# Geostatistical Inference Under Preferential Sampling

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June 1, 2009

#### Abstract

Geostatistics involves the fitting of spatially continuous models to spatially discrete data (Chilès and Delfiner, 1999). Preferential sampling arises when the process that determines the data-locations and the process being modelled are stochastically dependent. Conventional geostatistical methods assume, if only implicitly, that sampling is non-preferential. However, these methods are often used in situations where sampling is likely to be preferential. For example, in mineral exploration samples may be concentrated in areas thought likely to yield high-grade ore. We give a general expression for the likelihood function of preferentially sampled geostatistical data, and describe how this can be evaluated approximately using Monte Carlo methods. We present a model for preferential sampling, and demonstrate through simulated examples that ignoring preferential sampling can lead to misleading inferences. We describe an application of the model to a set of bio-monitoring data from Galicia, northern Spain, in which making allowance for preferential sampling materially changes the results of the analysis

**Key words:** environmental monitoring; geostatistics; log-Gaussian Cox process; marked point process; Monte Carlo inference; preferential sampling.

### 1 Introduction

The term geostatistics describes the branch of spatial statistics in which data are obtained by sampling a spatially continuous phenomenon  $S(x) : x \in \mathbb{R}^2$  at a discrete set of locations  $x_i : i = 1, ..., n$  in a spatial region of interest  $A \subset \mathbb{R}^2$ . In many cases, S(x) cannot be measured without error. Measurement errors in geostatistical data are typically assumed to be additive, possibly on a transformed scale. Hence, if  $Y_i$  denotes the measured value at the location  $x_i$ , a simple model for the data takes the form

$$Y_i = \mu + S(x_i) + Z_i : i = 1, ..., n$$
(1)

where the  $Z_i$  are mutually independent, zero-mean random variables with variance  $\tau^2$ , often in this context called the *nugget variance*. One interpretation of the  $Z_i$  in (1) is as measurement errors in the  $Y_i$ . Another, which explains the more colourful terminology, is as a device to model spatial variation on a scale smaller than the shortest distance between any two sample locations  $x_i$ . We adopt the convention that E[S(x)] = 0 for all x, hence in (1)  $E[Y_i] = \mu$ for all i. The model (1) extends easily to the regression setting, in which  $E[Y_i] = \mu_i = d'_i\beta$ , with  $d_i$  a vector of explanatory variables associated with  $Y_i$ . The objectives of a geostatistical analysis typically focus on prediction of properties of the realisation of S(x) throughout the region of interest A. Targets for prediction might include, according to context: the value of S(x) at an unsampled location; the spatial average of S(x) over A or sub-sets thereof; the minimum or maximum value of S(x); or sub-regions in which S(x) exceeds a particular threshold. Chilès and Delfiner (1999) give a comprehensive account of classical geostatistical models and methods.

Diggle, Moyeed and Tawn (1998) introduced the term model-based geostatistics to mean the application of general principles of statistical modelling and inference to geostatistical problems. In particular, they added Gaussian distributional assumptions to the classical model (1) and re-expressed it as a two-level hierarchical linear model, in which S(x) is the value at location x of a latent Gaussian stochastic process and, conditional on  $S(x_i) : i = 1, ..., n$ , the measured values  $Y_i : i = 1, ..., n$  are mutually independent, Normally distributed with means  $\mu + S(x_i)$  and common variance  $\tau^2$ . Diggle, Moyeed and Tawn (1998) then extended this model, retaining the Gaussian assumption for S(x) but allowing a generalized linear model (McCullagh and Nelder, 1989) for the mutually independent conditional distributions of the  $Y_i$  given  $S(x_i)$ .

As a convenient shorthand notation to describe the hierarchical structure of a geostatistical model, we use  $[\cdot]$  to mean "the distribution of," and write  $S = \{S(x) : x \in \mathbb{R}^2\}$ ,  $X = (x_1, ..., x_n), S(X) = \{S(x_1), ..., S(x_n)\}$  and  $Y = (Y_1, ..., Y_n)$ . Then, the Diggle, Moyeed and Tawn (1998) model implicitly treats X as being deterministic and has the structure  $[S, Y] = [S][Y|S(X)] = [S][Y_1|S(x_1)][Y_2|S(x_2)]...[Y_n|S(x_n)]$ . Furthermore, in (1) the  $[Y_i|S(x_i)]$  are univariate Gaussian distributions with means  $\mu + S(x_i)$  and common variance  $\tau^2$ .

As presented above, and in almost all of the geostatistical literature, models for the data treat the sampling locations  $x_i$  either as fixed by design or otherwise stochastically independent of the process S(x). Admitting the possibility that the sampling design may be stochastic, a complete model needs to specify the joint distribution of S, X and Y. Under the assumption that X is independent of S we can write the required joint distribution as [S, X, Y] = [S][X][Y|S(X)], from which it is clear that for inferences about S or Y we can legitimately condition on X and use standard geostatistical methods. We refer to this as non-preferential sampling of geostatistical data. Conversely, preferential sampling refers to any situation in which  $[S, X] \neq [S][X]$ .

We contrast the term *non-preferential* with the term *uniform*, the latter meaning that, beforehand, all locations in A are equally likely to be sampled. Examples of designs which are both uniform and non-preferential include completely random designs and regular lattice designs (strictly, in the latter case, if the lattice origin is chosen at random). An example of a non-uniform, non-preferential design would be one in which sample locations are an independent random sample from a prescribed non-uniform distribution on A. Preferential designs can arise either because sampling locations are deliberately concentrated in sub-regions of Awhere the underlying values of S(x) are thought likely to be larger (or smaller) than average, or more generally when X and Y together form a marked point process in which there is dependence between the points, X, and the marks, Y.

We emphasise at this point that our definition of preferential sampling involves a stochastic dependence, as opposed to a functional dependence, between the process S and the sampling design X. For example, a model in which the mean of S and the intensity of X share a dependence on a common set of explanatory variables does not constitute preferential sampling. In most geostatistical applications it is difficult to maintain a sharp distinction between the treatment of variatio S(x) as deterministic or stochastic because of the absence of independent replication of the process under investigation. Our pragmatic stance is to represent by a stochastic model the portion of the total variation in S that cannot be captured by extant explanatory variables.

Curriero, Hohn, Liebhold and Lele (2002) evaluated a class of non-ergodic estimators for the covariance structure of geostatistical data, which had been proposed by Isaaks and Srivastava (1988) and Srivastava and Parker (1989) as a way of dealing with preferential sampling. They concluded that the non-ergodic estimators "possess no clear advantage" over the traditional estimators that we describe in Section 3.1 below. Schlather, Ribeiro and Diggle (2004) developed two tests for preferential sampling, which treat a set of geostatistical data as a realisation of a marked point process. Their null hypothesis is that the data are a realisation of a random field model. This model assumes that the sample locations X are a realisation of a point process  $\mathcal{P}$  on A, that the mark of a point at location x is the value at x of the realisation of a random field S on A, and that  $\mathcal{P}$  and S are independent processes. This is therefore equivalent to our notion of non-preferential sampling. Their test statistics are based on the following idea. Assume that S is stationary, and let  $M_k(h) = \mathbb{E}[S(x)^k | x, x + h \in \mathcal{P}]$ . Under the null hypothesis that sampling is non-preferential, the conditioning on  $x + h \in \mathcal{P}$  is irrelevant, hence  $M_k(h)$  is a constant. Schlather, Ribeiro and Diggle (2004) propose as test statistics the empirical counterparts of  $M_1(h)$  and  $M_2(h)$ , and implement the resulting tests by comparing the observed value of each chosen test statistic with values calculated from simulations of a conventional geostatistical model fitted to the data on the assumption that sampling is non-preferential. Guan and Afsharatous (2007) avoid the need for simulation and parametric model-fitting by dividing the observation into non-overlapping sub-regions that can be assumed to provide approximately independent replicates of the test statistics. In practice, this requires a large data-set; their application has a sample size n = 4358.

In this paper, we propose a class of stochastic models and associated methods of likelihoodbased inference for preferentially sampled geostatistical data. In Section 2 we define our model for preferential sampling. In Section 3 we use the model to illustrate the potential for inferences to be misleading when conventional geostatistical methods are applied to preferentially sampled data. Section 4 discusses likelihood-based inference using Monte Carlo methods and suggests a simple diagnostic for the fitted model. Section 5 applies our model and methods to a set of biomonitoring data from Galicia, northern Spain in which the data derive from two surveys of the same region, one of which is preferentially sampled the other not. Section 6 is a concluding discussion.

# 2 A shared latent process model for preferential sampling

Recall that S denotes an unobserved, spatially continuous process on a spatial region A, X denotes a point process on A and Y denotes a set of measured values, one at each point of X. The focus of scientific interest is on properties of S, as revealed by the data (X, Y), rather than on the joint properties of S and X, but we wish to protect against incorrect inferences that might arise because of stochastic dependence between S and X.

To clarify the distinction between preferential and non-preferential sampling, and the inferential consequences of the former, we first examine a related situation considered by Rathbun (1996), in which S and X are stochastically dependent but measurements Y are taken only at a different, pre-specified set of locations, i.e. independently of X. Then, the joint distribution of S, X and Y takes the form

$$[S, X, Y] = [S][X|S][Y|S].$$
(2)

It follows immediately on integrating (2) with respect to X that the joint distribution of S and Y has the standard form, [S, Y] = [S][Y|S]. Hence, for inference about S it is valid, if potentially inefficient, to ignore X, i.e. to use conventional geostatistical methods. Models analogous to (2) have also been proposed in a longitudinal setting, where the analogues of Y and X are a time-sequence of repeated measurements at pre-specified times and a related timeto-event outcome, respectively. See, for example, Wulfsohn and Tsiatis (1997) or Henderson, Diggle and Dobson (2000).

In contrast, if Y is observed at the points of X, the appropriate factorisation is

$$[S, X, Y] = [S][X|S][Y|X, S].$$
(3)

Even when, as is typical in geostatistical modelling, (3) takes the form

$$[S, X, Y] = [S][X|S][Y|S(X)],$$
(4)

the algebraic form of [Y|X, S] reduces to [Y|S(X)], an important distinction between preferential anbd non-preferential sampling is that in (??) the functional dependence between Sand X in the term [Y|S(X)] cannot be ignored, because the implicit specification of [S, Y]resulting from (??) is non-standard. Conventional geostatistical inferences which ignore the stochastic nature of X are therefore potentially misleading. The longitudinal analogue of (??) arises when subjects in a longitudinal study provide measurements at time-points which are not pre-specified as part of the study design; see, for example, Lipsitz, Fitzmaurice, Ibrahim, Gelber and Lipshultz (2002), Lin, Scharfstein and Rosenheck (2004) or Ryu, Sinha, Mallick, Lipsitz and Lipshultz (2007).

We now define a specific class of models through the following additional assumptions;

A1. S is a stationary Gaussian process with mean 0, variance  $\sigma^2$  and correlation function  $\rho(u; \phi) = \text{Corr}\{S(x), S(x')\}$  for any x and x' a distance u apart;

A2. conditional on S, X is an inhomogeneous Poisson process with intensity

$$\lambda(x) = \exp\{\alpha + \beta S(x)\};\tag{5}$$

A3. conditional on S and X, Y is a set of mutually independent Gaussian variates with  $Y_i \sim N(\mu + S(x_i), \tau^2)$ .

It follows from A1 and A2 that, unconditionally, X is a log-Gaussian Cox process (Møller, Syversveen and Waagepetersen, 1998). If  $\beta = 0$  in (4), then it follows from A1 and A3 that the unconditional distribution of Y is multivariate Gaussian with mean  $\mu \mathbf{1}$  and variance matrix  $\tau^2 I + \sigma^2 R$ , where I is the identity matrix and R has elements  $r_{ij} = \rho(||x_i - x_j||; \phi)$ .

Ho and Stoyan (2008) discuss essentially the same construction as A1 to A3 viewed as a model for a marked point process of locations X and marks Y, and derive its first and second moment properties.

We do not suggest that this model will be adequate for all applications. However, it is sufficiently rich to provide a vehicle for investigating the consequences of preferential sampling, and for the application described in Section 5 of the paper.

## 3 Impact of preferential sampling on geostatistical inference

We have conducted a simulation experiment in which we simulated data on A the unit square from an underlying stationary Gaussian process which we then sampled, with additive Gaussian measurement error, either non-preferentially or preferentially according to each of the following sampling designs. For the *completely random* sampling design, sample locations  $x_i$ were an independent random sample from the uniform distribution on A. For the *preferential* design, we first generated a realisation of S, then a realisation of  $X = \{x_i : i = 1, ..., n\}$ conditional on S using the model defined by equation (4) with parameter  $\beta = 2$ , and finally a realisation of  $Y = \{y_i : i = 1, ..., n\}$  conditional on S and on X using the conditional model



Figure 1: Sample locations and underlying realisations of the signal process for the model used in the simulation study. The left-hand panel shows the completely random sample, the centre-panel the preferential sample and the right-hand panel the clustered sample. In each case, the background image represents the realisation of the signal process, S(x), used to generate the associated measurement data. The model parameter values are  $\mu = 4$ ,  $\sigma^2 = 1.5$ ,  $\phi = 0.15$ ,  $\kappa = 1$ ,  $\beta = 2$ ,  $\tau^2 = 0$ 

defined by A3 above. For the *clustered* design, we used the same procedure to generate a realisation of X, but then generated a realisation of Y on the locations X using a second, independent realisation of S, so that the resulting Y is a realisation of the standard geostatistical model (1). This gives a non-preferential design with the same marginal properties for X and Y as the preferential design.

The model for the spatial process S was stationary Gaussian, with mean  $\mu = 4$ , variance  $\sigma^2 = 1.5$ , and Matérn correlation with scale parameter  $\phi = 0.15$  and shape parameter  $\kappa = 1$ . In each case, we set the nugget variance,  $\tau^2 = 0$ , hence the data  $y_i$  consisted of the realised values of  $S(\cdot)$  as the sample locations  $x_i$ .

The Matérn (1986) class of correlation functions takes the form

$$\rho(u;\phi,\kappa) = \{2^{\kappa-1}\Gamma(\kappa)\}^{-1}(u/\phi)^{\kappa}K_{\kappa}(u/\phi) : u > 0,$$

where  $K_{\kappa}(\cdot)$  denotes the modified Bessel function of the second kind, of order  $\kappa > 0$ . This class is widely used because of its flexibility. Although  $\kappa$  is difficult to estimate without extensive data, the integral part of  $\kappa$  determines the degree of mean square differentiability of the corresponding process  $S(\cdot)$ , giving both a nice interpretation and, in at least some contexts, a rationale for choosing a particular value for  $\kappa$ . The special case  $\kappa = 0.5$  gives an exponential correlation function,  $\rho(u; \phi) = \exp(-u/\phi)$ .

The three panels of Figure 1 show a realisation of each of the three sampling designs superimposed on a single realisation of the process S. The preferential nature of the sampling in the centre-panel of Figure 1 results in the sample locations falling predominantly within the darker shaded areas.

### 3.1 Variogram estimation

The theoretical variogram of a stationary spatial process Y(x) is the function  $V(u) = \frac{1}{2} \operatorname{Var} \{Y(x) - Y(x')\}$  where u denotes the distance between x and x'. Non-parametric estimates of V(u) are widely used in geostatistical work, both for exploratory data analysis and for diagnostic checking.

Consider a set of data  $(x_i, y_i) : i = 1, ..., n$ , where  $x_i$  denotes a location and  $y_i$  a corresponding measured value. The *empirical variogram ordinates* are the quantities  $v_{ij} = (y_i - y_j)^2/2$ . Under non-preferential sampling, each  $v_{ij}$  is an unbiased estimator for  $V(u_{ij})$ , where  $u_{ij}$  is the distance between  $x_i$  and  $x_j$ . A scatterplot of  $v_{ij}$  against  $u_{ij}$  is called the *variogram cloud*. A smoothed version of the variogram cloud can be used to suggest appropriate parametric models for the spatial covariance structure of the data; in what follows, we use simple binned estimators. For more information on variogram estimation, see for example Cressie (1985; 1991, Section 2.4), Chilès and Delfiner (1999, Section 2.2) or Diggle and Ribeiro (2007, Chapter 5).

The two panels of Figure 2 show simulation-based estimates of the point-wise bias and standard deviation of smoothed empirical variograms, derived from 500 replicate simulations of each of our three sampling designs. With regard to bias, the results under both uniform and clustered non-preferential sampling designs are consistent with the unbiasedness of the empirical variogram ordinates; although smoothing the empirical variogram ordinates does induce some bias, this effect is small in the current setting. In contrast, under preferential sampling the results show severe bias. With regard to efficiency, the right-hand panel of Figure 2 illustrates that clustered sampling designs, whether preferential or not, are also less efficient than uniform sampling. The bias induced by preferential sampling is qualitatively unsurprising; the effect of the preferential sampling is that sample locations predominantly span a reduced range of values of S(x), which in turn reduces the expectation of pair-wise squared differences at any given spatial separation. Note, incidentally, that the sample variogram has substantially smaller variance under preferential than under non-preferential clustered sampling. However, this is of little practical interest in view of its severe bias under preferential sampling. In general, the implicit estimated of the empirical variogram is the variance of Y(x) - Y(x') conditional on both x and x' belonging to X, which under preferential sampling differs from the unconditional variance; see, for example, Wälder and Stoyan (1996) or Schlather (2001).

### 3.2 Spatial prediction

Suppose that our target for prediction is  $S(x_0)$ , the value of the process S at a generic location  $x_0$ , given sample data  $(x_i, y_i)$ : i = 1, 2, ..., n. The widely used ordinary kriging predictor estimates the unconditional expectation of  $S(x_0)$  by generalised least squares, but using plug-in estimates of the parameters that define the covariance structure of Y. In classical geostatistics, these plug-in estimates are obtained either subjectively (Chilès and Delfiner, 1999, Section 2.6) or by non-linear least squares (Cressie, 1991, Section 2.6.2). We used maximum likelihood estimation under the assumed Gaussian model for Y.

Each simulation yields an estimate of the bias,  $\hat{S}(x_0) - S(x_0)$ , and the mean square error,



Figure 2: Estimated bias and standard deviation of the sample variogram under random (thick solid line), preferential (dotted line) and clustered (dashed line) sampling. The left-hand panel shows pointwise means plus and minus two pointwise standard errors, the right-hand panel pointwise standard deviations. See text for detailed description of the simulation model.

 $\{\hat{S}(x_0) - S(x_0)\}^2$ , for the ordinary kriging predictor  $\hat{S}(x_0)$ . The first two lines of Table 1 show approximate 95% confidence intervals, calculated as means plus and minus two standard errors over 500 replicate simulations, for the bias and root mean square error at the prediction location  $x_0 = (0.49, 0.49)$ .

The bias is large and positive under preferential sampling with  $\beta > 0$ . This prediction bias is a direct consequence of the bias in the estimation of the model parameters, which in turn arises because the preferential sampling model leads to the over-sampling of locations corresponding to high values of the underlying process S. The correct predictive distribution for Sis [S|Y, X] which, with known parameter values, takes a standard multivariate Gaussian form whether or not sampling is preferential. The two non-preferential sampling designs both lead to approximately unbiased prediction, as predicted by theory. The substantially larger mean square error for clustered sampling by comparison with completely random sampling reflects the inefficiency of the latter, as already illustrated in the context of variogram estimation.

In a second set of simulations, we set the values of the model parameters to correspond to the maximum likelihood estimates obtained in the analysis of the 1997 Galicia bio-monitoring data reported in Section 5 below, hence  $\mu = 1.515$ ,  $\sigma^2 = 0.138$ ,  $\phi = 0.313$ ,  $\kappa = 0.5$ ,  $\beta = -2.198$ ,  $\tau^2 = 0.059$ . The results are qualitatively as expected, but the differences amongst the three sampling designs are much smaller for two reasons. Firstly, the degree of preferentiality is much weaker; a measure of this is the product  $\beta\sigma$ , which takes the values 3 and 0.815 for the first and second simulation models, respectively. Secondly, the effect is further diluted by the inclusion of a non-zero nugget variance.

Table 1: Impact of sampling design on the bias and mean square error of the ordinary kriging predictor  $\hat{S}(x_0)$ , when  $x_0 = (0.49, 0.49)$  and each sample consists of 100 locations on the unit square. Each entry in the table is an approximate 95% confidence interval calculated from 500 independent simulations. See text for detailed description of the simulation models 1 and 2.

Model	Sampling design					
	Completely random	Preferential $(\beta = 2)$	Clustered			
1	bias	(-0.014, 0.055)	(0.951, 1.145)	(-0.048, 0.102)		
1	root mean square error	(0.345, 0.422)	(1.387, 1.618)	(0.758, 0.915)		
2	bias	(0.003, 0.042)	(-0.134, -0.090)	(-0.018, 0.023)		
2	root mean square error	(0.202, 0.228)	(0.247, 0.292)	(0.214, 0.247)		

## 4 Fitting the shared latent process model

#### 4.1 Monte Carlo maximum likelihood estimation

For the shared latent process model (3), the likelihood function for data X and Y can be expressed as

$$L(\theta) = [X, Y] = E_S [[X|S][Y|X, S]],$$
(6)

where  $\theta$  represents all of the model parameters and the expectation is with respect to the unconditional distribution of S. Evaluation of the conditional distribution [X|S] strictly requires the realisation of S to be available at all  $x \in A$ . In practice, we approximate the spatially continuous realisation of S by the set of values of S on a finely spaced lattice to cover A, and replace the exact locations X by their closest lattice points. We then partition S into  $S = \{S_0, S_1\}$ , where  $S_0$  denotes the values of S at each of n data-locations  $x_i \in X$ , and  $S_1$  denotes the values of S at the remaining N - n lattice-points.

To evaluate  $L(\theta)$  approximately, a naive strategy would be to replace the intractable expectation on the right hand side of (5) by a sample average over simulations  $S_j$ . This strategy fails when the measurement error variance  $\tau^2$  is zero, because unconditional simulations of S will then be incompatible with the observed Y. It also fails in practice when the measurement error is small relative to the variance of S, which is the case of most practical interest.

We therefore re-write the exact likelihood (5) as the integral

$$L(\theta) = \int [X|S][Y|X,S] \frac{[S|Y]}{[S|Y]} [S] dS.$$
(7)

Now, write  $[S] = [S_0][S_1|S_0]$  and replace the term [S|Y] in the denominator of (6) by  $[S_0|Y][S_1|S_0, Y] = [S_0|Y][S_1|S_0]$ . Note also that  $[Y|X, S] = [Y|S_0]$ . Then, (6) becomes

$$L(\theta) = \int [X|S] \frac{[Y|S_0]}{[S_0|Y]} [S_0] [S|Y] dS$$

$$= E_{S|Y} \left[ [X|S] \frac{[Y|S_0]}{[S_0|Y]} [S_0] \right]$$
(8)

and a Monte Carlo approximation is

$$L_{MC}(\theta) = m^{-1} \sum_{j=1}^{m} \left[ [X|S_j] \frac{[Y|S_{0j}]}{[S_{0j}|Y]} [S_{0j}] \right],$$
(9)

where now the  $S_j$  are simulations of S conditional on Y. Note that when Y is measured without error,  $[Y|S_{0j}]/[S_{0j}|Y] = 1$ . To reduce the Monte Carlo variance, we also use antithetic pairs of realisations, i.e. for each j = 1, ..., m/2 set  $S_{2j} = 2\mu_c - S_{2j-1}$ , where  $\mu_c$  denotes the conditional mean of S given Y. The use of conditional simulation in (8) by-passes the difficulty with the naive strategy by guaranteeing that the simulated realisations of S are compatible with the data Y.

To simulate a realisation from [S|Y], we use the following construction. Recall that the datalocations  $X = \{x_1, ..., x_n\}$  constitute a sub-set of the  $N \ge n$  prediction locations,  $X^* = \{x_1^*, ..., x_N^*\}$  say. Define C to be the n by N matrix whose *i*th row consists of N - 1 zeros and a single 1 to identify the position of  $x_i$  within  $X^*$ . Note that, unconditionally,  $S \sim \text{MVN}(0, \Sigma)$ and  $Y \sim \text{MVN}(\mu, \Sigma_0)$  with  $\Sigma_0 = C\Sigma C' + \tau^2 I$ . Then, if Z denotes an independent random sample of size n from  $N(0, \tau^2)$  and y denotes the observed value of Y, it follows that

$$S_c = S + \Sigma C' \Sigma_0^{-1} (y - \mu + Z - CS)$$
(10)

has the required multivariate Gaussian distribution of S given Y = y (Rue and Held, 2005, Chapter 2; Eidsvik, Martino and Rue, 2006). Hence, for conditional simulation when N is large, we need a fast algorithm for unconditional simulation of S. We have used the circulant embedding algorithm of Wood and Chan (1994) applied to a rectangular region containing the region of interest, A. The subsequent calculations for  $S_c$  then involve only the relatively straightforward inversion of the  $n \times n$  matrix  $\Sigma_0$  and simulation of the n independent Gaussian random variables that make up the vector Z in (9).

### 4.2 Goodness-of-fit

We have already noted in Section 1 the availability of tests for preferential sampling; see, for example, Schlather, Ribeiro and Diggle (2004) or Guan and Afsharatous (2007). Here, we suggest a way of assessing the goodness-of-fit of the preferential sampling model described in Section 2, by comparing the sample locations with realisations of the fitted Cox model for their unconditional distribution.

A standard diagnostic tool for stationary spatial point processes is the reduced second-moment measure, or K-function (Ripley, 1977), defined by  $\lambda K(s) = E[N_0(s)]$  where  $N_0(s)$  denotes the number of points of the process within distance s of an arbitrary origin of measurement, conditional on there being a point of the process at the origin, and  $\lambda$  is the expected number of points of the process per unit area. Under the preferential sampling model, the marginal model for the sample locations X is a log-Gaussian Cox process with stochastic intensity  $\Lambda(x) = \exp\{\alpha + \beta S(x)\}$ . For this process, the K-function is of the form

$$K(s) = \pi s^{2} + 2\pi \int_{0}^{s} \gamma(u) u du,$$
(11)

where  $\gamma(u) = \exp\{\beta^2 \sigma^2 \rho(u; \phi)\} - 1$  is the covariance function of  $\Lambda(x)$  (Diggle, 2003, Section 5.5) To assess the goodness-of-fit informally, we compare the estimated K-function of the data with the envelope of estimates obtained from sets of sample locations generated from simulated realisations of the fitted model. For a formal Monte Carlo test, we use a goodness-of-fit statistic that measures the discrepancy between estimated and theoretical K-functions, as described in Section 5.2.2. Note that this aspect of the model is not considered explicitly in the fitting process described in Section 4.1.

### 5 Heavy-metal bio-monitoring in Galicia

Our application concerns bio-monitoring of lead pollution in Galicia, northern Spain, using the concentrations in moss samples, in  $\mu g/g$  dry weight, as the measured variable. An initial survey was conducted in the spring of 1995 "to select the most suitable moss species and collection sites" (Fernández, Rey and Carballeira, 2000). Two further surveys of lead concentrations in samples of the moss species (*Scleropodium purum*) took place in October 1997 and July 2000. Figure 3 shows the sampling locations used in these two surveys. Note that some locations appear to lie outside Galicia, and others in the sea to the north. The boundary shown in Figure 3 is both approximate and imperfectly registered; however, it plays no role in the analysis and is included only to add context to the map.

In the 1997 survey, sampling was conducted more intensively in sub-regions where large gradients in lead concentrations were anticipated, in line with suggestions in Ruhling (1994). The resulting design was highly non-uniform and potentially preferential. The second survey used an approximately regular lattice design, which is therefore non-preferential; gaps in the lattice arose only where a different species of moss was collected. For further details, see Fernández, Rey and Carballeira (2000) and Aboal, Real, Fernández and Carballeira (2006). In particular, Fernández, Rey and Carballeira (2000) studied the changes in heavy metal concentrations between the two years ignoring the corresponding spatial distributions. Our objective in analysing these data is to estimate, and compare, maps of lead concentrations in 1997 and 2000.

The measured lead concentrations included two gross outliers in 2000, each of which we replaced by the average of the remaining values from that year's survey. Table 2 gives summary statistics for the resulting 1997 and 2000 data. Note that the mean response is higher for the 1997 data than for the 2000 data, which would be consistent either with the former being preferentially sampled near potential pollutant sources, or with an overall reduction in pollution levels over the three years between the two surveys. Also, the log-transformation eliminates an apparent variance-mean relationship in the data and leads to more symmetric distributions of measured values (Figure 4).



Figure 3: Sampling locations for 1997 (solid dots) and 2000 (open circles). The unit of distance is 100km. Two outliers in the 1997 data were at locations (6.50, 46.90) and (6.65, 46.75).

Table 2: Summary statistics for lead pollution levels ( $\mu g/g$  dry weight) measured in 1997 and 2000.

	untransformed		log-transformed		
	1997	2000	1997	2000	
Number of locations	63	132	63	132	
Mean	4.72	2.15	1.44	0.66	
Standard deviation	2.21	1.18	0.48	0.43	
Minimum	1.67	0.80	0.52	-0.22	
Maximum	9.51	8.70	2.25	2.16	

### 5.1 Standard geostatistical analysis

For an initial analysis, we assumed the standard Gaussian model (1) with the underlying signal S(x) specified as a zero-mean stationary Gaussian process with variance  $\sigma^2$  and Matérn correlation function  $\rho(u; \phi, \kappa)$ , and Gaussian measurement errors,  $Z_i \sim N(0, \tau^2)$ , and fitted



Figure 4: Empirical distributions of log-transformed lead concentrations in the 1997 (solid line) and 2000 (dashed line) samples.

this model separately to the 1997 and 2000 data.

Figure 5 shows, for each of 1997 and 2000, smoothed empirical variograms and theoretical variograms with parameters fitted by maximum likelihood. Based on the general shape of the two empirical variograms, we used a fixed value  $\kappa = 0.5$  for the shape parameter of the Matérn correlation function. The estimated variograms differ in some respects, notably the absence of a nugget component (i.e.  $\hat{\tau}^2 \approx 0$ ) in the variogram estimated from the 2000 data; however, this parameter is poorly identified because of the lattice-like arrangement of the 2000 sampling design, whereas the inclusion of close pairs of locations in the 1997 sampling design enables better estimation of  $\tau^2$ . Other features of the two fitted variograms are similar; for example, the height of the asymptote (i.e.  $\hat{\tau}^2 + \hat{\sigma}^2$ ) and the approximate range (i.e.  $\hat{\phi}$ ). These observations support the idea that a joint model for the two data-sets might allow at least some parameters in common between the two years. The generalised likelihood ratio test statistic (GLRTS, Cox and Hinkley, 1974, Section 9.3) to test the hypothesis of common  $\sigma$ ,  $\phi$  and  $\tau$ , under the admittedly dubious assumption that neither sample is preferential, was 7.66 on 3 degrees of freedom (p = 0.054). We re-visit this question in the next sub-section.

### 5.2 Analysis under preferential sampling

#### 5.2.1 Parameter estimation

We now investigate whether the 1997 sampling is indeed preferential. We used the Nelder-Mead simplex algorithm (Nelder and Mead, 1965) to estimate the model parameters, increas-



Figure 5: Smoothed empirical (open circles) and fitted theoretical (lines) variograms for 1997 (left-hand panel) and 2000 (right-hand panel) log-transformed lead concentration data.

ing the number of Monte Carlo samples, m, progressively to avoid finding a false maximum. With m = 100,000, the Monte Carlo standard error in the evaluation of the log-likelihood ratio was reduced to approximately 0.3 (the actual value varies over the parameter space) and the approximate GLRTS to test  $\beta = 0$  was 27.7 on 1 degree of freedom (p < 0.001).

We then fitted a joint model to the two data-sets, treating the 1997 and 2000 data as preferentially and non-preferentially sampled, respectively. To test the hypothesis of shared values for  $\sigma$ ,  $\phi$  and  $\tau$ , we fitted the model with and without these constraints, obtaining a GLRTS of 6.2 on 3 degrees of freedom (p = 0.102). The advantage of using shared parameter values when justified is that the parameters in the joint model are then estimated more efficiently and the model is consequently better identified (Altham, 1984). This is particularly important in the geostatistical setting, where the inherent correlation structure of the data reduces their information content by comparison with independent data having the same sample size.

Table 3 shows the Monte Carlo maximum likelihood estimates together with estimated standard errors and correlations for the model with shared  $\sigma$ ,  $\phi$  and  $\tau$ . Standard errors and correlations were evaluated by fitting a quadratic surface to Monte Carlo log-likelihoods by ordinary least squares. Parameter combinations were initially set as a 3<sup>6</sup> factorial design centred on the Monte Carlo maximum likelihood estimates, with parameter values chosen subjectively after examining the trajectories through the parameter space taken by the various runs of the Nelder-Mead optimisation algorithm. The quadratic surface was then re-fitted after augmenting this design with a 2<sup>6</sup> factorial on a more closely spaced set of parameter values, to check

Table 3: Monte Carlo maximum likelihood estimates of parameters in the joint model for the 1997 and 2000 Galicia biomonitoring data. Approximate standard errors and correlations are computed from a quadratic fit to the Monte Carlo log-likelihood surface (see text for details)

Parameter	Estimate	Standard error	Correlation matrix					
$\mu_{97}$	1.515	0.136	1.000	0.023	0.095	-0.243	-0.222	0.167
$\mu_{00}$	0.762	0.110		1.000	0.230	-0.229	-0.281	0.342
$\log(\sigma)$	-0.992	0.049			1.000	-0.217	-0.744	0.469
$\log(\phi)$	-1.163	0.075				1.000	0.604	-0.675
$\log(\tau)$	-1.419	0.042					1.000	-0.652
$\beta$	-2.198	0.336						1.000

the stability of the results. Each evaluation of the log-likelihood used m = 10,000 conditional simulations. The non-negative parameters  $\sigma$ ,  $\phi$  and  $\tau$  are estimated on a log-transformed scale, to improve the quadratic approximation to the log-likelihood surface.

Note that the expectation of  $S(\cdot)$  shows a substantial fall between 1997 and 2000, and that the preferential sampling parameter estimate is negative,  $\hat{\beta} = -2.198$ . The latter finding is both counter-intuitive, because the over-sampled northern half of the region is more industrialised than the under-sampled southern half, and critically dependent on our allowing the two mean parameters to differ. Otherwise, because the observed average pollution level is substantially higher in 1997 than in 2000, we would have been forced to conclude that the 1997 sampling was preferential with a positive value of  $\beta$ . One piece of evidence against this alternative interpretation is that, within the 1997 data, the observed pollution levels are lower in the over-sampled northern half of the region (n = 47, mean log-concentration= 1.38, SD = 0.49) than in the under-sampled southern half (n = 16, mean= 1.62, SD = 0.40), consistent with a negative value of  $\beta$ .

#### 5.2.2 Goodness-of-fit

Figure 6 shows the estimated K-function for the 1997 sampling locations together with the envelope of 99 simulations of the fitted Cox process, and the theoretical K-function. The estimate lies within the simulation envelope for distances up to 0.22 (22km). For a formal Monte Carlo goodness-of-fit test, we define the test statistic

$$T = \int_0^{0.25} [\{\hat{K}(s) - K(s)\}^2 / v(s)] ds$$

where K(s) is given by (10) and v(s) is the variance of K(s), estimated from the simulations of the fitted Cox process. This gives p = 0.03. The Cox model slightly under-estimates the extent of spatial aggregation in the data-locations.



Figure 6: Estimated K-function of the 1997 sample locations (solid line) and envelope from 99 simulations of the fitted log-Gaussian Cox process (dashed lines).

#### 5.2.3 Prediction

What impact does the acknowledgement of preferential sampling make on the predicted 1997 pollution surface? Figure 7 shows the predicted surfaces  $\hat{T}(x) = E[T(x)|X, Y]$ , where  $T(x) = \exp\{S(x)\}$  denotes lead concentration on the untransformed scale, together with the pointwise differences between the two. Each surface is a Monte Carlo estimate based on m = 10,000 simulations, resulting in Monte Carlo standard errors of 0.026 or less. The predictions based on the preferential sampling model have substantially wider range, over the lattice of prediction locations, than those that assume non-preferential sampling (1.310 to 7.654 and 1.286 to 5.976, respectively). The difference surface also covers a relatively large range (-0.715 to 3.693) and shows strong spatial structure. The size of the difference between the two predicted surfaces is at first sight surprising, as both are partially constrained by the observed concentrations. However, in the presence of a nugget effect the predictions are not constrained to interpolate the data. Also, ignoring the preferential nature of the sampling leads to biased parameter estimates. Finally, the effect of the back-transformation from log(Pb) to Pb, this being the scale on which predictions are required, is to amplify the differences.

Using the conditional expectation as a point predictor is conventional, but questionable when, as here, the measurement process has a highly skewed distribution. As an alternative summary, Figure 8 compares pointwise 5%, 50% and 95% limits of the plug-in predictive distribution of lead concentrations under preferential and non-preferential modelling assumptions, holding the model parameters fixed at their estimated values. The differences between the two are smaller than in Figure 7, but still non-negligible.

Finally, in Figure 9, we show kernel density estimates of the plug-in predictive distributions for the areal proportion of Galicia in which 1997 lead concentrations exceed 3, 5 or  $7\mu g/g$  dry weight. In all three cases, recognition of the preferential sampling results in a pronounced shift in the predictive distribution. Note, however, that these plug-in predictive distributions



Figure 7: Predicted surface of lead concentrations in 1997 under preferential (left-hand panel) and non-preferential (centre panel) assumptions, together with the pointwise difference between the two (right-hand panel). All three surfaces are plotted on a common scale, as shown.

do not account for parameter uncertainty

Our overall conclusion is that the preferential sampling has made a material difference to our predictive inferences for the 1997 pollution surface.

### 6 Discussion

We have shown that conventional geostatistical models and associated statistical methods can lead to misleading inferences if the underlying data have been preferentially sampled. We have proposed a simple model to take account of preferential sampling and developed associated Monte Carlo methods to enable maximum likelihood estimation and likelihood ratio testing within the proposed class of models. The resulting methods are computationally intensive, each run taking several hours of CPU time. The computations reported in the paper were run on a DELL Latitude D620 lap-top PC, using the R software environment (R Development Core Team, 2008, see also www.r-project.org) and associated CRAN packages fields, geoR and splancs. The data and R code are available from www.lancs.ac.uk/staff/diggle/. There is undoubtedly very considerable scope to improve the efficiency of the authors' code. In particular, we did not seek to automate the progressive increase in the number of Monte Carlo samples as we explored the log-likelhood surface.

The computation of the Monte Carlo likelihood uses direct simulation, as in Diggle and Gratton (1984), rather than Markov chain Monte Carlo. Hence, issues concerning convergence of the simulations do not arise, and the variability between replicate simulations gives a direct estimate of the size of the Monte Carlo error.

We have described an application to a set of environmental bio-monitoring data from Galicia,



Figure 8: Pointwise 5%, 50% and 95% limits of the predictive distribution of lead concentrations in 1997 under preferential (upper row) and non-preferential (lower row) assumptions. All six surfaces are plotted on a common scale, as shown.

northern Spain. An important feature of these data is that they are derived from two surveys of the region of interest, the first of which used a spatially irregular set of sampling locations and has been shown to be preferentially sampled. In the second survey the sampling locations formed a near-regular grid over the study-region and we have therefore taken it to be nonpreferentially sampled. This, coupled with our finding that several of the model parameters can be assumed to take a common value for the two samples, led to a better identified joint model for the two surveys. To illustrate this point, we also fitted the preferential sampling model to the 1997 data alone. Although, as reported earlier, the value of the maximised loglikelihood was obtained relatively easily, the subsequent quadratic fitting method to estimate the standard errors of the maximum likelihood estimates proved problematic. Using a  $3^5 + 2^5$ factorial design analogous to the earlier  $3^6 + 2^6$  design for the model fitted to the 1997 and 2000 data jointly, and with 10,000 simulations for each log-likelihood evaluation as before, the quadratic fit explained only 72% of the variation in the Monte Carlo log-likelihoods, compared with 93% for the joint model, the implied estimate of  $\partial^2 L/\partial\beta^2$  was not significantly different from zero, and the ratio of largest to smallest eigenvalues of the Hessian matrix was 34.5, compared with 22.3 for the joint model.

Alternative strategies for dealing with poorly identified model parameters could include treat-



Figure 9: Predictive distributions for the areal proportion of Galicia in which 1997 lead concentrations exceed 3, 5 and  $7\mu g/g$  dry weight (left-hand, centre and right-hand panels, respectively) under preferential (solid line) and non-preferential (dashed line) assumptions.

ing the preferential sampling parameter  $\beta$  as a sensitivity parameter, since its value is typically not of direct scientific interest, or using Bayesian methods with informative priors.

A natural response to a strongly non-uniform sampling design is to ask whether its spatial pattern could be explained by the pattern of spatial variation in a relevant covariate. Suppose, for the sake of illustration, that S is observed without error, that dependence between X and S arises through their shared dependence on a latent variable, U, and that the joint distribution of X and S is of the form

$$[X,S] = \int [X|U][S|U][U]dU, \qquad (12)$$

so that X and S are conditionally independent given U. If the values of U were to be observed, we could then legitimately work with the conditional likelihood, [X, S|U] = [X|U][S|U] and eliminate X by marginalisation, exactly as is done implicitly when conventional geostatistical methods are used. In practice, "observing" U means finding explanatory variables which are associated both with X and with S, adjusting for their effects and checking that after this adjustment there is little or no residual dependence between X and S. If so, the analysis could then proceed on the assumption that sampling is no longer preferential. Note, in this context, that any of the existing tests for preferential sampling can be applied, albeit approximately, to residuals after fitting a regression model for the mean response.

The value of seeking relevant explanatory variables to contribute to a spatial statistical model cannot be over-stated. We hold the view that in most geostatistical applications, spatial correlation reflects, at least in part, smooth spatial variation in relevant, but unobserved, explanatory variables rather than being an inherent property of the phenomenon being studied; an example to the contrary would be the spatial distribution of the prevalence of an infectious disease during an epidemic where, even for a uniformly distributed population in a completely homogeneous environment, the process of transmission from infectious to susceptible individuals would induce spatial correlation in the prevalence surface. This in turn leads us to emphasise that our paper is not a plea for uniform sampling, but rather for ensuring that any model for a set of data should respect whatever sampling design has been used to generate the data; for a thorough discussion that also uses point process models of sampling designs, albeit in a very different setting, see McCullagh (2008).

Returning to the geostatistical setting, and specifically to the application described in Section 5, Fernández et al (2005) give a practitioner's perspective on the ways in which different sampling designs can materially affect any analysis of spatial variation. To meet their primary objective of mapping concentration surfaces, they favour regular lattice designs but note that for other purposes, "additional sampling in areas of anomalously high concentrations of contaminants makes good sense." We agree. Our paper formalises this idea, whilst acknowledging that it necessarily complicates the subsequent modelling and inference.

## Acknowledgements

This work was supported by the UK Engineering and Physical Sciences Research Council through the award of a Senior Fellowship to Peter Diggle.

We thank the Ecotoxicology Group, University of Santiago de Compostela, for permission to use the Galicia data and, in particular, José Angel Fernández, for helpful discussions concerning the data.

We also thank Havard Rue for advice on efficient conditional simulation of spatially continuous Gaussian processes.

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