A comparison of design-based and model-based approaches for finite population spatial sampling and inference.

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3 Abstract

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- 1. The design-based and model-based approaches to frequentist statistical inference rest on fundamentally different foundations. In the design-based approach, inference relies on random sampling. In the model-based approach, inference relies on distributional assumptions. We compare the approaches in a finite population spatial context.
- 2. We provide relevant background for the design-based and model-based approaches and then study their performance using simulated data and real data. The real data is from the United States Environmental Protection Agency's 2012 National Lakes Assessment. A variety of sample sizes, location layouts, dependence structures, and response types are considered. The population mean is the parameter of interest, and performance is measured using statistics like bias, squared error, and interval coverage.
 - 3. When studying the simulated and real data, we found that regardless of the strength of spatial dependence in the data, the generalized random tessellation stratified (GRTS) algorithm, which explicitly incorporates spatial locations into sampling, tends to outperform the simple random

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- sampling (SRS) algorithm, which does not explicitly incorporate spatial locations into sampling. We also found that model-based inference tends to outperform design-based inference, even for skewed data where the model-based distributional assumptions are violated. The performance gap between design-based inference and model-based inference is small when GRTS samples are used but large when SRS samples are used, suggesting that the sampling choice (whether to use GRTS or SRS) is most important when performing design-based inference.
- 4. There are many benefits and drawbacks to the design-based and model-based approaches for finite population spatial sampling and inference that practitioners must consider when choosing between them. We provide relevant background contextualizing each approach and study their properties in a variety of scenarios, making recommendations for use based on the practitioner's goals.

44 Keywords

- Design-based inference; Finite population block kriging (FPBK); Generalized
- random tessellation stratified (GRTS) algorithm; Local neighborhood variance
- estimator; Model-based inference; Restricted maximum likelihood (REML) esti-
- 48 mation; Spatially balanced sampling; Spatial covariance

49 1. Introduction

- When data cannot be collected for all units in a population (population units),
- data are collected on a subset of the population units this subset is called a
- sample. There are two general approaches for using samples to make frequentist
- 53 statistical inferences about a population: design-based and model-based. In the
- design-based approach, inference relies on randomly assigning some population

units to be in the sample (random sampling). Alternatively, in the model-based approach, inference relies on distributional assumptions about the underlying data-generating stochastic process (superpopulation). Each paradigm has a deep 57 historical context (Sterba, 2009) and its own set of benefits and drawbacks (Brus and De Gruijter, 1997; Hansen et al., 1983). In this manuscript, we compare 59 design-based and model-based approaches for finite population spatial sampling and inference. Spatial data are data that have some sort of spatial index (usually specified via coordinates). De Gruijter and Ter Braak (1990) and Brus and DeGruijter (1993) give early comparisons of design-based and model-based approaches for spatial data, quashing the belief that design-based approaches could not be used for spatially correlated data. Since then, there have been several general comparisons between design-based and model-based approaches for spatial data Brus and De Gruijter, 1997; Brus, 2021; Ver Hoef, 2002; Ver Hoef, 2008). Cooper 68 (2006) reviews the two approaches in an ecological context before introducing a 'model-assisted" variance estimator that combines aspects from each approach. In addition to Cooper (2006), there has been substantial research and development 71 into estimators that use both design-based and model-based principles (see e.g., Sterba (2009) and Cicchitelli and Montanari (2012), and for Bayesian approaches, 73 see Chan-Golston et al. (2020) and Hofman and Brus (2021)). While comparisons between design-based and model-based approaches have 75 been studied in spatial contexts, our contribution is comparing design-based approaches specifically built for spatial data to model-based approaches. Though 77 the broad comparisons we draw between design-based and model-based approaches generalize to finite and infinite populations, we focus on finite populations. A finite population contains a finite number of population units (we assume the finite number is known) – an example is lakes (treated as a whole

- with the lake centroid representing location) in the conterminous United States.
- An infinite population contains an infinite number of population units an
- 84 example is locations within a single lake.
- The rest of the manuscript is organized as follows. In Section 1.1, we introduce
- and provide relevant background for design-based and model-based approaches
- 87 to finite population spatial sampling and inference. In Section 2, we describe
- 88 how we intend to compare performance of the approaches using simulated and
- 99 real data. The real data is from the United States Environmental Protection
- 90 Agency's 2012 National Lakes Assessment (NLA) (USEPA, 2012). In Section 3,
- 91 we present analysis results for the simulated data and real data. And in Section
- 92 4, we end with a discussion and provide directions for future research.

93 1.1. Background

- The design-based and model-based approaches incorporate randomness in
- ₉₅ fundamentally different ways. In this section, we describe the role of randomness
- 96 for each approach and the subsequent effects on statistical inferences for spatial
- 97 data.

98 1.1.1. Comparing Design-Based and Model-Based Approaches

- The design-based approach assumes the population is fixed. Randomness is
- incorporated via the selection of population units according to a sampling design.
- A sampling design assigns a probability of selection to each sample (subset of
- 102 population units). Some examples of commonly used sampling designs include
- simple random sampling, stratified random sampling, and cluster sampling.
- 104 The inclusion probability of a population unit is calculated by summing each
- sample's probability of selection over all samples that contain the population unit.
- 106 Inclusion probabilities are often used when selecting samples and estimating
- population parameters.

When samples are chosen in a manner such that the layout of sampled units 108 reflects the layout of the population units, we call the resulting sample spatially 109 balanced. By "reflecting the layout of the population units," we mean that if 110 population units are concentrated in specific areas, the units in the sample should 111 be concentrated in the same areas. Because spatially balanced samples reflect 112 the layout of the population units, they are not necessarily spread out in space 113 in some equidistant manner. One method of selecting spatially balanced samples 114 is the generalized random tessellation stratified (GRTS) algorithm (Stevens and 115 Olsen, 2004), which we discuss in more detail in Section 1.1.2. To quantify the spatial balance of a sample, Stevens and Olsen (2004) proposed loss metrics 117 based on Voronoi polygons (i.e., Dirichlet Tessellations). Fundamentally, the design-based approach combines the randomness of the 119 sampling design with the data collected via the sample to justify the estimation and uncertainty quantification of fixed, unknown parameters of a population (e.g., 121 a population mean). Treating the data as fixed and incorporating randomness 122 through the sampling design yields estimators having very few other assumptions. 123 Confidence intervals for these types of estimators are typically derived using 124 limiting arguments that incorporate all possible samples. Sample means, for 125 example, are asymptotically normal (Gaussian) by the central limit theorem 126 (under some assumptions). If we repeatedly select samples from the population, 127 then 95% of all 95% confidence intervals constructed from a procedure with 128 appropriate coverage will contain the true fixed population mean. Särndal et al. (2003) and Lohr (2009) provide thorough reviews of the design-based approach. 130 The model-based approach assumes the population is a random realization of a

data-generating stochastic process. Randomness is formally incorporated through

distributional assumptions on this process. Strictly speaking, randomness need

not be incorporated through random sampling, though Diggle et al. (2010)

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warn against preferential sampling. Preferential sampling occurs when the 135 process generating the data locations and the process being modeled are not 136 independent of one another. To guard against preferential sampling, model-137 based approaches can implement some form of random sampling. It is common, 138 however, for model-based approaches to sample non-randomly. When model-139 based approaches do implement random sampling, the inclusion probabilities are 140 ignored when analyzing the sample (in contrast to the design-based approach, which relies on these inclusion probabilities to analyze the sample). 142 Instead of estimating fixed, unknown population parameters, as in the designbased approach, often the goal of model-based inference is to predict the value 144 of a realized variable. For example, suppose the realized mean of all population units (the realized population mean) is the variable of interest. Instead of a fixed, 146 unknown mean, we are predicting the value of the mean, a random variable. Prediction intervals are then derived using assumptions of the data-generating 148 stochastic process. If we repeatedly generate realizations from the same process and select samples, then 95% of all 95% prediction intervals constructed from a 150 procedure with appropriate coverage will contain their respective realized means. 151 Cressie (1993) and Schabenberger and Gotway (2017) provide thorough reviews 152

approach with a fixed population and random sampling to the model-based approach with random populations and non-random sampling.

of model-based approaches for spatial data. In Fig. 1, we provide a visual

comparison of the design-based and model-based approaches (Ver Hoef (2002)

and Brus (2021) provide similar figures). Fig. 1 contrasts the design-based

1.1.2. Spatially Balanced Design and Analysis 158

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We previously mentioned that the design-based approach can be used to 159 select spatially balanced samples. Spatially balanced samples are useful because 160 parameter estimates from these samples tend to vary less (be more precise) 161

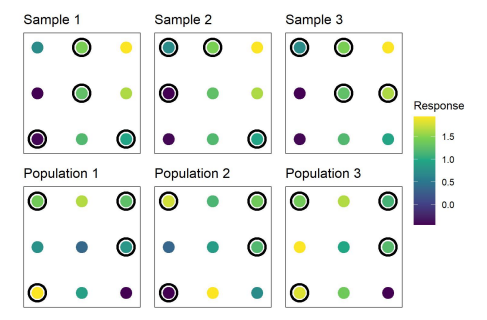
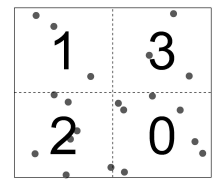


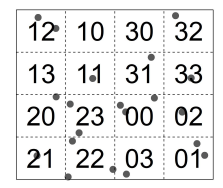
Figure 1: A visual comparison of the design-based and model-based approaches. In the top row, the design-based approach is highlighted. There is one fixed population with nine population units and three random samples of size four (points circled are those sampled). The response values at each site are fixed. In the bottom row, the model-based approach is highlighted. There are three realizations of the same data-generating stochastic process that are all sampled at the same four locations. The response values at each site are random.

than parameter estimates from samples lacking spatial balance (Barabesi and

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Franceschi, 2011; Benedetti et al., 2017; Grafström and Lundström, 2013; Robert-163 son et al., 2013; Stevens and Olsen, 2004; Wang et al., 2013). The first spatially 164 balanced sampling algorithm to see widespread use was the generalized random 165 tessellation stratified (GRTS) algorithm (Stevens and Olsen, 2004). After the 166 GRTS algorithm was developed, several other spatially balanced sampling algo-167 rithms emerged, including stratified sampling with compact geographical strata 168 (Walvoort et al., 2010), the local pivotal method (Grafström et al., 2012; Graf-169 ström and Matei, 2018), spatially correlated Poisson sampling (Grafström, 2012), balanced acceptance sampling (Robertson et al., 2013), within-sample-distance 171 sampling (Benedetti and Piersimoni, 2017), and Halton iterative partitioning sampling (Robertson et al., 2018). In this manuscript, we select spatially bal-173 anced samples using the GRTS algorithm because it is readily available in the spsurvey R package (Dumelle et al., 2022) and naturally accommodates finite 175 and infinite sampling frames, unequal inclusion probabilities, and replacement 176 units. Replacement units are additional population units that can be sampled 177 when a population unit originally selected can no longer be sampled. A couple 178 of reasons why an originally selected site can no longer be sampled include its 179 location being physically inaccessible or it is on private land that the researcher 180 does not have permission to access. 183 The GRTS algorithm selects samples by utilizing a particular mapping 182 between two-dimensional and one-dimensional space that preserves proximity 183 relationships. First, the bounding box of the domain is split up into four 184 distinct, equally sized squares called level-one cells. Each level-one cell is randomly assigned a level-one address of 0, 1, 2, or 3. The set of level-one 186 cells is denoted by A_1 and defined as $A_1 \equiv \{a_1 : a_1 = 0, 1, 2, 3\}$. Within each 187 level-one cell, the inclusion probability for each population unit (which is pre-188





(a) Assignment of level-one cells to the spa-(b) Assignment of level-two cells to the tial domain. Grey circles indicate popula- spatial domain. Grey circles indicate population units.

Figure 2: Assignment of level-one and level-two cells to the spatial domain. In (a), each level-one cells is randomly given a level-one address of 0, 1, 2, or 3. In (b), each level-two cell within each level-one cell is randomly given a level-two address of 0, 1, 2, or 3.

specified) is summed, and if any of these sums are one or greater, a second level 189 of cells is added. Then each level-one cell is split into four distinct, equally sized squares called level-two cells. Each level-two cell is randomly assigned 191 a level-two address of 0, 1, 2, or 3. The set of level-two cells is denoted by 192 A_2 and defined as $A_2 \equiv \{a_1 a_2 : a_1 = 0, 1, 2, 3; a_2 = 0, 1, 2, 3\}$. The inclusion 193 probabilities within each level-two cell are summed, and if any of these sums 194 are one or greater, a third level of cells is added. This process continues for 195 k steps, until all level-k cells have inclusion probability sums less than one. 196 Then $A_k \equiv \{a_1...a_k : a_1 = 0, 1, 2, 3; ...; a_k = 0, 1, 2, 3\}$. Figure 2 provides some 197 intuition regarding the assignment of level-one and level-two cells. 198 After determining A_k , the set is placed into hierarchical order. Hierar-199 chical order is a numeric order that first sorts A_k by the level-one addresses 200 from smallest to largest, then sorts A_k by the level-two addresses from small-201 est to largest, and so on. For example, A_2 in hierarchical order is the set 202 $\{00, 01, 02, 03, 10, ..., 13, 20, ..., 23, 30, ..., 33\}$. Because hierarchical ordering sorts

by level-one cells, then level-two cells, and so on, population units that have similar hierarchical addresses tend to be nearby one another in space. Next, each 205 population unit is mapped to a one-dimensional line in hierarchical order where 206 each population unit's inclusion probability equals its line-length. If a level-k 207 cell has multiple population units in it, they are randomly placed within the 208 cell's respective line segment. A uniform random variable is then simulated in 209 [0,1] and a systematic sample is selected on the line, yielding n sample points for 210 a sample size n. Each of these sample points falls on some population unit's line 211 segment, and thus that population unit is selected in the sample. For further 212 details regarding the GRTS algorithm, see Stevens and Olsen (2004). 213

After selecting a sample and collecting data, unbiased estimates of population means and totals can be obtained using the Horvitz-Thompson estimator (Horvitz and Thompson, 1952). If τ is a population total, the Horvitz-Thompson estimator for τ , denoted by $\hat{\tau}_{ht}$, is given by

where z_i is the value of the *i*th population unit in the sample, π_i is the inclusion

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$$\hat{\tau}_{ht} = \sum_{i=1}^{n} z_i \pi_i^{-1},\tag{1}$$

probability of the ith population unit in the sample, and n is the sample size. An 215 estimate of the population mean is obtained by dividing $\hat{\tau}_{ht}$ by N, the number 216 of population units. 217 It is also important to quantify the uncertainty in $\hat{\tau}_{ht}$. The Horvitz-Thompson 218 (Horvitz and Thompson, 1952) and Sen-Yates-Grundy (Sen, 1953; Yates and 219 Grundy, 1953) variance estimators are often used to estimate $Var(\hat{\tau}_{ht})$, but 220 these estimators have two drawbacks. First, they rely on calculating π_{ij} , the 221 probability that population unit i and population unit j are both in the sample 222 this quantity can be challenging if not impossible to calculate analytically for 223 GRTS samples. Second, these estimators tend to ignore the spatial locations of

the population units. To address these two drawbacks simultaneously, Stevens 225 and Olsen (2003) proposed the local neighborhood variance estimator. The local 226 neighborhood variance estimator does not rely on π_{ij} and estimates the variance 227 of $\hat{\tau}$ conditional on the random properties of the GRTS sample – the idea being 228 that this conditioning should yield a more precise estimate of τ . They show that 229 the contribution from each sampled population unit to the overall variance is 230 dominated by local variation. Thus the local neighborhood variance estimator 231 is a weighted sum of variance estimates from each sampled population unit's 232 local neighborhood. These local neighborhoods contain the sampled population 233 unit itself and its three nearest neighbors (among all other sampled population 234 units). For more details, see Stevens and Olsen (2003).

236 1.1.3. Finite Population Block Kriging

Finite population block kriging (FPBK) is a model-based approach that 237 expands the geostatistical Kriging framework to the finite population setting 238 (Ver Hoef, 2008). Instead of developing inference based on a specific sampling design, we assume the data are generated by a spatial stochastic process. We 240 summarize some of the basic principles of FPBK next – see Ver Hoef (2008) for technical details and see Higham et al. (2021b) for an extension to cases of 242 imperfect detection among population units. Let $\mathbf{z} \equiv \{z(s_1), z(s_2), ..., z(s_N)\}$ be 243 an $N \times 1$ response vector at locations s_1, s_2, \ldots, s_N that can be measured 244 at the N population units. Suppose we want to use a sample to predict some 245 linear function of the response variable, $f(\mathbf{z}) = \mathbf{b}'\mathbf{z}$, where \mathbf{b}' is a $1 \times N$ vector of weights (e.g, the population mean is represented by a weights vector whose 247 elements all equal 1/N). Denoting quantities that are part of the sampled population units with a subscript s and quantities that are part of the unsampled 249 population units with a subscript u, let 250

$$\begin{pmatrix} \mathbf{z}_s \\ \mathbf{z}_u \end{pmatrix} = \begin{pmatrix} \mathbf{X}_s \\ \mathbf{X}_u \end{pmatrix} \beta + \begin{pmatrix} \delta_s \\ \delta_u \end{pmatrix}, \tag{2}$$

where \mathbf{X}_s and \mathbf{X}_u are the design matrices for the sampled and unsampled population units, respectively, $\boldsymbol{\beta}$ is the parameter vector of fixed effects, and $\boldsymbol{\delta} \equiv [\boldsymbol{\delta}_s \ \boldsymbol{\delta}_u]'$, where $\boldsymbol{\delta}_s$ and $\boldsymbol{\delta}_u$ are random errors for the sampled and unsampled population units, respectively.

FPBK assumes δ in Equation (2) has mean-zero and a spatial dependence structure that can be modeled using a covariance function. This covariance function is commonly assumed to be non-negative, second-order stationary (depending only on the separation vector (e.g., distance) between population units), and isotropic (independent of direction) (Cressie, 1993). Henceforth, it is implied that we have made these same assumptions regarding δ . Chiles and Delfiner (1999), pp. 80-93 discuss covariance functions that are not second-order stationary, not isotropic, or not either. A variety of flexible covariance functions can be used to model δ (Cressie, 1993) – one example is the exponential covariance function. Cressie (1993) provides a thorough list of spatial covariance functions. The i, jth element of the exponential covariance matrix, $cov(\delta)$, is

$$cov(\delta_{i}, \delta_{j}) = \begin{cases} \sigma_{1}^{2} \exp(-h_{i,j}/\phi) & h_{i,j} > 0\\ \sigma_{1}^{2} + \sigma_{2}^{2} & h_{i,j} = 0 \end{cases}$$
(3)

where σ_1^2 is the variance parameter that quantifies the spatially dependent (correlated) variability, σ_2^2 is the variance parameter the quantifies that spatially independent (not correlated) variability, ϕ is the distance parameter that measures the distance-decay rate of the covariance, and $h_{i,j}$ is the Euclidean distance between population units i and j. In geostatistical literature, σ_1^2 is called the partial sill, σ_2^2 is called the nugget, and ϕ is called the range. We denote θ as the

vector of covariance parameters that composes $\boldsymbol{\delta}$. In Equation 3, $\boldsymbol{\theta} = \{\sigma_1^2, \sigma_2^2, \phi\}$.

The parameters in Equation 2 can be estimated using a variety of techniques, but we focus on restricted maximum likelihood (REML) (Harville, 1977; Patterson and Thompson, 1971; Wolfinger et al., 1994). REML is preferred over maximum likelihood (ML) because ML estimates can be badly biased for small sample sizes, due to the fact that ML makes no adjustment for the simultaneous estimation of β and θ (Patterson and Thompson, 1971). Minus twice the REML log-likelihood of the sampled sites is given by

$$\ln |\mathbf{\Sigma}| + (\mathbf{z}_s - \mathbf{X}_s \tilde{\boldsymbol{\beta}})^T \mathbf{\Sigma}_{ss}^{-1} (\mathbf{z}_s - \mathbf{X}_s \tilde{\boldsymbol{\beta}}) + \ln |\mathbf{X}_s^T \mathbf{\Sigma}_{ss}^{-1} \mathbf{X}_s| + (n - p) \ln(2\pi), \quad (4)$$

where $\tilde{\beta} = (X_s^T \Sigma_{ss}^{-1} X_s)^{-1} X_s^T \Sigma_{ss}^{-1} z_s$ and Σ_{ss} is the covariance matrix of the sampled sites. Minimizing Equation 4 yields $\hat{\theta}_{reml}$, the REML estimates of 263 $\boldsymbol{\theta}$. Then $\hat{\boldsymbol{\beta}}_{reml}$, the REML estimate of $\boldsymbol{\beta}$, is given by $(\boldsymbol{X}_s^T \hat{\boldsymbol{\Sigma}}_{ss}^{-1} \boldsymbol{X})^{-1} \boldsymbol{X}_s^T \hat{\boldsymbol{\Sigma}}_{ss}^{-1} \boldsymbol{z}_s$ where $\hat{\Sigma}_{ss}$ is Σ_{ss} evaluated at $\hat{\theta}_{reml}$. 265 With the model formulation in Equation 2, the best linear unbiased predictor (BLUP) of $f(\mathbf{b}'\mathbf{z})$ and its prediction variance can be computed. While details 267 of the derivation are in Ver Hoef (2008), we note here that the predictor and 268 its variance are both moment-based, meaning that they do not rely on any distributional assumptions. Distributional assumptions are used, however, when 270 constructing prediction intervals. Other approaches, such as k-nearest-neighbors (Fix and Hodges, 1989; Ver 272 Hoef and Temesgen, 2013) and random forest (Breiman, 2001), among others, could also be used to obtain predictions for a mean or total from finite population 274 spatial data. Compared to the k-nearest-neighbors and random forest approach, 275 we prefer FPBK because it is model-based and relies on theoretically-based 276

variance estimators leveraging the model's spatial covariance structure, whereas

k-nearest-neighbors and random forests use ad-hoc variance estimators (Ver Hoef

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and Temesgen, 2013). Additionally, Ver Hoef and Temesgen (2013) compared FPBK, k-nearest-neighbors, and random forest in a variety of spatial data contexts, and FPBK tended to perform best.

282 2. Materials and Methods

In this section we describe how we used simulated and real data to investigate performance between simple random sampling (SRS) and GRTS sampling as well as performance between design-based (DB) and model-based (MB) inference. In SRS and GRTS sampling, all population units had equal inclusion probabilities and were selected without replacement. The important distinction between SRS and GRTS is that SRS ignores spatial locations while sampling but GRTS explicitly incorporates them. Together, the two sampling plans (SRS and GRTS) combined with the two inference approaches (DB and MB) yielded four sampling-inference combinations: SRS-DB, SRS-MB, GRTS-DB, and GRTS-MB. For SRS-DB, the Horvitz-Thompson estimator (1) was used to estimate means and the commonly-used SRS variance formula (Lohr, 2009; Särndal et al., 2003) was used to estimate variances. This variance formula is given by

$$\frac{f[\sum_{i=1}^{n} (z_i - \bar{z})^2]}{n(n-1)},\tag{5}$$

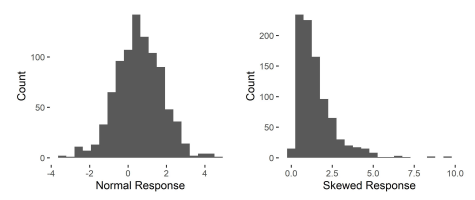
where z_i is the response value of the *i*th sampled population unit, \bar{z} is the mean of all z_i , n is the sample size, N is the population size, and f = (1 - n/N) (f is often called the finite population correction factor). For GRTS-DB, the Horvitz-Thompson estimator was used to estimate means and the local neighborhood variance was used to estimate variances. For SRS-MB and GRTS-MB, FPBK was used to estimate means and variances using restricted maximum likelihood. SRS, GRTS sampling, and design-based inference were implemented using the spsurvey \mathbf{R} package (Dumelle et al., 2022). FPBK was implemented using the

291 sptotal **R** package (Higham et al., 2021a).

The simulated and real data were used for distinct objectives. The simulated data was used to compare the sampling-inference combinations across many realized populations (from the same data-generating stochastic process) and random samples. The real data was used to compare the sampling-inference combinations within a single realized population but across random samples. With the simulated data, we were in control of the data-generating stochastic process and the random sampling process. With the real data, we were only in control of the random sampling process (which is typically the case in practice).

300 2.1. Simulated Data

We evaluated performance of the four sampling-inference combinations in 301 36 different simulation scenarios. The 36 scenarios resulted from the crossing of 302 three sample sizes, two location layouts (of the population units), two response 303 types, and three proportions of dependent random error (DRE). The three sample 304 sizes (n) were n = 50, n = 100, and n = 200. Samples were always selected from 305 a population size (N) of N = 900. The two location layouts were random and 306 gridded. Locations in the random layout were randomly generated inside the 307 unit square ($[0,1] \times [0,1]$). Locations in the gridded layout were placed on a fixed, 308 equally spaced grid inside the unit square. The two response types were normal and skewed. For the normal response type, the response was simulated using 310 mean-zero random errors with the exponential covariance (Equation 3) for three 31 proportions of dependent random error (DRE): 0% DRE, 50% DRE, and 90% 312 DRE. Recall the proportion of DRE is represented by $\sigma_1^2/(\sigma_1^2 + \sigma_2^2)$, where σ_1^2 313 and σ_2^2 are the DRE variance and independent random error (IRE) variance from 314 Equation 3, respectively. The total variance, $\sigma_1^2 + \sigma_2^2$, was always 2. The distance 315 parameter was always $\sqrt{2}/3$, chosen so that the correlation in the DRE decayed 316 to nearly zero at $\sqrt{2}$, the largest possible distance between two population units



(a) Histogram of a single realized popula-(b) Histogram of a single realized population for the normal response.

Figure 3: Histograms of single realized populations simulated for the normal and skewed respones using the random layout and 50% DRE.

in the domain. For the skewed response type, the response was first simulated using the same approach as for the normal response type, except that the total variance was 0.6931 instead of 2. The response was then exponentiated, yielding a skewed random variable whose total variance was 2. The skewed responses were used to evaluate performance of the sampling-inference approaches for data that were not normally distributed but were still estimated using REML, which relies on a normal log-likelihood. Figure 3 shows an example of a realized population for the normal and skewed responses using the random location layout and 50% DRE.

In each of the 36 simulation scenarios, there were 2000 independent simulation trials. Within each trial, a population was simulated according to the specifications of the particular simulation scenario (for the random location layout, locations were simulated separately for each trial). Next, a random SRS sample and a random GRTS sample were selected. Then, design-based and model-based inferences were used to estimate (design-based) or predict (model-based) the realized mean and construct 95% confidence (design-based) or

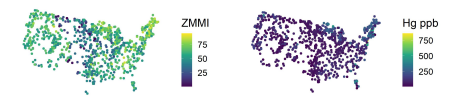
95% prediction (model-based) intervals. With model-based inference, covariance 334 parameters and the realized mean were estimated (using REML) separately for 335 each trial. After all 2000 trials, we summarized the long-run performance of the 336 sampling-inference combination in each scenario by calculating mean bias, root-337 mean-squared error, and interval coverage. Mean bias was taken as the average 338 deviation between each trial's estimated (or predicted) mean $(\hat{\mu}_i)$ and its realized 339 mean (μ_i) : $\frac{1}{n}\sum_{i=1}^{2000}(\hat{\mu}_i-\mu_i)$, where i indexes the simulation trials. Because each trial had a different realized population, μ_i changed with i. Root-mean-squared 34: error was taken as the square root of the average squared deviation between each trial's estimated (or predicted) mean and its realized mean: $\sqrt{\frac{1}{n}\sum_{i=1}^{2000}(\hat{\mu}_i-\mu_i)^2}$. 343 Interval coverage was taken as the proportion of simulation trials where the realized mean was contained in its 95% confidence (or prediction) interval. These 345 intervals were constructed using the normal distribution – justification comes from the asymptotic normality of means via the central limit theorem (under 347 some assumptions). Quantifying these metrics is important because together, they give us an idea of the accuracy (mean bias), spread (RMSE), and validity 349 (interval coverage) of the sampling-inference combinations. 350

2.2. National Lakes Assessment Data

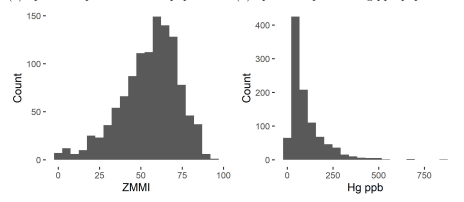
The United States Environmental Protection Agency (USEPA), states, and tribes periodically conduct National Aquatic Research Surveys (NARS) to assess the water quality of various bodies of water in the conterminous United States. One component of NARS is the National Lakes Assessment (NLA), which measures various aspects of lake health and water quality. We focus on analyzing zooplankton multi-metric indices (ZMMI) and mercury concentrations in parts per billion (Hg ppb) from the 2012 NLA. For ZMMI, data were collected at 1035 unique lakes. At less than 10% of lakes, two ZMMI replicates were collected. These were averaged for the purposes of our study so that each lake had one

measurement for ZMMI. For Hg ppb, data were collected at 995 unique lakes (there were no replicates). The ZMMI and Hg ppb data are shown as spatial maps and as histograms in Figure 4. The ZMMI data tend to be highest near the 363 coasts, lowest in the Central United States, are relatively symmetric, and have a mean of 55.05. The Hg ppb data tend to be highest in the Northeastern United 365 States, lowest elsewhere, are skewed, and have a mean of 103.16 ppb. Also in Figure 4 are separate spatial semivariograms for ZMMI and Hg ppb. The spatial 367 semivariogram quantifies the halved average squared differences (semivariance) 368 of responses whose separation (distance) falls within a separation class. The spatial semivariance is closely related to the spatial covariance, and spatial 370 semivariograms are often used to gauge the strength of spatial dependence in data. Both ZMMI and Hg ppb seem to have moderately strong spatial 372 dependence (Figure 4), as the empirical semivariance increases steadily with distance (meaning that observations near one another tend to be more similar 374 than observations far apart from one another). 375

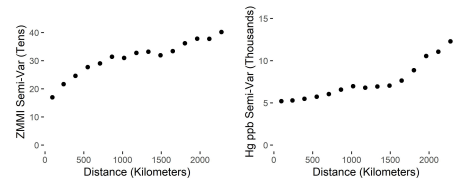
We studied performance of the four sampling-inference combinations by 376 selecting 2000 SRS and GRTS samples of size n = 50, n = 100, and n = 200377 from the realized ZMMI and Hg ppb populations and then analyzing the samples 378 using MB and DB inference. In total, there were six separate scenarios (two 379 responses crossed with three sample sizes). Within each SRS and GRTS sample, design-based and model-based inferences were used to estimate or predict the 381 population mean and construct 95% coverage intervals. With model-based inference, the exponential covariance was assumed, and covariance parameters 383 and the population mean were estimated using REML (separately for each SRS and GRTS sample). We used the same evaluation metrics as for the simulated data: mean bias, RMSE, and interval coverage. Mean bias was taken as the average deviation between each sample's estimated (or predicted) mean $(\hat{\mu}_i)$ and 387



(a) Spatial map of the ZMMI population. (b) Spatial map of the Hg ppb population.



(c) Histogram of the ZMMI population. (d) Histogram of the Hg ppb population.



(e) Empirical semivariogram of the ZMMI (f) Empirical semivariogram of the Hg ppb population.

Figure 4: Exploratory graphics representing populations for the zooplankton multi-metric indices (ZMMI) and mercury concentration in parts per billion (Hg ppb) in the 2012 National Lakes Assessment (NLA) data.

the population mean (μ) (of ZMMI or Hg ppb): $\frac{1}{n} \sum_{i=1}^{2000} (\hat{\mu}_i - \mu)$, where i indexes the simulation trials. Because each trial had the same realized population, μ did not change with i (in contrast to the simulated data, where the realized mean changed with i). Root-mean-squared error was taken as the square root of the average squared deviation between each sample's estimated (or predicted) mean and its population mean: $\sqrt{\frac{1}{n}\sum_{i=1}^{2000}(\hat{\mu}_i - \mu)^2}$. Interval coverage was taken as the proportion of simulation trials where the population mean was contained in its 95% confidence (or prediction) interval. These intervals were constructed using the normal distribution.

397 3. Results

399

3.1. Simulated Data

36 scenarios, so we omit a more detailed summary of those results here. Tables 400 for mean bias in all 36 simulation scenarios are provided in the supporting 401 information. 402 We define the relative RMSE as a ratio with numerator given by the RMSE 403 for a sampling-inference combination and the denominator given by the RMSE 404 for SRS-DB. Relative RMSEs for the random location layout are provided in 405 Fig. 5. When there is no spatial covariance (Fig. 5, "DRE%: 0%"), the four 406 sampling-inference combinations have approximately equal RMSE. In these 407 scenarios, using GRTS sampling or model-based inference does not generally 408 increase efficiency compared to SRS-DB. When there is spatial covariance (Fig. 409 5, "DRE%: 50%" and "DRE%: 90%"), GRTS-MB tends to have the lowest 410 RMSE, followed by GRTS-DB, SRS-MB, and finally SRS-DB. As the strength 411 of spatial covariance increases, the gap in RMSE between SRS-DB and the other sampling-inference combinations widens. Finally we note that when there is 413

Mean bias is nearly zero for all four sampling-inference combinations in all

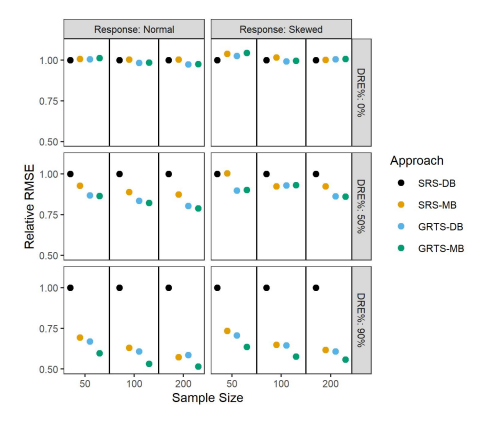


Figure 5: Simulated data relative RMSE for the four sampling-inference combinations and three sample sizes in the random location layout. The rows indicate the proportion of dependent error and the columns indicate the response type. The solid, black lines separate the sample sizes.

- spatial covariance, SRS-MB has a much lower RMSE than SRS-DB, suggesting
- that the lack of efficiency from SRS is largely mitigated by model-based inference.
- These RMSE conclusions are similar to those observed in the grid location
- 417 layout, so we omit a figure and discussion regarding the grid location layout here.
- 418 Tables for RMSE in all 36 simulation scenarios are provided in the supporting
- 419 information.
- 95% interval coverage for each of the four sampling-inference combinations
- in the random location layout is shown in Fig. 6. Within each simulation
- scenario, all sampling-inference combinations tend to have fairly similar interval

coverage, though when n = 50 or n = 100, GRTS-DB coverage is usually a 423 few percentage points lower than the other combinations, which suggests that 424 the local neighborhood variance estimate may be slightly too small for small n. 425 Coverage in the normal response scenarios is usually near 95%, while coverage in 426 the skewed response scenarios usually varies from 90% to 95% but increases with 427 the sample size. At a sample size of 200, all four sampling-inference combinations 428 have approximately 95% interval coverage in both response scenarios for all DRE 429 proportions. These interval coverage conclusions are similar to those observed in 430 the grid location layout, so we omit a figure and discussion regarding the grid 43 location layout here. Tables for interval coverage in all 36 simulation scenarios 432 are provided in the supporting information.

3.2. National Lakes Assessment Data

Mean bias is nearly zero for all four sampling-inference combinations in all
six scenarios, so we omit a more detailed summary of those results here. Tables
for mean bias in all six simulation scenarios are provided in the supporting
information.

The relative RMSE of both ZMMI (symmetric response) and Hg ppb (skewed response) for all four sampling-inference combinations are shown in Fig. 7. GRTS-MB has the lowest RMSE, followed by GRTS-DB, SRS-MB, and then SRS-DB. The difference in RMSE among GRTS-MB and GRTS-DB tends to be quite small. When n=50, SRS-MB RMSE is approximately evenly between SRS-DB RMSE and GRTS-MB RMSE, but for the larger sample sizes (n=100, n=200), SRS-MB RMSE is closer to GRTS-MB RMSE. Lastly we note that GRTS-MB, GRTS-DB, and SRS-MB all have noticeably lower RMSE than SRS-DB. Tables for RMSE in all six scenarios are provided in the supporting information.

95% interval coverage of both ZMMI and Hg ppb for all four samplinginference combinations is shown in Fig. 8. When n=50, interval coverage for

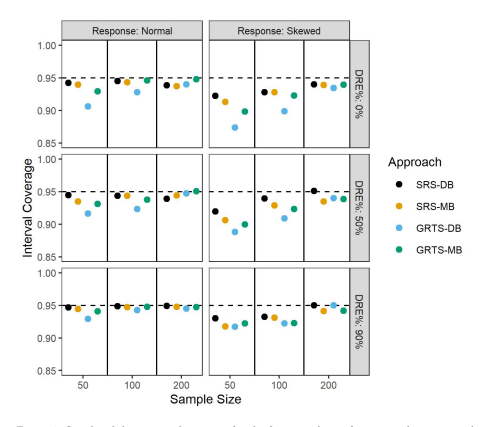


Figure 6: Simulated data interval coverage for the four sampling-inference combinations and three sample sizes in the random location layout. The rows indicate the proportion of dependent error and the columns indicate the response type. The solid black lines separate the sample sizes and the dashed black lines represent 95% coverage.

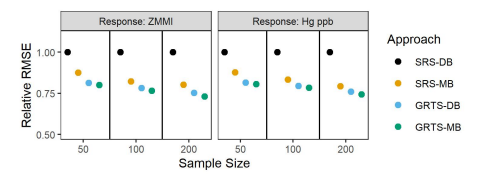


Figure 7: NLA data relative RMSE for the four sampling-inference combinations. The columns indicate the response type. The solid, black lines separate the sample sizes.

both responses is too low, though interval coverage is higher for ZMMI (symmetric 450 response) than for Hg ppb (skewed response). When n = 100, ZMMI interval 451 coverage is approximately 95% except for GRTS-DB, which has coverage around 452 92%, while Hg ppb interval coverage ranges from approximately 90% (GRTS-DB) 453 to 93% (GRTS-MB). When n = 200, ZMMI interval coverage is approximately 454 95% while Hg ppb interval coverage ranges from approximately 93% (GRTS-DB) 455 to 95% (GRTS-MB). As with the simulated data, coverages for the NLA data 456 tend to increase with the sample sizes, coverages tend to be higher for symmetric 457 responses than for skewed responses, and the local neighborhood variance was slightly too small for small n, yielding slightly lower interval coverages than the 459 other sampling-inference combinations. Recall that model-based inference defines interval coverage properties across realized populations. With the simulated data, 461 we evaluated interval coverage across realized populations, but for the NLA data, we evaluated interval coverage within a single realized population. We did find 463 that model-based coverages were similar to the design-based coverages, however, 464 suggesting that for some realized populations it is reasonable to heuristically 465 view data from separate random samples as being from approximately separate 466 realized populations. But generally, if model-based intervals constructed from 467 many random samples of a single realized population show improper coverage, 468 this does not necessarily imply a deficiency in model-based inference. Tables for 469 interval coverage in all six simulation scenarios are provided in the supporting 470 information. 471

472 4. Discussion

The design-based and model-based approaches to frequentist statistical inference rest on fundamentally different foundations. Design-based approaches rely on random sampling to estimate population parameters. Model-based approaches

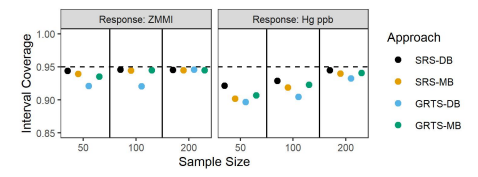


Figure 8: NLA data interval coverage for the four sampling-inference combinations. The columns indicate the response type. The solid black lines separate the sample sizes and the dashed black lines represent 95% coverage.

rely on distributional assumptions to predict realized values of a data-generating 476 stochastic process. Though model-based approaches do not rely on random 477 sampling, random sampling can still be beneficial as a way to guard against pref-478 erential sampling. While design-based and model-based approaches have often 479 been compared in the literature from theoretical and analytical perspectives, 480 our contribution lies in studying them for finite population spatial data while 481 implementing GRTS sampling and the local neighborhood variance estimator. 482 Aside from the theoretical differences described throughout the manuscript, a 483 few analytical findings from the simulated and real data studies were particularly 484 notable. All sampling-inference combinations had approximately zero mean bias. Independent of the inference approach, the GRTS samples yielded lower RMSE 486 than their SRS counterparts. Though GRTS-DB and GRTS-MB generally had 487 very similar RMSE, SRS-MB tended to have much lower RMSE than SRS-DB, 488 suggesting that the model-based inference mitigated much of the inefficiency in 489 RMSE from SRS. As the proportion of dependent random error in the simulated 490 data increased, SRS-MB, GRTS-DB, and GRTS-MB become increasingly more 491 efficient (lower RMSE) than SRS-DB. Interval coverage tended to be higher for 492 the symmetric responses than skewed responses and tended to increase with the

sample size. At a sample size of n=200, generally all interval coverages were near the desired value of 95%.

There are several benefits and drawbacks of the design-based and model-based 496 approaches for finite population spatial sampling and inference. Some we have 497 discussed, but others we have not, and they are worthy of discussion. First, we 498 discuss advantages of the design-based approach. Design-based inference is often 499 computationally efficient, while model-based inference can be computationally 500 burdensome, especially for likelihood-based estimation methods like REML that 501 rely on the inverse of a covariance matrix. Design-based inference easily handles binary data through a straightforward application of the Horvitz-Thompson 503 estimator. In contrast, analyzing binary data using model-based inference generally requires a logistic mixed regression model, the parameters of which 505 can be difficult to estimate and interpret (Bolker et al., 2009). An advantage of design-based inference is that interval coverage is valid (has the proper coverage 507 rate) as long as 1) the sample is sufficiently large to ensure the statistic's sampling 508 distribution is approximately normal and 2) the variance estimator is consistent 509 (Brus and De Gruijter, 1997; Särndal et al., 2003). This is because with the 510 design-based approach, the sampling plan and inclusion probabilities are specified 511 directly by the researcher. An advantage of SRS-DB not previously mentioned 512 is that it is likely to be valid given the consistency of its variance estimator 513 (Särndal et al., 2003). With the model-based approach, however, interval coverage 514 is unlikely to be valid if the model assumptions made do not not accurately reflect reality. Whether model assumptions accurately reflect reality can be a 516 challenging and sometimes impossible question to answer definitively. 517

Now, we discuss advantages of the model-based approach. The modelbased approach can more naturally quantify the relationship between covariates (predictor variables) and the response variable than design-based approaches.

Model-based inference also yields estimated spatial covariance parameters, which help better understand the dependence structure of the process in study. Model 522 selection is also possible using model-based inference and criteria such as cross 523 validation, likelihood ratio tests, or AIC (Akaike, 1974). Model-based inference 524 is capable of more efficient small-area estimation than design-based inference 525 because model-based inference can leverage distributional assumptions in areas 526 with few observed population units. Model-based approaches also accommodate 527 unit-by-unit predictions at unobserved locations that can be used to construct 528 informative visualizations like smoothed maps. Brus and De Gruijter (1997) provide a more thorough discussion regarding the benefits and drawbacks of the 530 two approaches. In short, when deciding whether the design-based or modelbased approach is more appropriate to implement, these benefits and drawbacks 532 should be considered alongside the particular goals of the study.

There are many extensions of this research worthy of future consideration that include sampling with unequal inclusion probabilities, using different spatially balanced sampling approaches (instead of GRTS), using different spatial data configurations, using different spatial domains like stream networks (Ver Hoef and Peterson, 2010), using different response or covariance structures, and using spatial or external mean trends (which can be defined through covariates).

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552 Conflict of Interest Statement

There are no conflicts of interest for any of the authors.

554 Author Contribution Statement

- All authors conceived the ideas; All authors designed the methodology; MD and MH performed the simulations and analyzed the data; MD and MH led the writing of the manuscript; All authors contributed critically to the drafts and
- 558 gave final approval for publication.

559 Data and Code Availability

- This manuscript has a supplementary **R** package that contains all of the
- data and code used in its creation. The supplementary R package is hosted on
- 562 GitHub. Instructions for download are available at
- https://github.com/michaeldumelle/DvMsp.
- If the manuscript is accepted, this repository will be archived in Zenodo.

565 Supporting Information

- In the supporting information, we provide tables of summary statistics for
- ⁵⁶⁷ all 36 simulation scenarios and all six real data scenarios.

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