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Grid-spacing and the quality of abundance maps for species that show spatial autocorrelation and zero-inflation



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ABSTRACT

The effect of grid-spacing on the quality of species abundance maps is explored for species that show zero-inflation and spatial auto-correlation. Using a zero-inflated Poisson mixture model multiple fields of the prevalence parameter π and the intensity parameter μ were simulated. A selected field was sampled by grid-sampling with 200, 400, 800, 1600, and 3200 m grid-spacing and used to predict at a fixed set of validation locations by simple kriging with an external drift. The external drift variables were silt, silt squared and altitude. The estimated sampling distribution of MSE against grid-spacing shows that beyond a spacing of 1600 m the mean of MSE increases at a much faster rate. Based on these findings the 1600 m grid which consists of 446 locations for our study area of 2400 km² gives a compromise between sampling costs and prediction accuracy.

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1. Introduction

The relationship between species and their environment is generally described by species distribution models; in particular, by habitat suitability or environmental niche models (Guisan and Thuiller, 2005). Such models are constructed using survey data available at a limited set of sampling locations

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and allow one to create predictive species distribution maps on the basis of environmental data which are usually available for a much larger set of locations (Guisan and Zimmermann, 2000). The number of sampling locations is known to affect the accuracy of the species distribution models and maps (Stockwell and Peterson, 2002; Wisz et al., 2008). Knowledge of the trade-off function between number of sampling locations and accuracy of the predictions is usually not obtained a priori. If number of samples is too low, accuracy will suffer by an unknown amount: if sampling intensity is too high, the design will be unnecessarily costly. (Caughlan and Oakley, 2001; Reynolds et al., 2011).

Several studies evaluated effects of sample size on the accuracy of species distribution models (Stockwell and Peterson, 2002; Pearson et al., 2007; Wisz et al., 2008; Hanberry et al., 2012). For example, Stockwell and Peterson (2002) assessed sample size requirements for modelling bird species in Mexico by random sampling between 1 and 100 locations. Wisz et al. (2008) considered three sample sizes (10, 30, and 100 locations) to evaluate the quality of model predictions using data for 46 species obtained from natural history collections. Finally, Hanberry et al. (2012) used sample sizes ranging from 30 to 2500 locations to model tree species in northeastern Minnesota. All these studies consider presence—absence maps, but often, predictive species abundance maps in the form of numerical or biomass density are to be preferred, because they are more informative than presence—absence maps (Vieira et al., 2012; Cozzi et al., 2013). Fortin et al. (1989) constructed such maps using sugar-maple tree density data gathered in southwestern Québec. The authors evaluated the ability to predict spatial patterns using different sample sizes and designs. They considered two sample sizes of 50 and 64 points, both derived from a 200-point dataset.

Using real datasets only, as Fortin et al. (1989) did, limits comparison between the effects of different sample designs and sample number sizes as well as uncertainties in the model's parameter values. These limitations were recognized in recent studies, such as those by Perner and Schueler (2004), Rachowicz et al. (2006), Bijleveld et al. (2012) and Foster et al. (2014). An updated approach is to first simulate a spatial field resembling reality as much as possible, a pseudo-reality, which is then subsequently sampled using different sampling designs. The performance of sampling designs is then compared by confronting predictions with simulated values that serve as ground-truth. Zurell et al. (2010) call this the virtual ecologist approach. Following this approach, Bijleveld et al. (2012) used the results of an existing intertidal benthic monitoring programme to construct various spatial models with an exponential spatial autocorrelation function. With these models they simulated virtual populations with a Normal distribution and sampled these populations using different sampling designs. They provided a trade-off function between sampling distance and prediction error which was rather flat for those virtual species that hardly showed spatial autocorrelation, but much steeper for species with strong spatial autocorrelation. The assumed normality of the data was clearly violated by the empirical data because of the many zero observations.

The assumption of normality is a common practice, because when dealing with species abundance (count) data the more obvious Poisson distribution is rarely applicable. Ecological count data have two properties that ask for a specific treatment, other than relying on the classical assumption of independent and normally distributed data. These properties are zero-inflation (Martin et al., 2005; Clarke and Green, 1988; Lewis et al., 2011) and spatial autocorrelation, i.e. nearby observations are more similar than observations far apart, even when environmental conditions do not differ. Hitherto most studies have dealt with these two properties, but only one at a time (Tyre et al., 2003; Bijleveld et al., 2012).

For example, the first property was accounted by Tyre et al. (2003) who considered the zero-inflated negative binomial model, but ignored autocorrelation. Ignoring spatial autocorrelation in simulation studies on how sampling designs affect the accuracy of estimates of population or model parameters or the accuracy of spatial predictions, may lead to biased estimates of this accuracy (Legendre et al., 2002).

Contrary to Tyre et al. (2003), the Bijleveld et al. (2012) study took account of the autocorrelation by using a stochastic model that included spatial autocorrelation of the error. But, as mentioned earlier, they simulated normally distributed data. Clearly, there is a need to integrate both properties in a single study and to examine how zero-inflation and autocorrelation may affect recommendations for the optimal sampling design, sample size, and distance between samples.

Studies that simultaneously address zero-inflation and autocorrelation for species abundances (see e.g. Recta et al., 2012; Boyd et al., 2015) do not treat the question of optimal sampling design. We

attempt to fill this gap by following a paper by Lyashevska et al. (2016) in simulating fields with zero-inflated, spatially autocorrelated count data, and sampling the fields repeatedly with different sampling designs. More specifically, we will sample the fields by grid-sampling with a varying spacing. The aim of this paper is to quantify the trade-off between grid-spacing and accuracy of predictions of species-abundance model parameters on a fine grid for mapping, for species that show zero-inflation and spatial autocorrelation. Most species will show these two properties (see Martin et al., 2005, and references therein).

2. Materials and methods

2.1. Data

Data used in this paper were zero-inflated (66% are zeros) and autocorrelated counts of a benthic species *Macoma balthica* (Fig. 1(a)) that were collected in the yearly Synoptic Intertidal Benthic Surveys (SIBES) monitoring programme conducted in the Dutch Wadden Sea (Bijleveld et al., 2012; Compton et al., 2013). The study area, bordered by the barrier islands on the north and by the mainland coast on the south, is formed by intertidal and subtidal mudflats and gullies. The monitoring network consists of 3451 permanent locations on intertidal mudflats at the nodes of a 500 m grid. The square grid is supplemented by 578 locations. These locations were selected by first selecting 578 out of the 3451 grid-points by simple random sampling without replacement. Then at each selected grid-point one point was selected at a uniformly distributed distance between 0 and 250 m distance from the grid-point, in a direction randomly chosen from the four directions defined by the grid-lines (Bijleveld et al., 2012). The total sample size was 4029 locations.

The most important determinants of habitat structure used for mapping the abundance were sediment texture characteristic, more specifically the mass fraction of silt, and altitude (Amsterdam Ordnance Datum, Rijkswaterstaat¹). To be used as a predictor in mapping, the covariate must be known everywhere in the study area. Therefore the mass fraction of silt was interpolated by inverse distance weighting in ArcGIS 10.0.

2.2. Overview of evaluation method

The starting point of the our procedure for evaluating the sampling designs is a model for the spatial distribution of the zero-inflated and autocorrelated count data. This spatial distribution is modelled through a spatial zero-inflated Poisson mixture model (ZIP) (Lambert, 1992; Agarwal et al., 2002):

$$P(Y_i = y | \eta_i) = \begin{cases} \pi_i + (1 - \pi_i) \exp(-\mu_i) & y = 0\\ (1 - \pi_i) \frac{\exp(-\mu_i) \mu_i^y}{y!} & y = 1, 2, 3, \dots \end{cases}$$
(1)

where Y_i is the count at location i, η_i is spatially dependent random effect, π_i the probability of a Bernoulli zero at location i, and $1 - \pi_i$ is the probability of a Poisson count, either zero or non-zero. The intensity (mean number of individuals) of the Poisson process at location i is μ_i . The first part of the model is the overall probability of zero (Hilbe and Greene, 2007).

The parameters π_i and μ_i at location *i* are random variables modelled by the following submodels:

$$\log(\pi_{i}) = \log\left(\frac{\pi_{i}}{1 - \pi_{i}}\right) = \mathbf{x}_{B,i}^{T} \boldsymbol{\beta}_{B} + \eta_{B,i}$$

$$\log(\mu_{i}) = \mathbf{x}_{P,i}^{T} \boldsymbol{\beta}_{P} + \eta_{P,i}$$
(2)

¹ www.rijkswaterstaat.nl.

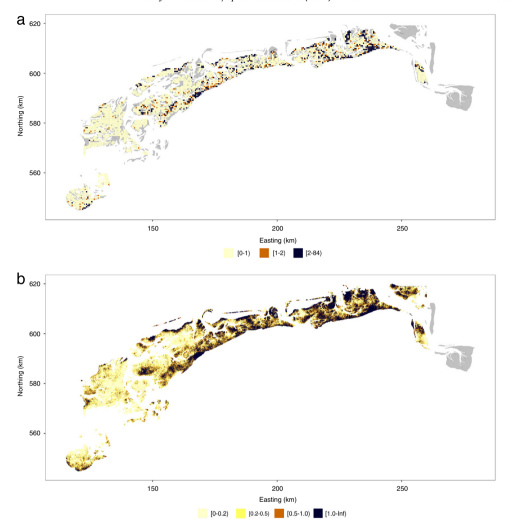


Fig. 1. Empirical species abundance map of *Macoma balthica* (a) and unconditional intensity map (b) conditionally simulated to the nodes of 100 m grid.

with $\mathbf{x}_{B,i}$ and $\mathbf{x}_{P,i}$ vectors with covariates at location i, $\boldsymbol{\beta}_B$ and $\boldsymbol{\beta}_P$ vectors with regression coefficients, and $\eta_{B,i}$, $\eta_{P,i}$ residuals of the spatial trend. Note that the model parameters can be modelled by different sets of covariates.

The residuals $\eta_{B,i}$, $\eta_{P,i}$ at any location i are random variables. The probability distribution of the residuals at all locations in the study area was modelled as

$$\begin{bmatrix} \boldsymbol{\eta}_{\mathrm{B}} \\ \boldsymbol{\eta}_{\mathrm{P}} \end{bmatrix} \sim \mathcal{N} \begin{pmatrix} \begin{bmatrix} \mathbf{0} \\ \mathbf{0} \end{bmatrix}, \begin{bmatrix} \mathbf{C}_{\mathrm{B}} & \mathbf{0} \\ \mathbf{0} & \mathbf{C}_{\mathrm{P}} \end{bmatrix}$$
 (3)

with C_B and C_P covariance matrices. So note that we assumed that the Bernoulli and Poisson residuals were independent. Testing for this assumption we revealed a weak correlation of 0.3. For both random residuals we further assumed isotropy, so that the covariance of the residuals at any two locations was modelled as a function of the distance h between the two locations. For instance, for the Bernoulli

residuals, the covariance was modelled as

$$C_{\rm B}(h) = \sigma_{\rm B}^2 \rho_{\rm B}(h;\phi_{\rm B}) + \tau_{\rm B}^2 \tag{4}$$

with σ_B^2 the partial sill, ϕ_B the range (distance parameter), τ_B^2 the nugget, and ρ_B the correlation function, for instance exponential or spherical (Webster and Oliver, 2007).

The two submodels in (2) are generalized linear mixed models, as they are the sum of a linear combination of covariates describing a spatial trend (fixed effect) and a spatially autocorrelated residual (random effect). Diggle and Ribeiro (2007) name this type of models as generalized linear *geostatistical* models.

Following Diggle and Ribeiro (2007), the sum of the trend and residual, representing the transformed model parameter, is referred to as the signal S, for instance $S_{B,i} = \mathbf{x}_{B,i}^T \boldsymbol{\beta}_B + \eta_{B,i}$. For convenience, all the parameters in one model, including the type of correlation function, are collected in a vector: $\boldsymbol{\theta}_B = (\boldsymbol{\beta}_B, \phi_B, \tau_B^2, \sigma_B^2, \rho_B)$ and $\boldsymbol{\theta}_P = (\boldsymbol{\beta}_P, \phi_P, \tau_P^2, \sigma_P^2, \rho_P)$. To avoid confusion the model parameters $\boldsymbol{\theta}_B$ and $\boldsymbol{\theta}_P$ are referred to as hyperparameters; with model parameters we mean the parameters $\boldsymbol{\pi}$ and $\boldsymbol{\mu}$.

The aim of evaluating the sampling strategy is to map the prevalence parameter π of the Bernoulli distribution and the intensity parameter μ of the Poisson distribution. Please note that the objective is not to predict the species abundance counts, but to use the observed counts in the sample to estimate, at the desired sites i, the *expected* counts conditional on the values \mathbf{x}_i of the covariates and the random effects η_i that express spatial dependence. We believe that predicting the counts themselves is not feasible in our situation, and not of practical relevance.

Our evaluation procedure is as follows. The SIBES data are used to estimate the parameters of a ZIP model. Several steps are involved in estimation. First, a ZIP model is fitted by maximum likelihood assuming that both residuals $\eta_{B,i}$ and $\eta_{P,i}$ are spatially independent. The fitted model parameters are then used to classify a zero count either as a Bernoulli or a Poisson zero, and to construct two datasets: the Bernoulli dataset with zeros (absent) and ones (present), and the Poisson subset, containing the SIBES locations with a one in the Bernoulli dataset, with counts. In the next step these two datasets are used to fit the two submodels for the parameters π_i and μ_i , but now accounting for spatial autocorrelation. This is done by simulating a large sample of signals S_B and S_P at the SIBES locations by Markov chain Monte Carlo (MCMC) using initial estimates of the hyperparameters, followed by Monte Carlo Maximum Likelihood (MCML) estimation of the hyperparameters. The final MCML estimates of the hyperparameters are used to simulate signals S_B and S_P at the SIBES locations conditional on the observations at these locations.

The fitted hyperparameters are then used to simulate the Bernoulli signal (S_B) and Poisson signal (S_P) at the nodes of a very fine square grid with a spacing of 100 m covering the study area. This grid is extended with 1000 randomly selected validation locations in between the grid-points.

In the next step the simulated signals at the grid-nodes and validation points are used to simulate fields with count data. One field with statistics closest to those of the SIBES data is selected, and underlying S_B and S_P fields are repeatedly sampled by grid-sampling. A range of grid-spacings is applied. Each selected grid is used to predict the model parameters π_i and μ_i at the validation locations. By comparing these predictions with the true (original) model parameters at the validation points the quality of the predictions is assessed. More details on all steps but the first (estimation of ZIP model parameters) are given below. For details on the first step we refer to Lyashevska et al. (2016).

- 1. The simulated signals S_B at the nodes of 100 m grid extended with 1000 validation locations are backtransformed using the inverse of the link function in Eq. (2) to give 100 fields of the prevalence parameter π of the Bernoulli distribution. The same procedure is followed for the S_P to give 100 fields of the intensity parameter μ of the Poisson distribution.
- 2. Apart from the field with zero-inflated counts, the two underlying fields with simulated prevalence parameter values π and simulated intensity parameter values μ are selected, as these are needed in the validation. Fig. 1 shows a map of the product of π and μ , representing the unconditional intensity (unconditional expected count), and a map of the SIBES count data. There is a clear resemblance between the two maps.
- 3. The two underlying fields are sampled on a grid with a spacing of 200, 400, 800, 1600, and 3200 m. For each grid-spacing 100 samples are randomly selected. The corresponding number of grid-points was on average 28 505, 7130, 1783, 446, and 110, respectively. An overlay is made of the

selected grids which are then used in prediction of the prevalence parameter π and the intensity parameter μ at the validation locations.

4. The model parameters π and μ at the 1000 validation locations are predicted by the same method as used in simulating our pseudo-reality, being simple kriging with an external drift, using silt, silt-squared and altitude as external drift variables. Ideally, for each grid-sample the hyperparameters are estimated from the 'pseudo-observations' of zero-inflated counts at the grid-points with Markov chain Monte Carlo maximum likelihood (MCML). Using these hyperparameters signals S_B and S_P should be simulated again conditional on the simulated counts (pseudo-observations). However, this is not feasible due to the computing time involved. Therefore, in predicting from the selected grid-points to the validation points we used the hyperparameters that were also used to simulate all 100 fields. These hyperparameters (referred to hereafter as the parent-hyperparameters) were estimated by MCML from the SIBES data. In practice these hyperparameters are unknown, so that by using the unknown parent-hyperparameters we ignore the contribution of uncertainty about the hyperparameters to the uncertainty about the predictions.

To obtain a rough idea about this contribution, per grid-spacing four grid-samples are selected that are used to estimate the hyperparameters. As a consequence the hyperparameters are not fixed but vary between the four samples of a given grid-spacing. The hyperparameters are not estimated from the pseudo-observations of zero-inflated counts at the selected grid-points, but from the Bernoulli and Poisson signals at these points. In doing so we avoid the time-demanding MCML estimation. By using the simulated signals as observations the hyperparameters can be estimated by Residual Maximum Likelihood (REML). We are aware that this estimation procedure does not reflect practice either, and that the contribution of uncertainty about the hyperparameters will be underestimated, but we see it as a first attempt within reasonable computing time.

2.3. Quality measures

The quality of the predicted prevalence parameters was quantified by the Mean Squared Error (MSE); for instance for the prevalence parameter π this MSE equals:

$$MSE(\pi) = \frac{1}{n} \sum_{i=1}^{n} \left\{ \hat{\pi}_i - \pi_i \right\}^2$$
 (5)

with n the number of validation points (n=1000), $\hat{\pi}_i$ the predicted prevalence parameter at location i and π_i the 'pseudo-observed' intensity parameter. For intensity parameter μ MSE is computed from the subset of validation points with a simulated value of 1 for the presence/absence indicator (species present). This subset contains 211 points. MSE was also calculated for the product of π and μ , representing unconditional intensity (intensity not conditioned on presence). For this product again all 1000 validation points are used.

For each grid-spacing we have 100 grid-samples. All 100 grid-samples are used in prediction with the fixed parent-hyperparameters $\theta_{\rm B}$ and $\theta_{\rm P}$ (no sample-specific estimation), leading to 100 estimates of ${\rm MSE}(\pi)$, ${\rm MSE}(\mu)$ and ${\rm MSE}(\pi \cdot \mu)$. The distribution of these 100 estimates is an estimate of the sampling distribution of the estimated mean quality of model-based predictions of the model parameters π and μ . Only five grid-samples are used in prediction with variable sample-specific estimates of the hyperparameters $\theta_{\rm B}$ and $\theta_{\rm P}$, so these five estimates give a very rough estimate of the sampling distribution only.

3. Results

3.1. Prevalence

The mean of the 100 MSEs (1 MSE per grid of a given spacing) of the predicted species prevalence parameter π increased with increasing grid-spacing (Fig. 2(a)). Using the fixed parent-hyperparameters the increase was from 0.006 at 200 m to 0.008 at 3200 m. The variance of MSE between the 100 grid samples was small for all grid-spacings.

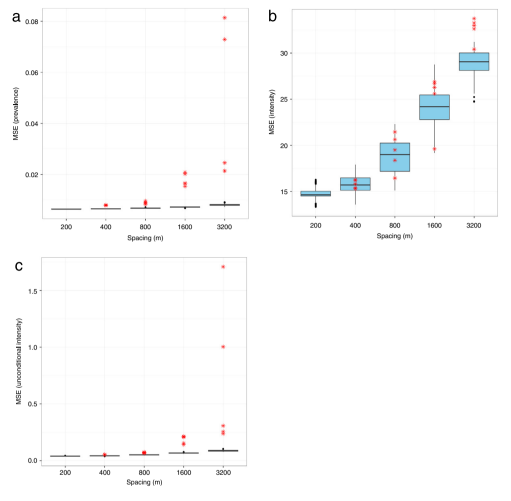


Fig. 2. The MSE for predicted prevalence (π) (a), predicted intensity (μ) (b) and predicted unconditional intensity $(\pi \cdot \mu)$ (c) as a function of grid-spacing for 200, 400, 800, 1600, and 3200 m. Predictions were obtained by simple kriging with an external drift with fixed parent-hyperparameters (blue) and hyperparameters estimated from a sample (red). All 100 grid-samples were used in prediction with fixed parent-hyperparameters and only five grid-samples were used in prediction with variable sample-specific estimates. The first grid-spacing (200 m) with estimated hyperparameters could not be estimated due to memory constraints. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

The mean of the five MSE values using hyperparameters estimated from the grid-samples, was for all spacings larger than the mean of MSE using the fixed parent-hyperparameters, especially for the largest spacing of 3200 m. This shows that the contribution of uncertainty about the hyperparameters to the uncertainty about the predictions was substantial. Remarkable is the strong increase of the mean MSE beyond a spacing of 1600 m.

3.2. Intensity

Similar to the prevalence parameter π , the mean of MSE of the predicted intensity parameter μ increased with increasing grid-spacing (Fig. 2(b)). Using fixed parent-hyperparameters the increase was from 14.73 at 200 m to 28.84 at 3200 m (Fig. 2(b)). Using hyperparameters estimated from the

grid-samples, the mean of MSE increased even more, from 15.89 for 400 m (compared with 15.73 at 400 m for fixed hyperparameters) to 32.32 at 3200 m.

The graph of the MSE for the product of π and μ , the unconditional intensity, is very similar to the graph for the prevalence parameter π . For the first four spacings the increase of the mean MSE was very modest, but beyond a spacing of 1600 m, the increase was much stronger (Fig. 2(c)).

4. Discussion and conclusions

The aim of this paper was to quantify the effect of grid-spacing on the quality of spatial predictions of the abundance of species that show zero-inflation and spatial autocorrelation. We proposed an approach in which multiple fields of the prevalence parameter π and the intensity parameter μ of a zero-inflated Poisson mixture model are simulated by generalized linear geostatistical models. These fields were used to simulate pseudo-realities of zero-inflated autocorrelated counts. One pseudo-reality was then selected with summary statistics that were close to summary-statistics of the available data. This pseudo-reality was then sampled by the sampling design under study, in this study grid-sampling at various grid-spacings. A selected sample was used to predict at a fixed set of validation locations and to compute the MSE of the predictions of π and μ . By repeating the selection of samples and the prediction at the validation points, an estimate of the sampling distribution of MSE is obtained.

To construct the graph with the estimated sampling distribution of MSE against grid-spacing the hyperparameters that were used to simulate the pseudo-reality were also used in spatial prediction at the validation points. As a consequence, this graph shows the effect of the grid-spacing given the model. This is common practice in designing spatial samples for mapping by kriging. McBratney and Webster (1981) optimized the spacing of grids using as an evaluation criterion the maximum kriging variance as obtained with ordinary kriging. Uncertainty about the variogram used in ordinary kriging is not accounted for. van Groenigen et al. (1999) optimized the spatial coordinates of a given number of sampling locations for ordinary kriging, also assuming that the variogram is known. Brus and Heuvelink (2007) did the same for kriging with an external drift (KED). This study resembles the study described here, apart from that a linear mixed model is used instead of a generalized linear mixed model. The variance as computed with KED does account for uncertainty about the trend coefficients, but does not account for uncertainty about the variogram parameters, so also in this study the contribution of uncertainty about the variogram parameters is ignored.

Alternatively, in order to account for the uncertainty about the hyperparameters in estimating the MSE, the hyperparameters that are used in prediction are estimated from the samples. By repeatedly selecting samples with a given sampling design, estimating the hyperparameters, and predicting at the validation points, a sampling distribution of the MSE is obtained with a mean that will be larger than that of the sampling distribution of MSE obtained with the fixed parent-hyperparameters. The difference in the sampling distributions of MSE reflects the contribution of the uncertainty about the hyperparameters to uncertainty about the predictions due to sampling errors in the estimated hyperparameters.

This procedure for evaluating sampling designs is relatively simple and versatile. However, the applicability of this approach in our case study was hampered by the computing time involved in estimating the hyperparameters from a sample. In the proposed model for the zero-inflated counts spatial dependency is introduced at the level of the model parameters π and μ , which cannot be directly observed. Besides, both model parameters are non-linearly related to environmental covariates. The parameters of such model can be estimated by Monte Carlo Maximum Likelihood (MCML), which involves repeated simulation of long Markov chains, which is time consuming. As an approximation we estimated the hyperparameters by REML from the unobservable model parameters π and μ at the selected sampling locations. Most likely this approximation underestimates the contribution of the uncertainty about the hyperparameters to the uncertainty about the predictions, but it is a first attempt within reasonable computing time. To improve the quality of the estimated MSEs at various grid-spacings a more efficient procedure for estimating the hyperparameters is needed. We welcome research into estimation of the hyperparameters of a ZIP mixture model by integrated nested Laplacian approximation, as proposed by Rue et al. (2009).

For the time being we must base our decisions on the graphs as presented in Fig. 2(a)–(c). The graphs of the prevalence parameter π and of the unconditional intensity $\pi \cdot \mu$ show that beyond a spacing of 1600 m the mean of MSE starts increasing at a much faster rate. This is especially true for the MSEs obtained with hyperparameters estimated from the samples. Based on these findings the 1600 m grid which consists of 446 locations for our study area of 2400 km² seems to be a good compromise between sampling costs and prediction accuracy. The increase in accuracy of the 800 m grid does not seem to outweigh the fourfold increase in number of sampling units taken. But of course the optimal spacing can only be determined when sampling costs and accuracy are defined in a common unit.

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Supporting information

Data and R-code can be found on https://github.com/lyashevska/GridSpacing-Paper.git.

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