Linearization, Bootstrap and Jackknife Variance Estimation for Ratio Estimators Under PPS Sampling

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Abstract

This paper mainly focuses on variance estimation for generalized ratio estimators under three probability proportional to size (PPS) sampling designs: randomized systematic PPS, PPS with replacement, and Poisson sampling. This paper also discusses three estimation methods, which are linearization, Bootstrap, and Delete-1 Jackknife, and how they are adapted according to each sampling design. A simulation study is also conducted to evaluate the performance of variance estimation under each method, with a special focus on the confidence interval. The study shows that PPS sampling with replacement consistently yields lowest variance and narrower confidence intervals. These findings highlight the practical considerations when applying model-assisted estimators under unequal probability sampling designs.

1 Introduction

A key challenge in probability sampling design is how to use the limited sample to draw inferences about the population parameters, such as the population mean. One possible way is to employ model-assisted estimators such as ratio estimators. When the sample is drawn using simple random sampling without replacement (SRSWOR), the application of ratio estimators is relatively straightforward, and the resulting estimators are referred to as simple ratio estimators. However, the use of ratio estimators becomes considerably more complex when the sample is obtained through an unequal probability sampling design, such as Probability Proportional to Size (PPS) sampling. In that case, the ratio estimator is called the generalized ratio estimator (Wu & Thompson, 2020, pp. 100–103). This paper will mainly focus on how to use a generalized ratio estimator under PPS sampling by implementing three variance estimation methods: linearization, Bootstrap, and delete-1 Jackknife.

For probability sampling design, there are two general types of estimation methods: model-based prediction and model-assisted estimation. In model-based prediction methods, the population outcome and auxiliary information follow a superpopulation model ξ , and survey design is ignored. Even though the estimators derived from specified superpopulation model are alwyas efficient; however, the model-based methods may generate misleading results when the underlying model is misspecified. In contrast, design-based inference does not rely on any model assumptions, which is more robust to model misspecification. This motivates the model-assisted estimators, which integrate design-based inference through a model-assisted approach. Specifically, an estimator is called model-assisted if it is unbiased under the assumed model and approximately unbiased under the probability sampling design (Wu &

Thompson, 2020, p. 95). This approach is particularly powerful, as it leverages the strengths of model-based estimation while preserving the validity of design-based inference.

This paper is structured as follows. Section 2 will review the simple ratio estimator, which can only be used under SRSWOR. In addition, Section 3 will introduce three PPS sampling methods used in this paper. The generalized ratio estimator and how to estimate its variance using linearization will be discussed in Section 4. Section 5 will introduce how the resampling methods Bootstrap and Jackknife can be applied to a generalized ratio estimator. Furthermore, A simulation study is conducted in Section 6, and the results will be presented as well. Finally, all the findings will be discussed in Section 7.

2 Simple Ratio Estimator

Simple ratio estimator is one of the model-assisted estimator to estimate the population mean μ_y . For simple ratio estimator, it is derived from the superpopulation model ξ such that $y_i = \beta x_i + \varepsilon_i$, i = 1, ..., N where ε_i are independent with $E_{\xi}[\varepsilon_i] = 0$ and $\mathrm{Var}_{\xi}(\varepsilon_i) = x_i \sigma^2$. Equivalently, the population mean μ_y can be derived as $\mu_y = \beta \mu_x + \bar{\varepsilon}_N$ where $\bar{\varepsilon}_N = N^{-1} \sum_{i=1}^N \varepsilon_i$. In addition, the sample \mathbf{S} under simple ratio estimator is restricted to be taken by Simple Random Sampling Without Replacement SRSWOR with survey data $\{y_i, x_i : i \in \mathbf{S}\}$ and assumes μ_x known. Formally, the simple ratio estimator is written as

$$\hat{\mu}_{y\mathbf{R}} = \hat{\beta}\mu_x = \frac{\bar{y}}{\bar{x}}\mu_x$$

where $\hat{\beta} = \frac{\bar{y}}{\bar{x}}$ is the weighted least square estimator of β under ξ . Denote E_{ξ} and E_{p} as taking expectation over ξ and the probability sampling design respectively, it can be shown that $E_{\xi} \left(\hat{\mu}_{yR} - \mu_{y} \right) = 0$ and $E_{p} \left(\hat{\mu}_{yR} \right) \doteq \mu_{y}$, which corresponds to two properties of model-assited estimators.

The proof of $E_p\left(\hat{\mu}_{y\mathrm{R}}\right) \doteq \mu_y$ relies on the technique called linearization, which can be further used to estimate the variance as well. In general, linearization simplifies a complex estimator by Taylor expanding it around its true population value and keep only the first-order term. This transforms the nonlinear estimator into an approximately linear one whose variance is easier to estimate. By linearization, it can be shown that the variance of the simple ratio estimator is given by (see Appendix A.1)

$$\operatorname{Var}(\hat{\mu}_{y\mathbf{R}}) = \left(1 - \frac{n}{N}\right) \frac{1}{n} \left(\sigma_y^2 + R^2 \sigma_x^2 - 2R\sigma_{xy}\right)$$

In addition to linearization, the variance of the simple ratio estimator can also be estimated using Bootstrap and delete-1 Jackknife resampling methods. The key difference between the two methods is that the Bootstrap resamples the entire sample with replacement, while the delete-1 Jackknife constructs replicates by systematically removing one observation at a time. The general algorithms for both methods are provided in the Appendix A.2. We will demonstrate how the three methods are applied in a more detailed way later in the generalized ratio estimator.

3 PPS Sampling methods

The main sampling method discussed in this paper is the Probability Proportional to Size (PPS) sampling. In PPS sampling, the key thing is the size variable (denoted by z) which is positively correlated to the outcome y and is available at the survey design stage. For instance, the previous income (z) is proportion to the expenses today (y). An important feature under PPS sampling design is that the first order inclusion probability π_i is proportional to the size variable, i.e., $\pi_i \propto z_i$. On the other hand, since the inclusion probabilities follows that $\sum_{i=1}^N \pi_i = n$, it follows that $\pi_i = n \frac{z_i}{T_z}$, or simply $\pi_i = n z_i$ if z is normalized. This paper will discuss three PPS sampling methods: randomized systematic PPS (RSPPS) sampling, PPS sampling with replacement (PPS-WR), and Poisson sampling.

3.1 Randomized systematic PPS sampling

Randomized systematic PPS sampling is a refined version of systematic PPS (SPPS) sampling. In the SPPS sampling, the number of sample n is assumed to be fixed and to be larger than 2. In addition, the size variable is assumed to be normalized, $\sum_{i=1}^{N} z_i = 1$, so that the first-order inclusion probability is given by $\pi_i = nz_i$ such that $\sum_{i=1}^{N} \pi_i = n$. The SPPS sampling contains two main steps:

1. Construct cumulative inclusion probabilities based on the given order of units in the sampling frame

$$b_0 = 0, b_1 = \pi_1, b_2 = \pi_1 + \pi_2, \dots, b_N = \pi_1 + \dots, \pi_N = n$$

2. Generate a random starting point $r \sim \text{Unif}(0,1)$, and select the unit i into the sample S if

$$b_{i-1} < r+k \leq b_i, \quad k \in \{0,1,2,\dots,n-1\}$$

Systematic PPS sampling implies two key properties. First, it implies the first-order inclusion probability $\pi_i = p(i \in \mathbf{S})$ is indeed equal to nz_i . This can be shown by considering two cases: when the interval $(b_{i-1}, b_i]$ lies entirely within a single unite interval [l, l+1], and when $(b_{i-1}, b_i]$ spans across an integer point l. The latter case is more complicated as there are two possible starting values r. Moreover, the second order inclusion probability, $\pi_{ij} = p(i, j \in \mathbf{S})$, may be zero for some pairs (i, j). This occurs because the randomized start r may cause certain pairs to be mutually exclusive in the final sample. For example, if $b_1 = 0.1, b_2 = 0.6$ but r = 0.5, then units 1 and 2 cannot be selected together as the r falls outside the range to include both.

To address the limitation of systematic PPS sampling regarding the second-order inclusion probabilities, one possible solution is to randomize the order of the N units on the sampling frame. By this extra step, it preserves the first-order inclusion probabilities that $\pi_i = nz_i$, but this ensures π_{ij} is strictly positive for all possible pairs (i, j). This refined systematic PPS sampling is referred as randomized systematic PPS sampling.

Due to the strictly positive π_{ij} , this ensures unbiased variance estimation for the Horvitz-Thompson estimator (Wu & Thompson, 2020, p. 74). Let $\hat{T}_{y\text{HT}} = \sum_{i \in \mathbf{S}} \frac{y_i}{\pi_i}$ be the H-T estimator of T_y , then

$$E(\hat{T}_{u\mathrm{HT}}) = T_u$$

Consequently, the variance of $\hat{T}_{y\text{HT}}$ can be expressed as

$$V\left(\hat{T}_{yHT}\right) = \sum_{i=1}^{N} \sum_{j=1}^{N} \left(\pi_{ij} - \pi_{i}\pi_{j}\right) \frac{y_{i}}{\pi_{i}} \frac{y_{j}}{\pi_{j}}$$

and an unbiased estimator of the variance is given by

$$v\left(\hat{T}_{yHT}\right) = \sum_{i \in \mathbf{S}} \sum_{j \in \mathbf{S}} \frac{\pi_{ij} - \pi_i \pi_j}{\pi_{ij}} \frac{y_i}{\pi_i} \frac{y_j}{\pi_j}$$

The above proof can be found at Appendix A.3.

3.2 PPS sampling with replacement

Another PPS sampling method is PPS sampling with replacement. In this approach, a unit is selected from the population with probabilities (z_1, \dots, z_N) such that $\sum_{i=1}^N z_i = 1$, and independently repeat it for n times with the same probabilities to have a sample of size n. Unlike randomized systematic PPS sampling, this method directly uses the size variables as selection probabilities rather than the inclusion probabilities. However, even it can be shown by Newton's formula that $\pi_i \doteq nz_i$, $\pi_i = P(i \in \mathbf{S}) = 1 - (1 - z_i)^n \doteq nz_i$, π_i is not equal to nz_i in PPS sampling with replacement.

To estimate the total T_y under PPS sampling with replacement, Hansen-Hurwitz (HH) estimator is used instead of Horitz-Thompson estimator. Formally, $\hat{T}_{y\rm HH}$ is defined as

$$\hat{T}_{y\text{HH}} = \sum_{i \in \mathbf{S}^*} \frac{y_i}{nz_i} = \frac{1}{n} \sum_{i=1}^n \frac{Y_i}{Z_i} = \frac{1}{n} \sum_{i=1}^n R_i$$

where S^* is the sample with the duplicated units (Wu & Thompson, 2020, p. 78). It can be shown that $\hat{T}_{y\text{HH}}$ is an unbiased estimator of T_y such that

$$E\left(\hat{T}_{\text{yHH}}\right) = T_y$$

with theoretical variance and variance estimator as

$$\begin{split} V\left(\hat{T}_{y\mathrm{HH}}\right) &= \frac{1}{n} \sum_{i=1}^{N} z_i \left(\frac{y_i}{z_i} - T_y\right)^2 \\ v\left(\hat{T}_{y\mathrm{HH}}\right) &= \frac{1}{n(n-1)} \sum_{i \in \mathbf{S}^*} \left(\frac{y_i}{z_i} - \hat{T}_{y\mathrm{HH}}\right)^2 \end{split}$$

All proofs can be found at Appendix A.4. Although nz_i is not exactly equal to π_i as in the Horvitz-Thompson estimator, Hansen-Hurwitz estimator $\hat{T}_{y\mathrm{HH}}$ can be viewed roughly as the $\hat{T}_{y\mathrm{HT}}$ for a large population. This is because, as population size N increases, the chance to have the duplicated units decreases. Therefore, the variance estimation $v(\hat{T}_{y\mathrm{HH}})$ can be used as an approximation to $v(\hat{T}_{y\mathrm{HT}})$. This approximation is advantageous because the variance estimator $v(\hat{T}_{y\mathrm{HH}})$ does not require the second-order inclusion probabilities which is very hard to compute in practice. We will further illustrate this in the next section.

3.3 Poisson Sampling

The last PPS sampling discussed in this paper is called the Poisson Sampling. In this approach, each unit is drawn independently from the population with $\pi_i = nz_i$, following the two key steps below:

1. Generate $r_i \sim \mathrm{Unif}(0,1)$ for each unit i and select the unit i to the sample if $r_i \leq \pi_i$.

2. Repeat step 1 for i = 1, ..., N independently.

Comparing to the previous two methods, Poisson sampling does not produce a fixed number of n; the actual sample size may be either larger or smaller than the set n. However, since the inclusion probabilities π_i is nz_i , the Horvitz-Thompson estimator can be applied to estimate the population total T_y . Nevertheless, as each unit is drawn independently, the variance of the HT estimator under Poisson sampling differs slightly from that under randomized systematic PPS sampling. As usual, $\hat{T}_{y\text{HT}}$ is defined as

$$\hat{T}_{y\text{HT}} = \sum_{i \in \mathbf{S}} \frac{y_i}{\pi_i}$$

which is unbiased for T_y . In addition, the theoretical variance of $\hat{T}_{y\mathrm{HT}}$ is

$$V\left(\hat{T}_{y\text{HT}}\right) = \sum_{i=1}^{N} \frac{1 - \pi_{i}}{\pi_{i}} y_{i}^{2} \tag{3.3.1}$$

and an unbiased estimator of this variance is

$$v\left(\hat{T}_{\text{yHT}}\right) = \sum_{i \in \mathbf{S}} \frac{1 - \pi_i}{\pi_i^2} y_i^2$$
 (3.3.2)

The proof can be found at Appendix A.5.

Comparing the three PPS sampling methods, each of them has distinct features in estimating the population total. Randomized systematic PPS sampling is the most general one as it includes the second-order inclusion probabilities. However, π_{ij} does not have a closed form formula and may be computed by simulation in practice (Thompson & Wu, 2008). In

contrast, PPS sampling with replacement and Poisson sampling involve independent draws and does not incorporate π_{ij} . However, the former one selects units based on the size variable, while the latter one selects based directly on the inclusion probabilities π_i . This is why the Horvitz-Thompson and Hansen-Huritz estimators are PPS sampling design-specific. We will explore how to estimate the variance of the ratio estimator under these three PPS sampling methods using linearization, Bootstrap, and Jackknife techniques.

4 Generalized Ratio Estimator

If the sample is drawn from either of the three PPS sampling method, the simple ratio estimator can not be simply applied to estimate the μ_y as it assumes the sample are drawn using SRSWOR. To deal with the sample that are drawn from unequal probability sampling design, we need another ratio estimator called generalized ratio estimator (GRE) to estimate the population mean. Like the simple ratio estimator, we still assumes the generalized ratio estimator works under the simple linear regression model, ξ . However, since the PPS sampling is an unequal probability sampling design, instead of setting $\hat{R} = \frac{\bar{y}}{\bar{x}}$, we need to use Horvitz-Thompson estimator or Hansen-Hurwitz estimator depends on the specific PPS sampling methods to estimate $R = \mu_y/\mu_x$.

4.1 Randomized Systematic PPS Sampling

For the most general unequal probability sampling design such as randomized systematic PPS sampling with $\pi_i = P(i \in \mathbf{S})$ and $\pi_{ij} = P(i, j \in \mathbf{S})$, the HT estimators for μ_y and μ_x

are written as

$$\hat{\mu}_{y\mathrm{HT}} = \frac{1}{N} \sum_{i \in \mathbf{S}} \frac{y_i}{\pi_i} = \frac{1}{N} \sum_{i \in \mathbf{S}} d_i y_i \quad \text{and} \quad \hat{\mu}_{x\mathrm{HT}} = \frac{1}{N} \sum_{i \in \mathbf{S}} \frac{x_i}{\pi_i} = \frac{1}{N} \sum_{i \in \mathbf{S}} d_i x_i$$

where $d_i = 1/\pi_i$ is called the basic design weights. Since $\hat{\mu}_{y\text{HT}}$ and $\hat{\mu}_{x\text{HT}}$ are unbiased with respect to μ_y and μ_x (Appendix A.3), denote \hat{R} as

$$\hat{R} = \frac{\hat{\mu}_{y\text{HT}}}{\hat{\mu}_{x\text{HT}}} = \frac{\sum_{i \in \mathbf{S}} d_i y_i}{\sum_{i \in \mathbf{S}} d_i x_i}$$

This implies the generalized ratio estimator as

$$\hat{\mu}_{y\text{GR}} = \hat{R}\mu_x = \left(\frac{\sum_{i \in \mathbf{S}} d_i y_i}{\sum_{i \in \mathbf{S}} d_i x_i}\right) \mu_x$$

where μ_x is known.

We can verify that the generalized ratio estimator is still a model-assisted estimator. Under the simple linear regression model, the expectation of HT estimator of y is

$$E_{\xi}\left(\hat{\mu}_{y\mathrm{HT}}\right) = \frac{1}{N} \sum_{i \in \mathbf{S}} \frac{E_{\xi}[y_i]}{\pi_i} = \frac{1}{N} \sum_{i \in \mathbf{S}} \frac{\beta x_i}{\pi_i} = \beta \hat{\mu}_{x\mathrm{HT}}$$

Also

$$E_{\xi}\left(\hat{\mu}_{y\text{GR}}\right) = \frac{\beta \hat{\mu}_{x\text{HT}}}{\hat{\mu}_{x\text{HT}}} \mu_{x} = \beta \mu_{x}$$

Since we also know that $E_{\xi}(\mu_y)=\beta\mu_x$, it implies $E_{\xi}\left(\hat{\mu}_{y\mathrm{GR}}-\mu_y\right)=0$.

To prove $\hat{\mu}_{y\text{GR}}$ is approximately unbiased under the sampling design. By linearization, \hat{R}

can be expressed as

$$\hat{R} = \frac{\hat{\mu}_{y\mathrm{HT}}}{\hat{\mu}_{x\mathrm{HT}}} = R + \frac{1}{\mu_x} \left(\hat{\mu}_{y\mathrm{HT}} - R \hat{\mu}_{x\mathrm{HT}} \right) + o_p \left(\frac{1}{\sqrt{n}} \right) \tag{4.1.1}$$

It follows that

$$\begin{split} E_p(\hat{R}) &= E_p \left(R + \frac{1}{\mu_x} \left(\hat{\mu}_{y\text{HT}} - R \hat{\mu}_{x\text{HT}} \right) + o_p \left(\frac{1}{\sqrt{n}} \right) \right) \\ &= R + \frac{1}{\mu_x} \left(\mu_y - R \mu_x \right) + o_p \left(\frac{1}{\sqrt{n}} \right) \\ &\doteq R \end{split}$$

Similarly, the variance of \hat{R} over the sampling desing can be written as

$$\begin{split} V_p(\hat{R}) &= V_p \left(R + \frac{1}{\mu_x} \left(\hat{\mu}_{y\text{HT}} - R \hat{\mu}_{x\text{HT}} \right) + o_p \left(\frac{1}{\sqrt{n}} \right) \right) \\ &\doteq \frac{1}{\mu_x^2} V_p \left(\hat{\mu}_{y\text{HT}} - R \hat{\mu}_{x\text{HT}} \right) \\ &= \frac{1}{\mu_x^2} V_p \left(\frac{1}{N} \sum_{i \in \mathbf{S}} d_i y_i - \frac{R}{N} \sum_{i \in \mathbf{S}} d_i x_i \right) \\ &= \frac{1}{\mu_x^2} V_p \left(\frac{1}{N} \sum_{i \in \mathbf{S}} d_i (y_i - R x_i) \right) \\ &= \frac{1}{\mu_x^2} V_p \left(\hat{\mu}_{e\text{HT}} \right) \end{split}$$

where $\hat{\mu}_{e\text{HT}} = N^{-1} \sum_{i \in \mathbf{S}} d_i e_i$ and $e_i = y_i - Rx_i$. Since $\hat{\mu}_{y\text{GR}} = \hat{R}\mu_x$, the above results imply that

$$E_{n}\left(\hat{\mu}_{\nu \text{GR}}\right) \doteq \mu_{\nu} \quad V_{n}\left(\hat{\mu}_{\nu \text{GR}}\right) \doteq V_{n}\left(\hat{\mu}_{e \text{HT}}\right)$$

In the general unequal probability sampling, the variance of $V_{p}\left(\hat{\mu}_{e\mathrm{HT}}\right)$ can be expressed as

$$V_p\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq V_p\left(\hat{\mu}_{e\mathrm{HT}}\right) = \frac{1}{N^2} \sum_{i=1}^N \sum_{j=1}^N \left(\pi_{ij} - \pi_i \pi_j\right) \frac{e_i}{\pi_i} \frac{e_j}{\pi_j} \tag{4.1.2}$$

and

$$v_p\left(\hat{\mu}_{y\text{GR}}\right) \doteq v_p\left(\hat{\mu}_{e\text{HT}}\right) = \frac{1}{\hat{N}^2} \sum_{i \in \mathbf{S}} \sum_{j \in \mathbf{S}} \frac{\pi_{ij} - \pi_i \pi_j}{\pi_{ij}} \frac{\hat{e}_i}{\pi_i} \frac{\hat{e}_j}{\pi_j} \tag{4.1.3}$$

where $\hat{N} = \sum_{i \in \mathbf{S}} \frac{1}{\pi_i} = \sum_{i \in \mathbf{S}} d_i$ and can be replaced by N if the population size is known (Wu & Thompson, 2020, pp. 100–103).

4.2 Other PPS Sampling Method

The linearization approach described above is applicable to general unequal probability sampling designs that involve second-order inclusion probabilities, such as RSPPS. However, as discussed in Sections 3.2 and 3.3, PPS sampling with replacement and Poisson sampling involve different estimation metrics for the population total and is not necessarily to have second order inclusion probabilities. Accordingly, the variance estimation for the generalized ratio estimator must also be adapted to remain consistent with the underlying sampling design.

In the case of Poisson sampling, each unit is drawn independently. As a result, the cross-product terms present in the Yates–Grundy–Sen form of the variance $V_p(\hat{\mu}_{e\mathrm{HT}})$ in equation 4.1.2 and 4.1.3 vanishes and is replaced by a simpler expression as shown in equation 3.3.1 and 3.3.2. On the other hand, in section 3.3, we have seen that $\hat{T}_{y\mathrm{HT}}$ under Poisson sampling is unbiased of T_y (Appendix A.5). This implies that \hat{R} in equation 4.1.1 is still approximately

design-unbiased of R. Following the same linearization procedures in section 4.1, the variance of generalized ratio estimator under Poisson sampling becomes

$$V_{p}\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq V_{p}\left(\hat{\mu}_{e\mathrm{HT}}\right) = \frac{1}{N^{2}} \sum_{i=1}^{N} \frac{1-\pi_{i}}{\pi_{i}} e_{i}^{2}$$

with the corresponding variance estimator given by

$$v_p\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq v_p\left(\hat{\mu}_{e\mathrm{HT}}\right) = \frac{1}{\hat{N}^2} \sum_{i \in \mathbf{S}} \frac{1 - \pi_i}{\pi_i^2} \hat{e}_i^2$$

Comparing to randomized systematic PPS and Poisson sampling, as discussed in section 3.2, PPS sampling with replacement requires the use of the Hansen–Hurwitz estimator rather than the Horvitz–Thompson estimator to estimate the population total. Since both $\hat{\mu}_{y\text{HH}}$ and $\hat{\mu}_{y\text{HH}}$ are unbiased for μ_y and μ_x respectively, the generalized ratio estimator under PPS sampling with replacement is constructed as

$$\hat{\mu}_{y\text{GR}} = \left(\frac{\hat{\mu}_{y\text{HH}}}{\hat{\mu}_{x\text{HH}}}\right) \mu_x = \left(\frac{\sum_{i \in \mathbf{S}} \frac{y_i}{nz_i}}{\sum_{i \in \mathbf{S}} \frac{x_i}{nz_i}}\right) \mu_x$$

Therefore, $\hat{R} = \frac{\hat{\mu}_{y\text{HH}}}{\hat{\mu}_{x\text{HH}}}$ is still approximately design-unbiased for R. Using linearization similarly as before, we can obtain

$$E_{p}\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq \mu_{y} \quad V_{p}\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq V_{p}\left(\hat{\mu}_{e\mathrm{HH}}\right)$$

where $\hat{\mu}_{e\text{HH}} = N^{-1} \sum_{i \in \mathbf{S}} w_i e_i$ with $w_i = 1/nz_i$ and $e_i = y_i - Rx_i$. The theoretical variance

of the generalized ratio estimator under this method is then

$$V_{p}\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq V_{p}\left(\hat{\mu}_{e\mathrm{HH}}\right) = \frac{1}{N^{2}}\frac{1}{n}\sum_{i=1}^{N}z_{i}\left(\frac{e_{i}}{z_{i}} - T_{e}\right)^{2}$$

with an unbiased estimator of the variance is given by

$$v_p\left(\hat{\mu}_{y\mathrm{GR}}\right) \doteq v_p\left(\hat{\mu}_{e\mathrm{HH}}\right) = \frac{1}{N^2} \frac{1}{n(n-1)} \sum_{i \in \mathbf{S}^*} \left(\frac{\hat{e}_i}{z_i} - \hat{T}_{e\mathrm{HH}}\right)^2$$

where $\hat{e}_i = y_i - \hat{R}x_i$ and \mathbf{S}^* denotes the sample with duplicated units.

5 Bootstrap and Jackknife in GRE

The previous section introduced the way to to estimate the variance of generalized ratio estimator using linearization. In addition to that, the variance can be also estimated using Bootstrap and Jackknife resampling methods.

For the bootstrap approach, suppose we have the sample $\mathbf{S}=(X_1,\dots,X_n)$ taken from the population using randomized systematic PPS sampling, the procedure to estimate the variance of $\hat{\mu}_{y\text{GR}}$ using Bootstrap works as follows:

- 1. Let S^* be the set of units selected including duplicated units from the S using SRSWR.
- 2. Obtain the bootstrap sample data S^* from the original data S such that $\{(y_i, x_i), i \in S^*\}$

3. Compute $\hat{\mu}_{y\text{GR}}$ using the bootstrap sample

$$\hat{\mu}_{y\text{GR}}^* = \left(\frac{\hat{\mu}_{y\text{HT}}^*}{\hat{\mu}_{x\text{HT}}^*}\right) \mu_x$$

where $\hat{\mu}^*_{y\text{HT}} = \sum_{i \in \mathbf{S}^*} d_i y_i$ and $\hat{\mu}^*_{x\text{HT}} = \sum_{i \in \mathbf{S}^*} d_i x_i$ with $d_i = 1/\pi_i$.

4. Repeat step 1 to 3 for B times to have $(\hat{\mu}_{yGR}^*(1), \dots \hat{\mu}_{yGR}^*(B))$, and then calculate the bootstrap variance estimator of $\hat{\mu}_{yR}$ by

$$v_B\left(\hat{\mu}_{y\mathrm{GR}}\right) = \frac{1}{B} \sum_{b=1}^{B} \left\{ \hat{\mu}_{y\mathrm{GR}}^*(b) - \hat{\mu}_{y\mathrm{GR}} \right\}^2$$

For Poisson sampling, the bootstrap procedure is identical to the above as the Horvitz—Thompson estimator is also used to estimate the population total. However, for PPS sampling with replacement, the procedure is slightly different due to the usage of Hansen–Hurwitz estimator. In particular, Step 3 changes to:

$$\hat{\mu}_{y\text{GR}}^* = \left(\frac{\hat{\mu}_{y\text{HH}}^*}{\hat{\mu}_{x\text{HH}}^*}\right) \mu_x$$

where
$$\hat{\mu}_{y\text{HH}}^* = \sum_{i \in \mathbf{S}^*} \frac{y_i}{nz_i}$$
, $\hat{\mu}_{x\text{HH}}^* = \sum_{i \in \mathbf{S}^*} \frac{x_i}{nz_i}$.

In addition to Bootstrap, the delete-1 Jackknife offers an alternative resampling method for estimating the variance of the generalized ratio estimator. However, unlike the standard Jackknife described in Appendix A.2.2, generalized ratio estimator now incorporates design weights through either the Hansen–Hurwitz or Horvitz–Thompson estimators. It is necessary to adjust for the weights within each replicate sample as well.

For the randomized systematic PPS sampling, it is suggested that the total weight for each Jackknife replicate preserves the original estimated total $\hat{N} = \sum_{i \in \mathbf{S}} d_i$ (Rust & Rao, 1996, pp. 287–289; Wolter, 2007). Specifically, if the unit i is excluded in a Jackknife replicate, the replicate weights should be defined as

$$\begin{cases} 0 & j=i \\ \frac{\hat{N}}{\hat{N}-d_i}d_j & j \neq i \end{cases}$$

which ensures $\sum_{j \in \mathbf{S}^*} d_j = \sum_{j \in \mathbf{S}^*} \frac{\hat{N}}{\hat{N} - d_i} d_j = \frac{\hat{N}}{\hat{N} - d_i} (\hat{N} - d_i) = \hat{N}$.

By the refined replicate weights, the complete Jackknife procedure works as follows

1. Compute the delete-1 Jackknife estimator

$$\hat{\mu}_{y\text{GR}[-i]} = \frac{\hat{\mu}_{y\text{HT}[-i]}}{\hat{\mu}_{x\text{HT}[-i]}} \mu_x, \quad i = 1, 2, \cdots, n$$

where

$$\begin{split} \hat{\mu}_{y\text{HT}[-i]} &= \frac{\sum_{j \neq i} d_j^{(-i)} y_j}{\sum_{j \neq i} d_j^{(-i)}}, \quad \hat{\mu}_{x\text{HT}[-i]} = \frac{\sum_{j \neq i} d_j^{(-i)} x_j}{\sum_{j \neq i} d_j^{(-i)}} \\ \text{s.t.} \quad d_j^{(-i)} &= \begin{cases} 0 & j = i \\ \frac{\hat{N}}{\hat{N} - d_i} d_j & j \neq i \end{cases} \end{split}$$

where
$$\hat{N} = \sum_{i \in \mathbf{S}} d_i$$
, $d_i = \frac{1}{\pi_i}$

2. Compute the Jackknife variance estimator as

$$v_J\left(\hat{\mu}_{y\mathrm{GR}}\right) = \frac{n-1}{n} \sum_{i=1}^n \left(\hat{\mu}_{y\mathrm{GR}[-i]} - \hat{\mu}.\right)^2$$

where
$$\hat{\mu}_{\cdot} = \frac{1}{n} \sum_{i=1}^{n} \hat{\mu}_{y GR[-i]}$$

As usual, Poisson sampling follows the same Jackknife algorithm as randomized systematic PPS sampling since both designs are based on first-order inclusion probabilities and HT-type estimators. For PPS sampling with replacement, the Hansen–Hurwitz estimator is used and the Jackknife procedure must be modified accordingly. In this case, Step 1 is adjusted to:

$$\hat{\mu}_{y\text{GR}[-i]} = \frac{\hat{\mu}_{y\text{HH}[-i]}}{\hat{\mu}_{x\text{HH}[-i]}} \mu_x, \quad i = 1, 2, \cdots, n$$

where $d_i = \frac{1}{nz_i}$ are the HH design weights.

6 Simulation

This simulation study is adopted from Changbao and Thompson (2020), where we firstly simulate the finite population data and then apply each of the three PPS sampling methods to generate corresponding samples. The objective of this study is to compare variance estimates obtained by the three proposed methods which are linearization, Bootstrap and Jackknife, under the three PPS sampling design respectively. In addition, this simulation study will also compute the confidence interval of μ_y based on the generalized ratio estimator, and evaluate their performance by comparing them to the true value of μ_y . All the R codes in the simulation can be found at Appendix A.6.

6.1 Simulation design

In this study, the covariates is assumed to be univariate and is generated from the uniform distribution $X_i \sim \text{Unif}(0,1)$ with size variables generated from an exponential distribution shifted by 0.5, $Z_i \sim \exp(1) + 0.5$. The outcome y_i is generated according to the model

$$y_i = 1 + 2x_i + 2.5z_i + \varepsilon_i$$

where ε_i is an independent error term drawn from a standard normal distribution, $\varepsilon_i \sim \mathcal{N}(0,1)$. In addition, the population size is set to N=5000, and the sample size n=500. The R codes for generating the population data can be found at Appendi A.6.1.

Table 1: Summary statistics of the population data for outcome y, covariate x and size z

Variable	Mean	SD	Min	Median	Max
y	5.767	2.714	-0.603	5.257	24.549
x	0.512	0.288	0.000	0.520	1.000
z	1.500	0.999	0.500	1.193	8.988

Table 1 presents the summary statistics for each variable in the population data. The true population mean μ_y is 5.767 with standard deviation 2.714. Later in this section, we estimate μ_y using the generalized ratio estimator using different methods and compare the results to the true value.

Table 2: Summary statistics of the outcome y in each of the three sample data.

Saample	SampleSize	Mean	SD	Min	Median	Max
RSPPS	100	7.229	3.794	2.445	6.059	18.713
Poisson	105	7.020	3.164	1.482	6.349	20.958
PPS-WR	100	7.560	3.711	1.122	6.966	20.083

Consequently, Table 2 presents the summary statistics of the outcome y in each of the three samples. Notably, the sample size under Poisson sampling is 105, which exceeds the expected sample size of 100. In addition, the estimated population mean $\hat{\mu}_y$ across all samples are all above 7, which deviates significantly from the true population mean 5.767. These observations all underscore the necessity of employing Horvitz-Thompson (HT) or Hansen-Hurwitz (HH) estimators to obtain unbiased estimation of the population mean.

6.2 Simulation metrics

As we discussed in Section 4 and 5, the way to estimate the variance for linearization, Bootstrap and Jackknife under each PPS sampling method is distinct due to different ways to draw the sample. In this simulation study, we follow the estimation procedures described in those sections precisely to estimate μ_y and its variance using generalized ratio estimator. However, the only exception is the linearization under randomized systematic PPS sampling. As mentioned in section 4.1, the estimation of second order inclusion probabilities π_{ij} is challenging

in practice. Although Thompson and Wu (2008) proposed a simulation-based approach to estimate π_{ij} , it requires long time to generate the full matrix of joint inclusion probabilities as the number of simulation is about 1,000,000 (Thompson & Wu, 2008). Therefore, this study will not consider to compute π_{ij} directly; instead, the variance is approximated using the Hansen–Hurwitz estimator under randomized systematic PPS sampling. This is also discussed in section 3.2.

To construct the confidence interval of $\hat{\mu}_y$, this paper proceed as follows. For linearization method, we adopt the approach recommended by Wu and Thompson (2020, p.103), to build the confidence interval as

$$\left(\hat{\mu}_{y\mathrm{GR}} - Z_{\alpha/2} \left[v_p \left(\hat{\mu}_{y\mathrm{GR}} \right) \right]^{1/2}, \quad \hat{\mu}_{y\mathrm{GR}} + Z_{\alpha/2} \left[v_p \left(\hat{\mu}_{y\mathrm{GR}} \right) \right]^{1/2} \right)$$

where $Z_{\alpha/2}$ is the $1-\alpha/2$ quantile of standard normal distribution, and α is the significance level. For the confidence interval of Bootstrap and Jackknife, this paper will use the Bootstrap percentile approach (Shao & Tu, 1995, pp. 132–133). That is, if $\alpha=0.05$, the 95% confidence interval is constructed by taking the 2.5th and 97.5th percentiles of the bootstrap or jackknife resampled estimates.

6.3 Simulation results

Table 3: Point Estimates and confidence intervals for μ_y via linearization for each sampling design

Method	Sampling	$v_p(\hat{\mu}_{y\mathrm{GR}})$	$\hat{\mu}_{y\mathrm{GR}}$	CI	Width
Linear.	RSPPS	0.211	5.596	(4.70, 6.50)	1.80
Linear.	Poisson	0.182	6.078	(5.24, 6.91)	1.67
Linear.	PPS_WR	0.177	5.732	(4.91, 6.56)	1.65

The R codes for this section can be found at Appendix A.6.2 to Appendix A.6.4. Table 3 shows the estimated variance of generalized ratio estimator of μ_y via linearization as described in section 5 under the three PPS sampling methods. It seems that the PPS sampling with replacement has the best performance among the three sampling methods as it has the lowest variance and the narrowest confidence interval. More importantly, this approach gives a very close point estimate $\hat{\mu}_y$ to the true population mean (5.732 vs. 5.767). For the rest two methods, the randomized systematic PPS sampling seems to have worst performance which has a estimated variance significantly larger than the other two.

In the case of Bootstrap and Jackknife, Figure 1 shows the distribution of $\hat{\mu}_{y\text{GR}}$ under the three PPS sampling designs. We can observed that the estimates based on PPS sampling with replacement consistently have the best performance among the three methods across both Bootstrap and Jackknife, as the estimates are closest to the true population mean (red dashed line). This aligns our observation from linearization. In contrast, estimates from Poisson sampling show the poorest performance, with greater deviation from the true

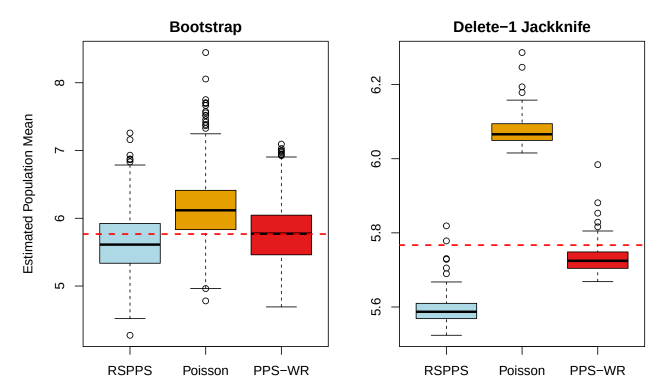


Figure 1: Booxplot of distribution of the estimated population mean using generalized ratio estimator via Bootstrap (left) and Delete-1 Jackknife (right) under the three PPS sampling method; red dashed line is the true population mean

population mean compared to randomized systematic PPS sampling. We can also observe the variance of $\hat{\mu}_{y\text{GR}}$ from Figure 1, where the estimated variance $v_p(\hat{\mu}_{y\text{GR}})$ for the three methods are approximately same in both Bootstrap and Jackknife as they have similar interquartile ranges IQR in the boxplots.

The patterns from Figure 1 can be verified from the Table 4, which presents the estimated variance of $\hat{\mu}_{y\text{GR}}$ and confidence intervals under both resampling methods. For each case, the estimated variance is around 0.2 with PPS sampling with replacement always return the lowest variance. As a result, PPS sampling with replacement also produces the narrowest confidence intervals across all comparisons. Comparing the three methods, the linearization under the PPS sampling with replacement seems to have the best performance among all.

Table 4: Point Estimates and confidence intervals for μ_y via Bootstrap and Jackknife for each sampling design

Method	Sampling	$v_p(\hat{\mu}_{y\mathrm{GR}})$	$\hat{\mu}_{y\mathrm{GR}}$	CI	Width
Bootstrap	RSPPS	0.200	5.596	(4.84, 6.59)	1.75
Bootstrap	Poisson	0.215	6.078	(5.31, 7.06)	1.75
Bootstrap	PPS_WR	0.192	5.732	(4.97, 6.69)	1.72
Jackknife	RSPPS	0.217	5.596	(4.68, 6.51)	1.83
Jackknife	Poisson	0.204	6.078	(5.19, 6.96)	1.77
Jackknife	PPS_WR	0.196	5.732	(4.86, 6.60)	1.73

7 Discussion

This paper shows how to estimate the population mean μ_y by the sample drawn from PPS sampling via a generalized ratio estimator, with the implementation of three variance estimation methods: linearization, Bootstrap, and Jackknife. From the simulation study, the performance of each method under each PPS sampling design is satisfied, especially the point estimate of μ_y for each method, which is much closer to the true population mean compared to the original sample mean from Table 2. PPS sampling with replacement has the best overall performance among all the methods. Notably, it returns the estimate $\hat{\mu}_y$, which is almost identical to the true population mean. However, each method has its drawbacks and must be addressed carefully.

In general, the variance estimates obtained using linearization are smaller than those produced by the bootstrap and delete-1 jackknife methods. This is expected as linearization is a parametric way to estimate the variance, while the other two are non-parametric. However, as noted earlier, linearization for randomized systematic PPS sampling requires second-order inclusion probabilities, which do not have a closed-form expression and are typically estimated through simulation-based methods (Thompson & Wu, 2008). In this study, we approximate variance estimation by using the Hansen–Hurwitz (HH) estimator from PPS sampling with replacement. Furthermore, the methods from PPS sampling with replacement have the best performance and are not coincidental. Because units are selected independently with probabilities proportional to their size measures, the sampling process repeatedly selects units with higher inclusion probabilities. This mechanism better preserves the underlying data variability than randomized systematic PPS or Poisson sampling designs.

Bootstrap will be more challenging to implement when we have non-iid samples. The main reason is that simply resampling the original data will break the data structure when the observations are dependent. As a result, the empirical distribution from the bootstrap sample will no longer be a good approximation to the true distribution, and it may also be a biased or inconsistent estimate of the sample distribution. In the randomized systematic PPS sampling design, the data are dependent on each other as it includes the second-order inclusion probabilities. This means that Bootstrap is theoretically not applicable here or can only be used when the fraction of sample size to population is very small. In our simulation design, the fraction is set to be 100/5000 = 0.02. While it is small, the dependence structure may still exist and potentially cause bias when implementing Bootstrap. We can actually

verify this from Table 4, where the variance generated from Bootstrap is higher than the other two. In contrast, PPS sampling with replacement is not affected by the sampling fraction, as units are drawn independently with replacement. This would preserve the validity of Bootstrap methods (Rao et al., 1992; Rao & Wu, 1988; Sitter, 1992).

In such cases, one possible solution is to apply a moving block bootstrap, which divides the data into blocks and resamples overlapping blocks of consecutive data with replacement to preserve local dependence (Liu & Singh, 1992; Shao & Tu, 1995, pp. 391–392). In addition, Shao and Tu also introduced the residual bootstrap method (Shao & Tu, 1995, pp. 395–396). Furthermore, the modern Bootstrap theory has few developments on complex surveys. For example, Rao and Wu (Rao & Wu, 1988) proposed the Rao-Wu bootstrap method, which deals with the stratified multi-stage design and with replacement. In addition, another method, known as the Mirror-Match Bootstrap, was proposed by Sitter (Sitter, 1992), which tried to mimic the original data structure.

In the case of delete-1 Jackknife, one of the main drawbacks is that the variance estimator v_J is only consistent when $\hat{\theta}$ is a smooth function of the sample mean or several sample means (Wu & Thompson, 2020, p. 241). The "smooth" estimator here is defined as the change of estimates that will not be so significant when removing one or some observation, such as sample mean. In contrast, "non-smooth" parameters such as median or other quantiles may change dramatically even when removing one observation (depending on the data structure). To deal with this, Shao and Tu suggested the more generalized delete-d Jackknife estimator v_{J-d} where d>1 and argue that it requires less stringent smoothness condition than the delete-1 Jackknife estimator (Shao & Tu, 1995, pp. 49–50). They also argue that the less

smooth the parameter, the higher order d is needed (Shao & Tu, 1995, p. 69).

Future research should focus on using the refined version of Bootstrap mentioned above in the PPS sampling design. It is also important to investigate the use of generalized delete-d Jackknife estimators in the context of PPS sampling, particularly to address issues related to non-smooth parameters. In addition, the estimation of second-order probabilities in PPS sampling remains challenging. Although Thompson and Wu (2008) proposed a simulationbased method, it is computationally intensive. Future research may focus on developing more efficient and practical methods for estimating second-order probabilities in practice.

A Appendix

A.1 Expectation and variance of simple ratio etimator using linearization

We can use linearization to estimate the variance of the simple ratio estimator. Since $\hat{\mu}_{yR} = \frac{\bar{y}}{\bar{x}}\mu_x$ with μ_x known, its variance can be expressed as $\mathrm{Var}_p(\hat{\mu}_{yR}) = \mu_x^2\,\mathrm{Var}_p(\frac{\bar{y}}{\bar{x}})$. Denote $R = \frac{\mu_y}{\mu_x}$ and $\hat{R} = \frac{\bar{y}}{\bar{x}}$. Using the linearization, we can express $\frac{\bar{y}}{\bar{x}}$ as

$$\hat{R} = \frac{\bar{y}}{\bar{x}} = \frac{\mu_y}{\mu_x} + \frac{1}{\mu_x} \left(\bar{y} - \mu_y \right) - \frac{\mu_y}{\mu_x^2} \left(\bar{x} - \mu_x \right) + \text{ higher order terms}$$

Since we know that $\bar{y} - \mu_y = O_p\left(\frac{1}{\sqrt{n}}\right)$ and $\bar{x} - \mu_x = O_p\left(\frac{1}{\sqrt{n}}\right)$, it can be further expressed as

$$\hat{R} = \frac{\bar{y}}{\bar{x}} = \frac{\mu_y}{\mu_x} + \frac{1}{\mu_x}(\bar{y} - R\bar{x}) + o_p\left(\frac{1}{\sqrt{n}}\right)$$

This implies $E_p\left(\hat{\mu}_{y\mathrm{R}}\right) \doteq \mu_y$. Consequently, the variance of $\mathrm{Var}(\hat{\mu}_{y\mathrm{R}})$ can be written as

$$\begin{split} \operatorname{Var}(\hat{\mu}_{y\mathbf{R}}) &= \mu_x^2 \operatorname{Var}(\frac{\bar{y}}{\bar{x}}) \\ &\doteq \mu_x^2 \frac{1}{\mu_x^2} V_p(\bar{y} - R\bar{x}) \\ &= \left(1 - \frac{n}{N}\right) \frac{1}{n} \left(\sigma_y^2 + R^2 \sigma_x^2 - 2R \sigma_{xy}\right) \end{split}$$

where $(1 - \frac{n}{N}) \frac{1}{n}$ comes from the SRSWOR.

A.2 The general algorithm for Bootstrap and delete-1 Jackknife

A.2.1 Bootstrap

Bootstrap is another non-parametric method that uses the sample we have, either draw by SRSWOR or other sampling methods, to buuld many re-samples and compute the estimates of a given estimator and then compute the variances. The general standard bootstrap procedures (Wu & Thompson, 2020, pp. 224–225) work as follows:

- 1. Take bootstrap samples (X_1^*, \dots, X_n^*) from the original sample (X_1, \dots, X_n) using SR-SWR.
- 2. Estimate the $\hat{\theta}$ using the bootstrap sample

$$\hat{\theta}^* = h\left(X_1^*, X_2^*, \cdots, X_n^*\right)$$

- 3. Repeat step 1 and 2 for B times independently to have $(\hat{\theta}_1^*, \dots, \hat{\theta}_B^*)$.
- 4. Use the $\{\hat{\theta}_b^*\}_{b=1}^B$ to obtain the bootstrap variance estimator for $\hat{\theta}$ by

$$v_B = \frac{1}{B} \sum_{b=1}^B \left(\hat{\theta}_b^* - \bar{\theta}^* \right)^2$$

where $\bar{\theta}^* = B^{-1} \sum_{b=1}^B \hat{\theta}_b^*$ or $\bar{\theta}^* = \hat{\theta}$. We can also replace B^{-1} by $(B-1)^{-1}$; however, the difference is ignorable for large B.

We can compute the variance of simple ratio estimator using bootstrap as well. It just replaces $\hat{\theta}$ by $\hat{\mu}_{yR}$ and have $\hat{\mu}_{yR}^* = \frac{\bar{y}^*}{\bar{x}^*} \mu_x$ in step 2. After we have B estimates $\hat{\mu}_{yR}$,

 $(\hat{\mu}_{uR}^*(1), \dots \hat{\mu}_{uR}^*(B))$, we can estimate its variance by

$$v_B\left(\hat{\mu}_{yR}\right) = \frac{1}{B} \sum_{b=1}^{B} \left\{ \hat{\mu}_{yR}^*(b) - \hat{\mu}_{yR} \right\}^2$$

A.2.2 Jackknife

The second resampling method is Jackknife, and we will only discuss the delete-1 Jackknife estimator. Compared to Bootstrap, which resamples the entire sample and has a new sample of size exactly n at each simulation, delete-1 Jackknife simply removes one unit every time across all units, resulting in n sub-samples of size n-1 (Wu & Thompson, 2020, p. 218). In addition, unlike Bootstrap, Jackknife is only applicable to estimator $\hat{\theta}$ that is the sample mean or a function of sample mean (Shao & Tu, 1995, p. 69); we will firstly discuss the delete-1 Jackknife estimator $\hat{\mu}$ and then extend to some general estimator $\hat{\theta}$.

For any general estimator $\hat{\theta}$, the general Jackknife procedures to compute the estimated variance $v(\hat{\theta})$ works as follows:

1. Compute the delete-1 estimators

$$\hat{\theta}_{-i} = h\left(X_1, \cdots, X_{i-1}, X_{i+1}, \cdots, X_n\right), \quad i = 1, 2, \cdots, n$$

2. Compute the delete-1 jackknife variance estimator by

$$v_J = \frac{n-1}{n} \sum_{i=1}^n \left(\hat{\theta}_{-i} - \hat{\theta}_{\cdot} \right)^2$$

where
$$\hat{\theta} = n^{-1} \sum_{i=1}^{n} \hat{\theta}_{-i}$$

We can extend this idea to simple ratio estimator, which follows as

1. Compute the delete-1 Jackknife estimator

$$\hat{\mu}_{-i} = \frac{\bar{y}_{[-i]}}{\bar{x}_{[-i]}} \mu_x, \quad i = 1, 2, \cdots, n$$

where
$$\bar{y}_{[-i]} = \frac{1}{n-1} \left(\sum_{j=1}^{n} y_j - y_i \right) = \frac{1}{n-1} \left(n\bar{y} - y_i \right)$$
 and similarly $\bar{x}_{[-i]} = \frac{1}{n-1} \left(n\bar{x} - x_i \right)$

2. Compute the Jackknife variance estimator as

$$v_{J}\left(\hat{\mu}_{yR}\right) = \frac{n-1}{n} \sum_{i=1}^{n} \left(\hat{\mu}_{-i} - \hat{\mu}_{\cdot}\right)^{2}$$

where
$$\hat{\mu}_{\cdot} = \frac{1}{n} \sum_{i=1}^{n} \hat{\mu}_{-i}$$

A.3 Hortitz-Thompson estimators in randomized systematic PPS sampling

Denote the inclusion indicator A_i such that $E(A_i)=\pi_i$. We can derive

$$E\left(\hat{T}_{y\mathrm{HT}}\right) = E\left(\sum_{i=1}^{N} A_i \frac{y_i}{\pi_i}\right) = \sum_{i=1}^{N} E(A_i) \frac{y_i}{\pi_i} = \sum_{i=1}^{N} y_i = T_y$$

$$V\left(\hat{T}_{y \text{HT}}\right) = V\left(\sum_{i=1}^{N} A_i \frac{y_i}{\pi_i}\right) = \sum_{i=1}^{N} \sum_{j=1}^{N} \text{Cov}(A_i, A_j) \frac{y_i}{\pi_i} \frac{y_j}{\pi_j} = \sum_{i=1}^{N} \sum_{j=1}^{N} (\pi_{ij} - \pi_i \pi_j) \frac{y_i}{\pi_i} \frac{y_j}{\pi_j}$$

$$\begin{split} E\left[v\left(\hat{T}_{y\text{HT}}\right)\right] &= E\left[\sum_{i \in S} \sum_{j \in S} \frac{\pi_{ij} - \pi_i \pi_j}{\pi_{ij}} \frac{y_i}{\pi_i} \frac{y_j}{\pi_j}\right] \\ &= E\left[\sum_{i = 1}^N \sum_{j = 1}^N A_i A_j \frac{\pi_{ij} - \pi_i \pi_j}{\pi_{ij}} \frac{y_i}{\pi_i} \frac{y_j}{\pi_j}\right] \\ &= \sum_{i = 1}^N \sum_{j = 1}^N \pi_{ij} \frac{\pi_{ij} - \pi_i \pi_j}{\pi_{ij}} \frac{y_i}{\pi_i} \frac{y_j}{\pi_j} \\ &= \sum_{i = 1}^N \sum_{j = 1}^N (\pi_{ij} - \pi_i \pi_j) \frac{y_i}{\pi_i} \frac{y_j}{\pi_j} \end{split}$$

A.4 Hansen–Hurwitz estimators in PPS sampling with replacement

Denote R_i as $\frac{Y_i}{Z_i}$ such that $E(R_i) = T_y$ and $V(R_i) = \sum_{i=1}^N \left(\frac{y_i}{z_i} - T_y\right)^2 z_i$, then we have

$$E\left(\hat{T}_{y\mathrm{HH}}\right) = \frac{1}{n} \sum_{i=1}^{n} E(R_i) = T_y$$

$$\begin{split} V\left(\hat{T}_{y\mathrm{HH}}\right) &= V\left(\frac{1}{n}\sum_{i=1}^{n}R_{i}\right) \\ &= \frac{1}{n^{2}}\sum_{i=1}^{n}V(R_{i}) \\ &= \frac{1}{n^{2}}\sum_{i=1}^{n}(y_{i}/z_{i}-T_{y})^{2}z_{i} \\ &= \frac{1}{n}\sum_{i=1}^{N}z_{i}\left(\frac{y_{i}}{z_{i}}-T_{y}\right)^{2} \end{split}$$

$$(\hat{T}_{z\mathrm{HH}}) = \frac{1}{n^{2}}\sum_{i=1}^{N}A_{i}\left(\frac{y_{i}}{z_{i}}-T_{y}\right)^{2} + \frac{1}{n^{2}}\sum_{i=1}^{N}A_{i}\left(\frac{y_{i}}{z_{i}}-T_{y$$

$$\begin{split} E\left(v\left(\hat{T}_{y\mathrm{HH}}\right)\right) &= \frac{1}{n(n-1)} \sum_{i=1}^{N} A_i \left(\frac{y_i}{z_i} - \hat{T}_{y\mathrm{HH}}\right)^2 \\ &= \frac{1}{n} \sum_{i=1}^{N} z_i \left(\frac{y_i}{z_i} - T_y\right)^2 \end{split}$$

A.5 Hortitz-Thompson estimators in Poisson sampling

Denote the inclusion indicator A_i such that $E(A_i) = \pi_i$. We can derive

$$\begin{split} E\left(\hat{T}_{y\mathrm{HT}}\right) &= E\left(\sum_{i=1}^{N}A_{i}\frac{y_{i}}{\pi_{i}}\right) = \sum_{i=1}^{N}E(A_{i})\frac{y_{i}}{\pi_{i}} = \sum_{i=1}^{N}y_{i} = T_{y} \\ V(\hat{T}_{y\mathrm{HT}}) &= V\left(\sum_{i=1}^{N}A_{i}\frac{y_{i}}{\pi_{i}}\right) \\ &= \sum_{i=1}^{N}V(A_{i})\left(\frac{y_{i}}{\pi_{i}}\right)^{2} \\ &= \sum_{i=1}^{N}\left(\frac{1-\pi_{i}}{\pi_{i}}\right)y_{i}^{2} \\ E(v(\hat{T}_{y\mathrm{HT}})) &= \sum_{i=1}^{N}A_{i}\frac{1-\pi_{i}}{\pi_{i}^{2}}y_{i}^{2} \\ &= \sum_{i=1}^{N}\left(\frac{1-\pi_{i}}{\pi_{i}}\right)y_{i}^{2} \end{split}$$

A.6 R codes

This section contains all the R codes used in this project. If you want to see how the results are implemented in the main text such as Table 1, please visit my GitHub page (here).

A.6.1 Generate the population data and draw samples under the three PPS sampling methods

The R codes for generating the population and sample data

```
# Generate the population data
set.seed(854)
N <- 5000</pre>
```

```
n <- 100
x <- runif(N)
z \leftarrow 0.5 + rexp(N)
y \leftarrow 1 + 2 * x + 2.5 * z + rnorm(N)
pdata <- cbind(y, x, z)</pre>
# 1. Randomized Systematic PPS Sampling Method
syspps <- function(x, n){</pre>
 N = length(x) # The population size
 U = sample(N,N) # Sample the index without replacement
 xx = x[U] # Find the corresponding size values
 z = rep(0,N) # Initial the size variable
 for(i in 1:N) z[i] = n * sum(xx[1:i]) / sum(x) # The grid of b
 r = runif(1)
 s = numeric()
 for(i in 1:N){
   if(z[i] >= r){
     s = c(s, U[i])
     r = r + 1
```

```
}
 return(s[order(s)])
}
set.seed(854)
sam = syspps(z, n) # The n units selected
ys = y[sam] # The y values in S, same as <math>ys=y[sam]
piN = n * z/sum(z) # The inclusion probabilities
\# sum(piN) \# Check: sum(pi) = n
pis = piN[sam] # pi_i for i in S
# We normalize the z here for the approximate of HT using HH later
zs \leftarrow z/sum(z)
zs <- zs[sam]
sdata_syspps = pdata[sam,] # The sample data matrix
sdata_syspps <- cbind(sdata_syspps, pis, zs)</pre>
# 2. PPS sampling with replacement
set.seed(854)
z <- z/sum(z) # normalized size variable
```

```
sam <- sample(N, n, replace = T, prob = z)</pre>
ys <- y[sam] # Values of y in the sample
zs <- z[sam] # Values of the size variable
sdata_wrpps <- pdata[sam,]</pre>
sdata_wrpps <- cbind(sdata_wrpps, zs)</pre>
sdata wrpps <- sdata wrpps[, !(colnames(sdata wrpps) == "z")]</pre>
# 3. Poisson sampling method
set.seed(854)
piN <- n * z / sum(z) # Inclusion probabilities
sam <- numeric()</pre>
for(i in 1:N){
 r = runif(1)
 if(r \le piN[i]) sam = c(sam,i)
}
ys = y[sam]
pis = piN[sam]
sdata_poisson = pdata[sam,]
sdata_poisson <- cbind(sdata_poisson, pis)</pre>
```

A.6.2 R codes for implementing linearization for each PPS sampling method

The R codes for linearization for each PPS sampling method

```
# HH weights & point estimate
w \leftarrow 1 / (n * z) # HH weight per draw
Ty <- sum(w * y)
Tx \leftarrow sum(w * x)
mu_GR <- (Ty / Tx) * mux</pre>
Rhat <- Ty / Tx
# Linearised residuals
e <- y - Rhat * x
# HH variance for residual total
Te_hat <- sum(w * e) # HH total of residuals</pre>
v_lin <- (1 / N^2) *
  (1 / (n * (n - 1))) *
  sum((e / z - Te_hat)^2)
# Confidence intervals
z \leftarrow qnorm(1 - (1 - conf)/2)
se <- sqrt(v_lin)</pre>
ci \leftarrow c(mu_GR - z * se,
        mu_GR + z * se)
```

```
return(list(point = mu_GR,
            var = v_lin,
            se = se,
            ci = ci,
            level = conf))
}
# 2. Estimating the variance and confidence interval for Poisson Sampling
# under linearization
linVarGR poisson <- function(sdata, mux, N, conf = 0.95) {</pre>
 y <- sdata$y
 x <- sdata$x
 pi <- sdata$pis # first-order inclusion probs</pre>
 d <- 1 / pi # HT weights
 n <- length(y)
 # Point estimate
 muY_HT <- sum(d * y) / N
 muX_HT <- sum(d * x) / N</pre>
 mu_GR <- (muY_HT / muX_HT) * mux</pre>
 Rhat <- muY_HT / muX_HT
```

```
# Linearised residuals
 e_hat \leftarrow y - Rhat * x
 # Poisson HT variance
 v_lin <- (1 / N^2) * sum((1 - pi) / pi^2 * e_hat^2)</pre>
 # Confidence intervals
 z < -qnorm(1 - (1 - conf)/2)
 se <- sqrt(v_lin)</pre>
 ci \leftarrow c(mu_GR - z * se,
         mu_GR + z * se)
 return(list(point = mu_GR,
            var = v_lin,
            se = se,
            ci = ci,
            level = conf))
}
# Use the functions to estimate the variance for each PPS sampling method
mux <- mean(pdata$x)</pre>
```

```
# 1. For radomized systematic PPS sampling
vlinear_syspps <- linVarGR_HH(sdata_syspps,</pre>
                               mux = mux,
                                  = 5000)
# vlinear_syspps
# 2. For Poisson Sampling
vlinear_poisson <- linVarGR_poisson(sdata_poisson,</pre>
                              mux = mux,
                                  = 5000)
# vlinear_poisson
# 3. For PPS sampling with replacement
vlinear wrpps <- linVarGR_HH(sdata wrpps,</pre>
                           mux = mux,
                               = 5000)
                           N
# vlinear_wrpps
```

A.6.3 R codes for Bootstrap under each PPS sampling Method

R codes for Bootstrap

```
# Read the generated data

pdata <- read.csv("data/pdata.csv")

sdata_syspps <- read.csv("data/sdata_syspps.csv")</pre>
```

```
sdata_poisson <- read.csv("data/sdata_poisson.csv")</pre>
sdata_wrpps <- read.csv("data/sdata_wrpps.csv")</pre>
# 1. Bootstrap function for randomized systematic PPS and
# Poisson sampling (use pi)
varBootGR <- function(sdata, mux, N, B = 1000, conf = 0.95){</pre>
 set.seed(3)
 ys <- sdata$y
 xs <- sdata$x
 pis <- sdata$pis</pre>
 n <- length(ys)
 # Hortitz-Thompson weights
 di <- 1 / pis
 # Point estimate
 muY_HT <- sum(di * ys) / N</pre>
 muX_HT <- sum(di * xs) / N</pre>
 muY GR <- (muY HT / muX HT) * mux
 muGR_star <- numeric(B)</pre>
```

```
for (b in seq_len(B)) {
  bsam <- sample.int(n, n, replace = TRUE)</pre>
  yb <- ys[bsam]</pre>
  xb <- xs[bsam]</pre>
  db <- di[bsam]</pre>
  muY_HT_star <- sum(db * yb) / N</pre>
  muX_HT_star <- sum(db * xb) / N</pre>
  muGR_star[b] <- (muY_HT_star / muX_HT_star) * mux</pre>
}
# Compute the variance of generalized ratio estimator
vB <- (B - 1) * var(muGR_star) / B</pre>
# Confidence interval
alpha <- 1 - conf
ci <- quantile(muGR_star,</pre>
                    probs = c(alpha/2, 1 - alpha/2),
                    names = FALSE)
return(list(var = vB,
     ci = ci,
```

```
level = conf,
      point = muY_GR,
      estimates = muGR_star))
}
# 2. Bootstrap function for PPS sampling with replacement (use z)
varBootGR_wrpps <- function(sdata, mux, N, B = 1000, conf = 0.95){</pre>
 set.seed(854)
 ys <- sdata$y
 xs <- sdata$x
 zs <- sdata$zs
 n <- length(ys)
 # Hansen-Hurwitz weights
 dHH <- 1 / (n * zs)
 # Point estimate
 Ty_HH <- sum(dHH * ys)
 Tx_HH <- sum(dHH * xs)</pre>
 muY_GR <- (Ty_HH / Tx_HH) * mux</pre>
```

```
muGR_star <- numeric(B)</pre>
for (b in seq_len(B)) {
  bsam <- sample.int(n, n, replace = TRUE)</pre>
  yb <- ys[bsam]</pre>
  xb <- xs[bsam]</pre>
  zb <- zs[bsam]</pre>
  dB <- 1 / (n * zb)
  Ty_star <- sum(dB * yb)</pre>
  Tx_star <- sum(dB * xb)</pre>
  muGR_star[b] <- (Ty_star / Tx_star) * mux</pre>
}
# Compute the variance
vB <- (B - 1) * var(muGR_star) / B</pre>
# Confidence interval of mu_y
alpha \leftarrow 1 - conf
ci <- quantile(muGR_star,</pre>
                 probs = c(alpha/2, 1 - alpha/2),
                 names = FALSE)
```

```
return(list(var = vB,
            ci = ci,
            level = conf,
            point = muY_GR,
            estimates = muGR_star))
}
mux <- mean(pdata$x)</pre>
N <- 5000
B <- 1000
vBoot_syspps <- varBootGR(sdata_syspps, mux, N, B)</pre>
# vBoot_syspps
vBoot_poisson <- varBootGR(sdata_poisson, mux, N, B)</pre>
# vBoot_poisson
vBoot_wrpps <- varBootGR_wrpps(sdata_wrpps, mux, N, B = 1000)</pre>
# vBoot_wrpps
```

A.6.4 R codes for delete-1 Jackknife under each PPS sampling Method

R codes for Delete-1 Jackknife

```
# Read the simulated data
pdata <- read.csv("data/pdata.csv")</pre>
sdata_syspps <- read.csv("data/sdata_syspps.csv")</pre>
sdata_poisson <- read.csv("data/sdata_poisson.csv")</pre>
sdata_wrpps <- read.csv("data/sdata_wrpps.csv")</pre>
# 1. The Delete-1 Jackknife variance estimation for H-T estimators
varJackGR <- function(sdata, mux, N, level = 0.95) {</pre>
 ys <- sdata$y
 xs <- sdata$x
 pis <- sdata$pis</pre>
 n <- length(ys)
 di <- 1 / pis
 Nhat <- sum(di)</pre>
 # Point estimate
 muY_HT <- sum(di * ys) / N</pre>
 muX_HT <- sum(di * xs) / N</pre>
       <- (muY_HT / muX_HT) * mux
 muGR
```

```
muGR_mi <- numeric(n)</pre>
for (i in seq_len(n)) {
  gamma <- Nhat / (Nhat - di[i])</pre>
  di_mi <- di * gamma
  di mi[i] <- 0
  muY_HT_mi <- sum(di_mi * ys) / sum(di_mi)</pre>
  muX HT mi <- sum(di mi * xs) / sum(di mi)</pre>
  muGR_mi[i] <- (muY_HT_mi / muX_HT_mi) * mux</pre>
}
# Jackknife variance
mu_dot <- mean(muGR_mi)</pre>
vJack <- (n - 1) / n * sum((muGR_mi - mu_dot)^2)</pre>
seJack <- sqrt(vJack)</pre>
# Confidence interval
alpha <- 1 - level
z <- qnorm(1 - alpha/2)</pre>
ci \leftarrow c(lower = muGR - z * seJack,
            upper = muGR + z * seJack)
return(list(point = muGR,
```

```
var = vJack,
            ci = ci,
            estimates = muGR_mi))
}
# 2. Delete-1 Jackknife estimator using HH estimators
varJackGR_wrpps <- function(sdata, mux, level = 0.95) {</pre>
 ys <- sdata$y
 xs <- sdata$x
 zs <- sdata$zs
 n <- length(ys)
 # HH weights
 z_i <- zs / sum(zs)</pre>
 wi <- 1 / (n * z_i)
 # Point estimate
 muY_HH <- sum(wi * ys) / sum(wi)</pre>
 muX_HH <- sum(wi * xs) / sum(wi)</pre>
 muY_GR <- (muY_HH / muX_HH) * mux</pre>
```

```
muGR_mi <- numeric(n)</pre>
for (i in seq_len(n)) {
  # Replicate weights
  wi <- wi
  wi_mi <- (n / (n - 1)) * wi
  wi_mi[i] <- 0
  muY_HH_mi <- sum(wi_mi * ys) / sum(wi_mi)</pre>
  muX_HH_mi <- sum(wi_mi * xs) / sum(wi_mi)</pre>
  muGR_mi[i] <- (muY_HH_mi / muX_HH_mi) * mux</pre>
}
# Jackknife variance
mu_dot <- mean(muGR_mi)</pre>
vJack <- (n - 1) / n * sum((muGR_mi - mu_dot)^2)</pre>
seJack <- sqrt(vJack)</pre>
alpha <- 1 - level
zcrit <- qnorm(1 - alpha/2)</pre>
ci <- c(lower = muY_GR - zcrit * seJack,</pre>
             upper = muY_GR + zcrit * seJack)
```

```
return(list(point = muY_GR,
            variance = vJack,
            ci = ci,
            estimates = muGR_mi))
}
mux <- mean(pdata$x)</pre>
  <- 5000
vJack_syspps <- varJackGR(sdata_syspps, mux, N)</pre>
# vJack_syspps
vJack_poisson <- varJackGR(sdata_poisson, mux, N)</pre>
# vJack_poisson
vJack_wrpps <- varJackGR_wrpps(sdata_wrpps, mux)</pre>
# vJack_wrpps
```

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