

A comparison of design-based and model-based approaches for spatial data.

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Abstract

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Potential Journals:

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1. Introduction

There are two general approaches for using data to make statistical inferences about a population: design-based approaches and model-based approaches. When data cannot be obtained for all units in a population (population units), data on a subset of the population units is collected in a sample. In the design-based approach, inferences about the underlying population are informed from a probabilistic process in which population units are selected to be in the sample. Alternatively, in the model-based approach, inferences are made from specific assumptions about the underlying process that generated the data. Each paradigm has a deep historical context (Sterba, 2009) and its own set of general advantages (Hansen et al., 1983).

Though the design-based and model-based approaches apply to statistical inference in a broad sense, we focus on comparing these approaches for spatial data. We define spatial data as variables measured at specific geographic locations. De Gruijter and Ter Braak (1990) give an early comparison of design-based and model-based approaches for spatial data, quashing the belief that design-based

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approaches could not be used for spatially correlated data. Thereafter, several comparisons between design-based and model-based for spatial data have been considered, but they tend to compare design-based approaches that ignore spatial locations to model-based approaches (Brus and De Gruijter, 1997; Ver Hoef, 2002; Ver Hoef, 2008). Cooper (2006) review 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 into estimators that use both design and model-based principles (see e.g. Cicchitelli and Montanari (2012), Chan-Golston et al. (2020) for a Bayesian approach, and Sterba (2009)). More recent overviews include Brus (2020) and Wang et al. (2012), but no numerical comparison has been made between design-based approaches that incorporate spatial locations and model-based approaches.

Lisa M.: Add paragraph describing contribution of manuscript.

The rest of this paper is organized as follows. In Section 2, we compare sampling and estimation procedures between the design-based approach and the model-based approach. In Section 3, we use simulated and real data to study the behavior of both approaches. And in Section 5, we end with a discussion and provide directions for future research.

2. Background

The design-based and model-based approaches incorporate randomness in fundamentally different ways. In this section, we describe the role of randomness and its effects on subsequent inferences. We then discuss specific inference methods for the design-based and model-based approaches for spatial data.

2.1. Comparing Design-Based vs. Model-Based

The design-based approach assumes the population is fixed. Randomness is incorporated in the selection of population units according to a sampling design. A sampling design assigns a positive probability of inclusion in the sample (inclusion probability) to each population unit. Some examples of commonly used sampling designs include independent random sampling (IRS), stratified random sampling, and cluster sampling. The goal is to use the sampling design and the sampled data to estimate population parameters like means and totals. These population parameters are traditionally assumed to be fixed but unknown.

Treating the data as fixed and incorporating randomness through the sampling design yields estimators having very few other assumptions. Confidence intervals for these types of estimators are typically derived using limiting arguments. Means and totals, for example, are asymptotically normally distributed by the Central Limit Theorem. Särndal et al. (2003) and Lohr (2009) provide thorough reviews of the design-based approach.

The model-based approach assumes the data are a random realization of a data-generating process. Randomness is often incorporated through distributional assumptions on this process. Instead of estimating fixed but unknown

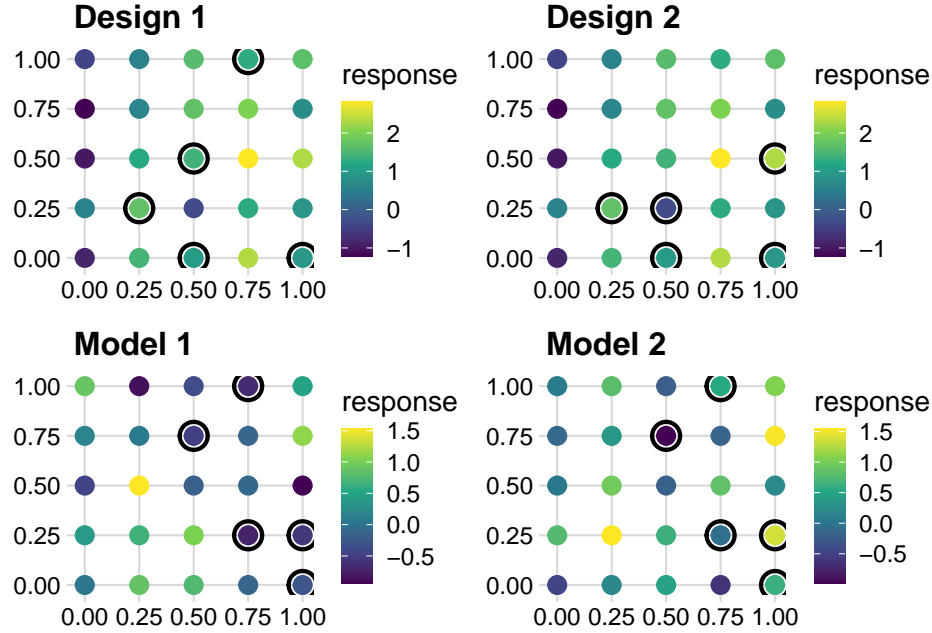


Figure 1: A comparison of sampling under the design-based and model-based frameworks. Points circled are those that are sampled. In the top row, we have one fixed population, and two random samples. In the bottom row, we have two realizations of the same spatial process sampled at the same locations.

parameters (as in the design-based approach), the goal of model-based inference in the spatial context is often *prediction* of an unknown quantity. For example, suppose the realized mean of all population units is the quantity of interest. Instead of *estimating* a fixed unknown mean, we are *predicting* the value of the mean, a random variable. We know that if we sampled all population units, we would have an exact prediction for the mean of our one realized process, without any uncertainty. But we are often not interested in the true, unknown mean of the underlying process.

Assuming the data is a realization of a specific data-generating process yields predictors that are linked to distributional assumptions. These distributional assumptions are used to derive prediction intervals. The distributional assumptions allow the prediction intervals to be more precise. Cressie (1993) and Schabenberger and Gotway (2017) provide reviews of model-based approaches for spatial data.

Description of Figure 1 goes here.

2.2. Spatially Balanced Design and Analysis

Lisa M.: Need a more precise definition of “miniature” in this context, and need an example.

97 The design-based approach can use spatial locations to obtain spatially
98 balanced samples. First we discuss spatial balance with respect to the population
99 (Stevens and Olsen, 2004). A sample is spatially balanced with respect to the
100 population if the sampled population units are a miniature of the population
101 units. A sample is a miniature of the population if the distribution of the sampled
102 population units mirrors the density of all population units. Spatial balance
103 with respect to the population is different than spatial balance with respect to
104 geography. A sample that is spatially balanced with respect to geography is
105 spread out in some type of equidistant manner over geographical space and is
106 not meant to be miniatures of the population. When we refer to spatial balance
107 henceforth, we mean spatial balance with respect to the population.

108 Spatially balanced samples are useful because they tend to yield estimates
109 that have lower variance than estimates constructed from sampling designs
110 lacking spatial balance (Barabesi and Franceschi, 2011; Benedetti et al., 2017;
111 Grafström and Lundström, 2013; Robertson et al., 2013; Stevens and Olsen,
112 2004; Wang et al., 2013). To quantify spatial balance, Stevens and Olsen (2004)
113 proposed loss functions based on Voroni polygons. The first spatially balanced
114 sampling algorithm that saw widespread use was the Generalized Random
115 Tessellation Stratified (Stevens and Olsen, 2004). Since GRTS was developed,
116 several other spatially balanced sampling algorithms have emerged, including
117 the Local Pivotal Method (Grafström et al., 2012; Grafström and Matei, 2018),
118 Spatially Correlated Poisson Sampling (Grafström, 2012), Balanced Acceptance
119 Sampling (Robertson et al., 2013), Within-Sample-Distance (Benedetti and
120 Piersimoni, 2017), and Halton Iterative Partitioning (Robertson et al., 2018).
121 We focus on the Generalized Random Tessellation Stratified (GRTS) algorithm
122 to select spatially balanced sampling because it has several attractive properties,
123 including Lisa M.: List major attractive properties, and detailed by Stevens and
124 Olsen (2004) and Dumelle et al. (2021).

125 The GRTS algorithm is used to sample from finite and infinite populations
126 and works by utilizing a mapping between two-dimensional and one-dimensional
127 space. The population units in two-dimensional space are divided into cells using
128 a hierarchical index. Population units are then mapped to a one-dimensional
129 line via the hierarchical indexing. The line length of each population unit equals
130 its inclusion probability. A systematic sample is conducted on the line and these
131 samples are linked to a population unit in two-dimensional space, which results
132 in the desired sample. Stevens and Olsen (2004) and Dumelle et al. (2021)
133 provide further details.

After collecting a sample using the GRTS algorithm, the data are used to
estimate population parameters. The Horvitz-Thompson estimator (Horvitz and
Thompson, 1952) yields unbiased estimates of population means and totals. For
example, if τ is a population total, then the Horvitz-Thompson estimator of τ
(denoted by $\hat{\tau}_{ht}$), is given by

$$\hat{\tau}_{ht} = \sum_{i=1}^n Z_i \pi_i^{-1}, \quad (1)$$

where Z_i and π_i are the observed value and inclusion probability of the i th population unit selected in the sample. A similar formula exists for estimating the mean, μ . Horvitz and Thompson (1952) and Sen (1953) provide variance estimators for $\hat{\tau}_{ht}$, but they have two drawbacks. First, they rely on calculating π_{ij} , the probability that population unit i and population unit j are included in the sample, and this can be very difficult to calculate. Second, they ignore the spatial locations of the population units. To address these drawbacks, Stevens and Olsen (2003) proposed a local neighborhood variance estimator. The local neighborhood variance estimator does not rely on π_{ij} , and it incorporates spatial locations by assigning higher weights to nearby observations. Stevens and Olsen (2003) show this variance estimator tends to reduce the estimated standard error of $\hat{\tau}$, yielding narrower confidence intervals for τ .

2.3. Finite Population Block Kriging

Finite Population Block Kriging (FPBK) is a model-based approach that expands the geostatistical Kriging framework to the finite population setting (Ver Hoef, 2008). Instead of basing inference off of a specific sampling design, we assume the data are generated by a spatial process. Ver Hoef (2008) gives details on the theory of FPBK, but some of the basic principles are summarized below. Let $\mathbf{z} \equiv \{z(s_1), z(s_2), \dots, z(s_N)\}$ be a response variable that can be measured at the N population units and is represented as an $N \times 1$ vector. Suppose we want to predict some linear function of the response variable, $f(\mathbf{z}) = \mathbf{b}'\mathbf{z}$, where \mathbf{b}' is a $1 \times N$ vector of weights. For example, if we want to predict the population total across all population units, then we would use a vector of 1's for the weights.

However, we often only have a sample of the N population units. Denoting quantities that are part of the sampled population units with a subscript s and quantities that are part of the unsampled population units with a subscript u ,

$$\begin{pmatrix} \mathbf{z}_s \\ \mathbf{z}_u \end{pmatrix} = \begin{pmatrix} \mathbf{X}_s \\ \mathbf{X}_u \end{pmatrix} \boldsymbol{\beta} + \begin{pmatrix} \boldsymbol{\delta}_s \\ \boldsymbol{\delta}_u \end{pmatrix}, \quad (2)$$

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 $\boldsymbol{\delta}_s$ and $\boldsymbol{\delta}_u$ are random errors for the sampled and unsampled population units, respectively. Denoting $\boldsymbol{\delta} \equiv [\boldsymbol{\delta}_s \ \boldsymbol{\delta}_u]'$, we assume the expectation of $\boldsymbol{\delta}$ equals $\mathbf{0}$.

We also assume that there is spatial correlation in $\boldsymbol{\delta}$, which can be modeled using a covariance function. It is common to assume the covariance function is second-order stationary and isotropic (Cressie, 1993), and that the spatial covariance decreases as the separation between population units increases. Many spatial covariance functions exist, but the primary function we use throughout the simulations and applications in this manuscript is the exponential covariance function: the i, j^{th} entry for $\text{cov}(\boldsymbol{\delta})$ is

$$\text{cov}(\delta_i, \delta_j) = \theta_1 \exp(-3h_{i,j}/\theta_2) + \theta_3 \mathbb{1}\{\mathbf{h}_{i,j} = 0\}, \quad (3)$$

where $h_{i,j}$ is the distance between population units i and j , and $\boldsymbol{\theta}$ is a vector of spatial covariance parameters of the partial sill θ_1 , the range θ_2 , and the

166 nugget θ_3 ; and, $\mathbb{1}$ is equal to 1 when distance $h_{i,j}$ is equal to 0, and equal to 0
 167 otherwise. However, any spatial covariance function could be used in the place of
 168 the exponential, including functions that allow for non-stationarity or anisotropy
 169 (Chiles and Delfiner, 1999, pp. 80–93).

170 Lisa M. : Include formulas. Perhaps, but, these are very heavy in notation
 171 and matrix algebra. We might consider, however, adding the formulas to an
 172 Appendix.

173 With the above model formulation, the Best Linear Unbiased Predictor
 174 (BLUP) for $f(\mathbf{b}'\mathbf{z})$ and its prediction variance can be computed. While details
 175 of the derivation are in (Ver Hoef, 2008), we note here that the predictor and its
 176 variance are both moment-based.

177 We note that we only use FPBK in this paper in order to focus more on
 178 comparing the design-based and model-based approaches. However, k-nearest-
 179 neighbors (Fix and Hodges, 1951; Ver Hoef and Temesgen, 2013), random forest
 180 (Breiman, 2001), Bayesian models (Chan-Golston et al., 2020), among others, can
 181 also be used to obtain predictions for a mean or total from spatially correlated
 182 responses in a finite population setting. We choose to use FPBK because it is
 183 faster than a Bayesian approach and random forest and because Ver Hoef and
 184 Temesgen (2013) showed that the method outperforms k-nearest-neighbors in
 185 many scenarios.

186 3. Numerical Study

187 There were several variables to alter in the simulations, and the names of
 188 the scenarios in future plots mirror these variables

- 189 • correlation type: dependent errors or independent errors
- 190 • error type:
 - 191 – normal: mean 0, variance 2
 - 192 – lognormal: log scale mean 0, log scale variance 2 (total variance 47)
 - 193 – lognormalbig: log scale mean 0, log scale variance 4 (total variance
 - 194 2,926)
- 195 • sample sizes: $n = 10, 50, 150$; $N = 900$
- 196 • layout: gridded vs random uniform population locations confined to a 1 x
- 197 1 unit square

198 So for example, the `inderror.normal.n50.randloc` is the simulation having
 199 independent random errors that are normal, a sample size of 50, and random
 200 population locations.

201 There were 2000 trials for each simulation. The original response (before
 202 exponentiating if applicable) for the dependent error cases was normally dis-
 203 tributed with an exponential covariance function with partial sill of 0.9, effective
 204 range of $\sqrt{2}$, and a nugget of 0.1. For the independent error cases, the partial
 205 sill was 0 and the nugget was 1.

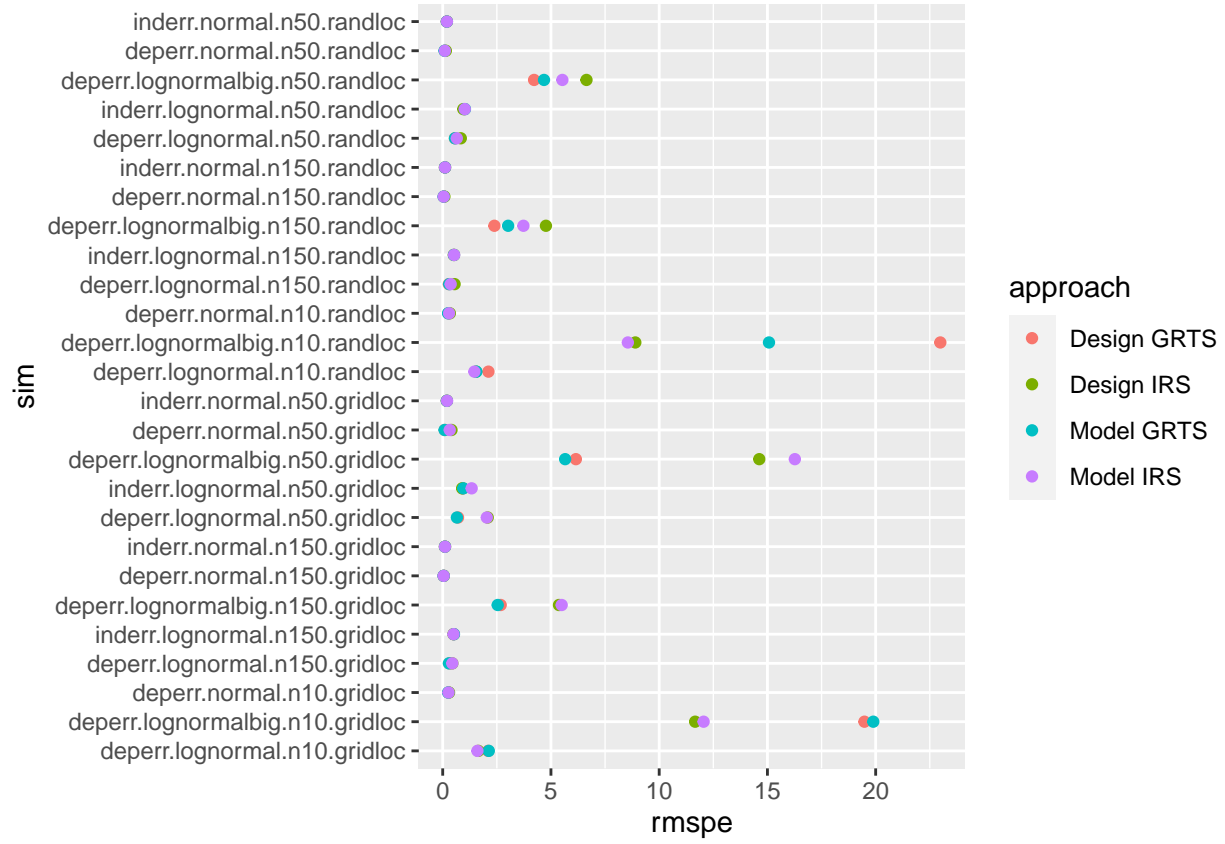
206 Lisa M. Notes: adding an intermediate level of spatial dependency? Transfer
 207 simulation scenarios to a table Explain what the effective range is Fill in the
 208 details of what exactly each approach means (perhaps a table would be a good
 209 way to do this) Reorder the sims in the first figure by some criterion. Think
 210 about what would be a “reasonable sample size” instead of 10. Define medae If
 211 the data has a large right-skew, wouldn’t one consider a transformation before
 212 the analysis? We should address this by stating that the BLUP for the log
 213 response does not mean that $e^{\log \text{BLUP}}$ is the BLUP for the response on the
 214 original scale.

215 In each simulated data set, a GRTS sample and an IRS sample were selected.
 216 Then for the GRTS sample, the design-based approach using the local neighbor-
 217 hood variance (Design GRTS) and a model-based approach were applied (Model
 218 GRTS). Then for the IRS sample, the design-based approach using the simple
 219 random sample variance (Design IRS) and a model-based approach were applied
 220 (Model IRS).

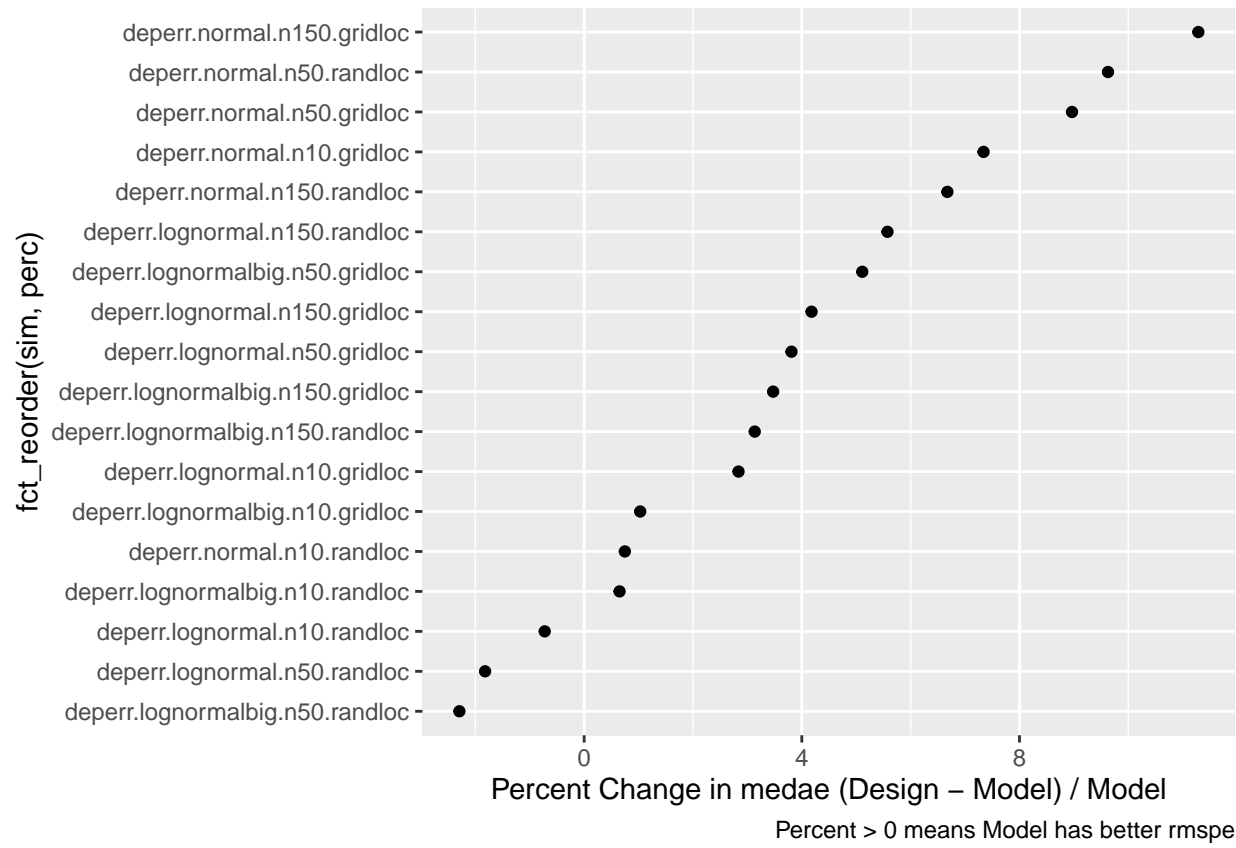
221 The GRTS algorithm and the local neighborhood variance estimator are
 222 available in the **R** package `spsurvey` (Dumelle et al., 2021). FPBK can be
 223 readily performed in **R** with the `sptotal` package (Higham et al., 2020). We
 224 use `sptotal` for both the simulation analysis and the application, estimating
 225 parameters with Restricted Maximum Likelihood (REML).

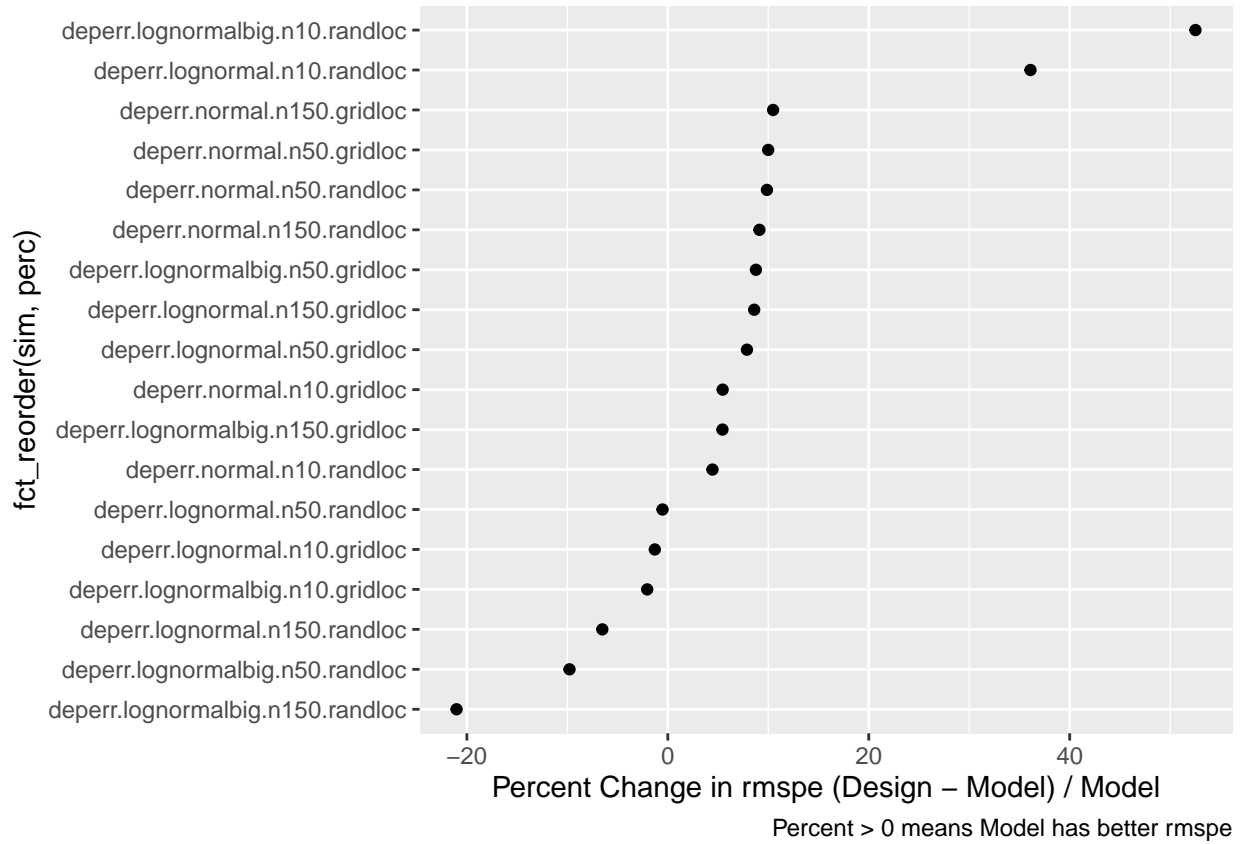
226 Major Points from August 3 Simulations:

- 227 1. In most of the dependent error simulation settings, either all four ap-
 228 proahces (IRS-Design, IRS-Model, GRTS-Design, and GRTS-Model) per-
 229 form equally or GRTS-Design and GRTS-Model outperform IRS-Design
 230 and IRS-Model. Exceptions to this are a couple of the settings with very
 231 small sample sizes ($n = 10$), in which the IRS does better than GRTS.
 232 In the independent error settings, it usually doesn’t matter much which
 233 approach is used, which makes sense.



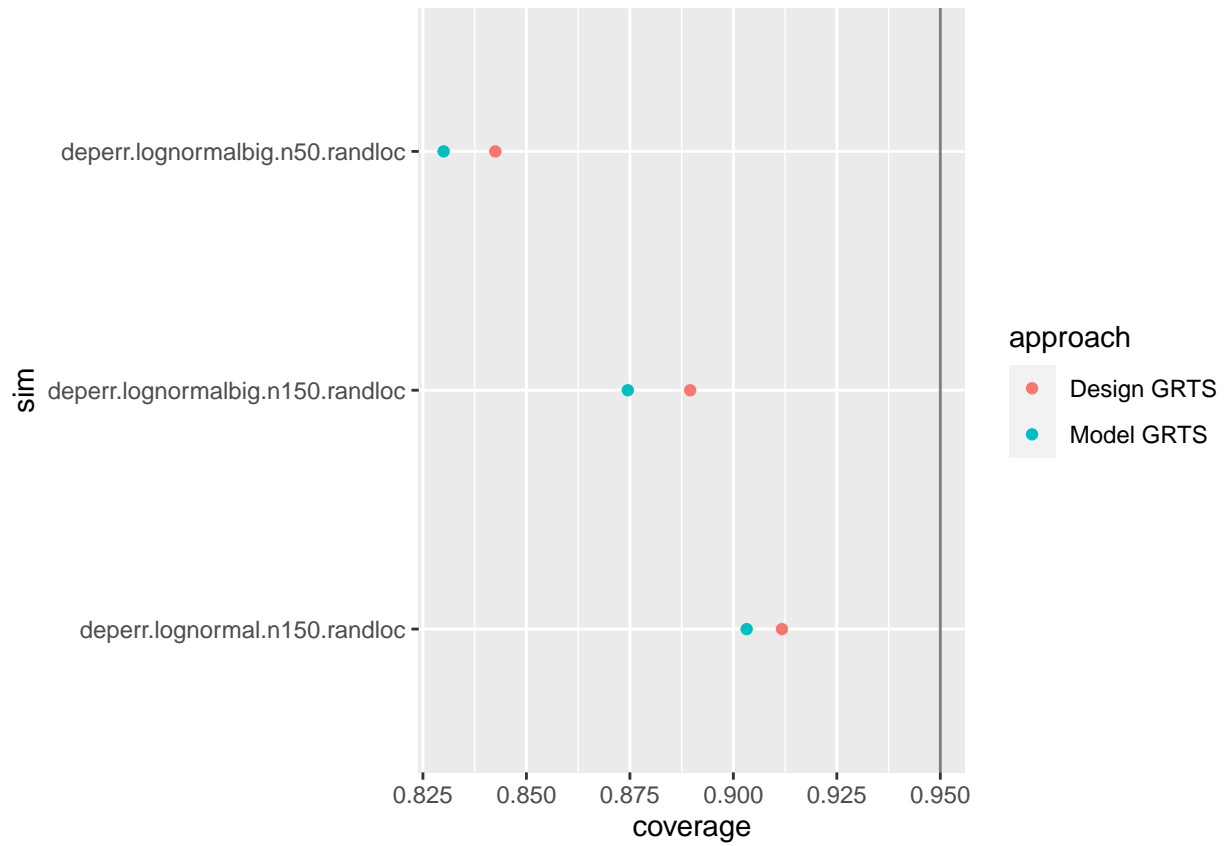
- We will now focus on comparing Design-GRTS to Model-GRTS, the two best approaches for any reasonable sample size. In the independent error settings, the two approaches perform very similarly, so those results are omitted in the following graph. In the dependent error settings, using rmspe as the performance criterion, Model-GRTS outperforms Design-GRTS in 12 of the 18 settings, the two approaches perform very similarly in 3 settings, and Design-GRTS outperforms Model-GRTS in 3 settings.





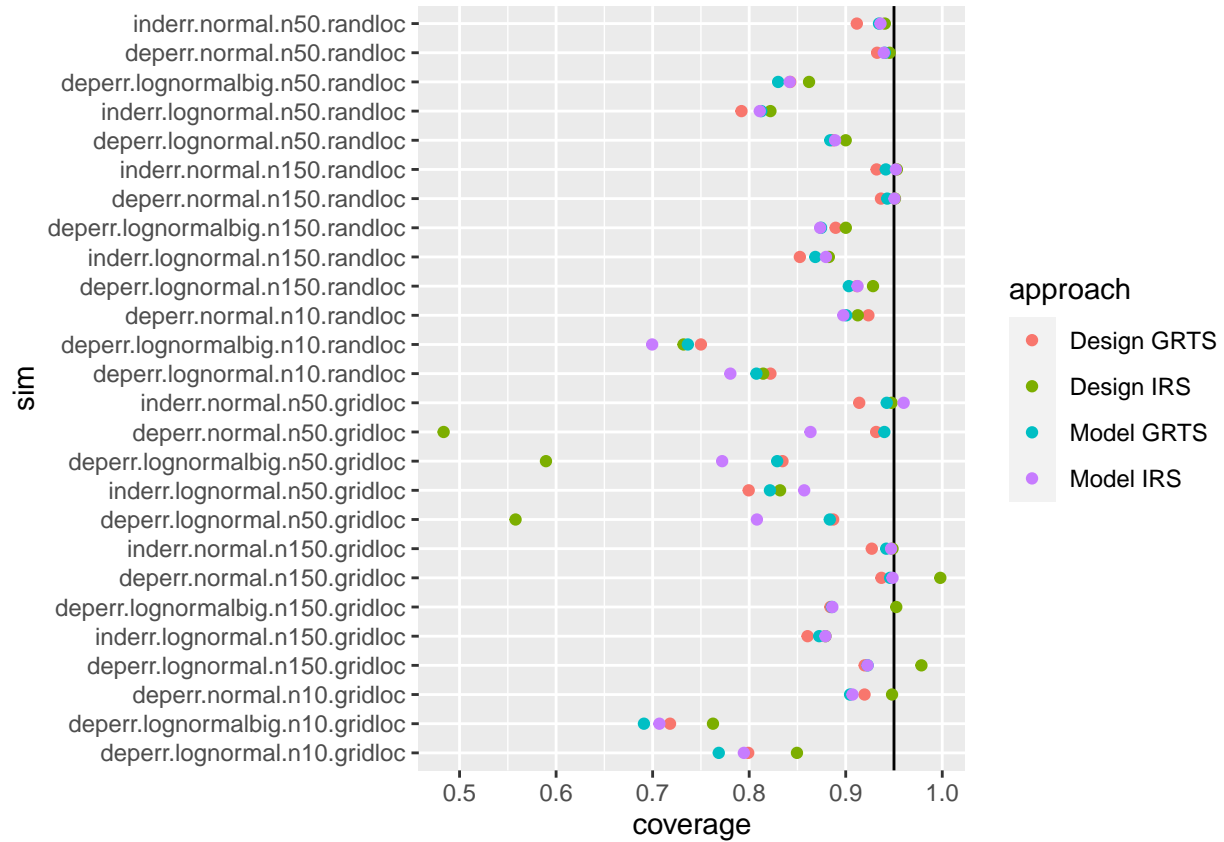
243

- 244 3. Focusing in on the three settings where Design-GRTS outperforms Model-
- 245 GRTS, we see that, in two of the settings, the log-normal response has a
- 246 large variance, corresponding to a large right-skew after exponentiation.
- 247 All three settings have sites in random locations. However, in only one
- 248 of these settings would we recommend actually using Design-GRTS. In
- 249 the other two settings, the data are sufficiently skewed that a practitioner
- 250 should not use either approach, though it is “safer” to use Design-GRTS.



4. Coverage

For Gaussian errors, coverage for all approaches tended to be near 0.95. There was less between-approach deviation in coverages for random locations compared to grid locations. Generally, the larger the skew, the worse the coverage, and the larger the sample size, the better the coverage. Design GRTS (local neighborhood variance) tended to slightly undercover, a result Tony was familiar with.



5. Take-home messages

- In terms of rmspe, a model-based analysis on a GRTS design yields an rmspe similar to or lower than a design-based analysis on a GRTS design, as long as the response variable is not “too skewed.”
- If the response variable is very skewed, then neither analysis is appropriate, but, the design-based analysis is better.
- a spatially balanced GRTS sample outperforms IRS in nearly all dependent error settings, as expected.
- methods that use spatial correlation generally perform better on random location points than they do on gridded points. This makes some intuitive sense because (1) on average, the minimum distance between an unobserved point and its nearest observed neighbor should be lower for random points and (2) the span of the study area is maximized for a grid based on the way that we set up the simulations (with the random points being drawn as uniform random variables within the boundary of the grid).
- comparison of Design-GRTS and Model-GRTS between two settings with different locations of points, but otherwise the same simulation parameters, should really be done on the same surface realization. One very strange

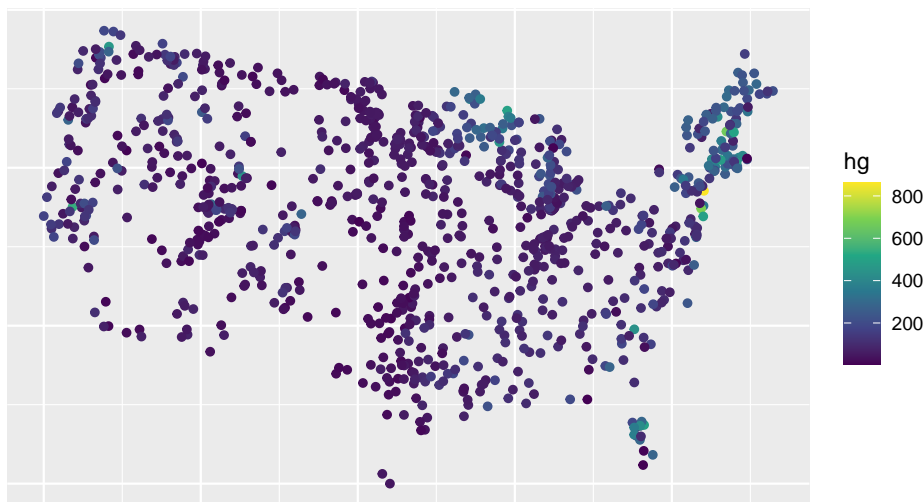


Figure 2: Population distribution of mercury concentration for 986 lakes in the contiguous United States. Thirty-five lakes were dropped from the analysis because they were missing mercury concentration.

277 realized response vector could drastically alter the results, especially on
 278 the exponentiated log data. In the same way that we compare the four
 279 approaches on the same realized data, we should also try to do the same
 280 with the locations, if they are of interest. (The realized mean won't be
 281 exactly the same but should be close).

282 4. Application

283 The Environmental Protection Agency (EPA), states, and tribes periodically
 284 conduct National Aquatic Research Surveys (NARS) in the United States to
 285 assess the water quality of various bodies of water. We will use the 2012 National
 286 Lakes Assessment (NLA), which measures various aspects of lake health and
 287 quality in lakes in the contiguous United States, to obtain an interval for mean
 288 mercury concentration. Although all lakes in the survey were measured in 2012,
 289 there may not always be enough time or money to do so. Therefore, we will
 290 explore whether or not we can still obtain an adequately precise estimate for the
 291 realized mean mercury concentration if we only take a sample of 100 of the 986
 292 lakes.

293 Figure 2 shows that mercury concentration is right-skewed, with most lakes
 294 having a low value of mercury concentration but a few having a much higher
 295 concentration. Mercury concentration exhibits some spatial correlation, with
 296 high mercury concentrations in lakes in the northeast and north central United
 297 States. Because we are considering these lakes to be our entire population, we
 298 know that the realized mean mercury concentration is 103.2 ng / g.

Table 1: Application of design-based and model-based approaches to the NLA data set on mercury concentration. The true mean concentration is 103.2 103.2 ng / g

Approach	Estimate	SE	95% LB	95% UB
Design IRS	112.7	8.8	95.4	129.9
Model IRS	110.5	7.9	95.0	125.9
Design GRTS	101.8	6.1	89.8	113.7
Model GRTS	102.3	5.9	90.8	113.9

Table 1 shows the application of a design-based analysis on an IRS, a model-based analysis on an IRS, a design-based analysis on a GRTS sample, and a model-based analysis on a GRTS sample. We see that, for all four analyses, the true realized mean mercury concentration is within the bounds of the 95% intervals. However, we should not generalize the results of this particular realization to any other data set or even to other potential samples of this data set.

But, we do note a couple of patterns. The design-based IRS analysis shows the largest standard error: a likely reason is that this is the only approach that does not use the spatial correlation in mercury concentration across the contiguous United States. We also see that, for the samples drawn, the both analyses with the GRTS sampling design have a lower standard error than the analyses with the IRS sampling design. We would expect this to be the case for most samples because mercury concentration exhibits spatial correlation so a spatially balanced sample should usually yield a lower standard error. If it is acceptable to have an interval for mean mercury concentration of about 25 ng / g and if we ignore the other variables that the EPA collects information on in these NLA surveys, then the EPA could consider sampling just 50 lakes to save time and money.

5. Discussion

- Pros of Design-Based (items we are not exploring): computationally efficient, few assumptions, more naturally handles binary data,
- Pros of Model-Based (items we are not exploring): covariate inference, more efficient small-area estimation, model selection?, estimated spatial parameters to better understand spatial structure, site-by-site predictions/prediction map

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