Model-Based Geostatistics the Easy Way

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Abstract

This paper briefly describes geostatistical models for Gaussian and non-Gaussian data and demonstrates the geostatsp and diseasemapping packages for performing inference using these models. Making use of R’s spatial data types, and raster objects in particular, makes spatial analyses using geostatistical models simple and convenient. Examples using real data are shown for Gaussian spatial data, binomially distributed spatial data, a log-Gaussian Cox process, and an area-level model for case counts.

Keywords: spatial statistics, geostatistics, R, INLA, Bayesian inference, kriging.

1. Introduction

In the past two decades spatial statistics has gradually become a mature and established branch of statistics with a suite of well defined models and proven inference methodologies capable of addressing a wide range of practical problems. The capability of R (R Core Team 2014) to store, manipulate, and display spatial data has similarly improved, and as a result spatial methodologies which were formerly only accessible to the specialist are available to the wider statistical community. This paper demonstrates model fitting for Gaussian, non-Gaussian, and point process data using the geostatsp and diseasemapping packages, with R’s spatial data classes being used to make spatial data analysis simple and the software intuitive.

1.1. Models and methods

Models and theory for Gaussian spatial data were first espoused by Matheron (1962) and popularized by Cressie (1993). Writing $U(s)$ as the value of a Gaussian random field $U$ at location $s$, the basic (stationary) geostatistical model is characterized by the joint multivariate normal distribution

$$[U(s_1) \ldots U(s_N)]^\top \sim \text{MVN}(0, \Sigma).$$
The entries of the covariance matrix $\Sigma$ are determined by a spatial correlation function $\rho$ with

$$\Sigma_{ij} = \text{cov}[U(s_i), U(s_j)] = \sigma^2 \rho[(s_i - s_j)/\phi, \theta].$$

Here $\phi$ is a scale parameter controlling the rate at which correlation decays with distance, and $\theta$ is a vector of possible additional parameters (controlling directional effects, for example). An isotropic process has correlation being a function of distance with $\rho[(s_i - s_j)/\phi] = \rho_0(||s_i - s_j||/\phi)$.

Various parametric functions have been used for $\rho$, and Stein (1999) makes a compelling case for the Matérn correlation function described in Appendix A. An isotropic process with a Matérn correlation has a single additional parameter $\kappa$ controlling the differentiability of the process. Two additional covariance parameters commonly used refer to geometric anisotropy, and comprise an angle of rotation indicating a preferred direction and a ratio parameter giving the ratio of the ranges on the two axes.

The parametrisation of the Matérn is different in each of the geoR (Ribeiro and Diggle 2001), RandomFields (Schlather, Malinowski, Menek, Oesting, and Strokorb 2015) and geostatsp packages. The specification of the Matérn in Appendix A, and in use in the geostatsp package, has the property that when varying $\kappa$ the correlation at a distance $\phi$ stays fairly close to 0.14, or $\rho([0, \phi]/\phi, \kappa) \approx 0.14$. A Matérn with $\kappa = \infty$ is a Gaussian density with $\phi$ being two standard deviations. The term ‘practical range’ is used at times to describe $\phi$ as defined here, interpreting $\phi$ as a distance beyond which correlation is ‘small’ is a manner analogous to interpreting the Gaussian density as being ‘small’ beyond two standard deviations.

The anisotropy angle refers to rotation of the coordinates anti-clockwise by the specified amount prior to calculating distances, which has the effect that the contours of the correlation function appear rotated clockwise by this amount. The anisotropy ratio is the amount the Y coordinates are divided by following rotation, with large values making the Y coordinates smaller and increasing the correlation in the Y direction (of the rotated coordinates).

**Gaussian data**

Data $Y_i$ observed at location $s_i$ with covariates $X(s_i)$ is often modelled with the linear geostatistical model (LGM):

$$Y_i|U(s_i) \sim N(\lambda(s_i), \tau^2)$$

$$\lambda(s_i) = \mu + \beta X(s_i) + U(s_i).$$

(1)

Although method-of-moments estimation of the covariance parameters $\phi$, $\sigma$ and $\tau$ is still common, Stein (1999) makes a thorough argument for using maximum likelihood estimates (MLEs). Writing $\psi = (\mu, \beta, \sigma, \tau, \phi)$, the MLEs $\hat{\psi}$ are the quantities which maximize the likelihood $pr(Y_1 \ldots Y_N; \psi)$. The $Y_i$ are jointly multivariate normal and the likelihood is tractable, albeit requiring the inversion of an $N$ by $N$ matrix, and numerical optimizers such as the optim function can be used to find $\hat{\psi}$.

Spatial prediction usually involves covering the study region with a large number of regularly spaced points $g_\ell; \ell = 1 \ldots L$ and mapping estimates of $\bar{U} = [U(g_1) \ldots U(g_L)]$ or $\bar{\lambda} = [\lambda(g_1) \ldots \lambda(g_L)]$. As the model is linear and Gaussian, the conditional distribution $[\bar{U}|Y]$ is multivariate normal with closed form expressions for the conditional mean and variance. The MLEs $\hat{\psi}$ are used to calculate these expressions, hence the uncertainty in these parameter estimates is ignored (see Diggle and Ribeiro 2006).
Non-Gaussian data

When the observed data \( Y_i \) are non-Gaussian, the model above is extended to the generalized linear geostatistical model (GLGM) used by Diggle, Moyeed, and Tawn (1998) and further described in Diggle and Ribeiro (2006). Consider a distribution \( f \) (i.e., Binomial or Weibull) with a mean parameter \( \lambda \) and possibly additional parameters \( \nu \). Writing \( g(\cdot) \) as a link function (i.e., log or logit), the GLGM takes the form

\[
Y_i | U(s_i) \sim f[\lambda(s_i), \nu] \\
g[\lambda(s_i)] = \mu + \beta X(s_i) + U(s_i) \\
\text{cov}[U(s_i), U(s_j)] = \sigma^2 \rho[(s_i - s_j)/\phi, \theta].
\]

The combination of non-Gaussian data and an unobserved latent variable make the likelihood function intractable and computing the MLEs difficult. Bayesian inference using Markov chain Monte Carlo (MCMC) algorithms has become the most common method for making statistical inference with GLGMs, as was done in Diggle et al. (1998). Bayesian inference requires specifying prior distributions for the model parameters \( \mu, \beta, \sigma \) and \( \phi \), with the posterior distributions \( \pi(\phi|Y) \) and \( \pi[U(s)|Y] \) forming the basis of inference.

The integrated nested Laplace approximation (INLA) algorithm of Rue, Martino, and Chopin (2009) is an alternative to MCMC for performing Bayesian with latent Gaussian models. MCMC’s principal drawback is the requirement that chains of posterior samples must be monitored and assessed for convergence and mixing, and obtaining a set of reliable posterior samples from a MCMC algorithm can be difficult and require a specialized skill set to accomplish. INLA is much easier to use in this regard, and although it’s maximisation step and numerical integration can sometimes require judicious choices of starting values and tuning parameters it is in general less labor-intensive to use than MCMC.

An additional recent development which has facilitated the implementation of the GLGM is the Markov random field approximation to the Matérn correlation function developed by Lindgren, Rue, and Lindström (2011). When the number of spatial locations \( N \) is large, inverting the variance matrix \( \Sigma \) can be time consuming or numerically unstable. Lindgren et al. (2011) use Gaussian Markov random fields (GMRF’s) to derive a simple expression for \( \Sigma^{-1} \) for Matérn correlations using various forms of stochastic partial differential equations. The \texttt{geostatsp} package makes use of the Matérn approximation of GMRF’s on grids of square cells with \( \kappa = 1 \) or 2. Although real datasets will rarely be sampled on a square lattice, the continuous surface \( U(s) \) can be well approximated by superimposing a fine lattice over the study region and assigning each data point to a cell. The fact that many (or most) of the cells will not have data observed in them is not problematic for INLA. This combination of INLA with the Markov random field approximation has been to estimate spatial variation in risk for Lupus in the city of Toronto, Canada from case incident locations by Li, Brown, Rue, al Maini, and Fortin (2012), and for assessing the effect of cancer risk of ambient radiation near a nuclear power facility using time-to-event data from a retrospective cohort in Jiang, Brown, Rue, and Shimakura (2014).

Lindgren et al. (2011) derive a GMRF approximation for the Matérn using an irregular lattice with triangular basis functions, which has a number of advantages over the grid cell approach. This approximation is implemented in the \texttt{INLA} software, and incorporation of this feature into \texttt{geostatsp} is work in progress.
1.2. Spatial statistics and R

The \texttt{sp} package (see Bivand, Pebesma, and Gómez-Rubio 2013) and \texttt{raster} package (Hijmans 2014) provide an excellent set of facilities for storing, manipulating, and visualising spatial data. The \texttt{sp} package provides \texttt{SpatialPointsDataFrame} and \texttt{SpatialPolygonsDataFrame} objects for storing point and polygon data respectively, and are compatible with many of the standard data formats most geographical information systems (GIS) uses. The \texttt{raster} package provides similar tools for raster data, which are pixelated images or rectangular lattices. The \texttt{rgdal} (Bivand, Keitt, and Rowlingson 2014) package provides a set of tools for reading spatial data from various formats into R, such as ESRI shapefiles for point and polygon data, and GeoTIFF files for raster data. These three packages (along with \texttt{spdep}, Bivand 2014, and others) have made R fully compatible with GIS software and R fulfils many of the criteria for it to be called a GIS in its own right.

The venerable \texttt{geoR} package (see Diggle and Ribeiro 2006) has provided tools for likelihood-based inference since 2000, and is one of the very few software packages for spatial analysis which accommodates all of: the Matérn correlation function; covariates; Maximum Likelihood Estimation; geometric anisotropy; and the Box-Cox transform. Since \texttt{geoR} predates the \texttt{sp} and \texttt{raster} packages, it has its own spatial data types.

For Bayesian inference, the excellent \texttt{INLA} (Rue, Martino, Lindgren, Simpson, and Riebler 2013) package developed by the authors of Rue et al. (2009) and Lindgren et al. (2011) implements INLA for a wide variety of models, including spatial Gaussian Markov random field models. \texttt{INLA} has been designed with flexibility of model specification being a priority, a job \texttt{INLA} accomplishes to an astonishing degree albeit at the cost rendering some tasks relatively complex in comparison to other packages. One such example is specifying a Matérn correlation function, with spatial locations being specified as grid cell indexes rather than coordinates. A considerable amount of code can sometimes be necessary for converting \texttt{INLA} results from a spatial model into a format which can be mapped.

The \texttt{geostatsp} package provides a set of user-friendly functions for Gaussian spatial models and an easy interface to \texttt{INLA} for fitting non-Gaussian models, resulting in a powerful set of tools for model-based geostatistical analyses in R. Response variables and covariates are specified with formulas, with data provided as \texttt{Raster} or \texttt{SpatialPointsDataFrame} objects. The interface to \texttt{INLA} has more complex set of routines underlying it, with observations being allocated to cells in a Markov random field and linear combinations of parameters and latent variables for predicted spatial surfaces being defined. The spatial predictions obtained from these packages are raster objects, making them easy to display and overlay on background maps.

2. Model-based geostatistics through examples

The \texttt{geostatsp} and \texttt{diseasemapping} packages described in this paper are available from the Comprehensive R Archive Network at \url{http://CRAN.R-project.org/} and R-Forge at \url{http://R-Forge.R-project.org/R/?group_id=312}. They both depend on the \texttt{INLA} package obtainable from \url{http://R-INLA.org/}. 
2.1. Maximum likelihood estimation and kriging

The Swiss rainfall dataset (see Diggle and Ribeiro 2006, 5.4.7) is a classic case study in Gaussian geostatistics. Loading of the \texttt{geostatsp} package and executing \texttt{data("swissRain")}

makes available the following objects: a \texttt{SpatialPointsDataFrame} named \texttt{swissRain} of rain values at a number of points, a \texttt{SpatialPolygonsDataFrame} named \texttt{swissBorder} of the border of Switzerland; and a \texttt{Raster} object \texttt{swissAltitude} containing elevation values for Switzerland. These three objects are plotted in Figure 1.

Using the linear geostatistical model in (1) with these data would have the rainfall measurements being the $Y_i$, elevation values being $X(s)$, and $\lambda(s)$ as the unknown true rainfall surface. Either Bayesian or Frequentist inference could be used to fit the model, with the former possible in a manner similar to the example in the subsequent section. Frequentist inference is accomplished with the \texttt{lgm} function in the \texttt{geostatsp} package, which in turn calls \texttt{likfitLgm} for estimating the model parameters and \texttt{krige} for computing conditional means and variances of $U(s)$ and $\lambda(s)$. The Swiss rainfall data is fit with the code below.

```r
R> names(swissRain)
[1] "ID" "rain"
R> names(swissAltitude)
[1] "CHE_alt"
R> swissFit <- lgm(rain ~ CHE_alt, swissRain, grid = 120,
+ covariates = swissAltitude, shape = 1, fixShape = TRUE, boxcox = 0.5,
+ fixBoxcox = TRUE, aniso = TRUE)
R> names(swissFit)
```

Figure 1: Swiss rainfall data (colored blue points and top legend) with elevation (background colors and bottom legend).
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| Parameter                  | Estimate | Std. error | CI 0.025 | CI 0.975 | Estimated |
|----------------------------|----------|------------|----------|----------|-----------|
| (Intercept)                | 4.86     | 1.29       | 2.32     | 7.39     | true      |
| Elev’n per 1000m           | 0.28     | 0.37       | -0.45    | 1.01     | true      |
| range, km                  | 0.06     | 0.03       | 0.11     | true     |           |
| sdNugget                   | 0.95     | 0.73       | 1.24     | true     |           |
| anisoAngleDegrees          | 37.00    | 31.74      | 42.27    | true     |           |
| anisoRatio                 | 7.48     | 3.94       | 14.19    | true     |           |
| shape                      | 1.00     | false      |          |          |           |
| boxcox                     | 0.50     | false      |          |          |           |
| sdSpatial                  | 2.97     | 1.89       | 4.68     | true     |           |

Table 1: Swiss rainfall parameter estimates, standard errors and confidence intervals obtained from a linear geostatistical model and the \texttt{lgm} function.

[1] "predict" "param" "varParam" "optim"

[5] "data" "model" "summary"

The data and covariates arguments contain the data required for fitting the model, with the fixed effects $\beta X(s)$ specified by formula. The variables listed in formula refer to names in either the \texttt{swissRain} or \texttt{swissAltitude} objects, and are not the names of the objects themselves. Variables in the right hand side of formula can refer to either: the name of a vector of values contained in the data argument; the name of a layer in a Raster object (a single layer, brick or stack) passed as covariates; or the name of one of the elements if covariates is a list of Raster objects. The latter is useful when covariate rasters have different resolutions and projections. If a covariate is a column in data, it will not be included in the predicted values for $\lambda(s)$.

The argument grid = 120 specifies that spatial prediction should be done on a raster with 120 cells in the X dimension, with this raster having square cells covering the bounding box of \texttt{swissRain}. The grid argument can alternatively be supplied as a Raster object. A Matérn spatial correlation function with shape parameter fixed at 1 and a Box-Cox transform with parameter fixed at 0.5 (a square-root transform) are used. The aniso = TRUE argument allows for geometric anisotropy in the correlation function. Additional function arguments are param and parscale, starting values and parameter scaling values passed from \texttt{lgm} to likfitLgm and ultimately the numerical optimizer optim. The Swiss data has spatial locations expressed in a UTM projection, with coordinates in metres and consequently a spatial range parameter likely to be in the hundreds of thousands. The default scaling of 1 in optim would be ineffective and arguments on the order of param = c(range = 10^{-5}) and parscale = c(range = 10^{-4}) are in order. The default starting value and scale which likfitLgm sets for the range parameter are 1/20 and 1/200 of the diagonal distance of the bounding box of data.

The \texttt{swissFit} object produced by \texttt{lgm} is a list with elements including predict, a RasterStack of spatial predictions and standard errors, and summary, a table of parameter estimates and confidence intervals. Table 1 shows the summary component, with the range parameter converted to kilometres. The standard deviation parameters $\sigma$ and $\tau$ are displayed in the sdSpatial and sdNugget rows respectively. Confidence intervals for the covariance parameters are derived from the observed information matrix, and will be missing if any of the
estimated parameters are on a boundary. Notice the ‘Estimated’ column indicating that the Matérn shape parameter and Box-Cox transformation parameter were not estimated from the data.

Spatial predictions of the rainfall surface \( \lambda(s) \) and the spatial random effect \( U(s) \) are contained in the RasterStack element of `swissFit$predict`, which has the following layers:

```
R> names(swissFit$predict)
[1] "space"   "random"   "predict.boxcox"
[4] "krigeSd"  "predict"
```

Using the notation in (1), these layers are (in the order given above): the predicted fixed effects \( \hat{\mu} + \hat{\beta} X(s) \); the kriged random effects \( E[U(s)|Y] \); the predicted rainfall surface \( E[\lambda(s)|Y] \) on the Box-Cox transformed scale; the prediction standard deviation \( sd[U(s)|Y] \); and predicted rainfall on the natural scale \( E\{\alpha \lambda(s) + 1\}^{1/\alpha}|Y \} \) with \( \alpha \) being the Box-Cox transformation parameter. Figure 2a shows the predicted rainfall values (on the natural scale), and results from the command `plot(swissFit$predict["predict"]))`. Notice the strong directionality is consistent with an angle of rotation of 37° and a ratio of the major to minor axes of 7.5.

Figure 2b shows the conditional probabilities that rainfall exceeds 30mm, computed with

```
R> exc30 <- excProb(swissFit, 30, nuggetInPrediction = TRUE)
```

The `excProb` function uses `pnorm` with means from the `predict.boxcox` layer and standard deviations from `krigeSd`, calculating probabilities of exceeding the Box-Cox transform of 30. The `nuggetInPrediction` argument can be set to `TRUE` to compute probabilities of new observations \( Y_i \) exceeding a threshold, with `FALSE` specifying exceedance probabilities for \( \lambda(s) \).

The data component of `swissFit` provides all the values necessary for further analysis such as conditional simulation or re-estimation of model parameters. This `SpatialPointsDataFrame` contains all covariates \( X(s_i) \), observed data \( Y_i \), and residuals \( Y_i - X(s_i)\hat{\beta} \) (the latter on the Box-Cox transformed scale if appropriate). Conditional simulation is required for making inference on non-linear functions of the latent process (such as total area above a threshold).
and is advisable when making inference on the latent process with a Box-Cox transformed model. Using a wrapper for the \texttt{RFsimulate} function in \texttt{RandomFields}, a sample from the conditional distribution \([U|Y]\) is obtained with

```r
R> oneSim <- geostatsp::RFsimulate(model = swissFit$param,
+ data = swissFit$data["resid"], err.model = swissFit$param["nugget"],
+ x = raster(extent(swissRain), nrow = 10, ncol = 10))
```

As a final note on the Gaussian geostatistical model, consider the comparison between the \texttt{geoR} package (see Diggle and Ribeiro 2006) and \texttt{geostatsp} below. The code below estimates the shape and Box-Cox parameters for an isotropic model. Notice the specification of scaling factors for parameters \texttt{myscale}, given as a \texttt{control} argument.

```r
R> swissRain$alt <- raster::extract(swissAltitude, swissRain)
R> library("geoR")
R> swiss2 <- as.geodata(swissRain, data.col = "rain", covar.col = "alt")
R> myscale <- c(range = 1000, shape = 1, boxcox = 1, nugget = 0.1)
R> geoRres <- likfit(swiss2, ini.cov.pars = c(1, 10000), kappa = 0.2,
+ trend = ~alt, lambda = 0.5, fix.lambda = FALSE, fix.nugget = FALSE,
+ fix.kappa = FALSE, lik.method = "REML", message = FALSE,
+ control = list(parscale = myscale["range", "nugget", "shape",
+ "boxcox"]))
```

The same model is fit with \texttt{lglm} with:

```r
R> swissFit2 <- lglm(rain ~ CHE_alt, swissRain, grid = 90,
+ covariates = swissAltitude, shape = 0.2, fixShape = FALSE,
+ boxcox = 0.5, reml = TRUE, fixBoxcox = FALSE, parscale = myscale)
```

The two sets of parameter estimates are comparable, as shown below.

|             | (Intercept) | CHE_alt | range | nugget | boxcox | shape | variance |
|-------------|-------------|---------|-------|--------|--------|-------|---------|
| geostatsp   | 6.57        | 0.000153| 54820 | 0      | 0.592  | 0.959 | 14.4    |
| geoR        | 6.63        | 0.000160| 52900 | 0      | 0.595  | 1.005 | 14.5    |

### 2.2. Generalized linear geostatistical models

The Loaloa data (see Diggle and Ribeiro 2006, 7.6.4) shown in Figure 3 contains the locations of villages where subjects were tested for a tropical disease, with the (binomially distributed) number of positive samples and total sample size being recorded. These data are accessible with \texttt{data("loaloa")} in the \texttt{geostatsp} package, which contains a \texttt{SpatialPointsDataFrame} (named \texttt{loaloa}), and raster images for elevation (\texttt{elevationLoa}), vegetation index (\texttt{eviLoa}) and land type (\texttt{ltLoa}). Land type is shown as background values in Figure 3.

The generalized linear geostatistical model from (2) would be suitable for these data with \(f\) being a binomial distribution and \(g\) being a logit link function. The surface \(X(s)\) is multivariate and have values for land type, vegetation index, and elevation. The model can be fit to these data using the \texttt{glgm} function in the \texttt{geostatsp} package.
As there is more than one covariate, the three rasters containing covariates are grouped together in a list. The elevation covariate is to be fit as a linear effect with a change point at 750m, with two variables `elLow` and `elHigh` being the negative and positive portions of elevation data minus 750. These two rasters are created with

```r
R> elevationLoa <- elevationLoa - 750
R> elevLow <- reclassify(elevationLoa, c(0, Inf, 0))
R> elevHigh <- reclassify(elevationLoa, c(-Inf, 0, 0))
```

Land types with a very small number of observations are merged with more populated land types, with: savannas (9) changed to woody savannas (8); wetlands (5) and mixed forests (11) changed to forest (2); and croplands (12) and urban areas (13) changed to crop/natural mosaic (14).

```r
R> rcl <- rbind(c(9, 8), c(5, 2), c(11, 2), c(12, 14), c(13, 14))
R> ltLoaRe <- reclassify(ltLoa, rcl)
R> levels(ltLoaRe) = levels(ltLoa)
```

The following code creates the corresponding list of rasters, note that they may have different extents, resolutions or projections.

```r
R> covList <- list(elLow = elevLow, elHigh = elevHigh, land = ltLoaRe,
+     evi = eviLoa)
```

The call to `g1gm` appears below. As with `1gm`, spatial predictions will be made on a grid as specified by the `grid` argument. It can be specified as a `raster` object though in this case a square grid with 150 cells in the X direction is used. The Markov random field implicitly assumes $U(s)$ takes values of zero outside of the study region, and this effect can be partially negated by adding a buffer (in this case of 50km) around the study region where $U(s)$ will be evaluated but the values in these cells are not returned. Variables listed in the `formula` argument can be contained in either the first argument or in `covariates`. The number of samples taken per village is passed as the `Ntrials` argument, and the $y$ variable in the formula is the number of positive samples. The argument `shape` specifies the (fixed) shape parameter $\kappa$ of the Matérn correlation function.
Table 2: Posterior expectations and quantiles of model parameters obtained by fitting a
generalized linear geostatistical model to the Loaloa dataset using the glgm function.

| Parameter                      | Mean     | 0.025 quantile | 0.975 quantile |
|--------------------------------|----------|----------------|----------------|
| (Intercept)                    | -2.18e+00| -3.50e+00      | -8.62e-01      |
| factor(land)Woody savannas     | -4.38e-01| -7.92e-01      | -8.92e-02      |
| factor(land)Cropland/natural   | -2.57e-01| -5.91e-01      | 7.20e-02       |
| evi                            | 2.63e-04 | -7.88e-06      | 5.35e-04       |
| elHigh                         | -3.55e-03| -4.88e-03      | -2.18e-03      |
| elLow                          | 2.74e-03 | 1.45e-03       | 3.94e-03       |
| range                          | 4.22e+04 | 2.67e+04       | 6.46e+04       |
| sd                             | 9.88e-01 | 7.80e-01       | 1.25e+00       |

Bayesian inference requires prior distributions, and the priors for the spatial covariance parameters are specified by the priorCI argument. Prior 95% intervals for $\sigma$ and $\phi$ are specified, and glgm creates gamma priors for the precision $1/\sigma^2$ and scaled range parameter $\phi/\delta$ (with $\delta$ being the cell size) having the 95% intervals specified. Priors other than the gamma are possible (though currently unimplemented in geostatsp). The INLA methodology requires priors to be continuous, but are otherwise unrestricted. The INLA software specifies that priors are set for log precisions, with prior distributions available including the log-gamma and normal. Incorporating additional priors into INLA or geostatsp would be relatively straightforward. Priors for the remaining parameters can be specified with inla arguments such as control.fixed = list(prec.intercept = 0.01).

The result of the glgm function is a list with elements: inla for the raw results from INLA; parameters containing parameter prior and posterior distributions; and raster containing the posterior means of the random effects and fitted values. Table 2 contains posterior means and quantiles of the model parameters, taken from the object loaFit$parameters$summary.

Markov chain Monte Carlo (MCMC) methods are an alternative (and more established) method for fitting spatial models to non-Gaussian data. The geoRglm (Christensen and Ribeiro 2002) package provides an excellent set of functions for fitting generalized linear geostatistical models with MCMC, with Matérn correlation functions and geometric anisotropy.
being available. MCMC is more labor intensive and computationally intensive to use than INLA, but is able to produce joint posterior samples. The geoRglm package does not use the GMRF approximation, which has advantages and disadvantages. Geometric anisotropy is straightforward without the GMRF approximation, and non-integer shape parameters can be used. Readers unfamiliar with MCMC methods are advised to skip ahead to Section 2.3, as the following paragraphs will presuppose a good deal of familiarity with MCMC.

The first step before fitting the Loaloa model using geoRglm is to create a new geodata object, copying over the values of the covariates extracted from the rasters by glgm.

```R
R> library("geoRglm")
R> loaNoMissing <- loaloa[as.integer(rownames(loaFit$inla$.args$data)), ]
R> loa2 <- as.geodata(loaNoMissing, data.col = "y")
R> loa2$covariate <- loaFit$inla$.args$data[, + c("evi", "elLow", "elHigh", "land")]
R> loa2$covariate$evi <- loa2$covariate$evi - 4000
```

Next, the model and prior distributions are specified. The model.glm.control function specifies the fixed effects portion of the model and the Matérn correlation. The prior for the range parameter is taken from the glgm output, though notice the difference in parametrisations for the range by factor of $\sqrt{8}$.

```R
R> model.10 <- model.glm.control(kappa = 1, cov.model = "matern", trend.d = + trend.spatial(~ 1 + elLow + elHigh + evi + factor(land), loa2))
R> phiSeq <- seq(1 * 1000, 100 * 1000, len = 1001)
R> phiValues <- approx(loaFit$param$range$prior, xout = phiSeq * sqrt(8))$y
```
Control parameters for the MCMC run are created next. The number of iterations, thinning and burn-in, and scaling parameters are specified by `mcmc.control`,

```r
R> mcmc.10 <- mcmc.control(S.scale = 0.004, n.iter = 4e+05,
+   S.start = raster::extract(loaFit$raster[["random.mode"]],
+     loaNoMissing), phi.start = 40 * 1000/sqrt(8), phi.scale = 10,
+   thin = 1000, burn.in = 10000)
R> mcmc.10$S.start[is.na(mcmc.10$S.start)] <- 0
```

The MCMC run is accomplished with the `binom.krige.bayes` function. The code below defines a function to run a single chain, and subsequently runs three chains in parallel. Trace plots for three of the model parameters are show in Figure 5, along with the posterior means and quantiles from INLA. The lack of mixing in the range parameter, despite thinning by a factor of 1000, gives an indication of the perseverance and skill often required when using MCMC for spatial problems.

```r
R> oneChain <- function(phiMult) {
+   set.seed(100 * phiMult)
+   mcmc.10$phiStart = mcmc.10$phiStart * phiMult
+   binom.krige.bayes(loa2, units.m = loaNoMissing$N,
+     model = model.10, prior = prior.10, mcmc.input = mcmc.10,
+     output = output.glm.control(messages = FALSE))
+ }
R> library("parallel")
R> test.10 <- mccollect(list(mcparallel(oneChain(0.8)),
+   mcparallel(oneChain(1)), mcparallel(oneChain(1.2))))
```
2.3. Log-Gaussian Cox processes

The log-Gaussian Cox process (LGCP) is closely related to the GLGM and is a model suitable for describing the data on murder locations in the city of Toronto, Canada in Figure 6. These data are from the years 1990 to 2013 and appear in the Toronto Star newspaper (http://www.thestar.com/news/crime/torontohomicidemap.html). The murder dataset in geostatsp contains these locations, as well as raster images for median household income (torontoIncome), population density (torontoPdens), and ambient light (torontoNight).

The LGCP (see Møller, Syversveen, and Waagepetersen 1998) is a spatial point process model with the event locations \( \{P_i; i = 1 \ldots N\} \) being independently distributed conditional on a random log-Gaussian spatial random field \( \lambda(s) \). Allowing for a vector of covariates \( X(s) \) at location \( s \) (in this case light, income, and population density), a LGCP model for the murder locations is

\[
\{P_i; i = 1 \ldots N\}|U(\cdot) \sim \text{Poisson process}[\lambda(\cdot)] \\
\log[\lambda(s)] = \mu + X(s)\beta + U(s) \\
\text{cov}[U(s_i), U(s_j)] = \sigma^2 \text{Matérn}(|s_i - s_j|/\phi; \kappa).
\]

Using a Gaussian Markov random field approximation for \( U(s) \), with \( U(s) \) being piecewise constant, reduces the inferential problem to modelling the count of points within cells with a Poisson distribution. An improved methodology for fitting LGCP’s, using the previously mentioned triangular bases on an irregular lattice, is available in INLA (see Rue et al. 2013). Currently this method is not implemented in geostatsp, and the results below use the GMRF on a square lattice.

The lgcp function in geostatsp operates very similarly to glgm. A list of covariates is first created, with income, population density, and ambient light transformed so as to make them roughly symmetrically distributed.

R> covList <- list(loginc = log(torontoIncome), logpop = log(torontoPdens), +   loglight = log(torontoNight))

Next, the lgcp function is called. Here murder is a SpatialPoints object, and the locations themselves are the response. The model formula is one sided, specifying the covariates. The
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| Parameter          | Mean   | 0.025 quantile | 0.975 quantile |
|--------------------|--------|----------------|----------------|
| (Intercept)        | -5.07  | -9.80          | -0.31          |
| log(income)        | -1.32  | -1.73          | -0.92          |
| log(pop'n)         | 0.76   | 0.63           | 0.90           |
| log(light)         | 0.74   | 0.51           | 0.97           |
| range              | 649.55 | 480.39         | 860.58         |
| sd                 | 0.83   | 0.71           | 0.94           |

Table 3: Posterior means and quantiles of model parameters obtained from fitting the murder data with lgcp.

Figure 7: Posterior means of spatial surfaces obtained from fitting the Toronto murder data with the lgcp function.

Parameter posterior distributions are shown in Table 3. The raster component of the results contains posterior distributions for $U(s)$ and $\log[\lambda(s)]$, with the posterior mean of $\lambda(s)$ contained in `murderFit$raster["predict.exp"]` and plotted in Figure 7.

As with the generalized linear geostatistical model in the previous section, MCMC is an alternative to using INLA for LGCP’s which avoids many of INLA’s limitations. The most accessible software for using MCMC with LGCP’s is the `lgcp` package described in Taylor, Davies, Rowlingson, and Diggle (2013). As was noted before, MCMC is more computationally intensive and labor intensive than INLA, and requires more than a moderate amount of specialist knowledge to use reliably. One situation where MCMC is required is the case where the point location data are not directly observable. Taylor, Davies, Rowlingson, and Diggle
(2015) show how the `lgcp` package allows for LGCP’s to be a latent variable in a hierarchical model, with the LGCP inference nested within a data augmentation algorithm.

### 2.4. The Besag York and Mollié model

The final model which will be demonstrated is the Besag, York, and Mollié (1991) model (BYM), useful for modelling disease case counts in polygons. Figure 8a shows the number of Larynx cancer cases in each county of Kentucky in a single year, and Figure 8b shows the count that should be expected given the population of each age and sex group in the counties and the US national rates for Larynx cancer. As the case count in a county is often zero or one, the Standardized Mortality Ratio (observed divided by expected) would be expected to be a poor estimator of underlying risk and a spatial random effects model with a Poisson-distributed response variable would be more useful. The BYM model models the case count $Y_i$ of region $i$, given the expected count $E_i$ and covariates $X_i$, as

\[
Y_i \sim \text{Poisson}(E_i \lambda_i)
\]

\[
\log(\lambda_i) = \mu + X_i \beta + U_i
\]

\[
U_i = W_i + V_i
\]

\[
V_i \sim \text{i.i.d. } \mathcal{N}(0, \tau^2)
\]

\[
W_i | \{W_j; j \neq i\} \sim \mathcal{N}(\text{mean}\{W_j; j \sim i\}, \sigma^2 / |j \sim i|)
\]

Here $W$ follows a Markov random field model on the irregular lattice of regions, with $j \sim i$ referring to regions $i$ and $j$ being neighbors (sharing a common boundary line). Including the spatially independent term $V$ in the model allows for flexibility in the spatial dependence of $U$, with $\tau$ being larger than $\sigma$ resulting in a rough surface and $\sigma$ being larger creating a smoother surface.

The `kentucky` dataset in the `diseasemapping` package contains a `SpatialPolygonsDataFrame` of the counties of Kentucky, and includes the population by age and sex group and the proportion of individuals living in poverty. The `larynx` object is a case file, with one row per individual with larynx cancer in a single year and a variable denoting their county of residence. The `cancerRates` and `getSMR` functions in `diseasemapping` can be used to generate observed and expected counts for each county with the following code. The observed counts and expected counts are shown in Figure 8.
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|       | Mean | 0.025 quantile | 0.975 quantile |
|-------|------|----------------|----------------|
| $\mu$ | 0.11 | −0.38          | 0.61           |
| poverty | 0.01 | −0.02          | 0.03           |
| $\sigma$ | 0.19 | 0.08           | 0.42           |
| $\tau$ | 0.20 | 0.08           | 0.41           |

Table 4: Posterior means and 95% credible intervals for model parameters obtained from fitting the Kentucky cancer data with the `bym` function.

Figure 9: Posterior means and probabilities obtained from fitting the Kentucky larynx cancer data with the `bym` function.

```
R> library("diseasemapping")
R> data("kentucky")
R> larynxRates <- cancerRates("USA", year = 1998:2002, site = "Larynx")
R> kentucky <- getSMR(kentucky, larynxRates, larynx, regionCode = "County")

The `bym` function performs Bayesian inference function for the BYM model. It takes as its arguments the `SpatialPolygonsDataFrame` containing the regional boundaries and variables, as well as the model formula and the prior 95% intervals for $\sigma$ and $\tau$.

```
R> kBYM <- bym(formula = observed ~ offset(logExpected) + poverty, data = + kentucky, priorCI = list(sdSpatial = c(0.1, 5), sdIndep = c(0.1, 5)))
```

The result has a component for the INLA results (`inla`), the parameter posterior distributions (`parameters`), and a `SpatialPolygonsDataFrame` with the spatial results. The posterior means and quantiles of the parameters are given in Table 4, obtained from `kBYM$parameters$summary`.

Figure 9a shows the posterior mean of relative risk $\lambda_i$ and can be produced with `spplot(kBYM$data, "fitted.exp")`. Figure 9b shows that posterior probability that each county has a cancer rate more than 30% in excess of the US national rates, obtained by numerically integrating the marginal posterior using the `excProb` function called below performs the integration using the `trapz` function in the `pracma` package (Borchers 2014).

```
R> kBYM$data$excProb <- excProb(kBYM$inla$marginals.fitted.bym, log(1.3))
```
Figure 10: Histograms of posterior samples (black bars) obtained from fitting the Kentucky cancer data with OpenBUGS (black bars), along with posterior densities from the bym function (blue lines).

Fitting the BYM model is much less computationally intensive than is the case for LGCP’s and the GLGM, and MCMC has a long history of being used with the BYM model (see Gilks, Richardson, and Spiegelhalter 1996). OpenBUGS (Sturtz, Ligges, and Gelman 2005) is a flexible and popular tool for running MCMC, and is able to fit the BYM model. The glmmBUGS and R2OpenBUGS packages (see Brown and Zhou 2010; Sturtz et al. 2005) can be used to fit the BYM model with a minimum amount of effort, providing a simple interface between R and OpenBUGS. First, model files are prepared and starting values computed. The priors argument creates gamma priors for the standard deviation parameters which are not entirely dissimilar from the priors used by the bym function. The glmmBUGS function requires priors to be specified for standard deviations, though it would be possible to edit the model file manually to set a posterior for the precision parameter in order to replicate INLA’s results.

```R
R> library("spdep")
R> kAdjMat <- poly2nb(kentucky, row.names = as.character(kentucky$County))
R> library("glmmBUGS")
R> kBYMbugs <- glmmBUGS(observed + logExpected ~ poverty,
+                      data = as.data.frame(kentucky), effects = "County", family = "poisson",
+                      spatial = kAdjMat, modelFile = "kentuckyBYM.txt", initFile = "kInit.R",
+                      priors = c(SDCountySpatial = "dggamma(5.46,42,0.555)",
+                      SDCounty = "dggamma(5.46,42,0.555)"))
```

Second, starting values loaded. The file kInit.R contains code for a function to generate random starting values, and users are encouraged to edit this file prior to sourcing it.

```R
R> startingValues <- kBYMbugs$startingValues
R> source("kBYMbugs$startingValues")
```

Finally, OpenBUGS is run.

```R
R> library("R2OpenBUGS")
R> kResult <- bugs(kBYMbugs$ragged, inits = getInits,
+                 model.file = "kentuckyBYM.txt", parameters = names(getInits()),
+                 n.chain = 3, n.iter = 1000, n.burnin = 200, n.thin = 200)
```
Figure 10 shows marginal posterior distributions for three of the model parameters. The spatial variance parameter has a slightly different posterior from $bym$, due to the differences in the prior distribution.

2.5. A short simulation study

This section aims to illustrate the ease with which spatial simulation studies can be carried out using the geostatsp package. Gaussian data is simulated and inference is carried out both with Maximum Likelihood Estimation using the likfit function, and Bayesian inference using glm.

Before simulating data, spatial covariates must be created and model parameters specified. The following code defines two simple covariates as sloping north to south and east to west respectively on a square area measuring 10 units across.

```r
R> covariates <- brick(xmn = 0, ymn = 0, xmx = 10, ymx = 10,
+    ncols = 200, nrows = 200, nl = 2)
R> values(covariates)[, 1] <- rep(seq(0, 1, len = nrow(covariates)),
+    ncol(covariates))
R> values(covariates)[, 2] <- rep(seq(0, 1, len = nrow(covariates)),
+    rep(nrow(covariates), ncol(covariates)))
R> names(covariates) <- c("cov1", "cov2")
```

Next, a spatial covariance structure is specified with $\sigma = 2$, $\phi = 2.5$, $\tau = 1/2$ and $\kappa = 2$ is specified.

```r
R> myModel <- c(intercept = 0.5, variance = 2^2, nugget = 0.5^2,
+    range = 2.5, shape = 2, cov1 = 0.2, cov2 = -0.5)
```

The RFsimulate function in geostatsp, a wrapper for the function of the same name in RandomFields (Schläfer et al. 2015), is used to simulate a $U(s)$ surface as a raster with the same resolution and dimension as cov1. An intercept and the two covariates are added to create a $\lambda(s)$.

Next, points in the study region are simulated at random. Observations at these locations are created by extracting values of $\lambda(s)$ and adding random normal noise with standard deviation 0.5.

```r
R> Npoints <- 50
R> myPoints <- SpatialPoints(cbind(runif(Npoints, 0, 10),
+    runif(Npoints, 0, 10)))
R> myPoints <- SpatialPointsDataFrame(myPoints,
+    as.data.frame(extract(covariates, myPoints)))
R> myPoints$fixed <- myModel["intercept"] + drop(as.matrix(data.frame(
+    myPoints))[, names(covariates)]) %*% myModel[names(covariates)]
R> myPoints$U <- RFsimulate(myPoints, model = myModel)$sim1
R> myPoints$y <- myPoints$fixed + myPoints$U + rnorm(length(myPoints),
+    0, sqrt(myModel["nugget"]))
```

MLEs are computed with geostatsp's lgm function,
Figure 11 shows the parameter estimates for 12 simulations. Figure 11a contains the posterior mean and 95% credible interval for the first $\beta$ coefficient obtained from Bayesian inference, and the MLE and 95% confidence interval obtained from Frequentist inference. Notice the estimates and intervals are nearly identical. Figure 11b shows the Bayesian posterior mean and credible interval for the range parameter $\phi$ along with the MLE. Confidence intervals for the range are not produced by lgm. The MLE and the posterior mean differ, sometimes substantially, but tend to identify the true value despite $\kappa$ being misspecified.

3. Discussion

The geostatsp and diseasemapping packages remove much of the drudgery involved in fitting spatial random effects models with INLA (Rue et al. 2013) or geoR (Diggle and Ribeiro 2013).
The creation of the lattice for the Markov Random Field in \texttt{INLA} in particular is time consuming to code, and translation of \texttt{INLA} results to maps and interpretable parameters is not always straightforward. The use of \texttt{Raster} objects for covariates and default values able to accommodate UTM spatial coordinates (with values in the thousands or hundreds of thousands) do away with the need to modify and reformat data prior to its analysis.

These packages provide a mechanism for fitting geostatistical models using \texttt{R} spatial data types: \texttt{SpatialPointsDataFrames}, \texttt{SpatialPolygonsDataFrames}, and \texttt{Rasters}. The advantage of working with these data types in place of matrices and vectors of coordinates is two-fold. First, data from various sources can be easily downloaded and included in these analyses. NASA provides a wide variety of satellite data, including elevation and vegetation indices, which can be loaded into \texttt{R} using the \texttt{raster} package. Census data are often available as Shapefiles which can be read using \texttt{rgdal}. Much of the geographic data from sources such as these have coordinates in a longitude-latitude projection, and geostatistical analyses involving Euclidean distances require coordinates on a metre-based (or UTM) projection. Converting coordinates is easily accomplished with the \texttt{spTransform} and \texttt{projectRaster} functions, and many additional functions in the \texttt{sp} and \texttt{raster} packages are available for data manipulation and processing.

A second advantage accruing from the use of \texttt{R} spatial objects is the ease with which results can be exported to GIS software or plotted with background map images in \texttt{R}. The maps presented in this paper have required projecting the result to a longitude-latitude coordinate reference system with the \texttt{projectRaster} function, downloading background layers from Openstreetmap.org with the \texttt{mapmisc} package, and obtaining city names and locations with \texttt{geonames}. While it is possible to improve on the maps presented here using GIS software, code for \texttt{R} generated maps can be incorporated in \texttt{Sweave} and \texttt{knitr} scripts thereby allowing any manual GIS map creation to be reserved for final drafts of documents.

The \texttt{geostastp} package could be improved by incorporating several more of the facilities in \texttt{INLA} and work towards this is ongoing. Replacing the grid of square cells in the MRF approximation with the irregular lattices in Lindgren et al. (2011) would increase speed and accuracy of the approximation, though spatial predictions could still be made on rasters and the inner workings of the approximation could remain hidden from the user. The \texttt{INLA} software allows for non-parametric effects of covariates, it would be possible to specify non-parametric effects in \texttt{glm} though they would not as of yet be included when making spatial predictions. Also, \texttt{INLA} can fit a variety of spatio-temporal models and a simple user-friendly interface to fitting spatio-temporal data would certainly be possible.

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A. Matérn correlation

There are several parametrisations of the Matérn correlation function, and the range parameter in \texttt{lgm} and \texttt{glgm} corresponds to \( \phi \) in

\[
\rho(h; \phi, \kappa) = \frac{1}{\Gamma(\kappa)2^{\kappa-1}} \left( \frac{\sqrt{8\kappa}\|h\|}{\phi} \right)^\kappa K_\kappa \left( \frac{\sqrt{8\kappa}\|h\|/\phi} \right).
\]

\( \Gamma(\cdot) \) is a gamma function and \( K_\kappa \) is a modified Bessel function of the second kind of order \( \kappa \).

Figure 12 shows plots of the Matérn for various values of \( \kappa \) and all with \( \phi = 1 \). Notice that, with the possible exception of \( \kappa = 0.1 \), the correlations intersect (more or less) at \( ||h|| = 1 \).

A not inaccurate interpretation of the range parameter \( \phi \) in this parametrization is it is the distance beyond which correlation is both fairly small (\(< 0.14\)), and decaying fairly slowly regardless of the shape parameter \( \kappa \).

The \texttt{geostatsp} package has a \texttt{matern} function which implements the parametrization above, though it may be helpful to consider the function below. Figure 12 is produced with this code.

```r
R> mymatern <- function(u, phi, kappa) {
+   uscale <- sqrt(8 * kappa) * u/phi
+   res <- (1/(gamma(kappa) * 2^(kappa - 1))) * uscale^kappa *
+         besselK(uscale, kappa)
+   res[u == 0] <- 1
+   res
+ }
```

Wikipedia (2013) and the ‘matern’ model in the \texttt{RandomFields} package define the range parameter as \( \phi_1 = \phi/2 \). Diggle and Ribeiro (2006), the \texttt{geoR} package, and the \texttt{whittle} model in \texttt{RandomFields} have a range parameter \( \phi_2 = \phi/\sqrt{8\kappa} \). It is also common to define the Matérn with a scale parameter in place of the range, with the scale parameter being \( \alpha = 1/\phi_2 \).

Lindgren \textit{et al.} (2011) use either the scale \( \alpha \) or the range \( \phi \). The \texttt{Range} parameter produced by \texttt{inla} is \( \phi\delta \), with \( \delta \) being the length of the sides of the grid cells, as confirmed below.

![Figure 12: Matérn correlation functions with \( \phi = 1 \) and various values of the shape parameter \( \kappa \) ](image.png)
R> c(loaFit$inla$summary.hyperpar["Range for space", "mode"] * 
+ xres(loaFit$raster), loaFit$par$summary["range", "mode"])

[1] 38489 38489

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