

<sup>2</sup> Cautions, Solutions, and a Comment on Ladle et al. (2016)

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## Summary

- 1. There are now many examples where ecological researchers used non-Euclidean distance metrics in geostatistical models that were designed for Euclidean distance, such as those used for kriging. This can lead to problems where predictions have negative variance estimates. Technically, this occurs because the spatial covariance matrix, which depends on the geostatistical models, is not guaranteed to be positive definite when non-Euclidean distance metrics are used.
- 2. I give a quick review of kriging and illustrate the problem with several fabricated examples, including locations on a circle, locations on a linear dichotomous network like streams, and locations on a linear trail or road network. I re-examine the linear network distance models from Ladle et al. (2016) and show that they are not guaranteed to have a positive definite covariance matrix.
  - 3. I introduce the reduced rank method, also called predictive process models, fixed-rank kriging, and spatial basis functions, for creating valid spatial covariance matrices with non-Euclidean distance metrics. It has an additional advantage of fast computation for large data sets.
  - 4. I re-analyze the data of Ladle et al. (2016), showing that their fitted models, which used linear network distance in a geostatistical model without any nugget effect, had poor predictive performance compared to a model using Euclidean distance with a nugget effect, and it also had improper coverage for the prediction intervals. The reduced rank approach using linear network distances had the best predictive performance and had proper coverage for the prediction intervals.

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8 KEY WORDS: spatial statistics, geostatistics, prediction, reduced-rank methods, predictive process

29 models

## 1 INTRODUCTION

- 32 The variety and sophistication of statistical methods in ecology is increasing rapidly (Touchon &
- McCoy, 2016). Occasionally, this leads to researchers making mistakes when proposing to extend
- 34 a method without fully realizing that certain foundations and assumptions of that method are
- violated. There are several examples in the ecological literature were non-Euclidean distances
- were used in kriging autocorrelation models developed under a Euclidean distance assumption.
- 37 My objective is to help ecologists understand the problem and avoid this mistake. In particular, I
- 38 comment on the problems with extending kriging to linear networks advocated by Ladle et al.
- (2016), and reanalyze their data to show a better method for kriging on linear networks.

## 40 A Quick Review of Kriging

- 41 Kriging is a method for spatial interpolation, beginning as a discipline of atmospheric sciences in
- Russia, of geostatistics in France, and appearing in English in the early 1960's (Gandin, 1963;
- 43 Matheron, 1963; Cressie, 1990). Kriging is attractive because it has both predictions and
- 44 prediction standard errors, providing uncertainty estimates for the predictions. Predictions and
- their standard errors are obtained after first estimating parameters of the kriging model. The
- 46 kriging model, like the familiar regression model, can be divided into two parts: 1) the
- 47 non-stochastic part (also called the fixed effects, which includes covariates and regression
- 48 parameters) and 2) the stochastic part (the random errors). The ordinary kriging model is,

$$Y_i = \mu + \varepsilon_i,$$
 eqn 1

where  $Y_i$  is a spatial random variable at location i, i = 1, 2, ..., n, with constant mean  $\mu$  (the

fixed effect) and random error  $\varepsilon_i$ . In classical statistics, such as regression, the random errors are

assumed to be independent from each other, with a single variance parameter. For kriging, the independence assumption is relaxed, and the spatial distance among locations is used to model 52 autocorrelation among random errors. Spatial autocorrelation is the tendency for spatial variables 53 to co-vary, either in a similar fashion, or opposite from each other. The most commonly observed spatial autocorrelation is when sites closer together tend to be more similar than those that are farther apart. These tendencies are captured in autocorrelation and covariance matrices. 56 Let R be an autocorrelation matrix among spatial locations. All of the diagonal elements of 57  $\mathbf{R}$  are ones. The ith row and ith column of the off-diagonal elements of  $\mathbf{R}$  are correlations, from 58 minus one to one, between site i and j. Then a covariance matrix  $\mathbf{C} = \sigma_{\mathbf{p}}^2 \mathbf{R}$  is just a scaled 59 autocorrelation matrix that includes an overall variance,  $\sigma_{\rm p}^2$ . In constructing kriging models, practitioners often include a "nugget" effect, which is an independent (uncorrelated) random 61 effect. Constructing a full covariance matrix for a kriging model generally yields

$$\boldsymbol{\Sigma} = \mathbf{C} + \sigma_0^2 \mathbf{I} = \sigma_\mathrm{p}^2 \mathbf{R} + \sigma_0^2 \mathbf{I}, \qquad \text{eqn 2}$$

where  $\sigma_{\rm p}^2 \geqslant 0$  is called the partial sill,  $\sigma_0^2 \geqslant 0$  is the nugget effect, and **I** is the identity matrix (a diagonal matrix of all ones). The total variance is  $\sigma_{\rm p}^2 + \sigma_0^2$ . The off-diagonal elements of **R** are obtained from models that generally decrease as distance increases. Several autocorrelation models (Chiles & Delfiner, 1999, p. 80–93), based on Euclidean distance,  $d_{i,j}$ , between sites i and

j, are

$$\begin{split} & \rho_{\rm e}(d_{i,j}) = \exp(-d_{i,j}/\alpha), \\ & \rho_{\rm s}(d_{i,j}) = [1 - 1.5(d_{i,j}/\alpha) + 0.5(d_{i,j}/\alpha)^3] \mathcal{I}(d_{i,j} < \alpha), \\ & \rho_{\rm g}(d_{i,j}) = \exp(-(d_{i,j}/\alpha)^2), \\ & \rho_{\rm c}(d_{i,j}) = 1/(1 + (d_{i,j}/\alpha)^2), \\ & \rho_{\rm h}(d_{i,j}) = (\alpha/d_{i,j}) \sin(d_{i,j}/\alpha) \mathcal{I}(d_{i,j} > 0) + \mathcal{I}(d_{i,j} = 0), \end{split}$$

where distances are scaled by  $\alpha \geqslant 0$ , called the range parameter.  $\mathcal{I}(a)$  is an indicator function, equal to one if the argument a is true, otherwise it is zero.

Examples of the autocorrelation models in eqn 3, scaled with a partial sill,  $\sigma_{\rm p}^2=2$ , and a 70 nugget effect,  $\sigma_0^2 = 1$ , are shown in Figure 1a. The exponential model,  $\rho_e(d_{i,j})$ , is a very popular 71 model, and a special case of the Matern model. It approaches zero autocorrelation asymptotically. The spherical model,  $\rho_{\rm s}(d_{i,j})$ , is also very popular, and attains exactly zero autocorrelation at  $\alpha$ . 73 Both the exponential and spherical models decrease rapidly near the origin, for short distances, whereas the Gaussian model,  $\rho_{g}(d_{i,j})$ , decreases more slowly near the origin. This is also a special 75 case of the Matern model, and creates very smooth spatial surfaces. The Cauchy model,  $\rho_{\rm c}(d_{i,j})$  is similar to the Gaussian, but approaches zero autocorrelation very slowly. Finally, The hole effect 77 model,  $\rho_h(d_{i,j})$  allows for negative autocorrelation in a dampened oscillating manner. These 78 models highlight different features of autocorrelation models, and they will be used throughout 79 this paper. Many more models are given in Chiles & Delfiner (1999, p. 80–93).

Kriging is often expressed as variograms and semivariograms. Semivariograms model the variance of the difference among variables. If  $Y_i$  and  $Y_j$  are random variables at spatial locations i and j, respectively, a semivariogram is defined as  $\gamma(d_{i,j}) \equiv \mathrm{E}(Y_i - Y_j)^2/2$ , where E is expectation.

84 All of the models in eqn 3 can be written as semivariograms,

$$\gamma_m(d_{i,j}) = \sigma_p^2 (1 - \rho_m(d_{i,j})),$$
 eqn 4

where m = e, s, g, c, or h for exponential, spherical, Gaussian, Cauchy, or hole effect, respectively. Figure 1b shows semivariograms that are equivalent to the models in Figure 1a. A matrix of

semivariogram values among spatial locations can be written in terms of eqn 2,

$$\mathbf{\Gamma} = (\sigma_0^2 + \sigma_p^2)\mathbf{I} - \mathbf{\Sigma}.$$

Autocorrelation needs to be estimated from data. Empirical semivariograms have been used since the origins of kriging. First, all pairwise distances are binned into distance classes,  $\mathcal{D}_k = [h_{k-1}, h_k), \text{ where } 0 \leq h_0 < h_1 \text{ and } h_{k-1} < h_k \text{ for } k = 1, 2, \dots, K, \text{ that partition the real line}$  into mutually exclusive and exhaustive segments that cover all distances in the data set. Then the empirical semivariogram is,

$$\hat{\gamma}(h_k) = \frac{1}{2N(\mathcal{D}_k)} \sum_{d_{i,j} \in \mathcal{D}_k} (y_i - y_j)^2,$$

for all possible pairs of i and j, and k = 1, ..., K, where  $y_1, ..., y_n$  are the observed data,  $h_k$  is a representative distance (often the average or midrange) for a distance bin  $\mathcal{D}_k$ , and  $N(\mathcal{D}_k)$  is the number of distinct pairs in  $\mathcal{D}_k$ . Empirical semivariograms have desirable estimation properties (it is an unbiased estimator, Cressie, 1993, p. 71) because, substituting eqn 1 into the semivariogram definition,  $\mu$  cancels, obviating the need to estimate it. To estimate autocorrelation, one of the models in eqn 3, in semivariogram form, eqn 4, can be fit to  $\hat{\gamma}(h_k)$  as a function of  $h_k$ , often using weighted least squares (Cressie, 1985). This concept is generalized by restricted maximum likelihood (REML, Patterson & Thompson, 1971, 1974), which can be used for autocorrelation in regression models with several covariates and regression coefficients (for REML applied to spatial models, see, e.g., Cressie, 1993, p. 93). In addition, REML eliminates the arbitrary binning of distances for variogram estimation. Although REML was originally derived assuming normality, REML can be viewed as unbiased estimating equations (Heyde, 1994; Cressie & Lahiri, 1996), so normality is not required to estimate covariance parameters. Later, I will use REML for estimation. Also, I focus on covariances, rather than variograms, because their interpretation is more readily understood in the broader context of statistical models.

After covariance parameters are estimated from the data, kriging is the spatial prediction
(interpolation) for spatial locations where data were not collected. Kriging provides best linear
unbiased predictions (BLUP) in the sense of minimizing the expected squared errors between the
data as predictors, and the predictand, subject to unbiasedness (on average). The ordinary
kriging prediction equations, in terms of the covariance matrix (Schabenberger & Gotway, 2005,
p.33), are

$$\hat{Y}_{n+\ell} = \hat{\mu} + \mathbf{c}' \mathbf{\Sigma}^{-1} (\mathbf{y} - \mathbf{1}\hat{\mu}),$$
 eqn 5

for M predictions with locations indexed by  $n + \ell$ ,  $\ell = 1, 2, ..., M$ . Here,  $\mathbf{1}$  is a vector of ones,  $\hat{\mu} = (\mathbf{1}'\mathbf{\Sigma}^{-1}\mathbf{y})/(\mathbf{1}'\mathbf{\Sigma}^{-1}\mathbf{1})$ , and  $\mathbf{c}$  has, as its ith element,  $\sigma_p^2 \rho_m(d_{i,n+\ell})$ , where m is the same model (one of those in eqn 3) that was used in  $\mathbf{\Sigma}$ . The prediction variance (the expected squared errors that were minimized) is given by

$$\operatorname{var}(\hat{Y}_{n+\ell}) = (\sigma_p^2 + \sigma_0^2) - \mathbf{c}' \mathbf{\Sigma}^{-1} \mathbf{c} + \frac{(1 - \mathbf{1}' \mathbf{\Sigma}^{-1} \mathbf{c})^2}{\mathbf{1}' \mathbf{\Sigma}^{-1} \mathbf{1}}$$
eqn 6

### 118 The Problem

One of the properties shared by all models in eqn 3 is that, when  $d_{i,j}$  is Euclidean distance, the covariance matrix in eqn 2 is guaranteed to be positive definite for all possible spatial configurations of points (in 3 dimensions or less) and all possible parameter values:  $\sigma_{\rm p}^2 \geqslant 0, \sigma_0^2 \geqslant 0$ , and  $\alpha \geqslant 0$  (one of  $\sigma_{\rm p}^2$  or  $\sigma_0^2$  must be greater than zero). It is important for  $\Sigma$  to be positive definite because many estimators and predictors in statistics are linear functions of the data, kriging being one of them. That is, let  $\omega$  be a vector of weights and  ${\bf y}$  be a vector of random variables with covariance matrix  $\Sigma$ . Then an estimator or predictor  $\hat{T} = \omega' {\bf y}$  will have variance

$$\operatorname{var}(\hat{T}) = \boldsymbol{\omega}' \boldsymbol{\Sigma} \boldsymbol{\omega},$$
 eqn 7

which is guaranteed to be positive only if  $\Sigma$  is positive definite (Guillot et al., 2014). Requiring  $\Sigma$  to be positive definite is the matrix analog of requiring a variance parameter to be positive. For example, Guillot et al. (2014) demonstrate that the triangle model (not given in eqn 3), which is only valid in one dimension, yields negative variances when used with Euclidean distances based on locations in two-dimensions.

The simplest way to check whether a matrix is positive definite is to check the eigenvalues of that matrix. A covariance matrix  $\Sigma$  should be composed of real values, and it should be symmetric. Then

$$\Sigma = \mathbf{Q}\Lambda\mathbf{Q}'$$
 eqn 8

is called the spectral decomposition of  $\Sigma$ , where each column of  $\mathbf{Q}$  contains an eigenvector, and the corresponding eigenvalue is contained in  $\Lambda$ , which is a diagonal matrix. Substituting eqn 8 into eqn 7 gives

$$\operatorname{var}(\hat{T}) = \mathbf{v}' \mathbf{\Lambda} \mathbf{v} = \sum_{i=1}^{n} v_i^2 \lambda_i$$

where  $\mathbf{v} = \mathbf{Q}'\boldsymbol{\omega}$ . Because  $v_i^2 \geqslant 0$ ,  $\operatorname{var}(\hat{T})$  is guaranteed to be positive as long as all  $\lambda_i$  are greater than zero and at least one  $v_i^2$  is greater than zero. So, if the smallest eigenvalue of  $\Sigma$  is greater than zero, then  $\Sigma$  is positive definite.

Now consider using the models in eqn 3 for cases where  $d_{i,j}$  is non-Euclidean. For example, 140 let 11 spatial locations occur at equal distances on a circle (Figure 2a). Let distance be defined as the shortest path distance, so that two adjacent points have distance  $2\pi/11$ , and the maximum 142 distance between any two points is  $10\pi/11$ . The  $11\times11$  distance matrix was used with 143 autocorrelation models in eqn 3, and the minimum eigenvalue is plotted in Figure 2b. Notice that 144 as the range parameter  $\alpha$  increases, the hole effect, Gaussian, and Cauchy models have a minimum eigenvalue that is less than zero, so for these values of  $\alpha$ , the matrix is not positive 146 definite, and cannot be a covariance matrix. This example points out a further problem. It 147 appears that the exponential model and spherical model are valid models for all range values; 148 however, this is only true for 11 points that are equidistant apart. There is no guarantee that the exponential and spherical model will provide positive definite covariance matrices for other 150 sample sizes and other spatial configurations. Later, I will discuss more general approaches for 151 developing models for all spatial configurations and all values of the range parameter. 152

Another example is provided by the spatial locations at the nodes of a dichotomous network (Figure 2c). The distance between each location and the nearest node is exactly one, and there are  $2^7 - 1$  locations. Again, let distance be defined as the shortest path between any two locations, so the maximum distance between two terminal locations is  $2 \times 6 = 12$ . Using the  $127 \times 127$  distance matrix with the autocorrelation models in eqn 3 for various  $\alpha$  values showed

that all models failed to consistently yield minimum eigenvalues below zero except the
exponential model (Figure 2d). The hole effect model illustrates how erratic the positive definite
condition can be, where small changes in  $\alpha$  causes wild swings on whether the covariance matrix
is positive definite. An argument on why the exponential model is always positive definite for the
dichotomous network situation is given by Ver Hoef & Peterson (2010).

Finally, consider the 25 locations in Figure 2e. This is representative of a road or trail 163 system on a perfectly regular grid. Again, consider the shortest path distance between any two 164 points. First, consider the situation where sites are only connected by the solid lines. In that case, 165 sites one and two are not connected directly, but rather the distance between them is 3 (through 166 sites 6 and 7). Using the  $25 \times 25$  distance matrix with the autocorrelation models in eqn 3 for 167 various  $\alpha$  values shows that none of the models are positive definite for all  $\alpha$  (Figure 2f). A 168 variation occurs if we let the sites with dotted lines be connected, as well as those with solid lines. 169 In this case, the exponential model remains positive definite for all values of  $\alpha$ , and an explanation is provided by Curriero (2006). 171

Figure 2 demonstrates that, in a variety of situations, models that guarantee positive 172 definite covariance matrices for any spatial configuration, and any range value  $\alpha > 0$ , when using 173 Euclidean distance, no longer guarantee positive definite matrices when using linear network 174 distances. Similarly, one might wonder why we do not use empirical covariances in  $\Sigma$ . That is, let 175 the i, j entry in  $\Sigma$  be  $(y_i - \hat{\mu})(y_j - \hat{\mu})$ , where  $\hat{\mu}$  is the average of all  $y_i$ . Again, there is no 176 guarantee that  $\Sigma$  will be positive definite. If it is not, then what is the analyst to do? Geostatistics has a long tradition of only considering models that guarantee positive definite 178 matrices (Journel & Huijbregts, 1978, p. 161). For example, Webster & Oliver (2007, p. 80) call 179 them "authorized" models, while Goovaerts (1997, p. 87) calls them "permissible" models. All of 180 the models in eqn 3 are permissible for Euclidean distance in three dimensions or less, but they

are clearly not generally permissible for linear network distances.

#### 33 Literature Review

Many authors have used autocovariance models, such as those in eqn 3, with non-Euclidean 184 distances, and they have been roundly criticized (Curriero, 2006). For example, for streams, 185 impermissible models have been used by Cressie & Majure (1997) and Gardner et al. (2003), who 186 substituted in-stream distance for Euclidean distance, and in fact this same idea was 187 recommended in Okabe & Sugihara (2012). Alternatively, permissible models that guarantee 188 positive-definite covariance matrices were developed (based on a spatial moving averages, a 189 spatially continuous analog of moving average models in times series) by Ver Hoef et al. (2006), 190 Cressie et al. (2006) and Ver Hoef & Peterson (2010). 191 For roads and trails, impermissible models have been used by Shiode & Shiode (2011), 192 Selby & Kockelman (2013) and Ladle et al. (2016), who substitute network-based distance for 193 Euclidean distance. However, the exponential is a permissible model for a perfect grid using 194 Manhattan distance (as described for Figure 2e); see Curriero (2006). I provide a more general 195 approach based on reduced-rank radial-basis functions below. 196 In estuaries, shortest-path distances were used to replace Euclidean distance in Little et al. 197 (1997), Rathbun (1998), and Jensen et al. (2006), which yields impermissible models. Instead, 198 permissible models based on reduced-rank radial-basis functions were given by Wang & Ranalli 199 (2007).200 There has been a great deal of interest in kriging over the surface of the earth, which is an 201 approximate sphere. Kriging on geographical coordinates can create distortions, yet such 202 applications have appeared (Ecker & Gelfand, 1997; Kaluzny et al., 1998), which have been 203 criticized (Banerjee, 2005). Most research has centered on geodesic, or great-circle distance. If

geodesic distance is substituted for Euclidean distance for the models in eqn 3, only the exponential and spherical models are permissible (Gneiting, 2013). Note that distance is measured in radians, and restricted to the interval  $[0, \pi]$ .

For an interesting ecological application, Bradburd et al. (2013) propose an extension of a
powered exponential, also called a stable geostatistical model, that combines Euclidean distance
with ecological or genetic distance. Then Guillot et al. (2014) show how the stable model can be
used with geodesic (great circle) distances, but only if the power parameter of the stable model is
restricted, and they also discuss ways of "gluing" geographical distances and environmental
distances to create permissible models.

The literature given above, with many examples, shows that replacing Euclidean distance with some other metric that makes more physical sense is intuitively appealing, but may lead to covariance functions that do not guarantee positive definite covariance matrices. I will discuss this further after a re-analysis of the data in Ladle et al. (2016).

# REANALYSIS OF LADLE ET AL. (2016)

Prior to a reanalysis of Ladle et al. (2016), I list several specific criticisms of their analysis. I then
review several general approaches to spatial models for non-Euclidean distance metrics. Finally, I
introduce the reduced rank method that I ultimately use on the data of Ladle et al. (2016).

# <sup>22</sup> Criticism of Ladle et al. (2016) Analysis

These criticisms only relate to spatial modeling and kriging used in Ladle et al. (2016). The
spherical variogram model used in Ladle et al. (2016) was incorrect. Fig. 2 in Ladle et al. (2016)
shows the spherical variogram going up and then back down. The correct spherical model reaches
an asymptote and remains constant, as shown in Fig. 1b, and virtually all textbooks on

geostatistics (Journel & Huijbregts, 1978; Isaaks & Srivastava, 1989; Cressie, 1993; Goovaerts, 1997; Chiles & Delfiner, 1999; Fortin & Dale, 2005; Webster & Oliver, 2007). Because Ladle et al. 228 (2016) fit an incorrect model, none of the results for the spherical variogram model are valid. 229 To compare variogram fits, Ladle et al. (2016) used AIC based on an assumption that the residuals of a nonlinear least-squares fit were independent and Gaussian. This is not valid. Every 231 point in the empirical variogram is binned, and re-uses the same location many times, both within 232 bins and among bins. This creates a complicated correlation structure that is not independent, 233 even if the spatial data are independent. If the data are normally distributed, then the squared 234 differences, under the best conditions, are chi-squared distributed, and not Gaussian. For a 235 review, see Cressie (1993). 236 Ladle et al. (2016) fit models without a nugget effect, justifying the decision without 237 examining the data and a prior belief that no nugget was present. Examination of Fig. 2 in Ladle 238 et al. (2016) would lead most spatial statistical modelers to include a nugget effect because a visual extrapolation to the origin leads to a discontinuity from zero. Moreover, when variograms 240 are fitted without a nugget effect, they should be checked carefully for fitting and prediction 241 instabilities. It is been well-known that models without nugget can lead to computational instability when inverting the covariance matrix (Diamond & Armstrong, 1984; Posa, 1989; 243 O'Dowd, 1991; Ababou et al., 1994). If the modeler insists on excluding the nugget effect (as 244 often occurs when using kriging to approximate deterministic computer models, e.g. Martin & 245 Simpson, 2005), a small nugget effect can be added to the diagonal (e.g.  $1 \times 10^{-6}$  was used in Booker et al. (1999)) to improve computational stability. Problems can occur due to model type (Gaussian autocorrelation is the worst) and the arrangement of the spatial locations, when "near 248 duplicate" locations can cause apparently singular matrices for computational purposes (Bivand 249

et al., 2008, p. 220).

The main objective of this paper, and my prior review, is that substitution of non-Euclidean 251 distance metrics into autocorrelation models derived for Euclidean distance can create covariance 252 matrices that are not positive definite. For the particular case of Ladle et al. (2016), using their 253 linear network distance matrix in the models given in eqn 3 showed that none of the models are permissible beyond a certain  $\alpha$  value (Figure 3a). On the other hand, using the Euclidean 255 distance matrix provided by Ladle et al. (2016), all models yield positive definite covariance 256 matrices at all values of  $\alpha > 0$  (Figure 3b), which simply verifies that they are permissible models. 257 Note that the fitted exponential model had  $\hat{\alpha} = 7620$  in Ladle et al. (2016) for motorised and  $\hat{\alpha} = 14245$  for nonmotorised variables, which yielded positive definite covariance matrices because 259  $\alpha < 28224$  had all positive eigenvalues (Figure 3a). The (incorrectly) fitted spherical models in 260 Ladle et al. (2016) had estimated range parameters > 40,000, which would not yield 261 positive-definite covariance matrices because  $\alpha > 15876$  had negative eigenvalues (Figure 3a).

#### Review of Non-Euclidean Distance Models

I will review two general approaches for creating spatial models in novel situations, whether for 264 non-Euclidean distances or other situations. The first is the spatial moving average, also called a 265 process convolution and autoconvolution. The spatial moving average approach is very similar to 266 a moving average model in time series, except that the random variables that are "smoothed" are 267 continuous in space (also known as a white noise process). This approach has been used for 268 flexible variogram modeling (Barry & Ver Hoef, 1996), multivariable (cokriging) models (Ver Hoef & Barry, 1998; Ver Hoef et al., 2004), nonstationary models (Higdon, 1998; Higdon et al., 1999), 270 stream network models (Ver Hoef et al., 2006; Cressie et al., 2006; Ver Hoef & Peterson, 2010), 271 models on the sphere (Gneiting, 2013), and spatio-temporal models (Wikle, 2002; Conn et al., 272 2015). Using the moving average approach requires solving integrals to obtain the autocorrelation

networks when purely dichotomous branching occurs (Ver Hoef et al., 2006), they are not 275 tractable for more general linear networks. 276 The second approach is a reduced rank idea, also called a dimension reduction (Wikle & 277 Cressie, 1999) and spatial radial basis (Lin & Chen, 2004; Hefley et al., 2016) method, which 278 handles non-Euclidean topology and has computational advantages. This is a very general 279 method, and the one that I will use to re-analyze the data of Ladle et al. (2016). It has been used 280 for shortest path distances in estuaries (Wang & Ranalli, 2007), but it is mostly featured as a 281 method for big data sets (e.g. Wikle & Cressie, 1999; Ruppert et al., 2003; Cressie & 282 Johannesson, 2008; Banerjee et al., 2008). I will use this method for models using linear network 283 distances, which I describe next. 284

function, or approximating the integrals. For example, the integrals are tractable for stream

#### 285 Reduced Rank Methods for Non-Euclidean Distances

Let **D** denote a matrix of Euclidean distances among locations and **L** denote a matrix of linear 286 network distances. Let  $\mathbf{R}_{m,\mathbf{A},\alpha}$  be a spatial autocorrelation matrix, where  $m=\mathrm{e,\,s,\,g,\,c,}$  or h, for 287 exponential, spherical, Gaussian, Cauchy, or hole effect, respectively, for one of the models in equ 288 3, **A** is a distance matrix, either **D** or **L**, and  $\alpha$  is the range parameter for one of the models in 289 eqn 3. For example,  $\mathbf{R}_{e,\mathbf{L},\alpha} = \exp(-\mathbf{L}/\alpha)$ . Then let  $\mathbf{R}_{m,\mathbf{A},\alpha}^r$  be the matrix where some of the 290 columns of  $\mathbf{R}_{m,\mathbf{A},\alpha}$  are kept as "knots", and all other columns have been removed; hence the term 291 "reduced rank." For example, for the Ladle et al. (2016) data,  $\mathbf{R}_{m,\mathbf{A},\alpha}$  is  $239 \times 239$ , but we will 292 reduce it to just 120 columns, so  $\mathbf{R}_{m,\mathbf{A},\alpha}^r$  is 239 × 120. 293 The reduced rank method requires the selection of knots. In general, knots can be placed 294 anywhere, and not only at the observed locations. I used K-means clustering (MacQueen, 1967) 295 on the spatial coordinates to create 120 groups. Because K-means clustering minimizes

within-group variance while maximizing among-group variance, the centroid of each group tends to be regularly spaced; i.e. it is a space-filling design (e.g. Ver Hoef & Jansen, 2015). Then, the knots were moved to the nearest observed location. The original knot locations are shown in blue, and then moved to the red circles in Fig. 4. It will be useful to have the matrix of Euclidean distances among knots only, which is a subset of the rows and columns of  $\mathbf{D}$ , and we denote the knot-to-knot distances as  $\mathbf{D}^k$ .

Now consider the random effects model,

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$$\mathbf{y} = \mathbf{1}\mu + [\mathbf{R}_{m,\mathbf{A},\alpha}^r]\boldsymbol{\gamma} + \boldsymbol{\varepsilon},$$
 eqn 9

where  $\gamma$  is a vector of zero-mean random effects, and  $var(\varepsilon) = \sigma_0^2 \mathbf{I}$ . The model in eqn 9 is just a generalization of eqn 1 in vector notation. It is a mixed model, often written as

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\boldsymbol{\gamma} + \boldsymbol{\varepsilon},$$
 eqn 10

where **X** is a design matrix with covariates,  $\beta$  is a vector of regression parameters, and **Z** is a random-effects design matrix. In statistical textbooks, Z in eqn 10 often contains dummy 307 variables (zeros or ones) that indicate some factor level of the random effect. However, Z can also 308 contain covariates, in which case  $\gamma$  would contain random effects for the slope of a line, illustrating that there are no restrictions on the types of values contained in Z. In eqn 9, I have replaced Z 310 with  $\mathbf{R}_{m,\mathbf{A},\alpha}^r$ , and there are no covariates in  $\mathbf{X}$ , so  $\mathbf{X}$  is a vector of ones. A broad introduction to 311 spatial basis functions, and rank reduction, for ecologists is given by Hefley et al. (2016). 312 For the linear mixed model, eqn 10, recall that  $var(\mathbf{y}) = \sigma_p^2 \mathbf{Z} \mathbf{C} \mathbf{Z}' + \sigma_0^2 \mathbf{I}$ , where  $\mathbf{C}$  is the 313 correlation matrix for  $\gamma$  and  $\sigma_{\rm p}^2$  is an overall variance for the random effects. Classically, for 314 mixed models, random effects are assumed independent, so C = I, and then 315

var( $\mathbf{y}$ ) =  $\sigma_{\mathrm{p}}^{2}\mathbf{Z}\mathbf{Z}' + \sigma_{0}^{2}\mathbf{I}$ . The innovations for reduced-rank spatial models in eqn 9 occur because: 1) we use correlation models of distance in the random effects design matrix, essentially  $\mathbf{Z} = \mathbf{R}_{m,\mathbf{A},\alpha}^{r}, \text{ and 2}) \text{ we also allow the random effects } \boldsymbol{\gamma} \text{ to be spatially autocorrelated using the}$ inverse covariance matrix from one of the models in eqn 3. The model in eqn 9 must have a
positive definite covariance matrix, so I assume Euclidean distance will be used for the distance
among knots. In that case, a general form of eqn 9 leads to,

$$\Sigma = \sigma_{\gamma}^{2} \mathbf{R}_{m,\mathbf{A},\alpha}^{r} [\mathbf{R}_{m,\mathbf{D}^{k},n}]^{-1} [\mathbf{R}_{m,\mathbf{A},\alpha}^{r}]' + \sigma_{\varepsilon}^{2} \mathbf{I}$$
 eqn 11

In fact, each model subscript m in eqn 11 could be different, and  $\mathbf{A}$  could be either  $\mathbf{D}$  or  $\mathbf{L}$ , or some other matrix based on any number of distance metrics. The construction eqn 11 is very flexible, and several comments are pertinent:

- 1. Strictly speaking, the covariance matrix in eqn 11 is guaranteed to be nonnegative definite. 325 The part  $\sigma_{\gamma}^2 \mathbf{R}_{m,\mathbf{A},\alpha}^r [\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1} [\mathbf{R}_{m,\mathbf{A},\alpha}^r]'$  will have zero eigenvalues equal in number to the 326 rank reduction. In practice, this is not a concern because for some set of weights,  $\omega$ , every 327 value of  $[\mathbf{R}_{m,\mathbf{A},\alpha}^r]'\omega$  would need to be zero to obtain a zero variance (negative variances are 328 still not possible), which is highly unlikely. This is no different than mixed models, eqn 10, 329 where recall that the variance was  $\mathbf{ZCZ'} + \sigma^2 \mathbf{I}$ . Note that the inverse of a positive definite 330 matrix will also be positive definite, so  $[\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1}$  is positive definite as long as Euclidean 331 distance  $\mathbf{D}^k$  is used. 332
  - 2. It might seem strange to model the covariance among the knots as the inverse  $[\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1}$ . Although any positive definite matrix could be used here, and the reasons for the inverse are complex (Banerjee et al., 2008), some intuition can be gained. Suppose that the reduced rank matrix is based on Euclidean distance, that is, let  $\mathbf{A} = \mathbf{D}$ , so we have  $\mathbf{R}_{m,\mathbf{D},\alpha}^r$ . Now,

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let the knots increase in number until the knots become exactly the same as the observed 337 locations. Then,  $\mathbf{R}_{m,\mathbf{D},\alpha}^r$  becomes  $\mathbf{R}_{m,\mathbf{D},\alpha}$ , the full covariance matrix, and  $[\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1}$ 338 becomes  $[\mathbf{R}_{m,\mathbf{D},\alpha}]^{-1}$ , the inverse of the full covariance matrix, and the inverse cancels one of 339 the full covariance matrices, so in eqn 11,  $\sigma_{\mathbf{p}}^2 \mathbf{R}_{m,\mathbf{D},\alpha} [\mathbf{R}_{m,\mathbf{D},\alpha}]^{-1} [\mathbf{R}_{m,\mathbf{D},\alpha}]' = \sigma_{\mathbf{p}}^2 \mathbf{R}_{m,\mathbf{D},\alpha}$ , which 340 is the  $n \times n$  symmetric covariance matrix without any reduction in rank. By using the 341 inverse, the formulation in eqn 11 allows us to recover a typical covariance matrix as the 342 knots become equal to the observed locations. However, it is also possible to let 343  $[\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1} = \mathbf{I}.$ 344

3. It is not necessary to use reduced rank. The full covariance matrices in eqn 11 could be used, but see the next item.

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4. In addition to allowing non-Euclidean distances in the random-effects design matrix, 347  $\mathbf{R}_{m,\mathbf{A},\alpha}^r$ , there is a computational advantage to using rank reduction in eqn 11. Notice that 348  $\Sigma$  is a 239  $\times$  239 matrix, and likelihood based methods (such as maximum likelihood, or 349 restricted maximum likelihood) require the inverse of  $\Sigma$ . Computing matrix inverses is 350 computationally expensive, and grows exponentially with the dimension of the matrix (as a 351 cube of the number of locations). However, the reduced rank formulation allows an inverse 352 of  $\Sigma$  that is reduced to the size of the rank reduction by using the 353 Sherman-Morrison-Woodbury result (Sherman & Morrison, 1949; Woodbury, 1950); see an 354 excellent review by Henderson & Searle (1981). In our case, if we choose 120 knots, then the 355 inverse would be for a  $120 \times 120$  matrix rather than a  $239 \times 239$  matrix. 356

In what follows, I will always choose a single model form across all 3 components of  $\mathbf{R}_{m,\mathbf{A},\alpha}^r[\mathbf{R}_{m,\mathbf{D}^k,\eta}]^{-1}[\mathbf{R}_{m,\mathbf{A},\alpha}^r]'$ , I will always use the linear network distance matrix  $\mathbf{L}$  for  $\mathbf{A}$ , but allow the autocorrelation parameter  $\alpha$  to be different from  $\eta$ . For example, the reduced rank

exponential model that uses linear network distance has a covariance matrix

$$\Sigma = \sigma_{\mathbf{p}}^{2} \mathbf{R}_{\mathbf{e}, \mathbf{L}, \alpha}^{r} [\mathbf{R}_{\mathbf{e}, \mathbf{D}^{k}, \eta}]^{-1} [\mathbf{R}_{\mathbf{e}, \mathbf{L}, \alpha}^{r}]' + \sigma_{0}^{2} \mathbf{I}.$$
 eqn 12

For this covariance matrix, there are 4 parameters to estimate;  $\sigma_p^2$ ,  $\alpha$ ,  $\rho$ , and  $\sigma_0^2$ . In what follows,

I fit all reduced rank models using REML.

The reanalysis of Ladle et al. (2016) is given in Table 1. The data were downloaded from the

## Reanalysis of the Ladle et al. (2014) Data

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Dryad Repository http://dx.doi.org/10.5061/dryad.62t17. The parameter estimates for the two 365 exponential models found in Ladle et al. (2016) for motorised and nonmotorised variables are 366 given in the first row. To evaluate models, I use four criteria, the first being AIC (Akaike, 1973; 367 Burnham & Anderson, 2002), which assumes that the data were distributed as a multivariate normal likelihood with a spatial covariance matrix (for an example using spatial models, see 369 Hoeting et al., 2006). 370 The rest of the criteria are based on leave-one-out crossvalidation. Let  $\mathbf{y}_{-i}$  be the vector of 371 observed data with the ith observation removed. Then, using  $\mathbf{y}_{-i}$  and the estimated covariance 372 matrix, the ith observation is predicted, denoted as  $\hat{y}_i$ , with eqn 5, and its prediction stardard 373 error, denoted as  $se(\hat{y}_i)$ , is estimated with eqn 6. The correlation was computed on the set  $\{y_i, \hat{y}_i\}$ 374 for all i and reported as Corr in Table 1. Root-mean-squared prediction error (RMSPE, Table 1) 375 was computed as the square root of the mean of  $(y_i - \hat{y}_i)^2$  for all i. The coverage of the 90% 376 prediction interval (CI90, Table 1) was the percentage of times that the interval 377  $[\hat{y}_i - 1.645se(\hat{y}_i), \ \hat{y}_i + 1.645se(\hat{y}_i)]$  contained the true value  $y_i$  for all i. 378 First, I consider the fitted exponential model reported in Ladle et al. (2016) (model Ladle in 379

Table 1). Note that Ladle et al. (2016) also used correlation between predicted and observed for leave-one-out crossvalidation. Using their model, I do not get exactly the same correlation for the 381 motorised variable as Ladle et al. (2016), where they report 0.472, and I obtained 0.491; however, 382 I obtain exactly the same correlation result for non-motorised (0.639). Of particular interest is the fact that the CI90 for the model in Ladle et al. (2016) covers the true value only 74.5% of the time for the motorised variable, and only 69.9% of the time for the non-motorised variable 385 (Table 1). This is due to the lack of a nugget effect. The covariance matrix is forcing high 386 autocorrelation among sites that are close together, assuming prediction is better than it really is, 387 which results in estimated prediction errors that are too small. 388 For the remaining fits, I used REML. The empirical semivariograms in Ladle et al. (2016) 380

clearly show that there should be a nugget effect in the model. I refit the exponential model with 390 linear network distance, but I added a nugget effect and used REML (model LinEN in Table 1). 391 The nugget effect was estimated to be substantial, being more than 50% of the partial sill (1.45/1.66 for motorised, and 1.19/1.75 for non-motorised). By every cross-validation metric, 393 model LinEN did a much better job at prediction than model Ladle (Ladle et al., 2016). The 394 correlation between observed and predicted was higher, the RMSPE was lower, and the 90% prediction interval covered the true value 89.1% of the time, much closer to the nominal 90%. 396 Note that this method is not recommended because linear network distance is not permissible in 397 models designed for Euclidean distance. It merely illustrates that a nugget effect should be 398 included in the models.

Fitting a model with Euclidean distance (model EucEN in Table 1) showed that it

performed slightly better than model LinEN for both motorised and non-motorised variables

based on AIC, Corr, and RMSPE, and much better than the original Ladle model. The 90%

prediction intervals appear to be very accurate, covering the true value 90% of the time in both

404 cases.

The final four models in Table 1 used the reduced rank approach, based on exponential, 405 spherical, Gaussian, and Cauchy autocorrelation models, labeled as RRexp, RRsph, RRgau, and 406 RRcau, respectively, using the covariance matrix shown in eqn 12. The estimated covariance parameters for each of the models are shown in Table 1 for both motorised and non-motorised 408 variables. For the motorised variable, RRcau had the highest Corr value and lowest RMSPE 400 among all models, although RRexp had the lowest AIC. For the non-motorised variable, RRcau 410 had the lowest AIC and RMSPE, and RRsph had the highest correlation. In general, the Ladle 411 model performed worst, with LinEN and EucEN better than Ladle and very similar to each other, 412 but the best models were RRexp, RRsph, RRgau, and RRcau. So not only were the reduced rank 413 models the best performers, they were all completely permissible and computationally faster than 414 the full rank models. There was little actual difference among the reduced rank models in 415 performance. 416

## 417 DISCUSSION AND CONCLUSIONS

If one is going to promote a statistical method, there are several things that are incumbent on the 418 author. First, the method should be shown to be better than the method it is supposed to 419 replace. In the case of the data in Ladle et al. (2016), there is no benefit to using linear network distance compared to Euclidean distance for models LinEN and EucEN, according to any of the 421 cross-validation statistics (Table 1). While linear network distance may make intuitive sense, if 422 the data exist, there is some obligation to do a comparison. For example, for stream networks, 423 several papers show linear distance models are better than Euclidean distance in a variety of ways (Peterson et al., 2013; Isaak et al., 2014; Rushworth et al., 2015). Secondly, an 425 estimator/predictor is intimately tied to a variance estimate of that estimator/predictor. 426

Statistics is a discipline for modeling uncertainty, and that uncertainty is captured by the
standard error estimate. The standard error estimate should appropriately reflect that
uncertainty. The model presented by Ladle et al. (2016) did not have proper prediction interval
coverage, whose actual coverage was between 70 and 75% for the 90% interval (Table 1). This is
easy to check with cross-validation. It is generally advisable to add a nugget effect to
geostatistical models and let the data decide how large it should be.

While it is possible to fit impermissible models such as Ladle and LinEN (Table 1) and then 433 check the fitted model to ensure that the covariance matrix is positive definite, this practice is 434 discouraged in traditional geostatistics. First, the fitting method itself may be susceptible to 435 irregularities. For example, the hole effect model in Fig. 2 oscillates wildly. An optimization 436 routine that depends on the inverse of the covariance matrix would behave erratically, and it 437 would be hard to constrain any optimization to  $\alpha$  (range) values that guaranteed a positive 438 definite covariance matrix. Also, note that models Ladle and LinEN (Table 1) happened to have positive definite covariance matrices for the specific set of locations and estimated  $\alpha$  values, 440 resulting in cross-validation predictions that had positive variance estimates. However, when 441 predicting at locations where data were not collected, a larger covariance matrix must be considered. Let  $\Sigma_{o,o}$  be the covariance matrix among the observed locations,  $\Sigma_{o,p}$  be the 443 covariance matrix between the observed and prediction locations, and  $\Sigma_{p,p}$  be the covariance 444 matrix among the prediction locations. Then 445

$$oldsymbol{\Sigma} = \left(egin{array}{cc} oldsymbol{\Sigma}_{o,o} & oldsymbol{\Sigma}_{o,p} \ oldsymbol{\Sigma}_{o,p} & oldsymbol{\Sigma}_{p,p} \end{array}
ight)$$

must be positive definite when making predictions at unobserved locations. This can be
computationally expensive or impossible to check if there are thousands of prediction locations, as

there were in Ladle et al. (2016) (it is computationally expensive to compute eigenvalues). It is much simpler, and safer, to choose permissible models/methods that guarantee positive definite covariance matrices for all spatial configurations and model parameter values.

I have shown that a reduced rank method can be used to create permissible models that 451 guarantee positive-definite covariance matrices for spatial models using linear network distance. 452 The reduced rank method is very flexible for various spatial topologies and distance metrics, and 453 also has computational advantages. For the data from Ladle et al. (2016), there was a small 454 benefit, by lowering RMSPE, for several of the linear network distance models (RRexp and 455 RRcau) over Euclidean distance (EucEN) for the motorised variable (Table 1), and a more 456 noticeable advantage for all reduced rank models for the non-motorised variable (Table 1). For 457 the reduced rank models, consideration must be given to the number and placement of knots 458 (Ruppert et al., 2003; Gelfand et al., 2012), which continues to be an area of active research. 459 The reduced-rank methods are not the only approach for developing models for 460 non-Euclidean distance metrics. Earlier, I mentioned the spatial moving average approach, also 461 called process convolutions. For continuous domains with irregular boundaries, soap film 462 smoothing (Wood et al., 2008) is another method. The larger point of Ladle et al. (2016) is important. Scientists are realizing that Euclidean distance may not represent ecologically-relevant 464 distance. New methods using non-Euclidean distance is exciting research, but it requires 465

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466

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statisticians and ecologists to ensure statistical models have appropriate properties.

product, or firm names does not imply an endorsement by the U.S. Government.

## 472 DATA AND CODE ACCESSIBILITY

- 473 Original data from Ladle et al. (2016) were made available at the Dryad Repository
- http://dx.doi.org/10.5061/dryad.62t17. An R (R Core Team, 2017) package called
- 475 KrigLinCaution was created that contains all data, code, and analyses. This manuscript was
- created using knitr (Xie, 2014, 2015, 2016), and the manuscript combining IATEX and R code is
- 477 also included in the package. The package can be downloaded at
- https://github.com/jayverhoef/KrigLinCaution.git, with instructions for installing the package.

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Table 1: Model fits and cross-validations statistics. The top part of the table is for the motorised data found in Ladle et al. (2016), and the lower part for the non-motorised. On the left of the table are parameter estimates using notation from eqn 2, eqn 3, and eqn 12. On the right are Akaike Information Criteria (AIC) and summary statistics from cross-validation, showing Corr, the correlation between true and predicted values, root-mean-squared prediction errors (RMSPE), and proportion of times that the 90% prediction interval covered the true value (CI90).

Model	$\sigma_p^2$	$\alpha$	$\eta$	$\sigma_0^2$	AIC	Corr	RMSPE	CI90
Wiodei	$U_p$	<u> </u>						
	Motorised							
$Ladle^a$	4.72	7620				0.491	1.850	0.745
${ m Lin}{ m EN}^b$	1.66	14806		1.45	968.78	0.552	1.705	0.891
$\mathrm{EucEN}^b$	2.05	18739		1.45	968.19	0.555	1.698	0.900
$RRexp^c$	1.51	9983	2123	1.55	967.06	0.564	1.686	0.874
$RRsph^c$	1.35	31164	7964	1.61	968.80	0.553	1.700	0.891
$RRgau^c$	1.17	15495	3954	1.62	969.16	0.549	1.706	0.891
$\mathrm{RRcau}^c$	1.41	7632	1753	1.55	967.80	0.565	1.685	0.883
	Non-motorised							
$\overline{\text{Ladle}^a}$	5.09	14245				0.639	1.594	0.699
${ m Lin}{ m EN}^b$	1.75	18676		1.19	899.11	0.662	1.498	0.883
$\mathrm{EucEN}^b$	1.73	11403		1.18	904.71	0.665	1.492	0.900
$RRexp^c$	1.58	12545	3368	1.32	899.61	0.674	1.475	0.891
$RRsph^c$	1.44	25962	9393	1.33	900.21	0.678	1.468	0.887
$RRgau^c$	1.20	10721	3586	1.35	903.76	0.671	1.481	0.883
$RRcau^c$	1.48	9768	3515	1.33	899.58	0.674	1.476	0.891

<sup>&</sup>lt;sup>a</sup>Model parameters reported in Ladle et al. (2016)

<sup>&</sup>lt;sup>b</sup>LinEN, EucEN are classical exponential models with a nugget effect, using linear network distance and Euclidean distance, respectively, fit using REML.

<sup>&</sup>lt;sup>c</sup>RRexp, RRsph, RRgau, RRcau are the reduced rank models using exponential, spherical, Gaussian, and Cauchy autocorrelation models, respectively, fit using REML.

## 647 FIGURES

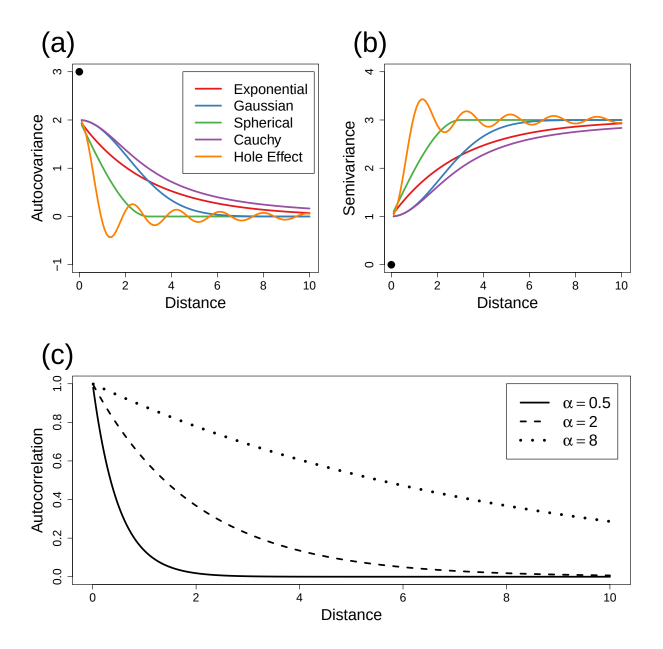


Figure 1: Autocorrelation models. (a) Autocovariance functions for various models, with a partial sill of 2 and a nugget effect of 1. (b) The same models as in (a), except represented as semivariogram models. (c) Effect of the range parameter  $\alpha$  on autocorrelation functions, where the exponential model was used as an example.

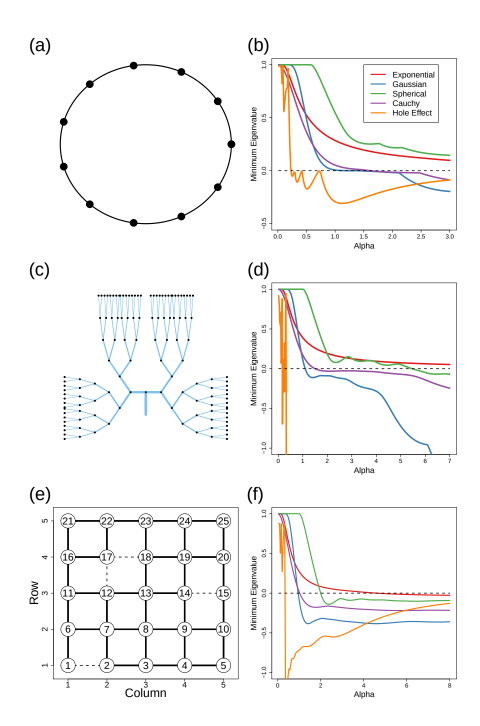
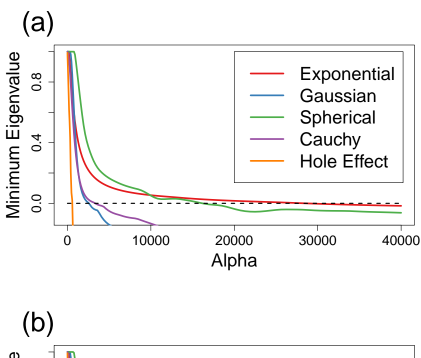


Figure 2: Cautionary examples. (a) 11 spatial locations on a circle are shown with solid circles. (b) Minimum eigenvalue for various autocorrelation models using distances on the circle. (c) A dichotomous branching network (stream) with 127 spatial locations at the node of each branch. (d) Minimum eigenvalue for various autocorrelation models using in-stream distance only. (e) 25 spatial locations on a grid network, where a perfect lattice includes the dashed line, but an irregular lattice includes only the solid lines. (f) Minimum eigenvalue for various autocorrelation models using shortest path distances along the irregular lattice.



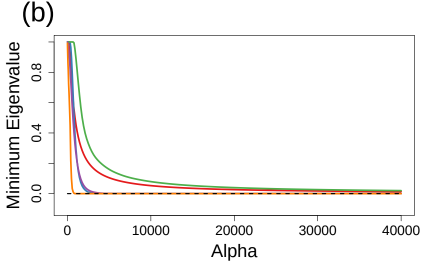


Figure 3: Minimum eigenvalues for various autocorrelation models for Ladle et al. (2016) data set. (a) Using linear distances among cameras. (b) Using Euclidean distances among cameras.

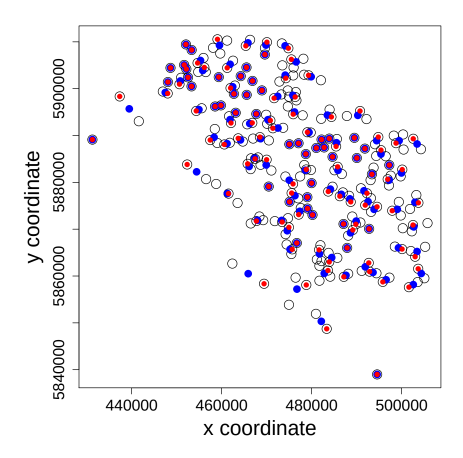


Figure 4: All spatial locations (open circles) and knot locations for reduced rank methods. Initially, k-means on x- and y-coordinates created 120 clusters with center locations given by solid blue circles, and then these were moved to nearest actual locations (solid red circles).