alone in the imputation model does not guarantee valid inference if important interactions between the weight and other covariates are ignored. In contrast, our proposed method results in nearly unbiased estimates and actual coverage that is closer to the nominal level under all three models, regardless of the misspecification. Substantial gains in terms of RMSE over the model-based method are also consistently observed in all scenarios considered (e.g. 0.303 vs. 0.220 for the slope under X only model, and 0.201 vs. 0.185 under the interaction model). This indicates that the ‘unweighting’ procedure has actually played dual roles in the process: its effect is not limited to untying the unequal probability selection and saving the effort of design-weighted analyses afterwards, but it also captures the interactions between the design and the survey variable of interest. Thus ignoring the design in the imputation model does no harm at all.

With a relevant design that is also a correlate of missingness (MAR_X,W), the imputation model will still require the use of the design variable (here the weight) to maintain an ignorable missing data mechanism. The fully parametric method behaves similarly to the case where the design is associated only with Y: failure to include the weight in the imputation model substantially biases all of the estimators considered, while including the weight as a covariate corrects for bias in the mean and intercept estimator but not in the slope. The synthetic model partially corrects for these biases by providing a correct estimate of the population distribution in the presence of missing data; however, unless the imputation model is correctly specified, some biases remain.
However, the synthetic model still substantially reduces RMSE relative to the fully parametric approach for the mean and intercept estimator when the weight is ignored in the imputation model. It also has reduced RMSE when estimating the slope when the weight is included as a covariate but the interaction between the slope and the probability of selection is ignored. Otherwise, the synthetic and fully parametric approaches have similar RMSE properties. The synthetic model also has slightly conservative coverage properties, in contrast to the anti-conservative coverage of the fully parametric estimator when the model is misspecified for the estimator of interest.

With an outcome irrelevant design (Table 2.4), there are very slight effects on the estimates when compared across methods and models. Including the irrelevant design variable in the imputation model results in negligible biases and introduces some modest inefficiencies, consistent with the findings in Reiter et al. (2006).

Figures 2.3 and 2.4 examine the properties of MI variance estimators. Figure 2.3 shows scatter plots of 100 estimated standard errors (SEs) of the mean from alternative imputation methods versus the empirical SEs from actual samples before deletion under three different imputation models, with outcome relevant design. The MI variance estimates under the proposed synthetic method are consistently lower than the fully parametric method. The triangles (representing synthetic MI) are closer than the circles (representing parametric MI) to the 45 degree straight line, and the contrast is most obvious under model 1. Similar results are observed with outcome irrelevant design (not being plotted here). Figure 2.4 plots the standard error (SE) versus empirical standard error (Emp.SE) from alternative MI methods, under all 12 combinations of simulation design and imputation model, for both the mean and the slope. While the variance
estimator for the parametric MI method (represented by red dots) either over- or under-
estimates when the imputation model is misspecified (i.e. under model 1 and model 2),
the blue triangles are always close to the 45 degree line. This indicates the approximate
unbiasedness of the variance estimator for the synthetic MI method as well as the
robustness of it to model misspecification.
Table 2.2 Before deletion study of the effects of the number of generated FPBB populations ($B$) on variance estimate

<table>
<thead>
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<th>Parameters Of Interest</th>
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Table 2.3 Performance of the proposed method in contrast to the fully parametric method under the relevant design condition

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Figure 2.3 Scatter plots of 100 estimated standard errors (SEs) of the mean from alternative imputation methods (x axis) versus the empirical SEs from actual samples before deletion (y axis), under three imputation models, with outcome relevant design.
2.5 Application to the Behavioral Risk Factor Surveillance System (BRFSS)

2.5.1 BRFSS Data

We next examine the effect of incorporating the survey weight in MI using data from one design stratum ($n=388$) of the 2009 Michigan BRFSS. This design stratum contains sampled households that belong to the medium-density (unlisted) telephone
numbers group. The BRFSS is a telephone survey conducted with a random sample of adults living in telephone-equipped households in the US. An independent sample of telephone numbers is used as the sampling frame; thus case weights are constructed to account for the fact that the probability of selection is proportional to the number of telephone lines and inversely proportional to the number of adults in a household. In addition, poststratification weights are used to adjust age-sex-race/ethnic distributions to Census totals. A mix of categorical and continuous variables is selected for analysis. These include health insurance coverage (yes/no), body-mass index (BMI) in kg/m², high blood pressure (yes/no), and five socio-demographic variables (age (in years), race (White vs. Nonwhite), annual household income (low: ≤ $25,000, medium: $25,000 - $75,000, high: >$75,000), gender (male/female), and employment status (yes/no/other) as well as a single design variable, the final weight. All survey variables except gender have certain degrees of missing data: income has the highest missing rate (16.5%), while others are missing 0%~6%.

2.5.2 BRFSS Imputation Method

We compare results from the fully parametric MI method with the proposed two-step semi-parametric MI method, using two imputation modeling strategies with each method: 1) assuming SRS, and 2) including the log of weights as a predictor in the model. We also include the weighted complete case analysis (CC). For the new method, we generated \( L = 100 \) Bayesian bootstrap (BB) samples and created \( B = 30 \) FPBB populations within each BB sample, with \( M = 5 \) multiple imputations performed for each FPBB population. Since we do not know the population size in advance, and the individual final weights sum up to nearly 200,000 cases which is unrealistic to generate, we assume that \( N = 4500 \) is large enough to be treated as a synthetic population. Since the degrees of
freedom is $L-I=99$, a normal distribution was used for inference.

### 2.5.3 BRFSS Analyses

We consider four different analyses: 1) the marginal distribution of income and health insurance accessibility (Table 2.5); 2) a linear regression model of BMI on key demographic variables as well as income (Table 2.5); 3) a loglinear model of a four-way contingency table defined by four categorical variables with no second-or-higher-order interactions (Table 2.6); and 4) a general location model for the joint distribution of the survey variables (Table 2.7).

Multivariate imputation by chained equations (mice) in R was used to impute the missing data under both MI methods. This technique is different from a joint modeling approach (Schafer, 1997a), which specifies a joint multivariate distribution for the missing data, this technique specifies the multivariate imputation model on a variable-by-variable basis by a set of conditional densities (Raghunathan et al., 2001; van Buuren & Groothuis-Oudshoorn, 2011). Therefore as a way to check the validity of the proposed method, we examine the joint distribution of all variables with missing data after imputation using a general location model (Olkin & Tate, 1961). However, due to sparse cells resulting from the complete cross-tabulation of all categorical variables in the original dataset, we are limited to looking at the joint distribution of BMI and age in a two-way contingency table defined by income (L,M,H) and gender (M,F). The parameters of interest are \( \theta = (P, \Gamma_c, \Omega_c), c = 1, \ldots, 6 \) where \( P \) is a vector of length 6, i.e. the contingency table cell probability, and the bivariate normal distribution of BMI \( (X_1) \) and age \( (X_2) \) within each cell is: \( X \sim N_c(\Gamma_c, \Omega_c) \), where \( \Gamma_c \) and \( \Omega_c \) denote the vector of means of BMI and age and the covariance matrix for the \( c^{th} \) cell, respectively.
We further collapse the medium and high income as one category for the loglinear analysis to avoid sparse cells.

### 2.5.4 BRFSS Results

Since the poststratification adjustment factor constitutes an important component of the final weight in BRFSS dataset, we presume that including the variables used to construct poststratification cells (age, race and gender in this case) in the imputation model should help in predicting the missing $Y$ variable. A linear regression of final weights on age, sex, and race shows that these covariates explain 40% of the variance of the weights, suggesting that some unknown design variables exist that contributes to the construction of survey weights. Thus we conclude that imputation approaches that condition only on the available design variables will be insufficient to account for the sampling weights.

Table 2.5 shows that under the fully parametric MI method, including survey weights in the imputation model has a large impact on the regression coefficients of BMI on income and gender (high income: -0.47 when weights are excluded vs. -0.33 with log(weights) included; female: 2.72 vs. 2.56). In fact, these differences are particularly significant for domain estimation for whites (medium income: -2.8 vs. -2.1; female: -0.13 vs. -0.68). Under the new method, however, all estimates are similar to those from the fully parametric method with weights accounted for. Moreover, there is essentially no difference whether or not we incorporate weights into the imputation model after the sample data are synthesized. This means, that the new method can adjust for the weight effects at the synthesizing step without the need to model survey weights at the imputation step. Similar conclusions can be made from the Table 2.6 results for the log-linear model: modeling the weight in the imputation model has major impacts on the
coefficients associated with income. For example, Low income x Has insurance: -0.37 under ‘excludes weights parametric MI’ vs. -0.33 under ‘includes log(weights) parametric MI’ vs. -0.31 under both modeling strategies of ‘synthetic MI’.

As both BMI and age only have a very small fraction of missing data, we see little difference between the SRS imputation model and the model that includes log(weight) as a predictor under either MI method in the estimation of the general location model (Table 2.7), either for overall estimation or within income by gender categories. Thus we only display the results for model-based MI with weights accounted for (which is considered as the correct imputation model to use), together with the synthetic MI with the SRS model and we contrast both of them with the complete case analysis (CC). In general, two-stage semi-parametric MI provides results that are similar to the fully parametric MI method. Both seem to improve efficiency for overall estimates of BMI and age relative to those obtained from CC.
Table 2.5 Estimation of marginal distributions for income and health insurance, and linear regression coefficients for the regression of BMI on income, age and gender

<table>
<thead>
<tr>
<th>Sample</th>
<th>Estimation</th>
<th>Variable</th>
<th>Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Complete Case</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Full Sample</td>
<td>Marginal</td>
<td>Low Income</td>
<td>0.50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medium Income</td>
<td>0.38</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Income</td>
<td>0.12</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No insurance</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>Regression</td>
<td>Intercept</td>
<td>27.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medium income</td>
<td>0.47</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High income</td>
<td>0.27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Age</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Female</td>
<td>2.29</td>
</tr>
<tr>
<td>Whites Domain</td>
<td>Marginal</td>
<td>Low Income</td>
<td>0.30</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medium Income</td>
<td>0.53</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Income</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No insurance</td>
<td>0.24</td>
</tr>
<tr>
<td></td>
<td>Regression</td>
<td>Intercept</td>
<td>31.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medium income</td>
<td>-1.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High income</td>
<td>-3.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Age</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Female</td>
<td>-1.7</td>
</tr>
</tbody>
</table>
Table 2.6 Estimation of log-linear model for four categorical variables (collapse categories for medium and high income)

<table>
<thead>
<tr>
<th>Methods</th>
<th>Complete Case</th>
<th>Parametric MI Exclude weights</th>
<th>Parametric MI Include log(weights)</th>
<th>Synthetic MI Exclude weights</th>
<th>Synthetic MI Include log(weights)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coef.</td>
<td>SE</td>
<td>Coef.</td>
<td>SE</td>
<td>Coef.</td>
</tr>
<tr>
<td><strong>Main effects</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low income</td>
<td>-0.01</td>
<td>0.12</td>
<td>0.08</td>
<td>0.13</td>
<td>0.04</td>
</tr>
<tr>
<td>Has insurance</td>
<td>0.61</td>
<td>0.12</td>
<td>0.64</td>
<td>0.12</td>
<td>0.62</td>
</tr>
<tr>
<td>White</td>
<td>-0.94</td>
<td>0.11</td>
<td>-1.0</td>
<td>0.10</td>
<td>-1.0</td>
</tr>
<tr>
<td>Male</td>
<td>-0.11</td>
<td>0.12</td>
<td>-0.09</td>
<td>0.10</td>
<td>-0.07</td>
</tr>
<tr>
<td><strong>Two-way Interactions</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low income x Has insurance</td>
<td>-0.36</td>
<td>0.12</td>
<td>-0.37</td>
<td>0.12</td>
<td>-0.33</td>
</tr>
<tr>
<td>Low income x White</td>
<td>-0.28</td>
<td>0.10</td>
<td>-0.20</td>
<td>0.09</td>
<td>-0.20</td>
</tr>
<tr>
<td>Low income x Male</td>
<td>-0.03</td>
<td>0.09</td>
<td>-0.03</td>
<td>0.09</td>
<td>-0.07</td>
</tr>
<tr>
<td>Has insurance x White</td>
<td>-0.13</td>
<td>0.12</td>
<td>-0.02</td>
<td>0.12</td>
<td>-0.02</td>
</tr>
<tr>
<td>Has insurance x Male</td>
<td>-0.01</td>
<td>0.13</td>
<td>-0.12</td>
<td>0.10</td>
<td>-0.14</td>
</tr>
<tr>
<td>White x Male</td>
<td>-0.08</td>
<td>0.09</td>
<td>-0.11</td>
<td>0.08</td>
<td>-0.11</td>
</tr>
</tbody>
</table>
Table 2.7 Estimation of general location model for joint distribution of BMI, age, income and gender after MI under alternative methods

(L_M = low-income male, M_M=medium-income male, H_M=high-income male; L_F = low-income female, M_F=medium-income female, H_F=high-income female)

<table>
<thead>
<tr>
<th></th>
<th>Proportion</th>
<th>SE(p)</th>
<th>bmi</th>
<th>SE(bmi)</th>
<th>age</th>
<th>SE(age)</th>
<th>Var(bmi)</th>
<th>Var(age)</th>
<th>Cov(bmi, age)</th>
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<tr>
<td>Complete Case Analysis (design-based analysis)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L_M</td>
<td>0.230</td>
<td>0.045</td>
<td>27.2</td>
<td>1.86</td>
<td>38.7</td>
<td>3.81</td>
<td>57.4</td>
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<td>-13.5</td>
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<tr>
<td>M_M</td>
<td>0.175</td>
<td>0.037</td>
<td>28.5</td>
<td>1.62</td>
<td>40.8</td>
<td>4.36</td>
<td>54.0</td>
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<td>51.8</td>
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<tr>
<td>H_M</td>
<td>0.068</td>
<td>0.023</td>
<td>29.7</td>
<td>1.78</td>
<td>49.2</td>
<td>2.76</td>
<td>32.0</td>
<td>122</td>
<td>1.3</td>
</tr>
<tr>
<td>L_F</td>
<td>0.269</td>
<td>0.034</td>
<td>30.7</td>
<td>1.26</td>
<td>43.2</td>
<td>2.48</td>
<td>86.9</td>
<td>313</td>
<td>4.5</td>
</tr>
<tr>
<td>M_F</td>
<td>0.201</td>
<td>0.028</td>
<td>30.5</td>
<td>1.08</td>
<td>46.2</td>
<td>1.97</td>
<td>57.9</td>
<td>197</td>
<td>-7.2</td>
</tr>
<tr>
<td>H_F</td>
<td>0.058</td>
<td>0.014</td>
<td>28.9</td>
<td>1.04</td>
<td>48.1</td>
<td>3.62</td>
<td>23.6</td>
<td>241</td>
<td>-13.7</td>
</tr>
<tr>
<td>Overall</td>
<td>-</td>
<td>-</td>
<td>29.3</td>
<td>0.70</td>
<td>43.0</td>
<td>1.56</td>
<td>61.0</td>
<td>278</td>
<td>8.0</td>
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<td>Model-Based MI log(weight) (design-based analysis)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L_M</td>
<td>0.237</td>
<td>0.042</td>
<td>26.8</td>
<td>1.46</td>
<td>38.3</td>
<td>3.49</td>
<td>48.2</td>
<td>301</td>
<td>4.8</td>
</tr>
<tr>
<td>M_M</td>
<td>0.155</td>
<td>0.036</td>
<td>28.3</td>
<td>1.43</td>
<td>41.5</td>
<td>4.48</td>
<td>47.7</td>
<td>361</td>
<td>45.6</td>
</tr>
<tr>
<td>H_M</td>
<td>0.077</td>
<td>0.029</td>
<td>27.7</td>
<td>1.78</td>
<td>41.4</td>
<td>6.07</td>
<td>33.7</td>
<td>299</td>
<td>45.0</td>
</tr>
<tr>
<td>L_F</td>
<td>0.268</td>
<td>0.035</td>
<td>30.5</td>
<td>1.05</td>
<td>44.0</td>
<td>2.50</td>
<td>76.9</td>
<td>356</td>
<td>5.97</td>
</tr>
<tr>
<td>M_F</td>
<td>0.204</td>
<td>0.032</td>
<td>30.1</td>
<td>0.96</td>
<td>44.5</td>
<td>2.42</td>
<td>54.6</td>
<td>254</td>
<td>-2.4</td>
</tr>
<tr>
<td>H_F</td>
<td>0.060</td>
<td>0.017</td>
<td>29.0</td>
<td>1.31</td>
<td>45.2</td>
<td>4.73</td>
<td>24.8</td>
<td>262</td>
<td>-10.4</td>
</tr>
<tr>
<td>Overall</td>
<td>-</td>
<td>-</td>
<td>28.9</td>
<td>0.58</td>
<td>42.1</td>
<td>1.42</td>
<td>55.5</td>
<td>312</td>
<td>14.9</td>
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<tr>
<td>Synthetic MI SRS (simple unweighted analysis)</td>
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</tr>
<tr>
<td>L_M</td>
<td>0.241</td>
<td>0.037</td>
<td>27.1</td>
<td>1.47</td>
<td>38.7</td>
<td>2.92</td>
<td>45.0</td>
<td>275</td>
<td>-0.41</td>
</tr>
<tr>
<td>M_M</td>
<td>0.161</td>
<td>0.033</td>
<td>28.2</td>
<td>1.40</td>
<td>41.4</td>
<td>4.09</td>
<td>43.0</td>
<td>321</td>
<td>40.1</td>
</tr>
<tr>
<td>H_M</td>
<td>0.070</td>
<td>0.022</td>
<td>28.2</td>
<td>1.67</td>
<td>43.8</td>
<td>5.30</td>
<td>26.6</td>
<td>211</td>
<td>26.0</td>
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<td>73.6</td>
<td>351</td>
<td>2.9</td>
</tr>
<tr>
<td>M_F</td>
<td>0.196</td>
<td>0.023</td>
<td>30.5</td>
<td>0.79</td>
<td>45.0</td>
<td>1.57</td>
<td>53.8</td>
<td>233</td>
<td>-4.2</td>
</tr>
<tr>
<td>H_F</td>
<td>0.056</td>
<td>0.013</td>
<td>29.0</td>
<td>1.00</td>
<td>46.8</td>
<td>3.22</td>
<td>25.5</td>
<td>231</td>
<td>-10.4</td>
</tr>
<tr>
<td>Overall</td>
<td>-</td>
<td>-</td>
<td>29.0</td>
<td>0.55</td>
<td>42.1</td>
<td>1.46</td>
<td>55.9</td>
<td>308</td>
<td>13.1</td>
</tr>
</tbody>
</table>
2.6 Discussion

Our primary goal was to propose a new method using the weighted finite population Bayesian Bootstrap to account for sampling weights in MI to deal with item-level missing data, and to evaluate the new method in a PPS sampling design setting. Our findings in the simulation study suggest that the new method does significantly reduce bias relative to the fully parametric methods, and with little loss in efficiency. Meanwhile, the weighted FPBB method maintains population-level multivariate relationships and potentially protects against model misspecification, for example, erroneous inclusion or exclusion of interactions between design variables and other covariates in the imputation model. A further advantage lies in that, unlike the fully parametric methods which include designs in the imputation model but still require complex survey packages to analyze the imputed datasets, the new method fully accounts for the unequal selection probabilities by unweighting them and restoring a population in a separate step. Therefore, only simple, unweighted complete-data analysis techniques are needed for inferences with the newly developed combining rules. This potentially allows a much wider variety of models to be considered with existing software, which, despite recent improvements, often cannot account straightforwardly for complex sample designs.

A limitation of the proposed method is the need to account for the design elements in the imputation procedure if the missingness mechanism requires their inclusion (e.g., if the probability of an item response is a function of the selection probability). As a practical matter, however, “undoing” the sample design to
generate the synthetic population may reduce the impact of misspecified missingness mechanisms by avoiding enhancement from misspecified data generation mechanisms.

The proposed two-stage semi-parametric multiple imputation approach has a number of possible extensions. The method developed here is designed for one-stage sample designs with independent selections and unequal probabilities of selection, as in the BRFSS sample design. Hence, extensions are required to account for multi-stage designs with clustering and stratification as part of the finite population Bayesian bootstrap. Another limitation is the assumption that no unit non-response occurs in the sample. Extensions that incorporate unit non-response provide another promising research opportunity. However, in public use samples that provide only final weights incorporating non-response adjustments, treating the final weight as a sampling weight (as we did in the BRFSS application) may be the only practical alternative.
3.1 Introduction

Survey data collected for social science and public health research is often clustered. Such clustered data often results from multi-stage sampling for cost and convenience reasons. For example, cluster sampling of students from schools is common in education surveys, where students are clustered by schools, or by classrooms within schools. In general population surveys, area probability sampling is considered a cost-effective way to select households because households are naturally clustered by geographic areas (e.g. PSUs/counties, blocks, etc.). In these and many other examples, it is realistic to assume that units in the same cluster tend to be more alike than units in different clusters. This leads to clustering effects, typically in the form of intraclass correlation (ICC) (Kish, 1965). Since a positive ICC implies an effective sample size less than the total sample, and thus increases variance estimates of statistics of interest, accounting for clustering effects is quite important.

Methods for doing this have been well-developed for complete-data approaches using either robust sandwich estimators based on Taylor Series approximations (Binder, 1993) or replication methods such as the balanced
repeated replication, jackknife, or bootstrap (Rust & Rao, 1996). These are now commonly available in statistical packages (e.g. the survey package in R, the svy suite of commands in Stata). A more model-based approach to the analysis of clustered data is multilevel modeling (Rabe-Hesketh & Skrondal, 2006), which allows the decomposition of overall variance estimates into components of variance due to different levels of sample selection. A good overview of software procedures for fitting linear mixed models is West & Galecki (2011); packages that fit nonlinear mixed models include R nlme, SAS nlmixed, and Stata gllamm (Li et al., 2011).

Concurrently with the development of methods and software for clustered sample designs, multiple imputation (MI) (Rubin, 1987) has become a standard method for dealing with item-level missing data in complex sample surveys. The need to incorporate sample design information into the imputation process has been recognized -- for example, Rubin (1996) in his MI review paper stated that any imputation model should minimally include major stratification and clustering indicators as well as design weights to make it proper. More generally, Schafer (1997a) also asserted that whether the estimation techniques ultimately applied to survey data are model-based or design-based, imputation models for missing values should incorporate important features of sample design. However, as this can be difficult to do in practice, imputation is typically performed under an assumption of simple random sampling.

This issue is considered in detail by Reiter, Raghunathan, and Kinney (2006). They attempt to incorporate sample design through simple modification
of existing imputation procedures such as SRMI in IVEware (Raghunathan et al., 2001), by modeling design variables (i.e. cluster membership indicators) as fixed effects in the imputation. Despite the simplicity of implementation, the fixed cluster effects model they propose is usually uncongenial to the analyst’s model in the sense of Meng (1994). It can become very inefficient when the number of clusters is large. Andridge (2011) also considered fitting a fixed cluster effects imputation model to data from cluster randomized trials. However, when she used such a modeling strategy, she found an upward bias in the MI variance estimator through both analytical proof and simulation studies.

An alternative approach consistent with standard model-based approaches is to impute data using a mixed/random effects model, where each cluster has its own intercept (usually assumed to be sampled from a normal distribution with zero mean and unknown variance). Software packages allowing the use of mixed effects models for multiple imputation include: 1) the R pan package (Schafer & Yucel, 2002; Zhao & Schafer, 2013), for imputing continuous variables in clustered or panel designs under a multivariate normal model. Details on the MCMC imputation method used in pan can be found in Schafer (1997b); and 2) the REALCOM-IMPUTE module of the multilevel model fitting software MLwiN (Carpenter, Goldstein & Kenward, 2011). This also adopts a joint modeling approach and MCMC-based solutions but assumes a multivariate latent normal model with random effects for mixed type response variables subject to missingness. While these imputation models achieve congeniality with respect to the cluster effects and result in more efficient estimates than a fixed effect
approach, their use in practice is limited due to several shortcomings: pan, for example, assumes normality on binary variables, and the linear mixed effects imputation model typically leads to implausible values that needs rounding (to the nearest 0 or 1) after imputation. This can cause biased estimates of parameters (Horton et al., 2003). With REALCOM-IMPUTE, uncongeniality remains an issue in the sense that conditional relationships among variables in the imputation model are typically assumed linear while the models of interest usually include non-linear relationships and/or interactions. Besides, the consequences are not clear of using a probit regression for imputation for binary data that are modeled by the analyst using a logistic regression. Moreover, both procedures are confined to relatively simpler data structure with no more than two levels.

On another front, attempts to extend the variable-by-variable imputation methodology to clustered designs, e.g. the sequential hierarchical regression imputation models (SHRIMP) for multivariate clustered data (Yucel & Raghunathan, 2006; Zhao & Yucel, 2009; Yucel, 2011) demonstrate problems with convergence. There is evidence of poor finite-sample repeated sampling properties in logistic mixed effect models with high ICC. Other problems with the fully parametric imputation methods above include: 1) imputations drawn from these models may perform poorly if the random effects follow a non-normal distribution instead; 2) the accommodation of both sample weights and cluster effects simultaneously is particularly problematic, especially in high-dimensional settings with many missing covariates. Such an accommodation requires the estimation of interactions between clusters and weights, as well as between
weights and other quantities of interest (e.g., regression parameters). This can be difficult to achieve using a standard parametric approach.

Thus the focus of this chapter is to develop an adaptation of the two-step semi-parametric MI procedure proposed in Zhou, Elliott, and Raghunathan (2013a) for disentangling clustering effects and sampling weight effects using a weighted finite population Bayesian bootstrap (FPBB) procedure developed in Dong, Elliott, and Raghunathan (2014). The new MI procedure is designed to produce draws from the posterior predictive distribution of the population that incorporate both clustering and weighting elements of the complex sample design. Item-level missingness is incorporated in these “uncomplexed” synthetic populations; these missing data can then be imputed under an IID assumption without explicit modeling of clustering and weight effects. We consider two-stage unbalanced cluster samples obtained from unequal probability of selection, and we assume a missing at random (MAR) missing data mechanism on a single survey outcome variable (Y). The parameters of interest are the population mean of Y and population regression parameters of Y on a covariate X based on the multiply imputed data.

Section 3.2 discusses MI under simple random sampling, together with standard fixed and random effects models to incorporate cluster effects. Section 3.3 develops the newly proposed weighted FPBB procedure. Section 3.4 conducts and discusses a series of simulation studies designed to assess the repeated sampling properties of MI under the various approaches discussed in Section 3.2 and 3.3. These simulation studies use different population models, differing
numbers of clusters, differing degrees of intraclass correlation, and differing MAR mechanisms (dependent only on the fully observed covariate independent of sample weights, or dependent on both the covariate and sample weights).

Section 3.5 applies the different MI procedures to the analysis of passenger vehicle injury data from the National Automotive Safety System – Crash Detection System (NASS-CDS) survey. Section 3.6 concludes with discussion and suggestions for next research steps.

3.2 Fully Parametric Imputation Methods for Clustered Sample Designs

3.2.1 Simple Random Sampling (SRS) Model

Let $Y_i$ be the survey outcome with missing values (assumed scalar for ease of exposition) and $X_i$ be a $p$-variate vector of other survey variables whose values are known for all cases in the sample, $i = 1, \ldots, n$. We assume that $Y$ is a member of the exponential family, and that the known transformation of $E(Y_i \mid X_i) = \mu_i$ can be modeled as a linear function of $X$: $g(\mu_i) = X_i^T \beta$, with $\text{Var}(Y_i \mid X_i) = \sigma_e V(\mu_i)$ for a known variance function $V$ and a possibly estimated scale parameter $\sigma_e$. (Examples of this include Gaussian regression with $g(\mu) = \mu$, $V(\mu) = 1$, and $\sigma_e$ as the [typically unknown] variance parameter; Poisson regression/log-linear model with $g(\mu) = \log(\mu)$, $V(\mu) = \mu$; and logistic regression with $g(\mu) = \log\left(\frac{\mu}{1-\mu}\right)$, $V(\mu) = \mu(1-\mu)$, and $\sigma_e = 1$.) For now, we assume independence of the $Y_i$ and equal probability of selection. A Gibbs Sampler can then be used to simulate the joint posterior of the unknown
parameters $\theta = (\beta, \sigma_\epsilon)$ and the missing data $y_{\text{mis}}$, Given starting values $\theta^{(0)}$ and $y_{\text{mis}}^{(0)}$, and let $f(\cdot)$ denote the probability density function, consider an iterative simulation algorithm in which the current version of the unknown parameters $\theta^{(r)} = (\beta^{(r)}, \sigma^{(r)}_\epsilon)$ and the missing data $y_{\text{mis}}^{(r)}$ are updated in two steps:

1. Draw model parameters from their posterior distributions conditional on the “filled-in data” (imputed plus observed) from the previous draw:

$$\theta^{(r+1)} \sim f(\theta \mid y_{\text{obs}}, y_{\text{mis}}^{(r)}) .$$

2. Draw missing values in $Y$ from its posterior given all other parameters drawn: $y_{\text{mis}}^{(r+1)} \sim f(y_{\text{mis}} \mid y_{\text{obs}}, \theta^{(r+1)}) .$

After a sufficient number of draws have been obtained to attain convergence of the Gibbs Sampler (Gelman & Rubin, 1992), $M$ widely separated draws of the missing data are combined with the observed data to form the completed data

$$y_{\text{comp}}^{(m)} = (y_{\text{obs}}, y_{\text{imp}}^{(m)}), m = 1, \ldots, M .$$

Inference about the population quantity of interest $Q$ is estimated using the combining rules of Rubin (1987):

$$\hat{Q} = M^{-1} \sum_{m=1}^{M} Q(y_{\text{comp}}^{(m)})$$

$$T^{1/2} (\hat{Q} - Q) \sim t_m$$

$$T = U + (1 + M^{-1})V_B$$

$$U = M^{-1} \sum_{m=1}^{M} \text{vár}(Q(y_{\text{comp}}^{(m)}))$$

$$V_B = (M - 1)^{-1} \sum_{m=1}^{M} \left( \hat{Q} - Q(y_{\text{comp}}^{(m)}) \right)^2$$

$$V_m = (M - 1) \left[ 1 + \frac{U}{(1 + M^{-1})V_B} \right]$$

where $Q(y_{\text{comp}}^{(m)})$ is the point estimate obtained from the $m^{th}$ completed dataset.
\( \gamma_{\text{comp}}^{(m)} \), \( U \) is the within imputation variance calculated as the average of variance estimates based on the \( M \) completed datasets, \( V_b \) is the between imputation variance, \( v_m \) is the degrees of freedom associated with the \( t \) reference distribution for inference about \( Q \).

### 3.2.2 Fixed Cluster Effects Model

We expand the simple random sampling model to \( g(\mu_i) = X_i^T \beta + Z_i^T \gamma \), for \( i = 1, \ldots, n \), where \( Z_i \) is a \((c - 1)\)-dimension vector of dummy variables for the \( c \) clusters selected in the sample. Alternatively we can further expand to include interactions between the fully-observed covariates and the cluster dummy variables:

\[ g(\mu_i) = X_i^T \beta + Z_i^T \gamma + X_i^T \otimes Z_i^T \eta \]. Imputation proceeds as in the SRS setting, with expanded parameter space \( \theta = (\beta, \gamma, \eta, \sigma) \).

### 3.2.3 Random Cluster Effects Model

Here we group the variables by cluster, for the \( i = 1, \ldots, c \) clusters with \( j = 1, \ldots, m_i \) observations. Our imputation model then becomes \( g(\mu_{ij}) = x_{ij}^T \beta + \omega_i \) where we assume that the random cluster effects are distributed as \( \omega_i \sim N(0, \sigma) \) independently for \( i = 1, \ldots, c \). The unknown parameters in the model are now \( \theta = (\beta, \sigma, \sigma_e) \) and \( \omega_i \). A Gibbs Sampler (for Gaussian regression imputations considered in this chapter) is then used to simulate the joint posterior of the unknowns. At current iteration \( t \), each parameter is drawn from its respective conditional posterior distribution as follows:
\[
\omega_i^{(r+1)} \sim f(\omega_i \mid y_{obs}^{(r)}, y_{mis}^{(r)}, \theta^{(r)}), \quad i = 1, \ldots, c.
\]
\[
\theta^{(r+1)} \sim f(\theta \mid y_{obs}, y_{mis}^{(r)}, \tilde{\omega}^{(r+1)}), \quad \tilde{\omega} = (\omega_1, \ldots, \omega_c)^T
\]
\[
y_{mis}^{(r+1)} \sim f(y_{mis} \mid y_{obs}, \tilde{\omega}^{(r+1)}, \theta^{(r+1)})
\]
where \( \omega_i \) is drawn from \( N \left( (1 + \sigma^2_{\omega})^{-1} (y_i - x_i \beta), \left( \sigma^{-2}_e + \sigma^{-2}_{\omega} \right)^{-1} \right) \); \( \sigma_{\omega} \) is drawn from \( \chi_2^{-2} \), \( \sigma_e \) is drawn from \( \chi_{\nu_2 + n - p}^{-2} \), where \( \nu_1 \) and \( \nu_2 \) are the prior specification of the hyper-parameters for the distribution of \( \sigma_e \) and \( \sigma_{\omega} \), respectively; \( \beta \) is drawn from \( N \left( \left( \sum_{i=1}^{c} x_i^T x_i \right)^{-1} \left( \sum_{i=1}^{c} x_i^T (y_i - \omega_i) \right), \sigma_e \left( \sum_{i=1}^{c} x_i^T x_i \right)^{-1} \right) \); and finally the missing data \( y_{j mis} \) are drawn from \( N \left( x_{j mis}^T \beta + \omega_i, \sigma_{\omega} + \sigma_e \right) \).

### 3.3 Multiple Imputation using the Weighted Finite Population Bayesian Bootstrap in Clustered and Weighted Sample Designs

#### 3.3.1 Overview

In this section, we develop a two-step multiple imputation methodology to account for complex sampling designs with two-stage cluster samples. The first step utilizes a weighted finite population Bayesian bootstrap to generate predictive draws of a population that capture the clustering and unequal probability of selection design features; the second step conducts the imputation within each of the predictive draws.

Define \( I \) as a (fully-observed) vector of sampling indicators for a target finite population of size \( N \). To develop the population posterior predictive distribution generation for the first step, let \( Y = [Y_s, Y_{ns}] \) be the survey outcome of interest where missing data occur, which divides into the sampled part \( Y_s \) (for
which \( I = 1 \) and the nonsampled part \( Y_{ns} \) (for which \( I = 0 \)). Define separate response mechanisms for the sampled and the nonsampled part of \( Y \):

\[
R = (R_s, R_{ns}), \text{ such that } Y_s = [Y_{s,obs}, Y_{s,mis}] \text{ further divides into the observed component } Y_{s,obs} \text{ and the missing component } Y_{s,mis}, \text{ corresponding to the sampled } Y \text{ values associated with } R_s = 1 \text{ and } R_s = 0 \text{ respectively, and similarly }
\]

\[
Y_{ns} = [Y_{ns,obs}, Y_{ns,mis}] \text{ divides into those that would have been observed had they been sampled (} R_{ns} = 1 \text{), and those that would have had missing values (} R_{ns} = 0 \text{).}
\]

Note that we can recombine as \( Y = [Y_{obs}, Y_{mis}] \) where \( Y_{obs} = [Y_{s,obs}, Y_{ns,obs}] \) and \( Y_{mis} = [Y_{s,mis}, Y_{ns,mis}] \). In a similar fashion, let \( X = [X_s, X_{ns}] \) be the complete covariate, and \( Z = (w, C) = [(w_s, C_s), (w_{ns}, C_{ns})] = [Z_s, Z_{ns}] \) be the complete design matrix which contains all the essential sample design features (here we restrict to sampling weights \( w \) and cluster membership indicators \( C \)). Note that \( w \) is a matrix \( \{(w_{ji}, w_{ij}), i = 1,...,n, j = 1,...,m_i\} \), where \( w_{ji} \) is the cluster-level weight for element \( j \) in cluster \( i \), \( w_{ij} \) is the element-level conditional weight for element \( j \) given cluster \( i \) is selected, and \( w_{ij} = w_{ji}w_{ij} \) is the overall sampling weight for element \( j \).

Assuming ignorable sampling, the joint predictive distribution of the nonsampled part of the population is given as (Rubin, 1987):

\[
p(Y_{ns}, X_{ns}, Z_{ns}, R_{ns} | Y_s, X_s, Z_s, R_s, I) \propto p(Y_{ns}, X_{ns}, Z_{ns}, R_{ns} | Y_s, X_s, Z_s, R_s),
\]

given \( p(I | Y, X, Z, R) \propto p(I | Y_s, X_s, Z_s, R_s). \) [3.1]

Explicit modeling for \( I \) can be avoided by incorporating design variables \( (Z) \) into a model for \( p(Y_{ns}, X_{ns}, Z_{ns}, R_{ns} | Y_s, X_s, Z_s, R_s) \) that conditionally eliminates
dependence on \( I \) (Gelman, 2007; Little, 2011). However, the practical 
implementation of this in a robust manner using standard parametric models can 
be daunting. Here we pursue a nonparametric approach using a weighted FPBB 
that combines a Polya sampling scheme developed by Ghosh and Meeden (1997) 
in the simple random sampling setting and extended by Meeden (1999) into a 
simple balanced cluster design setting with a second extension by Cohen (1997) 
to accommodate weighted data. These constitute a sample design reversing 
procedure as the first step.

The second step requires a correct imputation model that generates the 
posterior predictive distribution of missing values in the entire population, such 
that it is independent of the response mechanism for both sampled and 
nonsampled parts of the population. That is, we do not need an explicit model for 
\( R \) given ignorable missingness:

\[
 p(Y_{\text{mis}} \mid Y_{\text{obs}}, X, Z, R) \propto \\
\left\{ \begin{array}{ll}
 p(Y_{\text{mis}} \mid Y_{\text{obs}}, X), & \text{if } p(R \mid Y, X, Z) \propto p(R \mid Y_{\text{obs}}, X) \\
p(Y_{\text{mis}} \mid Y_{\text{obs}}, X, Z), & \text{if } p(R \mid Y, X, Z) \propto p(R \mid Y_{\text{obs}}, X, Z)
\end{array} \right. 
\]  

[3.2]

Here we will proceed with a standard parametric approach, modeling the missing 
\( Y \) conditional on the observed \( X \) and \( Z \). Note that the extent to which \( Z \) should be 
incorporated into the imputation model still depends on the specific form of the 
response mechanism, as shown in [3.2].

3.3.2 The Weighted-FPBB in Clustered and Weighted Sample Designs

In a simple two-stage cluster sampling (balanced case) with equal 
probability of selection at both cluster and element levels, Meeden (1999) 
proposed a “two-stage Polya posterior” approach to simulate draws that form an
entire population of clusters, and then an entire population of elements within each cluster. He showed that the posterior mean and variance are very close to standard design-based results. Cohen (1997) proposed a method to generate draws from a weighted Polya posterior using data obtained from weighted sample designs in a non-clustered setting. Here we extend the method of Meeden to a two-stage cluster sampling with unequal cluster sizes (unbalanced case) and combine it with the approach of Cohen to further handle unequal probability of selection.

Next, we propose two variations of a two-stage procedure to simulate draws from equation [3.1]. The first approach assumes that we know the sampling probabilities at both sampling stages, and uses a weighted FPBB at each stage to generate a population of clusters and then a population of elements (and hence is termed an “adapted two-stage Polya posterior” approach). The second approach requires only the final weights, as is typical in most public use surveys, and uses a Bayesian Bootstrap at the first stage to account for clustering effects, and a weighted FPBB using the final weights to generate the population of elements.

3.3.2.1 Double Weighted Finite Population Bayesian Bootstrap (SYN1)

We will call the first approach ‘double weighted FPBB’ . As indicated by the name, we propose to simulate the posterior predictive distribution in equation [3.1] by utilizing the weighted FPBB method at both the PSU (or cluster) level and the element level, such that a synthetic population of clusters is created at first and then a synthetic population of elements is created.
Let \( D = \{(Y_{ij}, X_{ij}, Z_{ij}, R_{ij}), i = 1, \ldots, N, j = 1, \ldots, M_i\} \) denote the population of values for element \( j \) within cluster \( i \), where \( N \) and \( M_i \) are the number of clusters and the number of elements within the \( i^{th} \) cluster in the population, respectively. Thus the population size is \( \sum_{i=1}^{N} M_i = M \). Let \( \{b^1, \ldots, b^r, \ldots, b^r, q = 1, \ldots, r\} \) be \( r \) (1 ≤ \( r \) ≤ \( N \)) distinct matrices of real numbers each of dimension \( |b^q_{row}| \times |b^q_{col}| \) (each column vector corresponds to a survey variable) with no row vectors in common. Each cluster in the population can take the form of one of \( b^q \)'s; let \( t \) be a vector of length \( N \), \( t_i = q \) when the elements in the \( i^{th} \) cluster take on the values of \( b^q \), for \( i = 1, \ldots, N \). Finally, let
\[
c_i(q) = \text{the number of } t_i's \text{ which equal } q, \text{ for } q = 1, \ldots, r, \text{ and}
\]
\[
c^i_{t,D}(k) = \text{the number of } (Y_{ij}, X_{ij}, Z_{ij}, R_{ij})'s \text{ which equal the } k^{th} \text{ row vector } b^i_k, \text{ for } k = 1, \ldots, |b^i_{row}|, i = 1, \ldots, N.
\]

Suppose a two-stage cluster sample is selected from the above finite population. Thus \( D \) divides into the sampled components \( D_s = (Y_s, X_s, Z_s, R_s) \) and the nonsampled components \( D_{ns} = (Y_{ns}, X_{ns}, Z_{ns}, R_{ns}) \). Let \( h \) and \( \tilde{h} \) each represents the collection of the sampled and nonsampled clusters, and thus we can break \( t \) into \( t = (t_h, t_{\tilde{h}}) \). Let \( n \) denote the number of sampled clusters and \( m_i \) the number of sampled elements within the \( i^{th} \) sampled cluster, and thus the total sample size is \( m = \sum_{i=1}^{n} m_i \). Assume \( n = r \), \( m_i = \|b^i_h\| \) the number of distinct row vectors in \( b^i_h \), and thus \( m = \sum_{i=1}^{n} \|b^i_h\| \) for convenience of exposition. Let
\[ w_{h} (i) = \text{the first-stage sample weight of the sampled cluster } i \in h \text{ which equal } b^{q}, \text{ for } i(\equiv q) = 1, \ldots, n(\equiv r). \]

\[ w_{h, D_{i}} (j) = \text{the second-stage sample weight of the } j^{th} \text{ sampled element in the } i^{th} \text{ sampled cluster which equal } b_{k}^{h}, \text{ for } j(\equiv k) = 1, \ldots, m_{i}(\equiv \| b_{k}^{h} \|). \]

Note that \( \sum_{i=1}^{n} w_{h} (i) = n \) and \( \sum_{j=1}^{m} w_{h, D_{i}} (j) = m_{i}. \) Let the set of distinct clusters in the observed sample \( (D_{s}) \) be \( \{b^{1}, \ldots, b^{q}, \ldots, b^{r}\}, \) and \( \zeta = \{\zeta_{1}, \zeta_{2}, \ldots, \zeta_{r}\} \) be the vector of probabilities that \( \Pr(D_{s, i} = b^{q} \mid \zeta) = \zeta_{q}, \text{ for } i = 1, 2, \ldots, n, \) and \( \sum_{q=1}^{r} \zeta_{q} = 1. \) Let \( c_{u} (q) \) and \( c_{r} (q) \) be the number of clusters taking value \( b^{q} \) in the sampled and the nonsampled part of the population, respectively, for \( q = 1, 2, \ldots, r, \) and

\[ \sum_{q=1}^{r} c_{u} (q) = n \text{ and } \sum_{q=1}^{r} c_{r} (q) = N - n. \] Let the set of distinct elements in the sampled clusters be \( \{d^{1}, \ldots, d^{k}, \ldots, d^{m}\}, \) and \( \lambda = \{\lambda_{1}, \lambda_{2}, \ldots, \lambda_{m}\} \) be the vector of probabilities that \( \Pr(D_{s, ij} = d^{k} \mid \lambda) = \lambda_{k}, \text{ for } i = 1, 2, \ldots, n, \) and \( j = 1, \ldots, m_{j}, \) and \( \sum_{k=1}^{m_{j}} \lambda_{k} = 1, \) and \( c_{i, D_{s}}^{i} (k) \) and \( c_{i, D_{u}}^{i} (k) \) be the number of elements taking value \( d^{k} \) in the sampled and the nonsampled part of the population, respectively, for \( k = 1, 2, \ldots, m, \) and

\[ \sum_{k=1}^{m} c_{i, D_{s}}^{i} (k) = m \text{ and } \sum_{k=1}^{m} c_{i, D_{u}}^{i} (k) = M - m = m'. \] Assuming a noninformative Haldane prior of \( \zeta : \zeta \sim \text{Dir}(0, \ldots, 0) \) together with multinomially distributed weighted counts of distinct clusters in the sample data

\[ p(w_{h} (1), \ldots, w_{h} (r) \mid \zeta) \propto \prod_{q=1}^{r} \zeta_{q}^{w_{h} (q)} \text{ yields a Dirichlet posterior distribution of } \zeta: \]
and hence the posterior predictive distribution of counts of distinct clusters in the nonsampled data follows a compound multinomial distribution:

\[ c_\zeta (1),..., c_\zeta (r) \mid w_h (1),..., w_h (r) \sim Mult (N - n; \zeta^*) , \] where \( \zeta^* \) is the parameter vector adjusted for unequal selection probability for clusters as in [3.4]. In a similar fashion, the posterior predictive distribution of counts of distinct elements in the nonsampled data follows a compound multinomial distribution:

\[ c_i^1 (1),..., c_i^1 (m) \mid w_{i, D_1} (1),..., w_{i, D_1} (m) \sim Mult (M - m; \lambda^*) , \] where \( \lambda^* \) is the parameter vector adjusted for unequal selection probability for elements as in [3.5]. We then have the “adapted two-stage Polya posterior” derived as follows (for simplicity of exposition and by an abuse of notation, we use \( t_\zeta \) and \( t_h \) to represent \( c_\zeta (1),..., c_\zeta (r) \) and \( w_h (1),..., w_h (r) \), respectively; similarly, we use \( D_{ns} \) and \( D_s \) for \( c_i^1 (1),..., c_i^1 (m) \) and \( w_{i, D_1} (1),..., w_{i, D_1} (m) \), respectively):
\[
p(D_{ns} \mid D_s) = \frac{p(D) \cdot p(D \mid t)}{p(D_s) \cdot p(D_s \mid t_h, t_\pi)}
\]
\[
= \frac{p(t_h, t_\pi) \cdot p(D_{ns} \mid D_s, t_h, t_\pi)}{p(t_h) \cdot p(D_s \mid t_h, t_\pi)}
\]
\[
= \frac{p(t_h) \cdot p(D_{ns} \mid D_s, t_h, t_\pi)}{p(t_h) \cdot p(D_s \mid t_h, t_\pi)}
\]
\[
= \frac{p(t_h) \cdot p(D_{ns}) \cdot p(D_s \mid t_h, t_\pi)}{p(t_h) \cdot p(D_s \mid t_h, t_\pi)}
\]
\[
= p(t_h) \cdot p(D_{ns} \mid D_s, t_h, t_\pi)
\]
\[
= \int_0^1 \ldots \int_0^1 \prod_{q=1}^{r-1} \xi_q w_{r,(q)-1} \cdot (1 - \sum_{q=1}^{r-1} \xi_q)^{w_{r,(r-1)-1}} \cdot d\xi_1 \ldots d\xi_{r-1}
\]
\[
\times \prod_{k=1}^{m-1} \lambda_k w_{r,s_{a_k}}(k)^{-1} \cdot (1 - \sum_{t=1}^{m-1} \lambda_t)^{-1} \cdot d\lambda_1 \ldots d\lambda_{m-1}
\]
\[
= \left\{ \prod_{q=1}^{r} \left\{ \Gamma(w_{r,(q)}) / \Gamma(w_{r,(q)}) \right\} / \{\Gamma(N) / \Gamma(n)\} \right\}
\]
\[
\times \left\{ \prod_{k=1}^{m} \left\{ \Gamma(w_{r,s_a}(k)) / \Gamma(w_{r,s_a}(k)) \right\} / \{\Gamma(M) / \Gamma(m)\} \right\}
\]  

[3.3]

where \( w_r(q) = w_{r,s_a}(q) + c_q(r) \), \( w_{r,s_a}(k) \) is the element-level weight, and

\( w_{r,s_a}(k) = w_{r,s_a}(k) + c_{s_a}(k) \).

The posterior distribution in [3.3] does not lend itself to direct calculation, and is approximated by a Monte Carlo simulation procedure described as follows:

**Step 1:** Apply the weighted FPBB to sampled clusters, that is, draw nonsampled clusters in the population based on the weighted Polya urn distribution. This step realizes the first factor in [3.3]. Denote the original sample of clusters by \( \{c_1, c_2, \ldots, c_n\} \). In generating \( N - n \) clusters \( \{c^*_1, c^*_2, \ldots, c^*_{N-n}\} \) from the original \( n \), we resample the sampled clusters with probability.
where \( l_{i,k-1} \) is the number of times that the \( i^{th} \) cluster has been sampled at the \((k-1)^{th}\) resampling, \( w_{n}(i) \) is the weight for the \( i^{th} \) cluster which is normalized to sum up to the total number of clusters, i.e. \( \sum_{i=1}^{n} w_{n}(i) = N \). Repeat step 1 \( L \) times to obtain \( L \) FPBB synthetic populations of clusters.

**Step 2:** For each repetition of step 1, form a population of clusters \( \{c_1,c_2,\ldots,c_n,c_1^*,c_2^*,\ldots,c_{N-n}^*\} \). Record the number of times each of the \( n \) clusters from the original sample appears in the FPBB population of clusters, denoted by \( \tau_{i}, i = 1,\ldots,n \) and \( \sum_{i=1}^{n} \tau_{i} = N \). Then update the within cluster element-level conditional weights to unconditional weights as follow: \( w_{jk}^* = w_{jk} \times \tau_{i}, i = 1,\ldots,n \), where \( w_{jk}^* \) is the inverse of the conditional probability that element \( j \) is sampled given cluster \( i \) sampled. (Note \( w_{jk}^* \) is the same as \( w_{l_{i,k-1}}(j) \) as previously defined.) Now each observed element in these \( n \) clusters is associated with an updated weight that represents all nonsampled elements in both the sampled cluster it belongs to and similar nonsampled clusters in the population. Note that the resulting sample has the same sample size of elements \( m \) but different survey weights than the original sample, and we term it a ‘FPBB sample’.

We then apply the weighted FPBB again to elements in the ‘FPBB sample’, and this realizes the second factor in equation \([3.3]\). In generating \( M-m \) units from the \( m \) elements in the FPBB sample, each of the elements in the FPBB
sample is resampled with probability

\[
\hat{\lambda}_{j|i}^* = \frac{w_{j|i}^* - 1 + I_{j|i,k-1} \times \left( \frac{M - m}{m} \right)}{M - m + (k-1) \times \left( \frac{M - m}{m} \right)}, k = 1, ..., (M - m + 1), j = 1, ..., m. \tag{3.5}
\]

where \( w_{j|i}^* \) is the updated conditional weight for the \( j^{th} \) element in the \( i^{th} \) cluster.

Note \( w_{j|i}^* \)'s inherit the information from the cluster-level selection probability \( \zeta_i^* \).

Again they are normalized to sum up to the total number of elements in the entire population, i.e. \( \sum_{j=1}^{m} w_{j|i}^* = M \). Thus we create a single synthetic population.

Repeat step 2 \( B \) times to obtain \( B \) FPBB synthetic populations of elements.

Such a procedure captures the sampling variance while untangling the sample weights at both cluster and element levels through the creation of synthetic populations. This also realizes our goal of resolving clustering effects (design effect due to cluster sampling, i.e. \( \text{deff}_c = 1 + (\bar{m} - 1) \rho \), where \( \bar{m} \) is the averaged sample size within each sampled cluster and \( \rho \) is the intraclass correlation) as a sampling phenomenon. A simple illustrative example goes as follows: suppose we obtain a sample of households (HHs) from a two-stage cluster sampling design, where all US metropolitan statistical areas (MSAs) are treated as clusters to be sampled at the first stage. Suppose Detroit is among such sampled clusters. By using the proposed procedure, we first replicate the number of all ‘Detroit-like MSAs’ based on the sample weight for Detroit and use it to update the conditional sample weights of sampled HHs within Detroit; then we replicate the number of all nonsampled HHs in Detroit as well as that in Detroit-like MSAs in the population. The same reasoning applies to other sampled MSAs that are different
from Detroit (e.g. LA and LA-like MSAs). Once we generate the population of HHs, HH-level inference becomes straightforward without the need to consider effects due to sampling at the MSA-level.

Because only final weights $w_{ij}$ are usually available in the sample data that are released to the public, we will often need to estimate both the cluster-level weight $w_{h}(i)$ in formulae [3.4] and the element-level conditional weight $w_{j|i}$ in formulae [3.5]. We propose estimating $w_{h}(i)$ by $\hat{w}_{i} = \sum_{j} w_{ij} / M_{i}$ and $w_{j|i}$ by $\hat{w}_{j|i} = w_{ij} / \hat{w}_{i}$, where $w_{ij}$ is the overall sample weight as defined in section 3.3.1.

### 3.3.2.2 Bayesian Bootstrap — Weighted FPBB (SYN2)

A variation of the procedure proposed in subsection 3.3.1.1 is to replace the first-stage weighted FPBB for clusters with regular Bayesian Bootstrap (BB), hence we name it “BB-weighted FPBB”. Operationally the difference is that, rather than obtaining a sample of clusters from a draw from a Polya posterior according to the cluster-level weights, we use a simple replication method assuming IID to capture the cluster-level sampling variance. The final sample weights ($w_{ij}$) instead of the adjusted element-level conditional weights (i.e. $w_{j|i}^{*}$) are then directly used as input in the second-stage weighted FPBB. We will show that this procedure not only serves as a handy approximation to the double weighted-FPBB due to easier implementation in practice, it can also deal with complex missing data problems as effective as the double weighted-FPBB where fully parametric MI fails. The full procedure is described as follows:
**Step 1:** Apply Bayesian Bootstrap (BB) (Rubin, 1981) to draw a sample of the clusters from the original sample of clusters. Unlike parametric model-based approaches, the BB can be used to obtain the distribution of a population parameter through the multinomial likelihood and an improper Haldane prior without assuming a fully parametric model for the response variable, therefore is particularly relevant and useful in the survey sampling context. See Aitkin (2008) for a comprehensive application of BB in finite population inference. In our case, we first use it to simulate the posterior for the proportions of distinct clusters in the population:

1) Draw \( n-1 \) i.i.d. random variates from \( \text{Unif}(0,1) \) and order them as 

\[ u_{(1)}, u_{(2)}, \ldots, u_{(n-1)}; \]

2) Calculating the gaps between \( u_{(i)} \)'s as \( g_i = u_{(i)} - u_{(i-1)}, i = 1, \ldots, n-1 \),

where \( u_{(0)} = 0 \) and \( u_{(n)} = 1; \)

3) Sample \( n \) clusters with replacement with the vector of probabilities 

\( g = (g_1, \ldots, g_n) \) to attach to the \( n \) distinct clusters in the parent sample, and record the number of times \( \tau_i \) each cluster is selected in the ‘Bayesian bootstrap (BB) sample’, \( \sum_{i=1}^{n} \tau_i = n. \)

4) Apply the ‘ultimate cluster principle’ (Wolter, 2007), that is, once a given cluster is taken into the bootstrap sample, all successive stage units are taken into the sample also. Modify the initial case weight in
the parent sample as follow:

\[ w_{ij}^* = \begin{cases} \tau_i \cdot w_{ij}, & \text{if } i^{th} \text{ cluster is selected or duplicated;} \\ 0, & \text{if } i^{th} \text{ cluster is not selected.} \end{cases} \]

Normalize \( w_{ij}^* \)'s to sum up to the population size. Note that the bootstrap sample size \( (m^*) \) is different from the parent sample size \( (m) \).

**Step 2**: Generate nonsampled elements of the population accounting for inclusion weights: Select a FPBB Pólya sample of size \( (M - m^*) \) from the compound multinomial distribution \( \text{mult}(M^*; p_1, \ldots, p_d) \), where \( p_j, j = 1, \ldots, d \), are the proportions of distinct values in the population and they are computed based on

\[
p_j = \frac{w_{ij}^* - 1 + l_{ij,k-1} \times \left( \frac{M - m^*}{m^*} \right)}{M - m^* + (k-1) \times \left( \frac{M - m^*}{m^*} \right)}, \quad k = 1, \ldots, M - m^* + 1, \quad j = 1, \ldots, d. \tag{3.6}
\]

Simulate several \( (B) \) copies of the nonsampled population.

**3.3.3 Multiply Imputing Missing Data**

Denote the \( L \times B \) FPBB “unweighed” synthetic populations generated by the weighted-FPBB by \( P^{\text{syn}} = \{P_{(b)}^{(l)}, b = 1, \ldots, B, l = 1, \ldots, L\} \), where

\[ P_{(b)}^{(l)} = (Y_{(b)\text{mis}}^{(l)}, P_{(b)\text{obs}}^{(l)}) \].

Having untied the sampling design, we are ready to perform conventional parametric MI under an IID assumption. Following the standard MI procedure or approximations such as SRMI (Raghunathan et al., 2001), we obtain draws from the posterior predictive distribution \( p(Y_{(b)\text{mis}}^{(l)} | P_{(b)\text{obs}}^{(l)}) \), where \( Y_{(b)\text{mis}}^{(l)} \) and
$P^{(l)}_{(b)obs}$ consist of the unobserved and observed data in the $lb^{th}$ FPBB dataset respectively. Without the need to include weights or cluster-level random effects in the imputation model, our task can now be concentrated on correctly modeling the covariates as well as interactions among them whenever necessary.

Point and variance estimate then proceeds as follows. Denote the imputed synthetic datasets as $P^{imp} = \{P^{(1)}_{(1)}, ..., P^{(1)}_{(M)}, ..., P^{(1)}_{(BM)}, ..., P^{(1)}_{(BM)}\}$. Note here $M$ denotes the number of imputations created and is different from the $M$ in the previous section which denotes population size of elements. By the standard Rubin (1987) MI combining rules, we have

$$Q \mid P^{imp} \sim t_{L-1}(\overline{Q}_L, (1 + L^{-1})V_L),$$

where $\overline{Q}_L = \frac{1}{L} \sum_{l} \tilde{Q}^{(l)}$, $V_L = \frac{1}{L-1} \sum_{l} (\tilde{Q}^{(l)} - \overline{Q}_L)^2$, and $\tilde{Q}^{(l)} = \lim_{M \to \infty} \frac{1}{BM} \sum_{b} \sum_{m} q^{(bm)}$, where $q^{(bm)}$ is an estimate of $Q$ obtained from the $m^{th}$ imputation of the $b^{th}$ synthetic population within $l^{th}$ (finite population) Bayesian Bootstrap sample, obtained as $\tilde{Q}^{(l)} = \frac{1}{BM} \sum_{b} \sum_{m} q^{(bm)}$. Note that the generation of the synthetic population sets the within imputation variance to 0 so that the posterior variance of $Q$ can be obtained using $V_L$ only; see Dong et al. (2014) and Zhou et al. (2013a).

Note a difference should be made to the small sample scenario when we apply these combining rules. Lo (1988) showed that the variance estimator for the FPBB mean in a simple random sample setting should be inflated by the factor $(\frac{n+1}{n-1})$ (here $n$ is the sample size of elements). Therefore, to get the variance...
estimate correct under the double-weighted FPBB, we should use \[ \frac{n+1}{n-1} (1+L^{-1}) V_L \]
(here \(n\) is the sample size of clusters).

3.4 Simulation Study

The simulation study considers two different clustered population data structures. For the first structure, we impose only clustering effects in the data model while leaving the weight independent of the data generating mechanism.

Structure 1: \( Y_{ij} \sim \alpha_0 + \alpha_i X_{ij} + u_i + \varepsilon_{ij} \), where \( u_i \sim N(0, \sigma_u^2), \varepsilon_{ij} \sim N(0, \sigma_e^2) \)

For the second structure, the data model depends on both clustering effects and the measure of size (MOS) \((Z)\) for second-stage sampling as well as its interaction with the covariate \((X)\). Under this model, the subsampling probability is correlated with the response variable even after conditioning on the covariate:

Structure 2: \( Y_{ij} \sim \beta_0 + \beta_1 X_{ij} + \beta_2 Z_{ij} + \beta_3 X_{ij} Z_{ij} + v_i + \varepsilon_{ij} \),
where \( v_i \sim N(0, \sigma_v^2), \varepsilon_{ij} \sim N(0, \sigma_e^2) \)

We conduct a separate factorial design for each population structure by considering the following simulation factors:

- **Intraclass correlation coefficients (ICC\(_{3,1}\))**: We set ICC\(_{3,1}\) to be 0.1 or 0.5. Note that the ICC\(_{3,1}\) we examine here is a conditional ICC which measures residual correlation after allowing for dependence on the \(X\) variables, which will in general be larger than ICC\(_Y\), the unconditional ICC of \(Y\). We also vary the cluster size \((M_i)\) to magnify the design effect. We purposely do not let the clustering occur on the covariates because the
impact of ignoring clustering effects in the imputation model may be abated by conditioning on $X$, which partially explains some of the unobserved clustering effects on $Y$ (Andridge, 2011).

- The number of selected clusters in the sample ($n$), given a fixed number of clusters ($N$) in the hypothetical population.

We set two values for the first stage sample size: $n = 10$ or 40. The purpose is to observe both small and large sample behaviors for alternative methods. Meeden argues that for the “two-stage Polya posterior” to yield sensible 0.95 credible interval estimates in a simple balanced two-stage cluster sample, the number of subunits (i.e. $m$) selected in each sampled cluster should be large enough. We follow his suggestion by setting $m = 20$ (or $\bar{m} = 20$ under an unbalanced design), and therefore we can focus on the effects of different PSU-level sample sizes on the performance of the methods.

- The response mechanism ($R$) for $Y$:

Much research that investigates methods for missing data imputation only looks at one type of MAR mechanism, namely where the response indicator depends on the regular covariate matrix $X$. However, it is reasonable to assume that the missingness is also related to sample design information available in the sample cases. For example, in the Survey of Income and Program Participation (SIPP), low-income populations are often oversampled, and thus low-income respondents tend to have smaller weights in the sample dataset. If these families or individuals are also less likely to respond to survey questions on the specific topic, then the response mechanism can be considered as being associated with
survey weights (W) (Andridge & Little, 2009). We will look at both types of response mechanisms (denoted as MAR_X and MAR_X,W, respectively) in our simulation for both data structures.

Based on the above stated factors, we conduct a full 2×2×2 factorial design for each data structure. A total of five MI methods are compared in the simulation study. They are described as follows:

- **SRS**: the linear model where clustering effects are ignored altogether
- **FX**: the linear fixed cluster effects model described in 3.2.2
- **RE**: the linear random cluster effects model described in 3.2.3
- **SYN1**: the two-step MI method with the double-weighted FPBB as 1st step
- **SYN2**: the two-step MI method with the BB-weighted FPBB as 1st step

The imputation tasks under SRS, FX, SYN1 and SYN2 are all implemented using R *mice* package (van Buuren & Groothuis-Oudshoorn, 2011). The imputation under RE is implemented using R *pan* package (Zhao & Schafer, 2013).

Specifically, we selected noninformative priors for regression parameters, and diffuse inverse-Wishart priors for variance components; we imputed the missing data 5 times taking 100 iterations between imputations, after a burn-in period of 1,000 iterations.

Finally, for imputation, we consider three model specifications:

- **X-only model**: \( y \sim N(\beta_0 + \beta_1 x + [b], \sigma^2) \)
- **Main effect model**: \( y \sim N(\beta_0 + \beta_1 x + \beta_2 w + [b], \sigma^2) \)
- **Interaction model**: \( y \sim N(\beta_0 + \beta_1 x + \beta_2 w + \beta_3 x w + [b], \sigma^2) \),
Note that the cluster effect term \([b]\) is only included under FX and RE. Note also that we will employ all three models for the three fully parametric MI methods although only the interaction model is considered as “appropriate”. For the two variations of synthetic MI methods, we will use X-only model for imputation throughout all scenarios.

3.4.1 Description of the Design

**Step 1: Population data generation**

We fix the number of clusters in the population as \(N = 400\) for all data structures. Varying cluster sizes are generated from the same uniform distribution \(M_j \sim \text{Unif}(50, 250), i = 1, \ldots, 400\), which sum up to a population size of \(N = 59501\) for population Structure 1 and \(N = 59230\) for population Structure 2.

For Structure 1, we let \(\alpha_0 = 1, \alpha_1 = 1.5\), and \(X_{ij} \sim N(0,1)\). For Structure 2,

\[
Z_{ij} \sim N(2,1), X_{ij} \sim N(0.1*Z,1), \quad \beta_0 = 0, \beta_1 = 0.2, \beta_2 = 0.6, \beta_3 = 0.5.
\]

The random cluster effects are set to be independent standard normal variates, i.e.

\(u_i, v_i \sim N(0,1)\). In order to generate hypothetical populations with varying ICC, we vary the random error terms as: \(e_{ij}^1 \sim N(0,1)\) and \(e_{ij}^2 \sim N(0,9)\), resulting in \(ICC_1 = 0.5\) and \(ICC_2 = 0.1\) respectively. These correspond to two hypothetical populations under each data structure.

**Step 2: Drawing replications of samples**

We draw clusters with probability proportional to the cluster size \(M_j\), and

\[
f_{ij} = n \cdot M_i / \sum_{k=1}^{N} M_k\; ; \text{The subsampling methods within each sampled cluster are}
\]
different for the two data structures: we select elements using SRSWOR with an equal rate of $f_2 = 0.1$ for the first data structure, while using PPSWOR with $Z$ as the MOS for the second data structure. As a result, the final weight is dominated by the cluster-level weight for the former and the element-level weight for the latter.

**Step 3: Imposing missingness**

For samples drawn from the first data structure: under MAR\_X, we use a logistic function as the deletion function,

$$\Pr(R = 1) = \expit(\lambda_0 + \lambda_i X), (\lambda_0 = 0.8, \lambda_i = -0.2);$$

under MAR\_X,W, we define a probit model $\Pr(R = 0| S) = \Phi(S)$ to generate missingness indicators, where the latent variable is $S = \eta_0 + \eta_i X + \eta_2 \log(\text{weight}) + e, e \sim N(0,1^2)$.

$(\eta_0 = -3.8, \eta_i = -0.8, \eta_2 = 1.)$ Similar forms of the probit model are used for data structure 2 with a different latent variable:

$$S = \xi_0 + \xi_i X + \xi_2 Z + \xi_3 X * Z + e, e \sim N(0,1^2),$$

where we set

$\xi_0 = -0.635, \xi_i = 0.4, \xi_2 = \xi_3 = 0$ under condition MAR\_X, and

$\xi_0 = -0.55, \xi_i = 0.4, \xi_2 = -0.5, \xi_3 = 0.4$ under condition MAR\_X,W. All scenarios result in a missing rate of approximately 30% on $Y$.

**Step 4: MI inference**

For each scenario, 200 simulations are obtained. The number of imputations under the fully parametric MI methods is $M = 5$ and we set $L = 100, B = 20$ and $M = 5$ for the synthetic MI methods. Our focus is on estimating the population mean of $Y$ and regression coefficients of $Y$ on $X$. The
analyst’s model is assumed \( Y = b_0 + b_1 X \).

Design-based analyses are performed for original samples before deletion (termed “BD”) and for fully parametric methods (SRS/FX/RE). Specifically, we use Horvitz-Thompson-type estimator for point estimation and ultimate cluster variance estimator for the within imputation variance estimation when estimating the population mean (Cochran, 1977), and we use weighted least squares estimators and sandwich variance estimators to estimate regression coefficients (Korn & Graubard, 1999).

In terms of interval estimation, since neither a sample size of \( n=10 \) or \( n=40 \) adequately allows for a normal approximation, we use \( t \) reference distribution instead to construct the 95\% confidence intervals for the mean and regression coefficients. Different degrees of freedom (df) are used for different methods: 1) For the actual samples BD, complete data df is used, i.e. \( v_{\text{com}} = n - p \) where \( p \) is the number of parameters; 2) For the fully parametric MI method, we use the small sample df derived in Barnard and Rubin (1999), i.e.

\[
\hat{\nu}_m = \left( \frac{1}{\hat{\nu}_m} + \frac{1}{\hat{\nu}_{\text{obs}}} \right)^{-1} = v_{\text{com}} \left[ \left( \hat{\lambda} (v_{\text{com}})(1 - \hat{\nu}_m) \right)^{-1} + \frac{v_{\text{com}}}{v_m} \right]^{-1}, \text{ where } v_m \text{ is the large-sample-finite-number-imputation df as defined in section 3.2.1, and } \hat{\nu}_m = (1 + M^{-1}) \frac{V_B}{T} \text{ is approximately the Bayesian fraction of missing information (FMI); and } \text{3) For the proposed synthetic MI method, because the FMI is not well-defined in the context of synthetic data in the current literature, and hence the df for constructing interval estimation for the small sample } t \text{-approximation synthetic data cannot be constructed using FMI, we propose using } v_{\text{syn}} = \min\{L-1, v_{\text{com}}\} \text{ as an}
approximation. This simultaneously accounts for finite number synthetic data generation \((L-1)\) and small sample size \((\nu_{com})\).

### 3.4.2 Results

Results pertaining to varying simulation conditions are summarized in Tables 3.1 - 3.6. The abbreviations for different methods are as previously described. We display the following five measures in the result tables: Pt.est. (point estimate) or Relbias (relative bias), SE (standard error of the estimate), Emp.SE (empirical standard error of the estimate), RMSE (root mean squared error) and Cov. (coverage rate of the nominal 95% confidence interval). We also display the interval widths (Intv.wth) in Tables 3.1 – 3.2.

- **Population structure 1**

  Under population structure 1, the final sampling weight is dominated by cluster-level weights and is uncorrelated with the outcome variable, so the estimator of \(Y\) is unbiased even when we exclude weight as a predictor in the imputation model (see Table 3.1 and 3.2). Thus, our focus is on the clustering effects which only impact the variance and hence coverage properties. The average estimated fraction of missing information (FMI) using the model-based MI methods (including SRS, FX and RE) is 11.5% for the mean and 32.5% for the slope. The estimated FMI differs by ICC when estimating the mean, which is 5.4% for ICC=0.5 and 17.6% for ICC=0.1.

  Despite MI’s ability to propagate imputation uncertainty, the standard imputations based on SRS where clusters are ignored altogether leads to even lower standard errors (SEs) than the actual before deletion samples, thus yielding the poorest CI coverage for the estimated mean. It also leads to larger than
expected SEs when estimating the slope, since the variability due to clustering is absorbed all in the residual variance which further inflates the variance of slope, yielding overly conservative intervals. On the other hand, the fixed effects model tends to overestimate the SEs for the mean (e.g. numbers in red), and thus results in intervals that are too conservative, especially when the ICC is low ($ICC=0.1$). This result replicates the findings of Andridge (2011). The results hold regardless of whether the response indicator is a function of $X$ alone (MAR$_X$), or both $X$ and $W$ (MAR$_{X,W}$).

The two newly proposed synthetic methods and the RE model have similar results when estimating the mean; their performances are generally better than FX and SRS (Table 3.1). Our synthetic methods demonstrate some advantages over RE when estimating the slope in the small sample scenario ($a=10$). While the use of t-corrected inference for small sample size under RE tends to overcorrect the coverage (e.g. numbers in blue), SYN1 and SYN2 yield approximately nominal coverage (Table 3.2). Again, results are similar for both the MAR$_X$ and MAR$_{X,W}$ missingness mechanisms.
Table 3.1 Performance of alternative MI methods for estimating the Mean: population 1, two-stage unbalanced sample design (tv=true value)

<table>
<thead>
<tr>
<th>Simulation Parameters</th>
<th>Method</th>
<th>MAR_X</th>
<th></th>
<th></th>
<th></th>
<th>MART</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
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<td>ICC=0.5, a=10</td>
<td>BD</td>
<td>1.02</td>
<td>0.34</td>
<td>0.36</td>
<td>0.36</td>
<td>1.54</td>
<td>95.5%</td>
<td>1.02</td>
<td>0.34</td>
<td>0.36</td>
</tr>
<tr>
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<td>0.36</td>
<td>1.33</td>
<td>90.50%</td>
<td>1.03</td>
<td>0.30</td>
<td>0.37</td>
</tr>
<tr>
<td></td>
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<td>0.37</td>
<td>0.36</td>
<td>0.36</td>
<td>1.69</td>
<td>96.50%</td>
<td>1.03</td>
<td>0.37</td>
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</tr>
<tr>
<td></td>
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<td>1.02</td>
<td>0.36</td>
<td>0.36</td>
<td>0.36</td>
<td>1.65</td>
<td>95.50%</td>
<td>1.02</td>
<td>0.37</td>
<td>0.37</td>
</tr>
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<td>0.36</td>
<td>0.36</td>
<td>1.59</td>
<td>95.50%</td>
<td>1.02</td>
<td>0.36</td>
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</tr>
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<td>95.50%</td>
<td>1.02</td>
<td>0.36</td>
<td>0.38</td>
</tr>
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<td>0.19</td>
<td>0.19</td>
<td>0.73</td>
<td>95.50%</td>
<td>1.04</td>
<td>0.18</td>
<td>0.19</td>
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<td>1.04</td>
<td>0.15</td>
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<td>0.43</td>
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<td>0.45</td>
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<td>98.50%</td>
<td>1.07</td>
<td>0.54</td>
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<td>0.45</td>
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<td>98.50%</td>
<td>1.08</td>
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<tr>
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<td>0.46</td>
<td>0.46</td>
<td>2.06</td>
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<td>97.00%</td>
<td>1.07</td>
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Table 3.2 Performance of alternative MI methods for the Slope: population 1, two-stage unbalanced sample design (tv=true value)

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</tbody>
</table>
• *Population structure 2:*

Under population structure 2, the final weight is dominated by the element-level weights and is associated with the outcome variable $Y$ after conditioning on $X$. Therefore the clustering effects on the inference for $Y$ are confounded by the weight effects.

Tables 3.3 - 3.4 display the results of SYN1 and SYN2 in the complete data context (SYN1_BD, SYN2_BD) for estimation of the mean and slope, respectively. Both versions of the proposed method result in point and variance estimates very close to that obtained from the actual replication samples (BD). This clearly indicates the ability of the synthetic procedure to untie both the weight and the clustering effects. There are even slight gains in efficiency (i.e. smaller RMSE) of the proposed methods over the design-based BD estimates.

Tables 3.5 - 3.6 display the results of different MI methods in the analysis of missing data for estimating the mean and slope, respectively. When estimating the mean under MAR$_X$ condition, all three fully parametric methods yield sizable biases and large RMSE in the point estimates under the $X$-only model, and at least the main effects model is required in this case for these biases to disappear. But if the missingness also relates to the weight (i.e. under MAR$_{X,W}$ mechanism), then only the interaction model gives approximately unbiased estimates. (For example, the relative biases are about 6.5% for SRS$_{x,w}$, FX$_{x,w}$ and RE$_{x,w}$.) For the slope, however, the correct (interaction) model is always required for obtaining unbiased estimates, regardless of the missingness
mechanisms. (For example, the relative biases are about 9.0% under MAR_X and 14.0% under MAR_X,W, for SRS_x,w, FX_x,w and RE_x,w.) As before, the SRS method underestimates the variance for the mean and overestimates the variance for the slope because the clustering effects are ignored. The FX method with small ICC overestimates the variance and hence the confidence interval coverage for MAR_X; and this problem becomes more pronounced for MAR_X,W. (For example, the SE./Emp.SE./Cov. are 0.584/0.538/99.0% for MAR_X and 0.638/0.567/99.5% for MAR_X,W, even under the correct model FX_x,w,x^w). With misspecified imputation models under all three fully parametric methods, the magnitude of biases and undercoverage is amplified by the extra association between the weight and the response mechanism in addition to its association with the outcome variable.

Our synthetic MI performs consistently well in all scenarios, and is superior over the fully parametric methods in this combination of population structure and sampling design. By using the simplest X-only imputation model, it not only yields comparable RMSE as the FX and RE using the appropriate interaction model, it also results in the smallest relative bias for point estimates. This is because, despite the operational and inferential similarity to the frequentist bootstrap (Efron, 1979), the (finite population) Bayesian bootstrap simulates the posterior distribution of the true parameter (\( \varphi \)) instead of the sampling distribution of a statistic estimating that parameter (\( \hat{\varphi} \)). As a result, the point estimates based on our method average closer to the true value than those obtained from fully parametric methods which more resemble the statistic based
on actual before deletion samples. There are slight underestimation for the variance (i.e. SE. < Emp.SE.) with SYN1 relative to SYN2 as well as the appropriate fully parametric models (i.e. FX_x,w,x\*w and RE_x,w,x\*w), but this does not affect the overall conclusion.

Furthermore, while the correct imputation model in the form of [3.2] is required for our semi-parametric method to break associations between the sample design (W) and response mechanism (R) under a MAR_X,W condition, the correct structure in the observed synthetic population data greatly reduce the effect of this association in the ultimate inference, even if the design is not included in the second-step imputation.
Table 3.3 Performance of alternative methods for estimating the mean: population 2, two-stage unbalanced sample design, before deletion results (tv=true value).

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<tr>
<th>Simulation Factor</th>
<th>Method</th>
<th>Summary Statistics</th>
</tr>
</thead>
<tbody>
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<td></td>
<td>Pt.est.</td>
</tr>
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<td>ICC =0.5, a=40; tv=1.542</td>
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<td></td>
<td>SYN1_BD</td>
<td>1.507</td>
</tr>
<tr>
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<td>SYN2_BD</td>
<td>1.514</td>
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<td>SYN2_BD</td>
<td>1.509</td>
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<td>1.511</td>
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<td>SYN2_BD</td>
<td>1.521</td>
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<td>SYN2_BD</td>
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Table 3.4 Performance of alternative methods for estimating the slope: population 2, two-stage unbalanced sample design, before deletion results (tv=true value).

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**Table 3.5** Performance of alternative MI methods for estimating the mean: population structure 2, two-stage unbalanced sampling design.

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<td>0.187</td>
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Table 3.6 Performance of alternative MI methods for estimating the slope: population structure 2, two-stage unbalanced sampling design.

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<td>Intv.wth</td>
<td>Cov.</td>
<td>Relbias</td>
<td>SE.</td>
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<td>13.6%</td>
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<td>Intv.wth</td>
<td>Cov.</td>
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<td>13.9%</td>
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<td>1.894</td>
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3.5 Application to NASS-CDS data

The National Automotive Sampling System Crashworthiness Data System (NASS-CDS) publishes micro-level crash record datasets that are obtained from a representative three-stage probability sample design: geographic region (county or groups of counties), police departments, and police-reported crashes. The design also has unequal probabilities of selection, with vehicles in more severe crashes much more likely to be sampled than vehicles in less severe crashes. The sample is selected annually from all police-reported crashes that resulted in at least one vehicle having to be towed from the scene for damage (http://www.nhtsa.gov/NASS).

A key feature of this car-crash data is the Delta-V measure (roughly defined as the “instantaneous” change in velocity). This measure of crash intensity has two qualities that make it quite useful as a tool for testing our imputation method. First, a very high proportion of car-crash cases are missing this variable; second, there exists a rich set of covariates which are potentially good predictors in the imputation for Delta-V. These include basic demographics, road condition, type of vehicle, injury condition of the driver, number of vehicles in the crash, etc. The survey variables selected for imputation and analysis purpose are summarized in Table 3.7. We estimate: 1) the mean of Delta-V as a continuous variable ($\bar{Y}_{\text{Delta-V}}$), and 2) the odds ratio of having severe injury or certain types of injury given varying levels of Delta-V.
Table 3.7 Summary of selected survey variables for imputation

<table>
<thead>
<tr>
<th>Selected Variables</th>
<th>Values and labels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delta V</td>
<td>Continuous</td>
</tr>
<tr>
<td>driver's age</td>
<td>Continuous</td>
</tr>
<tr>
<td>maximum ais injury</td>
<td>1-not injured/minor injury; 2-severe injury;</td>
</tr>
<tr>
<td>direction of force</td>
<td>1-frontal; 2-right side; 3-left side; 4-rear; 5-other</td>
</tr>
<tr>
<td>driver's gender</td>
<td>1-male; 2-female</td>
</tr>
<tr>
<td>light condition</td>
<td>1-daylight; 2-dark; 3-dark, lighted; 4-dawn/dusk</td>
</tr>
<tr>
<td>road surface condition</td>
<td>1-dry; 2-wet; 3-other</td>
</tr>
<tr>
<td>vehicle type</td>
<td>1-passenger car; 2-truck</td>
</tr>
<tr>
<td>model year</td>
<td>1-&gt;2006; 2&lt;=2006</td>
</tr>
<tr>
<td>vehicle make</td>
<td>1-american; 2-japanese; 3-korean; 4-european&amp;other</td>
</tr>
<tr>
<td>body region</td>
<td>1-head; 2-thorax; 3-lower extremity; 4-other</td>
</tr>
<tr>
<td>primary sampling unit</td>
<td>1,2,…,24.</td>
</tr>
<tr>
<td>final survey weight</td>
<td>Continuous</td>
</tr>
</tbody>
</table>

The analysis sample is obtained by merging four different data files at both occupant level and vehicle level. We retain only the driver data, and end up with $n = 7525$ cases in the final sample, where nearly half the sample cases (45%) are missing the Delta-V measure. There are 24 PSUs selected in the first sampling stage (out of a total of 1195 PSU’s in the population which were grouped into 12 strata based on geographic region and type, with at least two PSU’s selected from each stratum). We regard the stratum effects (at both first- and third-stage) as being reflected in the unequal selection probability or sample weights, and analyze the data using an ultimate-cluster approach with focus on the PSU-level clustering effects. This results in a moderate ICC of 0.1 on Delta-V among the complete cases.

As a preliminary analysis, we examined features of the design variables and the missing data pattern before imputation. The logistic regression coefficients of the response indicator on covariates are mostly significant, suggesting at least a MAR missing data mechanism. Therefore all these covariates are included as predictors for
imputation. A linear regression of final weights on all the survey variables indicates that several covariates as well as the Delta-V measure are strong predictors of final weights. Yet all together they explain only 10% of the variance in weights; therefore incorporating weights in the imputation is essential. Because the estimated population size based on the sum of the weights is in excess of 1 million, we assume for operational simplicity that $N = 80,000$, which is large enough for the proposed method to work properly.

Table 3.8 displays the results of applying different MI methods to the NASS-CDS Delta-V measure. Different modeling strategies are used under each method according to the extent the design information is modeled. Fully parametric imputation that does not include the weight appears to overestimate the mean Delta-V, as would be expected, since more severe crashes are overrepresented in the unweighted sample. When weights are included in the imputation, the fixed effect model $FX_X,W$ tends to result in wider confidence intervals. The FPBB method (SYN2) mimics the performance of the random effects model with weights in the imputation (RE_X,W). Results are broadly similar for estimates of the odds ratios of injury. Here the complete case model appears to underestimate the effect of Delta-V on injury risk, suggesting that injury cases are more likely to have high Delta-V measures missing. The SYN2 approach even enjoys some gains in precision when estimating the odds ratio of having head injury, where all imputation methods yield similar point estimates.
Table 3.8 Estimating mean Delta-V, odds ratio of severe injury given Delta-V, and odds ratio of head injury given Delta-V (high Delta-V=in excess of 35 kph; medium Delta-v=15-35 kph; low delta-V=less than 15 kph). CC=complete case; SRS=imputation under simple random sampling assumption; FX=imputation using fixed cluster effects; RE=imputation using random cluster effects; SYN2=Results from 2-stage finite population Bayesian bootstrap model using Bayesian bootstrap-finite population Bayesian bootstrap for synthetic population generation.

<table>
<thead>
<tr>
<th>Methods</th>
<th>CC</th>
<th>SRS</th>
<th>FX</th>
<th>RE</th>
<th>SYN2</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\bar{Y}_{\text{Delta-V}}$</td>
<td>CC</td>
<td>X</td>
<td>X, W</td>
<td>X</td>
<td>X, W</td>
</tr>
<tr>
<td>SE</td>
<td>0.88</td>
<td>0.88</td>
<td>0.96</td>
<td>1.12</td>
<td>1.05</td>
</tr>
<tr>
<td>OR (Severe Injury \text{high vs. low DV})</td>
<td>5.03</td>
<td>5.78</td>
<td>6.23</td>
<td>5.43</td>
<td>6.38</td>
</tr>
<tr>
<td>95% CI</td>
<td>[2.53, 9.99]</td>
<td>[3.16, 10.6]</td>
<td>[3.37, 11.5]</td>
<td>[2.99, 9.86]</td>
<td>[3.45, 11.8]</td>
</tr>
<tr>
<td>OR (Severe Injury \text{medium vs. low DV})</td>
<td>1.28</td>
<td>1.58</td>
<td>1.52</td>
<td>1.61</td>
<td>1.77</td>
</tr>
<tr>
<td>95% CI</td>
<td>[0.612, 2.68]</td>
<td>[0.838, 2.98]</td>
<td>[0.862, 2.68]</td>
<td>[0.917, 2.83]</td>
<td>[0.994, 3.15]</td>
</tr>
<tr>
<td>OR (Head Injury \text{high vs. low DV})</td>
<td>1.05</td>
<td>1.55</td>
<td>1.45</td>
<td>1.69</td>
<td>1.52</td>
</tr>
<tr>
<td>95% CI</td>
<td>[0.581, 1.90]</td>
<td>[0.76, 3.16]</td>
<td>[0.751, 3.16]</td>
<td>[0.904, 3.54]</td>
<td>[0.765, 3.02]</td>
</tr>
</tbody>
</table>
3.6 Discussion

Two-stage cluster sampling is a popular sampling scheme in survey research because it is cost effective and easy to administer. However, fully parametric MI does not work well with two-stage cluster sampling when sampling units at both stages are selected with unequal probabilities unless the sampling probabilities are properly accounted for. As shown in the simulation study, when model misspecification with respect to the sampling weights is present, both fixed effects and random effects imputations lead to biases in estimates, and hence invalid inferences. As Rabe-Hesketh and Skrondal (2006) put it, dealing with sampling weights in hierarchical/multilevel models can be challenging from either a computational or a modeling standpoint. We propose an alternative two-step MI approach, where the first step generates synthetic populations with missing data that account for weights and clusters, and the second step imputes under an IID assumption.

We propose two different approaches for the two-stage Bayesian nonparametric synthetic data generation from two-stage cluster sampling designs. The first (SYN1) uses a “FPBB-FPBB” approach, generating clusters from a finite-population Bayesian bootstrap and population elements from a finite-population Bayesian bootstrap from those FPBB-generated clusters. The “adapted two-stage Polya posterior” under this approach is different from the “two-stage Polya posterior” of Meeden (1999) in two ways. First, we extend Meeden’s simple two-stage cluster sampling (balanced case) with equal selection probability to an unbalanced case with unequal selection probabilities, for applications in the missing data context. Second, while Meeden considered a single variable ($Y$), we consider the joint distribution of several survey variables ($Y, X, Z, R$).
Our procedure correctly restores the population configuration of an outcome variable \((Y)\) that is built on complex relationships of the regular covariate \((X)\) and the sample design variables \((Z)\). Such population-level multivariate relationships are not guaranteed by Meeden’s procedure (according to results from a side simulation study not reported here). Our second approach (SYN2) uses a “BB-FPBB” procedure, resampling the clusters using a Bayesian bootstrap, and then generating the population elements from a finite-population Bayesian bootstrap. Simulation results show that both approaches are insensitive to either sample size or population structure. However, because SYN1 requires knowing both cluster-level weights and within cluster element-level conditional weights while SYN2 only needs the final weights, we recommend SYN2 for general purposes.

In this chapter, we focus on the mean structure of the imputation model for \(Y\) and use design-based analysis to account for weight and clustering effects after fully parametric MI. In our future research, we will investigate the robustness of the proposed new method in comparison with other MI methods against potentially incompatible normal assumptions on random cluster effects and random errors (Yucel & Demirtas, 2010). We will also consider extensions to stratified, clustered, and unequal probability of selection sample designs.
CHAPTER 4
A SYNTHETIC MULTIPLE IMPUTATION PROCEDURE FOR MULTI-STAGE COMPLEX SAMPLES

4.1 Introduction

Stratified multistage sampling is the most common type of sample design for large-scale surveys conducted by the U.S. federal statistical agencies. Examples of surveys using stratified multistage sample selection include the Current Population Survey (CPS) by the Census Bureau, the National Health Interview Survey (NHIS) and the National Health and Nutrition Examination Survey (NHANES) conducted by the National Center for Health Statistics, and the National Assessment of Educational Progress (NAEP) conducted by the National Center for Educational Statistics. This type of sample design combines the advantages of both stratification (for statistical efficiency) and cluster sampling (for cost and logistical efficiency). It thus may be as precise as a simple random sample or a single-stage stratified sample design, but costs significantly less (Murphy, 2008). Under this design, the primary sampling units (PSUs) are stratified in such a way that they are homogeneous with respect to a stratum-level aggregate of the variable(s) of interest. To permit a maximum degree of stratification and thus variance reduction, it is common practice to define a large number of strata where only a small number of PSUs are selected in each stratum.

On the one hand, such highly stratified multistage sample designs facilitate the data collection process while assuring broad representativeness of the target population.
On the other hand, because of the complexities involved, i.e. complex sample design features including stratification, clustering and unequal selection probability, these sample designs require sophisticated statistical methods at the analysis stage of survey data. When missing data are present, the analysis of complex survey data becomes particularly challenging. Taking the NAEP as an example, missing data may occur at two different levels: 1) survey nonresponse at the PSU level, if some sampled schools fail to participate in the entire survey or some school-level measures are missing; 2) survey nonresponse at the ultimate sampling unit level, if within participating schools, some students fail to provide responses to items in the survey questionnaire. We consider using multiple imputation (MI) (Rubin, 1976, 1987) to deal with item-level missing data at the ultimate sampling unit level. In particular, we consider the role of complex sample designs in the MI procedure.

Reiter, Raghunathan and Kinney (2006) demonstrated the importance of simultaneously accounting for stratum effects and clustering effects in multiple imputation. They showed that when design features were ignored in the imputation model, biases would occur on the estimated parameter, even if a design-based analysis method was applied on the imputed data. Current MI methods typically include dummy variables to represent strata as well as PSUs nested within each stratum in the imputation model. When necessary, they also identify statistically significant interactions between these dummies with other covariates through routine variable selection procedures such as stepwise regression (Reiter et al., 2006; Schenker et al., 2006). Such a modeling strategy is not only operationally burdensome but also inferentially inefficient when there are hundreds of strata in the sample design, and the sample in each stratum consequently
becomes sparse. For example, the Census Bureau’s CPS design groups 1768 nonself-representing PSUs into 220 strata.

A possibly better strategy is to consider clusters as random effects while treating strata as either fixed (using dummies) or random effects. However, many of the popular software packages that implement multiple imputation (e.g. SAS MI procedure, R packages mice or mi, and IVEware) cannot simply be adapted to such a mixed effects approach. While a few recent software modules (such as R package pan and MLwiN module REALCOM-IMPUTE) have started to consider mixed effects or multilevel modeling for imputation purposes, they typically assume normal or latent normal distribution for variables with missing data. Their performances for missing categorical variables (binary in particular) are unclear. Moreover, little research has formally investigated their use to incorporate strata as well as clusters.

To circumvent these problems with fully parametric model-based imputation techniques, we develop a modification of the two-step semi-parametric MI method proposed by Zhou, Elliott and Raghunathan (2013a, 2013b). The idea was to separate the need to account for complex sample designs from the treatment of missing data. In the first step, they reversed the sample designs through synthetic population data generation. They developed different variations of a weighted finite population Bayesian bootstrap (FPBB) (Cohen, 1997; Little & Zheng, 2007; Dong et al., 2014) for untying the sampling weights and clustering effects. In the second step, they imputed missing values in the created synthetic population based on a much simpler imputation model that assumes IID (identically independently distributed). To account for stratum effects, we propose an adapted version of their procedure in this chapter. The new procedure combines a
replication variance estimation method (Efron, 1979; Kovar, Rao, & Wu, 1988; Rao & Wu, 1988; Rao, Wu, & Yue, 1992; Rust & Rao, 1996) with the weighted FPBB. Under a standard missing at random (MAR) assumption (Little & Rubin 2002), our method requires neither complicated modeling of strata and clusters nor design-based analyses of the imputed data.

Although our method is applicable to multiple imputation in general settings, we focus in this chapter on the estimation of two quantities: quantile estimates for a continuous variable, and estimates of rare proportions and their associated logistic regression estimates. We consider a stratified two-stage sample design and investigate a full range of quantiles including tail behaviors. While design-based methods for quantile estimation from complex survey data have been developed (Francisco & Fuller, 1991; Woodruff, 1952), quantile estimation after imputation is rarely addressed in the literature, to our knowledge. This is despite the rapid development and increasing popularity of MI. We also consider MI for incomplete binary variables, with a focus on rare outcomes. It is well known that maximum likelihood estimation of logistic regression models typically suffers from small sample bias, the degree of which is strongly dependent on the number of sample cases in the less frequent of the two categories (King & Zeng, 2001). Thus when the dependent binary variable represents the occurrence of rare events, the logit coefficients can be substantially biased even with a simple IID data structure. Random effects logistic models are commonly used for fitting clustered binary data, however, these models rely heavily on asymptotic theory assumptions, which may not be met in sparse samples. All these issues might extend naturally to the missing data context. As shown by Zhao and Yucel (2009), sequential MI for binary data missing completely at
random in a multilevel setting suffers from severe bias and poor coverage in estimating probabilities that are close to 0 or 1, particularly when the intraclass correlation is high.

The objectives of this paper are: i) to develop an adapted version of the two-step synthetic MI method of Zhou et al. (2013a, 2013b) as a way to account for stratification, in addition to clustering and unequal inclusion probability; and ii) to demonstrate the effectiveness of the new method, with respect to quantile estimation and logistic regression for binary rare events data, as compared with existing fully parametric imputation strategies. Section 4.2 discusses the imputation strategies under three different models: simple random sample, fixed effects for clusters/strata, and random effects for cluster/strata. Section 4.3 introduces the newly proposed procedure and the MI inference rules for quantile estimation under this method. Section 4.4 presents a Monte Carlo simulation study as the validation tool to assess the repeated sampling properties of MI under the various approaches. Section 4.5 applies different MI procedures to the analysis of body mass index on youth data from the third National Health and Nutrition Examination Survey (NHANES III). Some concluding remarks follow in Section 4.6. We focus on the two-PSU-per-stratum design in this chapter, although the methods we develop can accommodate any number of PSUs per stratum.

4.2 Fully parametric imputation methods for the two-PSU per stratum design

Here we briefly describe fully parametric multiple imputation techniques with complex sample design features incorporated to different degrees. We assume the missing data $Y_i$ is a member of the exponential family, and that there are fully observed covariates $X_i$ (a $(p+1)$-dimension vector) such that $g(E(Y_i | X_i)) = X_i \beta$ for a known link function
\[ g(\bullet) \text{ (e.g. } g(u) = \log\left(\frac{u}{1-u}\right) \text{ for binary outcomes (logistic regression), } g(u) = \log(u) \text{ for count outcomes (Poisson regression), or } g(u) = u \text{ for continuous outcomes (Gaussian regression)).} \]

### 4.2.1 Standard regression model assuming SRS

Based on the maximum likelihood (ML) estimates \( \hat{\beta} \) and the associated asymptotic covariance matrix \( \hat{V}(\hat{\beta}) \) for the generalized linear model \( g(E(Y_i | X_i)) = X_i \beta \), the posterior predictive distribution of the parameters can be constructed, which is then used to impute the missing values (Rubin, 1987, pp. 169-170). The steps used to generate imputed values are summarized as follows:

1. At current iteration \( t \), draw new regression parameters \( \beta^{(t+1)} \) (a row vector with length \( (p+1) \)) from their normal approximation posterior predictive distribution \( N(\hat{\beta}, \hat{V}(\hat{\beta})) : \beta^{(t+1)} = \hat{\beta}^{(t)} + \hat{V}_c z \), where \( \hat{\beta}^{(t)} \) is the ML estimator of \( \beta \) based on the observed data \( X \) and \( Y^{\text{obs}} \) together with the filled-in \( Y^{\text{imp}(t)} \), \( \hat{V}_c \) is the upper triangular matrix in the Cholesky decomposition of covariance matrix \( \hat{V}(\hat{\beta}^{(t)}) \), and \( z \) is a vector of \( p+1 \) independent random normal deviates.

2. If the distribution of \( Y_i \) has an unknown scale parameter \( \sigma^2 \), draw \( \delta \sim \chi^2_{n-p-1} \) and compute \( \sigma^{2(t+1)} = \sum_i \left( Y_i - g^{-1}(X_i^T \beta^{(t+1)}) \right)^2 / \delta \).

3. For an observation with missing \( Y_i \), draw \( Y_{i\text{miss}} \) from the assumed distribution with mean \( g^{-1}(X_{i\text{miss}}^T \beta^{(t+1)}) \) and scale \( \sigma^{2(t+1)} \).
Point and variance estimates of the regression parameters can then be obtained using the usual MI combining rules (Rubin, 1987, p. 76). For the $p^{th}$ component of the regression parameter:

$$
\hat{\beta}_p = \frac{1}{M} \sum_{m=1}^{M} \hat{\beta}_{p,m},
$$

[4.1]

$$
\hat{V}(\hat{\beta}_p) = \frac{1}{M} \sum_{m=1}^{M} \hat{V}(\hat{\beta}_{p,m}) + \frac{M+1}{M(M-1)} \sum_{m=1}^{M} (\hat{\beta}_{p,m} - \hat{\beta}_p)^2
$$

[4.2]

and

$$
\left( \frac{\hat{\beta}_p - \hat{\beta}_p}{\sqrt{\hat{V}(\hat{\beta}_p)}} \right) \sim t_{v,v} = (M-1) \left( 1 + \frac{\sum_{m=1}^{M} \hat{\beta}_{p,m}}{(M+1) \sum_{m=1}^{M} (\hat{\beta}_{p,m} - \hat{\beta}_p)^2} \right)^2,
$$

[4.3]

where $m=1,...,M$ imputations are taken from draws widely separated to have practically eliminated autocorrelation. Multivariate combining rules for the joint distribution of $\hat{\beta}$ are available as well (Schafer, 1997a, pp. 112-118).

### 4.2.2 Appropriate fixed effects model (FX_APR)

Compared to the predictive model using standard generalized linear regression, we can add dummy variables indicating stratum and cluster memberships to account for stratification and clustering effects. Note we also need to include the log transformation of sampling weight as a predictor if the missing data mechanism depends on weights, to make the imputation model truly appropriate. The model takes the following form:

$$
g(E(Y_i | X_i)) = X_i\beta + D_i\gamma + E_i\eta + [\zeta \log(w_i)].
$$

[4.4]

where $D_i$ is a $1 \times (H-1)$ row vector of dummies representing the $H$ strata, and $E_i$ is a
$1 \times Q$ row vector of dummies representing the clusters nested within each stratum. Note that $Q = \sum_h Q_h - H$, where $Q_h$ is the number of clusters in each stratum; in the case of the two-PSU per stratum case, $Q = H$. The dummy part of the design matrix thus takes a block diagonal matrix form as follows:

$$
\begin{bmatrix}
X_1 & A_1 & 0 & \cdots & 0 & B_1 & 0 & \cdots & 0 \\
X_2 & 0 & A_2 & \cdots & 0 & 0 & B_2 & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\
X_H & 0 & 0 & \cdots & 0 & 0 & 0 & \cdots & B_H
\end{bmatrix}_{n \times (p + H + Q)}
$$

where $X_h$ is a $n_h \times (p+1)$ matrix containing the values of covariates for observations in stratum $h$, and $\sum_h n_h = n$, $h = 1, \ldots, H$. $A_h$ is a $n_h \times 1$ matrix containing a vector of ones for observations in stratum $h$, and $B_h$ is an $n_h \times (Q_h - 1)$ matrix containing vectors of dummy variables for the $Q_h$ PSUs in the $h^{th}$ stratum. To ensure identifiability, the dummy for the $H^{th}$ stratum and the $Q_h^{th}$ PSU are dropped. The parameter space under this model is expanded as $\theta = (\beta, \gamma, \eta, \zeta)$, and the steps for imputation are similar as in the SRS setting.

### 4.2.3 Appropriate mixed effects model (RE_APR)

As there are only two PSUs selected from each stratum, it is not feasible to model clusters as random effects separately within each stratum. Here we pool all $Q+H$ clusters in the sample, and model them using a single random effect term. The imputation model
is specified as follow:

\[ g \left( E(Y_j \mid X_j) \right) = X_j \beta + D_j \gamma + u_i + [\zeta \log(w_j)] , \]  \hspace{1cm} [4.5]

where \( u_i \sim N(0, \sigma_u^2) \) is a random intercept term representing cluster effects, for \( i = 1, \ldots, (Q + H) \), and \( \sigma_u^2 \) denotes the between cluster variance. Other terms are as previously defined. (In the two-PSU-per-stratum case, \( Q + H = 2H \).) This MI strategy obtains the imputed values in the following steps:

1) At current iteration \( t \), fit the above generalized linear mixed effects regression model using the observed data \( X \) and \( Y^{obs} \) together with the filled-in \( Y^{imp(t-1)} \). The inference about \( \theta^{(t)} = (\theta_0^{(t)}, \sigma_u^{(t)}) \), where \( \theta_0^{(t)} = (\beta^{(t)}, \gamma^{(t)}, \zeta^{(t)}) \), is based on the marginal likelihood function

\[
L = \prod_i \prod_j \int_{\mu_i} \int_{\sigma_u} p(y_{ij}^{(t-1)} \mid \theta^{(t)}) \varphi(u_i; 0, \sigma_u^{(t)}) \, du_i ,
\]

where \( y_{ij}^{(t-1)} \) is either the observed value or the filled-in value from the \((t-1)^{th}\) draw and \( \varphi(\cdot; \mu, \sigma^2) \) is the normal density with mean \( \mu \) and variance \( \sigma^2 \). When \( Y_i \) is non-normal, we employ adaptive Gaussian quadrature (Pinheiro & Bates, 1995) (with the number of quadrature points set as 10) to evaluate the integral and obtain the maximum likelihood estimates for the parameters \( (\hat{\theta}^{(t)}, \hat{\sigma}_u^{2(t)}) \) and the associated variance-covariance matrix of \( (\hat{\theta}^{(t)}, \hat{\sigma}_u^{2(t)})^T \) given by \( \hat{\Sigma}^{(t)} \) through numerical methods.

2) Obtain a draw from the normal approximation to the posterior predictive distribution of these parameters: \( \theta^{(t)} = \hat{\theta}^{(t)} + \begin{bmatrix} V_\phi \\ V_{\sigma_u} \end{bmatrix} \times z \), where \( z \) is a vector containing \( p + Q + H + 1 = p + 2H + 1 \) independent random normal variates,
are components of the cholesky root of $\hat{\Sigma}^{(t)}$ corresponding to the fixed
effects and random cluster effects, respectively.

3) Any residual scale parameter is drawn as in the SRS setting.

4) Finally, the missing values are drawn from the following distribution:

$$Y_{i\text{mis}}^{(t)} | X_{i\text{mis}}, G^{(i)}(\gamma_{j}^{(i)} + u_{j}^{(i)} + \xi^{(i)} \log(w_{i\text{mis}})) \sim g^{-1}(X_{i\text{mis}}, \beta^{(i)} + D_{j} \gamma^{(i)} + u_{j}^{(i)} + \xi^{(i)} \log(w_{i\text{mis}}))$$

Point and variance estimates and 95% confidence intervals are then obtained using the
standard Rubin MI combining rules as previously described.

### 4.3 Synthetic MI using the weighted FPBB for stratified samples

In this section, we extend the two-step multiple imputation methodology proposed
by Zhou et al. (2013a, 2013b) to a stratified two-stage sample design where a
combination of complex sampling techniques are considered, namely, stratification,
clustering and unequal inclusion probability. We develop methods for an unrestricted
number of clusters per stratum, but for our simulations and application we focus on the
special case of two primary sampling units (PSUs) selected per stratum, which mimics
the form of a public use dataset that is commonly released for analyses.

#### 4.3.1 Synthetic data generation to account for complex sample designs

Consider a finite population $P$, which is stratified into $H$ strata with $N_h$ PSUs in
the $h^{th}$ stratum, and hence the population size of PSUs is $\sum_{h=1}^{H} N_h = N$. For the $h^{th}$
stratum, select $n_h$ PSUs with/without replacement from some probability sampling plan,
independently across strata, and hence the total sample size of PSUs is $\sum_{h=1}^{H} n_h = n$. 

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Subsampling of $m_{hi}$ elements (treated as the ultimate sampling units in this example) from a total of $M_{hi}$ is then conducted within the $i^{th}$ sampled PSU of the $h^{th}$ stratum, for $i=1,...,n_{h}, h=1,2,...,H$. Hence the overall sample size and population size of elements are $\sum_{h=1}^{H} \sum_{i=1}^{n_{h}} m_{hi} = m$ and $\sum_{h=1}^{H} \sum_{i=1}^{N_{h}} M_{hi} = M$, respectively, where $m_{hi}$ and $M_{hi}$ are sample size and population size of elements for the $h^{th}$ stratum, respectively. The population consists of four types of survey variables: a single outcome $Y$, a single covariate $X$, a design matrix $Z = [S, C, w]$ including the stratum indicators ($S$), the cluster indicators ($C$) and the sample weight ($w$), and the response indicator $R$. Let $D = (D_{s}, D_{ns}) = \{ (Y_{hi}, X_{hi}, Z_{hi}, R_{hi}), h=1,...,H, i=1,...,N_{h}, j=1,...,M_{hi} \}$ denote the population of values measured on the survey variables, which is divided into the sampled component ($D_{s}$) and the nonsampled ($D_{ns}$) component.

We generate synthetic populations using a two-stage procedure. The first stage accommodates stratification and clustering, and the second weighting. We have two broad approaches. The first, which we term SYN1, assumes that first (cluster-level) and second (PSU-level) stage sample weights are available for the analysis, and implements a weighted finite population Bayesian bootstrap (FPBB) at each level to generate the synthetic population. The second, which we term SYN2, assumes that only final weights are available for the analysis, and uses a Bayesian bootstrap to account for stratification and clustering at the first stage, and the weighted FPBB to account for the final weight at the second stage.

4.3.1.1 Double Weighted Finite Population Bayesian Bootstrap (SYN1)

For the $h^{th}$ stratum, let $t_{s,h}$ and $t_{ns,h}$ index the sampled and nonsampled clusters,
respectively, and \( \{b^1, \ldots, b^q, \ldots, b^q, q = 1, \ldots, r_h \} \) be the \( r_h \) \((1 \leq r_h \leq N_h)\) distinct matrices of real numbers each of dimension \( |b^q_{row}| \times |b^q_{col}| \) with no row vectors in common. Each cluster in the stratum can take the form of one of \( b^q \)'s. Let \( t_{hi} = q \) when the \( i^{th} \) cluster takes on the values of \( b^q \), for \( i = 1, \ldots, N_h \). Assume \( n_h = r_h \) and \( m_h = \|b^q\| \) (the number of distinct row vectors in \( b^q \)) for convenience of exposition. Let \( w_{t_{hi}}(i) \) be the sample weight of the \( i^{th} \) sampled cluster in the \( h^{th} \) stratum which equals \( b^q \), for \( i = 1, \ldots, n_h \), and \( w_{t_{hi},D_{ih}}(j) \) be the sample weight of the \( j^{th} \) sampled element in the \( i^{th} \) sampled cluster which equals \( b^q \), for \( j = 1, \ldots, m_h \). Finally, let \( c_{t_{hi}}(q) \) and \( c_{t_{ih},D_{ih}}(k) \) be the number of sampled and nonsampled clusters that equal \( b^q \), and \( c_{t_{hi},D_{ih}}(k) \) be the number of sampled and nonsampled elements that equal \( b^q \).

Zhou et al. (2013b) showed that, within a stratum \( h \), the Polya posterior for the counts of distinct unobserved elements \( D_{ns,h} \) is given by

\[
p(D_{ns,h} \mid D_{s,h}) = \left\{ \prod_{q=1}^{s} \left\{ \Gamma(w^q_{t_{hi}}(q)) / \Gamma(w_{t_{hi}}(q)) \right\} \right\} / \{ \Gamma(N_h) / \Gamma(n_h) \} \\
\times \left\{ \prod_{k=1}^{m_h} \left\{ \Gamma(w_{t_{hi},D_{ih}}(k)) / \Gamma(w_{t_{hi},D_{ih}}(k)) \right\} \right\} / \{ \Gamma(M_h) / \Gamma(m_h) \}
\]

where \( w_{t_{hi}}(q) = w_{t_{hi}}(q) + c_{t_{ih}}(q) \) and \( w_{t_{hi},D_{ih}}(k) = w_{t_{hi},D_{ih}}(k) + c_{t_{ih},D_{ih}}(k) \), for

\[m_h = \sum_{k=1}^{m_h} c_{t_{ih}}(k) \] and \( m^h = M_h - m_h = \sum_{k=1}^{m_h} c_{t_{ih}}(k) \). The full posterior is then given by the product of the posteriors within each stratum, since these strata are independent and all strata in the population are in the sample:
\[ p(D_{ns} | D_s) = \prod_{h=1}^{H} p(D_{ns,h} | D_{s,h}) . \] \[4.7\]

A Monte Carlo procedure to simulate from this posterior distribution is then given as follows:

(i) Draw the \( N_h - n_h \) nonsampled clusters in the population based on the Polya posterior distribution independently for each stratum. Each of the sampled clusters is resampled with probability

\[
\zeta_{h,i} = \frac{w_{n,h} (i) - 1 + l_{n,h,k-1} \times \left( \frac{N_h - n_h}{n_h} \right)}{N_h - n_h + (k-1) \times \left( \frac{N_h - n_h}{n_h} \right)}, \quad k = 1, ..., N_h - n_h + 1 , \tag{4.8}
\]

where \( l_{n,h,k-1} \) is the number of times that the \( i^{th} \) cluster in the \( h^{th} \) stratum has been resampled at the \((k-1)^{th}\) resampling, and \( w_{n,h} (i) \) is the weight for the \( i^{th} \) sampled cluster in the \( h^{th} \) stratum which is normalized to sum up to the total number of clusters, i.e. \( \sum_{i=1}^{n_h} w_{n,h} (i) = N_h \).

(ii) From Step 1, form a population of clusters

\[ \{ c_{i1}, c_{i2}, ..., c_{in_h}, c_{i1}^*, c_{i2}^*, ..., c_{iN_h-n_h}^*, ..., c_{H1}, c_{H2}, ..., c_{Hn_h}, c_{H1}^*, c_{H2}^*, ..., c_{HN_H-n_n}^* \} . \]

Record the number of times each of the clusters from the original sample appears in the FPBB population of clusters, denoted by \( \tau_{hi}, i = 1, ..., n_h, h = 1, ..., H, \) and

\[ \sum_{h=1}^{H} \sum_{i=1}^{n_h} \tau_{hi} = N . \]

Then update the within cluster element-level conditional weights as follows: \( w_{j|hi}^* = w_{j|hi} \times \tau_{hi}, i = 1, ..., n_h, h = 1, ..., H, \) where \( w_{j|hi} \) is the inverse of the conditional probability that element \( j \) is selected given cluster \( i \) in stratum \( h \) is selected. Now pool all elements from these clusters together and treat
them as a single **FPBB sample** (i.e., as if they have no stratum or cluster boundaries). Note that this FPBB sample has the same sample size

\[
m = \sum_{h=1}^{H} \sum_{i=1}^{n_h} m_{hi}
\]

but different sampling weights than the original sample. We then apply the weighted FPBB again to these pooled elements to generate \(M - m\) units from the \(m\) units in the FPBB sample. We resample from each of the resampled clusters \(M - m\) elements, cycling through \(M - m\) times and resampling with probability

\[
\lambda_{jhi} = \frac{w_{jhi}^* - 1 + l_{hij,k-1} \times \left(\frac{M-m}{m}\right)}{M - m + (k-1) \times \left(\frac{M-m}{m}\right)}, \quad k = 1, \ldots, (M-m+1), \quad [4.9]
\]

where \(l_{hij,k}\) is the number of times that the \(j^{th}\) element in the \(i^{th}\) cluster in the \(h^{th}\) stratum has been resampled at the \(k^{th}\) resampling, and \(w_{jhi}^*\) is the updated conditional weight for the \(j^{th}\) element in the \(i^{th}\) cluster in the \(h^{th}\) stratum. Again, they are normalized to sum up to the total number of units in the entire population, i.e. \(\sum_{h=1}^{H} \sum_{i=1}^{n_h} \sum_{j=1}^{m_{hi}} w_{jhi} = M\). Thus we create a single synthetic population.

Repeat step 2 \(B\) times to obtain \(B\) FPBB synthetic populations.

**(iii)** Repeat steps 1-2 \(L\) times to obtain \(L\) bootstrap samples, yielding \(L \times B\) FPBB populations \(P_{(lb)}^{mn} = (P_{(lb)jdn}^{mn}, P_{(lb)mub}^{mn}), l = 1, \ldots, L, b = 1, \ldots, B,\) each of which consists of both responding elements and nonresponding elements on a vector of variables \(\{Y, X, Z, R\}\).
4.3.1.2 Bootstrap — Weighted Finite Population Bayesian Bootstrap (SYN2)

Because we often do not know the first- and second-stage weights in public-use dataset, we consider an alternative to the procedure proposed in subsection 4.3.1.1. Rather than obtaining a sample of clusters from a draw from a Polya posterior, we use replication methods (Rust & Rao, 1996) to capture the cluster-level sampling variance. The final sampling weights instead of the adjusted element-level conditional weights are then directly used as input in the second-stage weighted FPBB. We use Rao and Wu (1988)’s rescaling bootstrap, which is a generalized extension of McCarthy and Snowden (1985)’s “with replacement bootstrap”. The reason for choosing replication methods, particularly this special type of bootstrap, is three-fold: 1) replication methods in general are simple and direct to implement, and the formation of replicate samples further creates replicate weights that inherently represent the effects of a complex sample design in addition to reflecting the effects of a wide range of reweighting techniques such as calibration weighting; 2) among various replication methods, while the standard delete-1 jackknife is known to give an inconsistent variance estimator for a quantile as a classic example of nonsmooth statistics, the bootstrap yields sensible estimates for a variety of estimators. In fact, for stratified multistage samples, both consistency of the bootstrap variance estimators and confidence intervals for both smooth statistics and nonsmooth statistics have been established by researchers. Examples include functions of sample means by Rao and Wu (1988), and sample quantiles and sample low income proportions by Shao and Chen (1998). Asymptotic consistency of the Balanced Repeated Replication (BRR) and the bootstrap has also been established by Shao and Wu (1992) and Rust and Rao (1996). The bootstrap is adopted here because its application extends readily to more PSU sample allocations other than two-PSU-per-stratum design; 3) The Rao-Wu variant
of the conventional bootstrap yields adequate and stable variance estimates when the sample sizes are small, which occurs most often with stratified multistage sampling where only a small number of PSUs are selected within each stratum. Once the PSUs have been sampled, we continue with the weighted FPBB approach to complete the synthetic population data generation. The proposed procedure is as follows:

(i) Select a sample of $n_h^* = n_h - 1$ PSUs from the parent sample in each stratum via SRSWR sampling;

(ii) Apply the “ultimate cluster principle” (Wolter, 2007), that is, once a PSU is taken into the bootstrap replicate, all elements in that PSU are taken into the replicate also. Thus, we obtain our first bootstrap sample;

(iii) Repeat the previous steps $L$ times to obtain $L$ bootstrap samples

$$\{Boot_l, l = 1, ..., L\};$$

(iv) Within each bootstrap sample, update the element-level sampling weights as:

$$w_{hij}^* = w_{hij} \times \left(\frac{n_h}{n_h - 1}\right)^{n_h - 1}_h$$

where $w_{hij}$ is the weight of the $i$th PSU in the $h$th stratum.

As $w_{hij}$ itself implicitly carries over the strata and PSU information in addition to unequal inclusion probability, we can drop the subscripts $hi$ henceforth by pooling all elements in the bootstrap sample regardless of which stratum and PSU they originally came from. Normalize $w_j^*$’s to sum up to $m^*$:

$$\sum_{j=1}^{m^*} w_j^* = m^*,$$

where $m^*$ is the bootstrap sample size.

(v) For the $l$th bootstrap sample, $l = 1, ..., L$, apply the weighted FPBB algorithm to create an entire population $D = (D_{ns}, D_x^*)$ based on the posterior predictive
distribution of elements in the nonsampled population

\[ D_{ns} = \{ (Y_j, X_j, Z_j, R_j), j = m^* + 1, \ldots, M \} \]

given the elements in the bootstrap sample \( D^*_s = \{ (Y_j, X_j, Z_j, R_j), j = 1, \ldots, m^* \} \):

\[
p(D_{ns} | D^*_s) = \frac{\prod_{j=1}^{K} \lambda_j^{w_j^*+r_j-1} \left( 1 - \sum_{j=1}^{K-1} \lambda_j \right)^{w_j^*+r_j-1} \prod_{j=1}^{K-1} d\lambda_j \cdots d\lambda_{K-1}}{\Gamma(M) / \Gamma(m^*)},
\]

where \( r_j \) is the number of elements in the nonsampled FPBB population that take on the same value as the \( j^{th} \) element in the bootstrap sample, for \( j = 1, \ldots, K(\equiv m^*) \). Operationally, we draw a Polya sample of size \( M^* = M - m^* \) from \( \text{mult}(M^*; \lambda_1, \ldots, \lambda_K) \) where the selection probability \( \lambda_k, k = 1, \ldots, K. \) is a function of \( w_j^* \):

\[
\lambda_k = \frac{w_j^* - 1 + 1_{j,k-1} \times \left( \frac{M^*}{m^*} \right)}{M^* + (k-1) \times \left( \frac{M^*}{m^*} \right)}, \quad k = 1, \ldots, M^* + 1.
\]

Repeat Step (v) for \( B \) times to obtain \( L \times B \) FPBB populations.

### 4.3.2 Imputation of the synthesized populations

Once the set of FPBB synthetic populations \( P^{syn} = \{ P^{(l)}_{(b)}, l = 1, \ldots, L, b = 1, \ldots, B \} \),

where \( P^{(l)}_{(b)} = (Y^{(l)}_{(b)_{\text{mix}}}, Y^{(l)}_{(b)_{\text{obs}}}) \) are created using either SYN1 method or SYN2 method, we generate imputations \( P^{imp} = \{ P^{(l)}_{(b)_{a}}, l = 1, \ldots, L, b = 1, \ldots, B, a = 1, \ldots, A. \} \) from the posterior predictive distribution \( p(Y^{(l)}_{(b)_{\text{mix}}} | P^{(l)}_{(b)_{\text{obs}}}) \) based on a parametric model that does not
condition on sample design features, i.e. a model taking similar form as the SRS model stated in section 4.2.1. We consider imputations based on the covariate \( X \) only (SYN1_srs or SYN2_srs) or imputations that include the log of the sample weights in the linear predictors (SYN1_lwt or SYN2_lwt).

We obtain the MI inference by applying the combining rules developed in Zhou et al. (2013a) to the \( L \times B \times A \) estimates, based on both the observed set
\[
P_R = \{ P_{(b)obs}^{(l)}, b = 1, ..., B, l = 1, ..., L \}
\]
and the imputed set
\[
P_R = \{ Y_{(b)mis}^{(l)}, l = 1, ..., L, b = 1, ..., B, a = 1, ..., A \}
\]
of the synthetic populations, where \( R \) and \( \bar{R} \) represent responding and nonresponding, respectively. We estimate the population mean of \( Y \) by calculating the mean of the synthetic population
\[
\hat{Y}_{lba} = \frac{\sum_{i \in P_R} y_i + \sum_{j \in P_R} y_j}{M}, \tag{4.12}
\]
To estimate a generalized linear regression parameter, we solve the regression score function
\[
U(\beta)_{lba} = [\sum_{i \in P_R} x_i^T (y_i - g^{-1}(x_i^T \beta)) + \sum_{j \in P_R} x_j^T (y_j - g^{-1}(x_j^T \beta))], \tag{4.13}
\]
where \( g(\cdot) \) is the link function for the transformation of the mean, so that
\[
\hat{\beta}_{lba} = \{ \beta : U(\beta)_{lba} = 0 \}.
\]
For quantile estimation, we proceed by first obtaining the empirical distribution function based on the \( lba^{th} \) imputed synthetic population:
\[
\hat{F}_{lba}(y) = \frac{\sum_{i \in P_R} I(y_i < y) + \sum_{j \in P_R} I(y_j < y)}{M}, \tag{4.14}
\]
Then we estimate the \( \gamma^{th} \) quantile (\( q_\gamma \)) as:
\[
\hat{q}_{\gamma,lba} = (\hat{F}_{lba})^{-1}(\gamma) = \inf\{ y : \hat{F}_{lba}(y) \geq \gamma \}, \tag{4.15}
\]
The MI point estimator for the population statistic of interest \( Q \) (mean, regression
estimator, quantile) is then given by the mean of the $lba$th point estimators:

$$\hat{Q}_{MI} = \frac{1}{LBA} \sum_i \sum_b \sum_a \hat{Q}_{iba},$$  \hspace{1cm} [4.16]

The MI variance estimator is:

$$\hat{V}_{MI} = (1 + L^{-1})V_L = (1 + L^{-1}) \frac{1}{L-1} \sum_t (\hat{Q}_t - \hat{Q}_{MI})^2,$$

$$\text{where } \hat{Q}_t = \frac{1}{BA} \sum_b \sum_a \hat{Q}_{iba}, \hspace{1cm} [4.17]$$

We then construct the 95% interval estimate for quantiles based on $t$ reference distribution with degrees of freedom equal to $\min\{v_{com} = \sum_h n_h - H, v_{syn} = L - 1\}$. These results arise from the fact that, by the standard Rubin (1987) MI combining rules, we have

$$Q|\text{P}^{\text{imp}} \sim t_{L-1}(\bar{Q}_L, (1 + L^{-1})V_L),$$  \hspace{1cm} [4.18]

where $\bar{Q}_L = \frac{1}{L} \sum_t \check{Q}^{(i)}$, $V_L = \frac{1}{L-1} \sum_t (\check{Q}^{(i)} - \bar{Q}_L)^2$, and $\check{Q}^{(i)} = \lim_{BA \to \infty} \frac{1}{BA} \sum_b \sum_a \hat{Q}_{iba}$. Replacing $\check{Q}^{(i)}$ with its finite simulation estimator $\hat{Q}_t$ replaces $\bar{Q}_L$ with $\hat{Q}_{MI}$ and gives the results above.

Note that the generation of the synthetic population sets the within imputation variance to 0 so that the posterior variance of $Q$ can be obtained using the between-bootstrap variance only; see Dong et al. (2014) and Zhou et al. (2013a). The result assumes that $E(\hat{q}_{iba}) = Q$ – a result guaranteed by our Bayesian bootstrap estimator if the imputation model is also correct – as well as a sufficiently large sample size for the $t$ approximation to be reasonable.

Lo (1988) showed that the variance estimator for the FPBB mean in a simple random sample setting should be inflated by the factor $(\frac{n+1}{n-1})$. Thus in the double-weighted FPBB (SYN1) setting, a small sample correction to the variance estimate needs
to be used when the number of clusters per stratum is small. When \( n_h = a \) is a constant across all strata, we use \( \frac{n_h + 1}{n_h - 1} (1+L^{-1}) V_L \); otherwise we suggest \( \frac{\tilde{n}_h + 1}{\tilde{n}_h - 1} (1+L^{-1}) V_L \), where

\[
\tilde{n}_h = H^{-1} \sum_h n_h.
\]

4.4 Simulation Study

We conducted a simulation study to investigate the performance of the proposed method for incorporating stratified cluster sampling effects in multiple imputation. We targeted three population statistics: 1) population quantiles, 2) proportions of binary event data, and 3) logistic regression parameters relating the covariate to the binary data. The simulation is a 2×2 factorial design based on the following factors: 1) keeping the first stage sampling plan constant, we let the subsampling rate \( f_2 \) of elements within sampled clusters be a) independent of or b) dependent on the stratum effects, and 2) assume a) the missingness on the \( Y \)-variable (continuous or binary) depends only on the covariate \( X \) (MAR_X), or b) depends on both \( X \) and the final sampling weight \( W \) (MAR_X,W).

We focus on a two-PSU-per-stratum sample design, both because it is a common design, especially in public use settings, and because it is a “limiting case” in terms of the number of PSUs per stratum. In addition to the two variants of our synthetic MI estimators, we consider standard parametric MI under the SRS, appropriate fixed effect (FX_APR), and appropriate random effect (RE_APR) models.

4.4.1 Description of the Design

- Data generation
Let $i$ be the index for strata, $j$ be the index for clusters, and $k$ be the index for elements. Suppose there are 50 strata in the population. First, the number of PSUs in each stratum was randomly determined according to a uniform distribution, i.e. $C_i \sim \text{Unif}(2,54), i = 1,...,50$; second, the number of population elements within PSUs was randomly generated as $N_{ij} \sim \text{Unif}(20,80), i = 1,...,50, j = 1,...,C_i$. Thus we obtained a population of size $N = 67385$. The complete data for four survey variables $Y = (Y_1, Y_2, Y_3, Y_4)^T$ were generated from a super-population model according to a two-step process: In the first step, $Y_1$ and $Y_2$ were randomly selected from a bivariate linear mixed effects model; let $N_2(\cdot)$ denote a bivariate normal distribution function:

$$
\begin{pmatrix}
Y_{1ijk} \\
Y_{2ijk}
\end{pmatrix} \sim N_2(\mu, \Sigma), \quad \mu = \begin{bmatrix}
\beta_1 + S_i + u_{1ij} + e_{1ijk} \\
\beta_2 + u_{2ij} + e_{2ijk}
\end{bmatrix}, \quad \Sigma = \begin{bmatrix}
\sigma_{11} & \sigma_{12} \\
\sigma_{12} & \sigma_{22}
\end{bmatrix},
$$

[4.19]

Let $\beta_1 = \beta_2 = 15$ be the fixed covariate effects, $S_i = \frac{i}{5}$ be the fixed stratum effects, and let 

$$
\begin{bmatrix}
u_{1ij} \\
u_{2ij}
\end{bmatrix} \quad \text{and} \quad \begin{bmatrix}e_{1ijk} & e_{2ijk}
\end{bmatrix}^T
$$

be the random cluster effects and random error terms drawn from two independent bivariate normal distributions: $N_2 (0, \Sigma_u)$ and $N_2 (0, \Sigma_e)$. Elements of $\Sigma_u$ were set as: $\sigma_{u_1}^2 = 4, \sigma_{u_2}^2 = 4, \sigma_{u_1u_2} = 0.2$, and elements of $\Sigma_e$ were set as: $\sigma_{e_1}^2 = 4, \sigma_{e_2}^2 = 3, \sigma_{e_1e_2} = 1.732$. This results in conditional intraclass correlations (ICC) of $Y_1$ and $Y_2$ as $\rho_{Y_1} = 0.5$ and $\rho_{Y_2} = 0.25$ (note that the unconditional ICC for the two variables may be smaller than these values). In the second step, a random effects logistic regression model (Anderson & Aitkin, 1985; Stiratelli, Laird, & Ware, 1984) was used to simulate two binary outcome variables $Y_3$ and $Y_4$ as a function of $Y_2$. Under this model, a random effect is added to the linear part of the logistic regression model for each element.
in the cluster. The conditional mean of \( Y_{3ijk} \) and \( Y_{4ijk} \) is

\[
\pi_{ijk} = E(Y_{ijk} | Y_{2ijk}, u_{ij}) = \Pr(Y_{ijk} = 1 | Y_{2ijk}, u_{ij}) = \frac{e^{\alpha_0 + \alpha_1 Y_{2ijk} + u_{ij}}}{1 + e^{\alpha_0 + \alpha_1 Y_{2ijk} + u_{ij}}},
\]

[4.20]

where \( u_{3ij} \sim N(0, 6^2) \), \( u_{4ij} \sim N(0, 10^2) \) and \( \alpha = (\alpha_0, \alpha_1, \alpha_2)^T \) is the vector of fixed covariate effects. We fixed \( \alpha_2 = 1.5 \) and vary \( \alpha_0 \) and \( \alpha_1 \) to obtain two different binary variables \( Y_{3ijk} \) and \( Y_{4ijk} \), with either moderate (\( \alpha_0 = -5, \alpha_1 = -1.5 \)) or rare probabilities (\( \alpha_0 = -8, \alpha_1 = -6 \)). Given \( u_{ij} \), the \( Y_{ijk} \)'s in the cluster are independent Bernoulli variables, that is, \( Y_{ijk} \mid u_{ij} \sim \text{Bern}(\pi_{ijk}). \)

Figure 4.1 shows the correlations among variables in the simulated population, where the different shades of grey represent different degrees of association between any of the two variables. The darker shades indicate higher correlation. All survey outcome variables (\( Y_1, Y_3, Y_4 \)) have moderate to strong (0.2~0.8) stratum effect (\( H \) or \( \text{strID} \)) and clustering effect (\( U_1, U_3, U_4 \)), indicating that accounting for these effects in the analysis of missing data is essential.
The Sample Design

Within each stratum, we drew a two-stage cluster sample according to the following procedure: first, we drew a sample of 2 PSUs without replacement with probability proportional to the cluster size \( f_{ij} = \frac{2 \times N_{ij}}{\sum_j N_{ij}} \); second, we sampled elements from each sampled cluster by two different subsampling schemes: 1) sampling probability independent of \( S_i \) which was defined in [4.19]: SRS with an equal sampling fraction of \( f_{2kij} = 1/5 \); 2) sampling probability related to \( S_i \): SRS with varying sampling fractions across strata, i.e. \( f_{2kij} = \text{expit}(-0.8 - 0.12*S_i) \), where \( \text{expit}(x) = 1/(1+e^{-x}) \).

An average of 1122 elements are selected in each of the 200 simulation replications. The distributions of sampling weights are shown in Figure 4.2. The distributions of sampling weights under the two subsampling schemes are generally very similar with somewhat more skewness under subsampling scheme 2.
Throughout the simulation study, we assume that \( Y_2 \) is always completely observed and we impose missing values on \( Y_1, Y_3 \) and \( Y_4 \) independently according to the following deletion function conditional on \( Y_2 \) and/or log transformation of the weight:

\[
\Pr(R = 0 \mid Y_2, W) = \frac{\exp(\lambda_0 + \lambda_1 Y_2 + \lambda_2 \log(W))}{1 + \exp(\lambda_0 + \lambda_1 Y_2 + \lambda_2 \log(W))},
\]

where \( R \) is the response indicator and \( W \) is the overall sample weight. Setting \( \lambda_2 = 0 \), we obtain the first MAR mechanism (i.e. MAR_X, note we treat \( Y_2 \) as \( X \) here), under which we further set \( \lambda_0 = 3.42, \lambda_1 = -0.2 \) and \( \lambda_0 = -2.58, \lambda_1 = 0.2 \) for deleting values on \( Y_1 \) and \( Y_3, Y_4 \), respectively. Setting \( \lambda_2 = -0.6 \), we obtain the second MAR mechanism (i.e. MAR_X,W), under which we fix \( \lambda_1 = 0.2 \) and set two values on \( \lambda_0 (= -0.274 \) or \(-0.33) \).

**Imposing missingness**

Figure 4.2 Distribution of weights under the two subsampling schemes
for deleting values on all three outcome variables under subsampling scheme 1 and subsampling scheme 2, respectively. All deletion functions result in approximately 40% missingness.

- **Parametric Multiple Imputation**

  Both simple random sample SRS (including SRS, SYN1_srs and SYN2_srs) and fixed-effect model FX_APR can be implemented in R (R Core Team, 2013) using mice routines; for the logistic model associated with the binary outcome, the method ‘logreg’ must be specified. We use the pan package (Schafer, 1997b) in R for the mixed effects imputation (RE_APR) for the missing continuous outcome; logistic mixed effects imputation is programmed in SAS for the missing binary outcome, as there is no package readily available for use.

- **Parameters of interest and inference**

  We focus on inference for the following population parameters: the mean of the continuous variable $Y_1$, the mean of the binary variables $Y_3$ and $Y_4$ (i.e. Bernoulli proportions), linear regression coefficients of $Y_1$ on $Y_2$, logistic regression coefficients of $Y_3$ (or $Y_4$) on $Y_2$, and the population percentiles of the continuous variable $Y_1$.

  Weighted analyses and sandwich variance estimators accounting for strata and clusters are used to estimate smooth statistics (including proportions and regression parameters) under the three fully parametric MI methods. For estimating quantiles of the distribution of a continuous survey variable, we construct the sample-weighted point estimator with confidence intervals based on the test inversion method (Francisco & Fuller, 1991). We chose the test-inversion method instead of Woodruff’s method (Woodruff, 1952) despite the computational intensity, because the literature suggests that
it may outperform Woodruff in heavily stratified samples or in small-to-moderate-sized samples (Kovar et al., 1988), and these are the sample designs we are particularly interested in in this chapter. Based on the $a^{th}$ imputed dataset, the empirical distribution function can be written as

$$
\hat{F}^{(a)}(y) = \left[ \sum_{S_{\pi}} w_{hij} I(y_{hij}^{obs} < y) + \sum_{S_{\pi}} w_{hij} I(y_{hij}^{(a)} < y) \right] / \sum_{S} w_{hij}, \tag{4.22}
$$

where $S_{\pi}$ and $S_{\pi}'$ are subsets of the sample data $S$, consisting of respondents and nonrespondents, respectively. The estimator $\hat{F}(y)$ and its associated estimated variance $\nu(\hat{F}(y))$ can then be obtained using the variance estimator proposed by Francisco and Fuller (1991) together with standard Rubin combining rules as previously described. The sample $\gamma^{th}$ quantile estimator thus is $\hat{q}_{\gamma} = (\hat{F})^{-1}(\gamma)$, with 95% asymptotic confidence interval given by

$$
[L, U] = \left[ \hat{F}^{-1}\left( \gamma - t_{0.025} \sqrt{\text{var}(\hat{F}(q_{\gamma}))} \right), \hat{F}^{-1}\left( \gamma + t_{0.025} \sqrt{\text{var}(\hat{F}(q_{\gamma}))} \right) \right]. \tag{4.23}
$$

### 4.4.2 Results

Figures 4.3 through 4.6 plot the point estimates as well as their upper and lower confidence interval lines for 19 population quantiles (from 0.05 to 0.95 with an increment of 0.05). These were obtained from the two proposed finite population Bayesian bootstrap procedures (SYN1 and SYN2). These graphs demonstrate visually how the proposed methods work in incorporating complex sample design features for estimating population quantiles. From left to right in each figure, we compare the proposed methods in the absence of missing data (synthetic BD), the proposed methods in the presence of missing data under mechanism MAR_X (synthetic ADX) and under MAR_X,W.
(synthetic ADXW), with the design-based quantile estimation method based on the actual replication samples (Complete Data). The perfect overlapping of the red plot and black plot indicates that all point and interval estimates obtained from the synthetic BD (both SYN1 and SYN2) are identical to those using the design-based method. This provides good evidence that both proposed procedures were able to accommodate all sample design features to produce synthetic populations that behaved as simple random samples from the underlying true population. In the missing data setting, the point estimates are generally very close. The variance estimates increased as expected — note the green dashed lines always encompass the black dashed lines — due to the added noise from multiply imputing missing data. Hence, we further investigate the performance of the synthetic MI in comparison with fully parametric MI methods, by looking at several key summary measures under the four simulation conditions in Table 4.1.

Table 4.1 compares the average width \( \times 10^2 \) and average coverage rates of the 95% CI of \( q(\alpha) \), where \( \alpha = 0.05, 0.10, 0.25, 0.50, 0.75, 0.90 \) and 0.95, corresponding to seven selected population quantiles. Among all methods considered, the SRS imputation model yields the poorest coverage. This results from the compounding effects of biases and variance underestimation, due to ignoring stratum effects and clustering effects, respectively. As we increase the dependence of both the sampling mechanism and response mechanism on stratum effects and sampling weights, the performance of SRS gets even worse, as exhibited by the markedly increased RelBias and decreased coverage rates. In addition, ignoring stratum and/or weight effects that are highly relevant to either mechanism seems to impact the median and 2\(^{nd}\) and 3\(^{rd}\) quartiles more than the tail quantiles under SRS, as evident in the relatively lower coverage rates in the right block of
Table 4.1.

The FX_APR model (Reiter et al., 2006; Rubin, 1996; Schenker et al., 2006), generally performs fairly well in our simulation study with respect to estimation of population quantiles. There is some modest underestimation of the small percentile quartiles with the second stage sampling constant. The RE_APR model also performs well, with the exception of moderate to high overcoverage when the second stage sampling probability is associated with the stratum mean and the missingness mechanism.

In contrast, our synthetic MI (SYN2 in particular) compares favorably with all of its competitors, and in most cases yield comparable results to the RE_APR, which is regarded as a “gold standard” as it is compatible with the data generating mechanism. There is some undercoverage when the stratified double-weighted FPBB estimator (SYN1) is used, perhaps due to the fact that the Lo small-sample adjustment is not as accurate when $n_b = 2$. However, use of a stratified bootstrap-weighted FPBB estimator (SYN2) generally eliminates this issue. Although an imputation model assuming SRS suffices for the synthetic MI method in most scenarios, we need to include the sampling weight as a predictor when the outcome $Y$ and the response indicator $R$ are strongly associated with each other through the sampling mechanism $I$, as is the case with the second subsampling scheme, when both the missingness indicator and the second-stage sampling rate are functions of the stratum mean.

Tables 4.2 and 4.3 compare the absolute relative bias

$$
relbias = 100 \times \left| \frac{\hat{\theta} - \theta_{\text{complete}}}{\theta_{\text{complete}}} \right| \% ,
$$

RMSE and 95% nominal CI coverage for the estimated mean/proportions of $Y_1$, $Y_3$ and $Y_4$, and the slopes of the three outcome variables on $Y_2$. 

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respectively. \(\hat{\theta}_{\text{complete}}\) is the estimated parameter with complete data, and \(\hat{\theta}\) is the estimated parameter under one of the different MI methods.) As in the estimation of the quantiles, the SRS imputation model is biased and has poor coverage as it ignores stratum and cluster effects. Again, dependence of subsampling on stratum effects and dependence of response on sampling weights damage the performance of SRS even further.

**FX_APR** generally performs well in estimating the mean of a continuous variable \((Y_1)\) and a regular binary variable \((Y_3)\) with moderate probability as well as the slopes. However, it fails for proportion estimation for rare events data \((Y_4)\), yielding biased point estimates and less than nominal coverage throughout all scenarios. One interpretation might be that overfitting occurs when including too many dummies to account for fixed strata and cluster effects. This damages the predictive efficacy when the fitted model is used for drawing missing values. The problem is particularly prominent when the logistic fixed effects imputation model is used along with the current sampling design where an average of only 10 elements are selected per PSU within each stratum; and this results in even more substantial biases on \(\hat{Y}_4\) than the SRS model.

Compared with **FX_APR**, **RE_APR** avoids the overfitting issue through shrinkage effects: note that under **RE_APR**, we pooled all PSUs from all strata as if there were no strata bounds, and the stratum effects can be thought as being implicitly modeled in the random intercept term \(u_j = I_h + u_{h(j)}\). This in some sense alleviates the small sample issue though not a complete solution (e.g. the estimated \(\hat{Y}_4\) under **RE_APR** is still moderately biased).

As in the quantile estimation setting, our synthetic MI compares favorably with
all of its competitors, and in most cases yields comparable results to the RE_APR for estimation of means and logistic regression parameters. In the case of rare events data, our proposed new method increases the analytical size through generating synthetic population data thus is even superior to RE_APR, consistently yielding negligible biases and close to nominal coverage. The impact of ignoring the weights in the imputation (under MAR_X,W mechanism) is less than in the quantile estimation setting, with the exception of the estimation of the continuous mean $\bar{Y}$, where including the weight is required to obtain approximately correct coverage.

A disadvantage of the method lies in the relative inefficiency for estimating nonlinear parameters (regression coefficients) (e.g. the synthetic MI results in unbiased point estimates but a larger RMSE than the two model-based MI methods). This is typical in that nonparametric methods cannot typically compete with their fully parametric counterparts under the correct model, and is a tradeoff made to improve robustness to model misspecification, and, in our setting, simplicity in implementation.
Figure 4.3 Comparison of point and interval estimation for 19 population quantiles using the Stratified Boot-FPBB and the design-based complete data analysis for subsampling scheme 1 ($f_2 \propto \text{constant}$). (a) before deletion (b) imputation based on covariates under MAR_X (c) imputation based on covariates and sampling weights under MAR_X,W.

Figure 4.4 Comparison of point and interval estimation for 19 population quantiles using the Stratified Boot-FPBB and the design-based complete data analysis for subsampling scheme 2 ($f_2 \propto h(S_i)$). (a) before deletion (b) imputation based on covariates under MAR_X (c) imputation based on covariates and sampling weights under MAR_X,W.
Figure 4.5 Comparison of point and interval estimation for 19 population quantiles using the stratified double weighted-FPBB and the design-based complete data analysis for subsampling scheme1 ( $f_2 \propto$ constant ). (a) before deletion (b) imputation based on covariates under MAR_X (c) imputation based on covariates and sampling weights under MAR_X,W.

Figure 4.6 Comparison of point and interval estimation for 19 population quantiles using the stratified double weighted-FPBB and the design-based complete data analysis for subsampling scheme2 ( $f_2 \propto h(S_i)$ ). (a) before deletion (b) imputation based on covariates under MAR_X (c) imputation based on covariates and sampling weights under MAR_X,W.
Table 4.1 Comparison of average width $\times 10^2$ and 95% CI coverage rates of $q(\alpha)$ for $\alpha = 0.05, 0.10, 0.25, 0.50, 0.75, 0.90$ and 0.95.

<table>
<thead>
<tr>
<th>Sampling Scheme</th>
<th>Missingness Mechanism</th>
<th>Methods</th>
<th>Average width of 95% CI $\times 10^2$</th>
<th>95% CI coverage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Complete Data</td>
<td>Actual</td>
<td>170 144 123 106 116 142 165</td>
<td>90.5% 92.5% 94.5% 93.5% 95.5% 91.5% 91.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1BD</td>
<td>172 144 126 105 117 146 165</td>
<td>90.5% 90.0% 95.0% 94.0% 95.0% 94.0% 89.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2BD</td>
<td>182 154 132 113 122 150 171</td>
<td>94.0% 95.0% 96.0% 94.5% 96.5% 96.0% 92.5%</td>
</tr>
<tr>
<td></td>
<td>MAR_X</td>
<td>SRS</td>
<td>165 134 112 101 108 132 158</td>
<td>93.0% 91.5% 86.0% 82.5% 83.0% 89.0% 93.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX_APR</td>
<td>171 143 120 105 116 146 172</td>
<td>92.5% 90.5% 90.5% 92.5% 93.5% 94.0% 95.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RE_APR</td>
<td>184 154 131 115 125 156 186</td>
<td>93.5% 94.0% 93.0% 97.5% 95.5% 95.5% 97.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>171 145 123 109 122 148 165</td>
<td>91.0% 89.5% 92.5% 95.0% 90.5% 89.5% 94.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
<td>182 158 134 118 129 156 175</td>
<td>93.5% 93.0% 94.5% 96.5% 94.5% 94.5% 95.0%</td>
</tr>
<tr>
<td></td>
<td>MAR_X,W</td>
<td>SRS</td>
<td>178 146 120 109 110 139 163</td>
<td>89.0% 81.0% 70.5% 69.0% 80.0% 90.0% 91.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX_APR</td>
<td>186 153 126 115 125 155 190</td>
<td>89.5% 92.5% 93.5% 95.5% 92.5% 92.5% 96.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RE_APR</td>
<td>197 166 140 127 136 168 197</td>
<td>95.0% 97.0% 97.0% 98.0% 96.0% 95.0% 96.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>173 150 124 111 119 146 163</td>
<td>91.5% 92.0% 93.0% 91.5% 90.0% 94.0% 92.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
<td>183 160 134 119 123 153 172</td>
<td>93.5% 95.5% 96.5% 92.5% 92.0% 93.0% 95.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_lwt</td>
<td>174 151 126 115 124 148 166</td>
<td>90.0% 89.0% 93.0% 94.5% 90.5% 96.0% 94.0%</td>
</tr>
<tr>
<td></td>
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<td>Syn2_lwt</td>
<td>184 158 136 122 132 155 174</td>
<td>92.0% 93.0% 95.5% 96.0% 94.5% 96.0% 95.0%</td>
</tr>
<tr>
<td></td>
<td>Complete Data</td>
<td>Actual</td>
<td>170 143 120 110 121 148 169</td>
<td>92.5% 94.5% 95.0% 96.0% 92.5% 87.5% 87.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1BD</td>
<td>177 142 120 108 121 152 175</td>
<td>91.0% 92.5% 92.0% 94.5% 92.5% 87.5% 87.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2BD</td>
<td>182 152 128 116 126 154 178</td>
<td>95.0% 97.0% 96.0% 97.0% 94.5% 90.0% 90.5%</td>
</tr>
<tr>
<td></td>
<td>MAR_X</td>
<td>SRS</td>
<td>175 139 121 111 116 141 169</td>
<td>86.5% 73.0% 57.0% 48.5% 61.0% 72.0% 80.5%</td>
</tr>
<tr>
<td></td>
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<td>FX_APR</td>
<td>174 142 111 124 162 202</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>RE_APR</td>
<td>181 150 128 119 131 168 205</td>
<td>94.0% 96.5% 97.0% 96.5% 97.0% 94.0% 96.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>166 140 119 111 126 156 180</td>
<td>93.5% 94.0% 96.5% 92.5% 92.0% 91.0% 90.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
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<td>94.5% 95.5% 98.0% 96.5% 95.0% 93.5% 92.5%</td>
</tr>
<tr>
<td></td>
<td>MAR_X,W</td>
<td>SRS</td>
<td>191 157 127 117 122 147 168</td>
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</tr>
<tr>
<td></td>
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<td>FX_APR</td>
<td>186 153 125 119 138 179 227</td>
<td>96.5% 97.0% 93.5% 96.5% 97.0% 95.0% 94.5%</td>
</tr>
<tr>
<td></td>
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<td>RE_APR</td>
<td>190 161 135 131 148 184 220</td>
<td>98.0% 99.5% 97.5% 98.0% 98.5% 96.5% 95.0%</td>
</tr>
<tr>
<td></td>
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<td>Syn1_srs</td>
<td>168 146 124 114 128 155 174</td>
<td>94.0% 92.5% 84.0% 73.0% 76.0% 87.5% 88.0%</td>
</tr>
<tr>
<td></td>
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<td>Syn2_srs</td>
<td>184 160 135 122 134 160 179</td>
<td>95.0% 95.5% 88.0% 77.5% 79.0% 87.0% 89.0%</td>
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<tr>
<td></td>
<td></td>
<td>Syn1_lwt</td>
<td>168 143 121 113 130 158 176</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_lwt</td>
<td>178 155 131 122 138 166 185</td>
<td>96.0% 95.5% 95.5% 92.5% 95.5% 95.0% 93.0%</td>
</tr>
</tbody>
</table>

Note: $f_2 \propto \text{const.}$ and $f_2 \propto h(S_i)$
Table 4.2 Comparison of RelBias, RMSE and 95% CI coverage rates for the mean of Y1 and proportions of Y3 and Y4,
Population True Value: $Y^* = 20.4$, $P_{Y_1} = 0.608$, $P_{Y_2} = 0.117$

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<tr>
<th>Sampling Scheme</th>
<th>Missingness Mechanism</th>
<th>Methods</th>
<th>RelBias</th>
<th>RMSE</th>
<th>95% CI coverage</th>
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</thead>
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<td>Complete Data</td>
<td>Actual</td>
<td>$\bar{Y}_1$</td>
<td>$P_{Y_1}$</td>
<td>$P_{Y_2}$</td>
</tr>
<tr>
<td>Actual samples BD: $\bar{Y}_1 = 20.3$</td>
<td></td>
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<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>0.0%</td>
</tr>
<tr>
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<td>10.8%</td>
</tr>
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<tr>
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</tr>
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<td>$f_2 \propto h(S_i)$</td>
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<td>Actual</td>
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<td></td>
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<td>Actual samples BD: $\bar{Y}_1 = 20.4$</td>
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<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
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<td></td>
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<td>1.2%</td>
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Table 4.3 Comparison of RelBias, RMSE and 95% CI coverage rates for the regression coefficients of \( Y_1 \), \( Y_3 \) and \( Y_4 \) on \( Y_2 \).

Population True Value: \( \beta_{1,|y_2} = 0.488 \), \( \beta_{1,|y_2} = 0.227 \), \( \beta_{1,|y_2} = 0.083 \)

<table>
<thead>
<tr>
<th>Sampling Scheme</th>
<th>Missingness Mechanism</th>
<th>Methods</th>
<th>RelBias</th>
<th>RMSE</th>
<th>95% CI coverage</th>
</tr>
</thead>
<tbody>
<tr>
<td>( f_2 \propto \text{const.} )</td>
<td>Complete Data</td>
<td>Actual Data</td>
<td>- - -</td>
<td>0.103 0.065 0.098</td>
<td>98.0% 96.0% 90.0%</td>
</tr>
<tr>
<td></td>
<td>Syn1BD</td>
<td>0.4% 1.1% 1.9%</td>
<td>0.104 0.067 0.098</td>
<td>96.0% 93.5% 88.0%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Syn2BD</td>
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<td></td>
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<tr>
<td>( f_2 \propto h(S_i) )</td>
<td>Actual samples BD: ( \beta_{1,</td>
<td>y_2} = 0.481 ), ( \beta_{1,</td>
<td>y_2} = 0.232 ), ( \beta_{1,</td>
<td>y_2} = 0.086 )</td>
<td>MAR_X</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX/APR</td>
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<td>97.0% 97.0% 92.5%</td>
</tr>
<tr>
<td></td>
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<td>RE/APR</td>
<td>0.3% 2.1% 22.6%</td>
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<td>98.0% 95.5% 95.0%</td>
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<td>SRS</td>
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<td>0.121 0.070 0.100</td>
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<td>FX/APR</td>
<td>0.4% 1.7% 53.5%</td>
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<td></td>
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<td>RE/APR</td>
<td>0.2% 6.5% 22.9%</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>3.6% 2.7% 9.7%</td>
<td>0.123 0.076 0.105</td>
<td>94.5% 91.5% 91.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
<td>3.5% 0.5% 4.6%</td>
<td>0.121 0.076 0.107</td>
<td>96.5% 96.0% 93.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_lwt</td>
<td>1.8% 1.4% 2.8%</td>
<td>0.121 0.075 0.104</td>
<td>95.5% 93.0% 90.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_lwt</td>
<td>2.2% 1.5% 2.1%</td>
<td>0.120 0.075 0.106</td>
<td>96.5% 96.0% 96.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MAR_X, W</td>
<td>SRS</td>
<td>- - -</td>
<td>0.108 0.066 0.088</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX/APR</td>
<td>0.1% 0.6% 2.2%</td>
<td>0.109 0.068 0.089</td>
<td>95.0% 95.0% 93.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RE/APR</td>
<td>0.4% 2.9% 6.5%</td>
<td>0.109 0.069 0.090</td>
<td>95.0% 96.5% 96.0%</td>
</tr>
<tr>
<td></td>
<td>Actual samples BD: ( \beta_{1,</td>
<td>y_2} = 0.481 ), ( \beta_{1,</td>
<td>y_2} = 0.229 ), ( \beta_{1,</td>
<td>y_2} = 0.090 )</td>
<td>MAR_X</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX/APR</td>
<td>0.5% 0.6% 43.5%</td>
<td>0.114 0.069 0.079</td>
<td>93.5% 95.0% 97.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RE/APR</td>
<td>0.8% 2.5% 19.0%</td>
<td>0.111 0.061 0.065</td>
<td>95.0% 95.5% 97.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>0.4% 0.7% 5.6%</td>
<td>0.126 0.082 0.097</td>
<td>94.0% 92.0% 91.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
<td>0.0% 3.5% 2.7%</td>
<td>0.124 0.082 0.098</td>
<td>95.0% 94.0% 96.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MAR_X, W</td>
<td>SRS</td>
<td>17.6% 12.4% 69.5%</td>
<td>0.141 0.069 0.101</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FX/APR</td>
<td>0.4% 5.7% 42.2%</td>
<td>0.118 0.066 0.082</td>
<td>93.5% 95.5% 55.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RE/APR</td>
<td>1.7% 3.1% 30.1%</td>
<td>0.111 0.054 0.073</td>
<td>97.5% 98.0% 97.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_srs</td>
<td>6.7% 3.1% 23.0%</td>
<td>0.136 0.073 0.093</td>
<td>93.0% 94.0% 94.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_srs</td>
<td>7.4% 0.4% 19.0%</td>
<td>0.136 0.075 0.095</td>
<td>96.0% 97.5% 97.0%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn1_lwt</td>
<td>0.9% 0.9% 6.0%</td>
<td>0.130 0.075 0.092</td>
<td>93.0% 95.5% 93.5%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Syn2_lwt</td>
<td>1.7% 2.6% 3.3%</td>
<td>0.126 0.076 0.094</td>
<td>97.0% 98.0% 97.5%</td>
</tr>
</tbody>
</table>
4.5 Application to NHANES III

We apply our method to the National Health and Nutrition Examination Survey (NHANES) III (1988-1994), which is designed to provide national estimates of health and nutritional status of the civilian noninstitutionalized population of the United States aged 2 months and older (NCHS 1994). The data is obtained from a stratified, multistage area probability sampling design with oversampling of certain age and ethnicity groups. For confidentiality and computational reasons, the public use data provides two pseudo PSUs per stratum. Another unique feature of NHANES is that data are collected through both interview and actual physical examinations of the sampled persons. Both unit- and item-level nonresponse occurs in both components of the survey, and there is a particularly high missing rate on the body mass index (BMI) measure for youth data in the physical examination component (30%). As a popular measure of overweight status and obesity, the percentiles of BMI for children and youths are of particular interest for public health reasons. The upper percentiles and the lower percentiles are also closely monitored for overweight and underweight status, respectively. As a result, we restrict our analysis sample to children and youths 2 months to 16 years of age.

We estimate population quantiles (from 0.05 to 0.95 with an increment of 0.05 along with two extreme percentiles: 0.03 and 0.97) of BMI for children and youths by gender. We also estimate the proportion of such a population being covered by health insurance, overall and by race. To assure congenial inference, we include the following variables that are either of primary interest in the substantive analysis, or are important predictors for BMI measures in the imputation model: age, gender, race, education, mother’s BMI, father’s BMI and family income (Yuan & Little, 2007a). We compared
three different methods in treatment of the missing data: 1) Complete case analysis (CC) with design-based estimation; 2) fully parametric model-based MI using design-based estimation, within which we apply both an imputation model assuming SRS and the appropriate model conditional on all three sample design features (i.e. dummy variables indicating cluster and stratum memberships as well the log transformation of sampling weights); and 3) our proposed finite population Bayesian bootstrap method (using SYN2_lwt, since we do not have separate weights for the first and second stages of sampling). Estimates of the median BMI and the proportion of children with health insurance are given in Table 4.4. The CC method appears to overestimate both the median of the BMI measure and health insurance coverage for full sample and race domains relative to the MI approaches, and yields the widest confidence intervals or largest standard errors as a result of decreased sample size. On the other hand, the median of BMI obtained from synthetic MI is quite similar to that from the model-based MI, while demonstrating some advantages in efficiency by yielding shorter intervals. The generally lower health insurance coverage estimates under the synthetic MI relative to model-based MI might be attributable to the fact that the synthetic MI are able to capture certain interactions between the sample design variables and the regular covariate matrix which are not explicitly modeled in the fully model-based MI.
Table 4.4 Alternative methods in estimating the median of BMI and the health insurance coverage rate, for full sample and by gender and race, respectively

<table>
<thead>
<tr>
<th>Variable</th>
<th>Domain</th>
<th>CC</th>
<th>Model-based MI</th>
<th>Synthetic MI</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMI</td>
<td>Overall</td>
<td>17.2 [17.1, 17.4]</td>
<td>17.1 [16.9, 17.3]</td>
<td>17.0 [16.9, 17.2]</td>
</tr>
<tr>
<td></td>
<td>Male</td>
<td>17.2 [16.9, 17.4]</td>
<td>17.0 [16.7, 17.2]</td>
<td>17.0 [16.8, 17.2]</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>17.3 [17.0, 17.7]</td>
<td>17.1 [16.8, 17.4]</td>
<td>17.1 [16.8, 17.3]</td>
</tr>
<tr>
<td>Health</td>
<td>Overall</td>
<td>0.785 (0.020)</td>
<td>0.778 (0.019)</td>
<td>0.761 (0.019)</td>
</tr>
<tr>
<td>Insurance</td>
<td>White</td>
<td>0.822 (0.018)</td>
<td>0.815 (0.017)</td>
<td>0.799 (0.016)</td>
</tr>
<tr>
<td></td>
<td>Non-White</td>
<td>0.645 (0.036)</td>
<td>0.643 (0.033)</td>
<td>0.634 (0.036)</td>
</tr>
</tbody>
</table>

Figure 4.7 displays a visual comparison of the percentile estimation for the three methods under consideration. We look at how those methods perform in three different percentile ranges by gender domains: the middle percentiles from 0.5 to 0.75, the upper percentiles from 0.90 to 0.97 and the lower percentiles from 0.03 to 0.1. We chose these percentile ranges because: first, the extreme lower and upper percentiles of BMI are typically used to monitor under- and over-weight for children and youths; second, there is evidence that gender difference exists in these BMI percentile ranges (particularly when age is considered, i.e. growth patterns in BMI). In general, both MI methods result in very similar BMI estimates, and they are lower than those obtained from CC analysis. This makes sense, because by comparing the distributions of age for complete cases and for missing cases on the BMI measure, we found that younger children are more susceptible to missingness, and therefore CC analysis tends to overestimate BMI by excluding those younger missing cases. The inclusion of the age variable as a predictor in the imputation model corrects such an overestimation. The magnitude of this correction for boys is bigger than that for girls in estimating the lower percentiles (0.03, 0.05). By examining a report on BMI-for-age percentiles by gender released from CDC
(http://www.cdc.gov/nchs/data/series/sr_11/sr11_246.pdf), we find that baby boys (corresponding to the lower quantiles here) have relatively higher BMI, which might be the explanation. Therefore, it is convincing to believe that our synthetic MI method is comparable to the appropriate model-based MI in adjusting for potentially incorrect estimation under CC analysis.
Figure 4.7 Comparison of methods for quantile estimation of BMI, by gender
4.6 Discussion

While multiple imputation has become a popular option for the analysis of missing data, open issues remain in its practical application with complex sample survey data. The complex features of sampling compounded with nonresponse in survey data often result in a rather complicated data structure, which prevents the standard MI techniques (such as a multivariate normal model assuming simple random sampling) from being applied straightforwardly. In this paper, we develop a general purpose approach to account for various design features in a highly stratified two-stage sample, using the two-step synthetic MI framework proposed by Zhou et al. (2013a). We have focused on evaluating the performance of the new method compared with existing methods, with respect to several missing data issues frequently encountered in large population-based socioeconomic and epidemiological studies. These include: i) accommodating stratification and multi-stage sampling in the imputation process; ii) the employment of nonstandard or non-normal imputation models for estimating probabilities of rare events; and iii) the estimation of population quantiles with multiply imputed data.

We demonstrate that the coverage properties of the proposed method are fairly good for nonsmooth statistics. Specifically, our stratified variations of the weighted Polya posterior exhibits robustness to the loss function for estimating upper and lower tails of the distribution function where even the appropriate model-based method (i.e. FX_APR) fails. In contrast with existing fully parametric MI methods, most of which perform poorly when applied to rare outcome binary data, the proposed method yields quite stable parameter estimates regardless of the rarity of the outcome.
It is worth stressing that our method requires only the simplest form of imputation model and combining rules for inference, because the effects of complex sample designs and the effect of estimating the nuisance parameter (e.g. regression coefficient in the regression imputation scheme is an example of a nuisance parameter, since our main parameters of interest are the population mean/quantiles of $Y$) in imputation are both correctly reflected in the replication variance estimation given the design-reversed and multiply imputed synthetic populations. The simplicity of the imputation model and the inference rules mean that any higher-level and nonlinear interactions in the covariate data, including those with the weights, clusters, or strata, will automatically be captured in the synthesizing step. However, when the imputation is conducted parametrically, as we do here, such design variable interactions will still need to be considered if they are associated with the missingness mechanism, although the impact of misspecification will generally be attenuated. Similarly, not-missing-at-random mechanisms that are dependent on the missing values are not accommodated in this framework.

Future research will investigate the inferential properties of the proposed method in situations where auxiliary information on all population units is available, using a constrained version of the Polya posterior. Two other possible research directions include: (i) extending the two-step synthetic MI framework to deal with unit nonresponse problems, and (ii) extending it to deal with generating synthetic data for disclosure risk limitation.
CHAPTER 5
CONCLUSION AND FUTURE RESEARCH

5.1 Contribution

Complex sample survey estimation and multiple imputation (MI) for survey nonresponse are two important areas in survey research. Coming up with a proper imputation model that is attentive to all complex sample design features can be difficult in many practical settings. In this thesis, we modified the classical MI framework of Rubin (1987) by dividing up the need to account for both sample designs and missing data into two separate steps. In the first step, we reversed the sampling mechanism by utilizing a noninformative Bayesian approach to finite population sampling. In the second step, we imputed the missing values in the created pseudo-population by constructing a parametric Bayesian model for the missing data under an IID assumption. Thus, the new framework stays within the Bayesian fundamentals underlying the standard MI theory. Yet it also generalizes naturally to increased flexibility for imputation with complex sample design surveys, because once a population with missing values is synthesized in a fashion that accommodates the sample design, a variety of imputation methods not limited to model-based MI can then be applied, for example, nonparametric hot deck imputation (Andridge & Little, 2010).

Chapters 2 to 4 develop different procedures but are methodologically interrelated within the general conceptual framework of the proposed two-step MI, in the sense that the weighted finite population Bayesian bootstrap (FPBB), or equivalently the weighted
Polya posterior, serves as the basic approach to the different procedures developed in each chapter.

Chapter 2 derives the theoretical formulation of the two-step MI from the traditional MI, filling a gap in the literature resulting from the fact that most current methods cannot adequately accommodate sampling weights. Chapter 3 addresses the compound effects of clustering and unequal selection probability in the imputation process. It derives a posterior predictive distribution of the population that improves upon the “two-stage Polya posterior” of Meeden (1999). The posterior under the new procedure relaxes an inherent exchangeability assumption about the correlational structure among survey variables between the sampled clusters and nonsampled clusters not guaranteed by the population data generation mechanism, yet required by Meeden’s “two-stage Polya posterior”. It therefore allows for a wider range of population data structures to be considered. Chapter 4 further extends the two-step MI approach into the area of stratified, clustered, and unequal probability-of-selection sample designs. It also explores the potential of the new method to deal with quantile estimation, and the estimation of proportions for binary rare events in the presence of item nonresponse.

In Chapters 3 and 4, we considered two variations of a two-stage procedure to create a synthetic population with missing data. One applies the weighted FPBB at each level of sampling and hence requires known design weights for units at all stages of sampling. The other employs replication methods at the PSU level and only applies the weighted FPBB to the ultimate sampling units using final weights (products of the design weights at each stage). Since the final weights are often the only design information available in public-use databases, the latter version has important practical value. The
simulation results demonstrated good repeated sampling properties of the proposed methodology in several sampling design settings, including PPS, two-stage clustering and highly stratified two-stage cluster samples. The proposed method also performed well with respect to estimation of means, regression parameters, and quantiles.

The proposed MI framework provides advantages over the existing fully parametric model-based MI in three ways:

(i) **modeling**: the “untying” step of the proposed MI procedure recovers most of the information about the data generating mechanisms, including the data model, the sampling process and part of the response process. Thus a simple imputation model assuming IID suffices (in most cases) for the resulting pseudo population. This strikes a balance between *congeniality* and *sparsity* simultaneously required of the model, something difficult to attain with the fixed effects modeling strategy (as shown in the simulation results in chapter 4)

(ii) **computation**: the proposed algorithms can be easily programmed in the R language (example codes for their applications on the BRFSS data and the NHANES data are provided in the Appendix). To be quite conservative for the proposed method to work properly, we purposefully generated a very large number of synthetic populations (i.e. $L$ and $S$) for both the simulation study and the real data application. This required, for example, about 1.5 hours for imputing the missing BMI in the NHANES III data, with $L=50$ at the PSU level, $S=5$ at the element level and $M=5$ multiple imputations, on a 2.50 GHz Intel Core i5 laptop. Meanwhile, the proposed method does not need complex numerical integration methods as is the case with random effects
modeling. Hence, they offer a feasible alternative to the random effects model which is well-known to have convergence issues with high dimensional categorical data;

(iii) **analysis:** the weighted Polya sampling draws all variables jointly, and hence preserves the population-level multivariate relationships that can then be used for a variety of analyses (as evident by the real data applications). Since the relationships of the survey variables with the sample weights are also maintained in the draws, once we simulate sensible copies of the entire population, these analyses become fairly straightforward without appeal to design-based estimation (as is the case with quantile estimation).

### 5.2 Limitations

A full retrospective appreciation of the work done in this thesis also reveals several limitations of the proposed methods that suggest directions for further investigation. A major limitation lies in the fact that the simulation studies are restricted to one-stage element sampling or at most a stratified two-stage cluster sampling. Methods are not fully developed for *multistage designs* that involve more than two stages of sampling. For example, for the application on NHANES III, we considered the data as a two-stage sample, and assumed that lower-than-PSU-level sampling design effects could be neglected. In seeking for an improved solution, analysts should consider mapping multistage sampling designs to an “ultimate cluster design” (Kalton, 1979) before implementing the proposed procedure developed in this thesis.

Another limitation is the lack of development for the *small sample degrees of
freedom (df) under the newly derived combining rules. As the method is a combination of multiple imputation and synthetic data generation, we attempted to draw theoretical support from both literatures (Raghunathan et al., 2003; Reiter, 2003, 2004; Reiter & Raghunathan, 2007; Rubin, 1987). However, despite the small sample df derived in Barnard and Rubin (1999) that is aimed at MI inference at the sample-level, the fraction of missing information (FMI) is not well-defined in the context of synthetic data in the current literature. Hence df for constructing interval estimation for the small sample t-approximation synthetic data cannot be constructed using FMI. In this regard, a next step is to break down the variance estimator under the new MI combining rules into components that may be used to define a sensible FMI.

The efficiency loss under the proposed method in a complex sample design setting is a further, though expected, limitation, resulting from the gains in robustness to model assumptions by using a computational nonparametric technique. We should note that, the proposed method is typically not as efficient (with higher RMSE) as its fully parametric counterparts with correctly specified imputation model. Improvement might be made through the use of auxiliary information, which will be discussed in the next section.

In addition, the validity of our method relies strongly on sample size. In chapter 4, we circumvented this problem by assuming a relatively large number of strata (H=50) in the sample. Hence, the synthetic procedure was still based on a reasonably large sample scenario, even though only 2 PSUs were selected within each stratum. The method will not work when both the number of strata and the number of clusters within strata are quite small and if single PSU strata exist in the sample. In such cases, we need to consider adaptation of the method by either collapsing strata or splitting PSUs. See Rubin
for a discussion of inappropriateness in using bootstrap and Bayesian bootstrap.

5.3 Future Research

The two-step MI framework developed in this thesis has meaningful extensions in several directions.

Throughout the thesis, I consider only the statistical adjustments for item-level missing data, assuming no unit nonresponse in the selected sample. A natural extension of the method would be to adapt the two-step MI framework to incorporate unit nonresponse. Unit nonresponse is typically accounted for by weighting or weight adjustments, for which the sampling weight of non-respondents is spread over respondents using either weighting class adjustments or propensity modeling. When replication variance estimators (such as BRR and Jackknife) are used, it remains a practical question whether it is necessary to repeat every weighting step (including adjustments for unknown eligibility, unit nonresponse adjustments and post-stratification using auxiliary variables) separately for each replicate sample in order to produce consistent or approximately unbiased variance estimates (Valliant, 2004). Findings from previous research are mixed. Some evidence suggests the need to recalculate weights for each replicate (Lemeshow, 1979; Valliant, 1993); while others suggest that such extra efforts make little difference (Rust, 1987) or even lead to overestimation of the variance in small samples (Valliant, 2004). On all accounts, there is a need to develop an adapted version of the two-step MI method that propagates the uncertainty in unit nonresponse adjustments, given the fact that a replication method and hence a replication-type variance estimator is used under the proposed MI framework. It would also be interesting
to compare the variance estimator under such an adapted two-step MI method with that resulting from an expedient method that does not propagate this source of variability. A feasible way to achieve this in a single-stage sampling design would follow a three-step procedure: 1) generate a Bayesian bootstrap (BB) sample from the parent sample \( S \) (including the unit response indicator variable, the base weight \( w_{\text{base}} \), and the item-level missing data); 2) for each BB sample, apply regular unit nonresponse adjustments to the base weights for the respondent set \( S_R \), using available auxiliary variables for the entire sample, and obtain the nonresponse (NR)-adjusted weight \( w_{\text{NR}} \); and 3) for each NR-adjusted bootstrap sample, apply the weighted Polya posterior with \( w_{\text{NR}} \) as the input weight in the algorithm to create synthetic populations (with item-level missing data). The subsequent imputation method that assumes IID as well as the combining rules for inference would follow as described in chapter 2.

A second valuable extension would be to investigate \textit{domain/small area estimation}. Statistical agencies and survey organizations routinely produce estimates for subpopulations, called domains, to aid public and private sectors in effective policy making. Despite the growing demand, reliable domain estimates are difficult to obtain using standard frequentist approaches, primarily because domains are typically defined after a sample was selected. When cut across planned strata, they usually have very small associated sample size, leading to less precise estimates than those for the whole population. Complications of domain estimation and methods to tackle them are accounted in the complete data context to some degree (Cochran, 1977; Ghosh & Rao, 1994). When missing data are present, multiple imputation for domain estimation remains an issue, especially when the domain size is small and response rate is low (Kim,
Theoretically, we should include domain indicators $Z$ as well as their interactions with the survey weight $W$ in the imputation scheme for the MI variance estimator to be approximately unbiased. In practice, however, many domain estimates of interest in the substantive analysis will not be identified at the time that the imputation was carried out. Even if they do, to model the interaction term $W*Z$ for all possible domains is not feasible. In this case, the proposed two-step MI method is a logical choice because the association of the weights with survey variables in the population data can be resolved by generating synthetic populations. Once the numerous complete copies of the population are created and imputed, the simulated population-level estimates can be simply combined. This differs from the standard MI method where inference is based on explicit design-based domain estimators that may be lacking for certain estimands, e.g. the difference or ratio estimator of two domain medians.

Another related extension of the proposed MI procedure would be to incorporate auxiliary information at the first step of synthetic data generation. Auxiliary information is routinely used in finite population sampling to facilitate inference. For example, ratio and regression estimators are used when the population mean of an auxiliary variable is known a priori. The procedures developed in this thesis all assumed an absence of prior auxiliary information. Our future work will focus on adapting the current procedures by capitalizing on such information to improve the efficiency of imputation and inference as a whole. Lazar, Meeden, and Nelson (2008) showed that the Polya posterior can be easily adapted to incorporate different levels of prior information. In a similar fashion, I can restrict the weighted-FPBB to create pseudo-populations that satisfy certain constraints.
specified by the prior information, such as known population means or ranges about the auxiliary variable(s). This would in effect extend the work of Sangeneh et al. (2011) to the missing data context. On the other hand, since the essence of all small area estimation methods lies in the use of available auxiliary variables (Rao, 2003), I believe a constrained version of the weighted-FPBB would also provide a nonparametric Bayesian solution for small area estimation in both complete data and missing data settings. Yet another utilization of such auxiliary information lies in the aforementioned nonresponse adjustments in each BB sample.

We considered a special type of missing data mechanism which depends on the selection probability (or sample weights). On the one hand, the design-inversing step of the new method ensures a correct estimate of the population distribution in the presence of missing data and thus reduces the impact of misspecified missingness mechanisms by avoiding enhancement from misspecified data generation mechanisms. On the other hand, elimination of the weights from the self-weighting FPBB population does not obviate the need to account for the weights in the imputation process to attain valid inference. Future extensions of this work could use other imputation methods to account for such a weight-dependent MAR missingness, for example, the weighted hot deck by Andridge and Little (2009). In some cases, the missingness may also depend on the unobserved random clustering effects and thus becomes nonignorable. We do not consider such scenarios here, since our method is proposed to address item nonresponse under the MAR assumption. But the method may be adapted to deal with nonignorable missing data problem under a two-stage cluster sampling design, by combining with the work of Yuan and Little (2007a, 2007b, 2008), Andridge and Little (2011) and West and Little (2013).
APPENDIX

1. R code for using the proposed two-step MI method on BRFSS

```r
require(mice)
require(survey)
set.seed(123)

#dt: sample data set for analysis, with recoded variables;
#N: synthetic population size;
#Bt1: number of BB samples created for stage 1;
#Bt2: number of weighted FPBB populations created for stage 2;
#Mt: number of multiple imputations;
#ps.n: sample size of parent sample;

SynMI <- function(dt, N, Bt1, Bt2, Mt, ps.n) {
  ##Step 1: synthesize populations;
  #Stage 1: draw bootstrap samples from the parent sample;
  #Normalize the final weights to sum up to the synthetic population size;
  dt[, "X_FINALWT"] <- dt[, "X_FINALWT"] * N/sum(dt[, "X_FINALWT"])
  dsgn <- svydesign(ids = ~1, strata = NULL, nest = FALSE, data = dt, weights = ~ X_FINALWT)
  dsgn.r <- as.svrepdesign(design = dsgn, type = "subbootstrap", replicates = Bt1)
  repwt <- as.matrix(dsgn.r$repweights)
  repwt[repwt == 0] <- NA

  #Set up a data frame to store results from loglinear analysis;
  logcf <- matrix(0, 20, Bt1)

  #Stage 2: within each bootstrap sample, draw weighted FPBB synthetic populations;
  for (j in 1:Bt1){
    st.bb <- cbind(dt, repwt[, j])
    #Delete those units with zero replicate weights for each bootstrap sample;
    st.BB <- na.omit(st.bb)
    #Recode those 9999 back to NA for imputation;
    st.BB[st.BB == 9999] <- NA
    st.rpm <- st.BB[rep(1:nrow(st.BB), round(st.BB$repwt)),]
    ns <- ps.n - 1
    Samwts <- rep((ps.n/2), st.BB$X_FINALWT, round(st.BB$repwt))
  }
}
```

---

155
lcf <- matrix(0,20,Bt2)
#Write the algorithm for creating weighted FPBB populations based on Cohen;
for(boot in 1: Bt2){
  l <- rep(0,ns)
  for(k in 1:(N-ns)){
    l <- l+rmultinom(1,1,((Samwts-1)+1*((N-ns)/ns))/((N-ns)+(k-1)*((N-ns)/ns)))
  }
  income <- as.factor(c(rep(st.rp[,6],l),st.rp[,6]))
  bmi <- as.numeric(c(rep(st.rp[,10],l),st.rp[,10]))
  bphigh <- as.factor(c(rep(st.rp[,9],l),st.rp[,9]))
  hlthplan <- as.factor(c(rep(st.rp[,1],l),st.rp[,1]))
  age <- as.numeric(c(rep(st.rp[,2],l),st.rp[,2]))
  racew <- as.factor(c(rep(st.rp[,5],l),st.rp[,5]))
  educa <- as.factor(c(rep(st.rp[,8],l),st.rp[,8]))
  employ <- as.factor(c(rep(st.rp[,7],l),st.rp[,7]))
  gender <- as.factor(c(rep(st.rp[,3],l),st.rp[,3]))
  lgwt <- as.numeric(log(c(rep(st.rp[,4],l),st.rp[,4])))
  # Step 2: Multiple imputation of synthetized populations;
  # Imputation model ignores the design;
  temp1 <- data.frame(cbind(income, bmi, bphigh, hlthplan, age, racew, educa, employ, gender))
  # Imputation model includes log of final weights;
  temp2 <- data.frame(cbind(income, bmi, bphigh, hlthplan, age, racew, educa, employ, gender, lgwt))
  # Impute for each synthetic population;
  temp_impx <- mice(temp1)
  temp_impxw <- mice(temp2)
  mlx <- data.frame()
  mlxw <- data.frame()
  for (u in 1:Mt){
    mult <- as.vector(rep(u,nrow(temp1)))
    mlx <- rbind(mlx,cbind(complete(temp_impx, u),mult))
    mlxw <- rbind(mlxw,cbind(complete(temp_impxw, u),mult))
  }
  lcf <- matrix(0,20,Mt)

  ##Analysis of the imputed synthetic populations;
  for (v in 1:Mt){
    for (mult in v:v){
      stx <- mlx[mlx[,"mult"]==v,]
      stxw <- mlxw[mlxw[,"mult"]==v,]
    }
  }
dsignx <- svydesign(id=~1, strata= NULL, weights= NULL, data=stx)
dsgnxw <- svydesign(id=~1, strata= NULL, weights= NULL, data=stxw)

#loglinear analysis on the imputed data;
ax <- svyloglin(~income+hlthplan+racew+gender,dsgnx)
bx <- update(ax,~.^2)
axw <- svyloglin(~income+hlthplan+racew+gender,dsgnxw)
bxw <- update(axw,~.^2)
lcf[,v] <- c(coef(bx),coef(bxw))

#Calculate the synthetic MI point and variance estimates;
lgcfd .boot] <- apply(lcf,1,mean)
print(boot)

lgcf[j] <- apply(lgcfd,1,mean)
print(j)

smpm_log <- apply(logcf,1,mean)
smpv_log <- (1+1/Bt1)*apply(logcf,1,var)
stat <- cbind(smpm_log,smpv_log)
write.table(stat,file="D:\Dissertation\paper1\brfss_syn_loglin.csv",row.names=FALSE, sep=" ", )

###Example run;
##Need to collapse the medium and high income categories to avoid sparse cells;
dt$INCOME[dt$INCOME==3]<-2
#missing data were coded as ‘9999’ in data set brfss09.csv;
SynMI(dt=dt, N=4500, Bt1=100, Bt2=30, Mt=5, ps.n=388)
2. R code for using the proposed two-step MI method on NHANES III

```r
require(survey)
require(mice)
require(polyapost)
set.seed(seed #)

syn_bmi<-function(dt, N, Bt1, Bt2, Mt){

##Step 1: Generate synthetic populations with missing data;
#Stage 1: Create bootstrap samples from the parent sample;
dsgn <- svydesign(ids = ~predcl, strata = ~pstrat, nest = TRUE, data = dat,
weights = ~predwt)
dsgn.RW <- as.svrepdesign(design = dsgn, type = "subbootstrap", replicates = Bt1)
dim(dsgn.RW$repweights)
repwt<-as.matrix(dsgn.RW$repweights)
repwt[repwt==0]<-NA
dim(repwt)

##set up arrays to hold point estimates from bootstrap samples;
btm<-matrix(0,nrow=Bt1,ncol=3)
bqt<-matrix(0,nrow=Bt1,ncol=21)
bqt<--matrix(0,nrow=Bt1,ncol=21)
bqt<-matrix(0,nrow=Bt1,ncol=21)

for (j in 1:Bt1){
    st.bb<-cbind(dt,repwt[,j])
    #delete those units with zero weights for each bootstrap sample;
    st.BB<-na.omit(st.bb)
    #recode those 999 back to NA so that the mice package can be used for
    imputation;
    st.BB$pybmi[st.BB$pybmi==999]<-NA

    #need to calculate the replicate weights;
    Samwt<-st.BB[,9]*st.BB[,13]
    #normalize again the adjusted weights;
    Samwts<-Samwt*N/sum(Samwt)
    np<-nrow(st.BB)
    ids<-seq(np)
    ns<-N-np

    ##Stage 2: Create unweighted synthetic populations within each bootstrap sample;
    Set up arrays to hold point estimates from imputed unweighted synthetic populations;
    fbm<-matrix(0,nrow=Bt2,ncol=3)
    fbqt<-matrix(0,nrow=Bt2,ncol=21)
    fbqt<--matrix(0,nrow=Bt2,ncol=21)
```

fbqt<-matrix(0,nrow=Bt2,ncol=21)

for(boott in 1:Bt2){
  l<-vector()
  smp<-wtpolyap(ids, Samwts, ns)
  #input the adjusted weights in the weighted Polya sampling algorithm;
  for (k in 1:np){
    l<c(l,length(smp[smp==k]))
  }
  #check if the vector of l sum up to the number of synthetic population size;
  sum(l);

  predY1<-c(rep(st.BB[,1],l)) #bmi
  predY2<-c(rep(st.BB[,2],l)) #race
  predY3<-c(rep(st.BB[,3],l)) #gender
  predY4<-c(rep(st.BB[,4],l)) #income
  predY5<-c(rep(st.BB[,5],l)) #education
  predY6<-c(rep(st.BB[,6],l)) #mother’s bmi
  predY7<-c(rep(st.BB[,7],l)) #father’s bmi
  predY8<-c(rep(st.BB[,8],l)) #age
  predwt1<-c(rep(st.BB[,9],l))
  predlwt<-log(predwt1) #log of sample weight
  predCID<-c(rep(st.BB[,12],l)) #cluster ID
  predSTID<-c(rep(st.BB[,11],l)) #stratum ID

  ##Step 2: Multiple imputation of the unweighted synthetic populations;

  #use the imputation model including log of weight as a predictor (syn_lwt);
  temp1<-data.frame(cbind(predY1, predY2, predY3, predY4, predY5, predY6, predY7, predY8, predlwt))
  temp1_imp<-mice(temp1,method="norm", m=Mt)
  ml<-complete(temp1_imp, 'long')
  ml$bmit<-exp(ml$predY1) #back transform bmi to its normal scale
  mlmale<-subset(ml, predY3==1)
  mlfem<-subset(ml, predY3==2)
  multm<-cbind(as.vector(by(ml$bmit,ml$.imp,mean)),
               as.vector(by(mlmale$bmit,mlmale$.imp,mean)),
               as.vector(by(mlfem$bmit,mlfem$.imp,mean)))
  multqt<-sapply(with(ml,by(ml.$imp,function(x)quantile(x$bmit, c(0.03,seq(0.05,0.95,0.05),0.97)))),as.vector)
  multqtqm<-sapply(with(mlmale,by(mlmale.$imp,function(x)quantile(x$bmit, c(0.03,seq(0.05,0.95,0.05),0.97)))),as.vector)
  multqt<--sapply(with(mlfem,by(mlfem.$imp,function(x)quantile(x$bmit, c(0.03,seq(0.05,0.95,0.05),0.97)))),as.vector)

  fbm[boott,]<-t(apply(multm,2,mean))
fbqt[boott,]<-t(apply(multqt,1,mean))
fbqtm[boott,]<-t(apply(multqtm,1,mean))
fbqtf[boott,]<-t(apply(multqtf,1,mean))
print(boott)

btm[j,]<-t(apply(fbm,2,mean))
btqt[j,]<-t(apply(fbqt,2,mean))
btqtm[j,]<-t(apply(fbqtm,2,mean))
btqtf[j,]<-t(apply(fbqtf,2,mean))
print(j)

smpm<-apply(btm,2,mean)
smpv<-((1+1/Bt1)*apply(btm,2,var))
smpse<-sqrt(smpv)
smpqt<-apply(btqt,2,mean)
smpqtv<-((1+1/Bt1)*apply(btqt,2,var))
smpqts<-sqrt(smpqtv)
smpqtm<-apply(btqtm,2,mean)
smpqtvm<-((1+1/Bt1)*apply(btqtm,2,var))
smpqtsm<-sqrt(smpqtvm)
smpqtf<-apply(btqtf,2,mean)
smpqtvf<-((1+1/Bt1)*apply(btqtf,2,var))
smpqtsf<-sqrt(smpqtvf)

tt<-cbind(smpqt,smpqtm,smpqtf,smpqts,smpqtsm,smpqtsf)
ss<-cbind(smpm,smpse)
write.table(tt,file="D:/Dissertation/paper3/nhanes/synbmiqt_lwt.csv",row.names=FALSE,sep=",")
write.table(ss,file="D:/Dissertation/paper3/nhanes/synbmimn_lwt.csv",row.names=FALSE,sep=",")

###########;
#Example run;
syn_bmi(dt=dt, N=100000, Bt1=50, Bt2=5, Mt=5)
dt<-read.csv("D:/Dissertation/paper3/nhanes/synbmi.csv")
#Set the synthetic population size about 10 times the sample size;
N<-100000
#Normalize the weights to sum up to the assumed synthetic population size;
dt[,"predwt"]<-dt[,"predwt"]*N/sum(dt[,"predwt"])
sum(dt$predwt)
#Recode the missing values to 999;
dat[is.na(dat)]<-999

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[http://www.lse.ac.uk/statistics/events/SpecialEventsandConferences/CarpenterJR.pdf](http://www.lse.ac.uk/statistics/events/SpecialEventsandConferences/CarpenterJR.pdf).


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