ngspatial: A Package for Fitting the Centered Autologistic and Sparse Spatial Generalized Linear Mixed Models for Areal Data

by John Hughes

Abstract Two important recent advances in areal modeling are the centered autologistic model and the sparse spatial generalized linear mixed model (SGLMM), both of which are reparameterizations of traditional models. The reparameterizations improve regression inference by alleviating spatial confounding, and the sparse SGLMM also greatly speeds computing by reducing the dimension of the spatial random effects. Package **ngspatial** ('ng' = non-Gaussian) provides routines for fitting these new models. The package supports composite likelihood and Bayesian inference for the centered autologistic model, and Bayesian inference for the sparse SGLMM.

Background and introduction

The traditional autologistic model (Besag, 1972) and areal GLMM (Besag et al., 1991) have enjoyed widespread popularity: they have been applied thousands of times in many fields, e.g., epidemiology, marketing, agriculture, ecology, forestry, geography, and image analysis. But it was recently discovered that both models are spatially confounded (Caragea and Kaiser, 2009; Clayton et al., 1993). This confounding can cause bias and/or variance inflation in the estimators of regression coefficients, leading to erroneous regression inference. This is a serious drawback because many spatial modelers are interested in regression (rather than, or in addition to, prediction).

To address the confounding of the traditional autologistic model, Caragea and Kaiser (2009) devised the centered autologistic model, so named because it replaces the traditional model's autocovariate with a centered autocovariate (see below for details).

The confounding of the mixed model was first addressed by Reich et al. (2006) using a technique now known as restricted spatial regression (Hodges and Reich, 2010). This technique alleviates spatial confounding and also yields a faster mixing Markov chain, but the computational burden remains high because the dimension of the spatial random effects is reduced only slightly relative to the traditional model. By using the so called Moran operator, Hughes and Haran (2013) were able to reparameterize the mixed model in a way that not only improves regression inference but also dramatically reduces the dimension of the random effects. The resulting model, which we will call the sparse SGLMM, can be fitted so efficiently that even the largest areal datasets can be analyzed quickly.

These promising new models cannot be applied using existing software, and so we have provided support for the models in version 1.0 of R package **ngspatial**, the subject of this article. First we discuss the two models in some detail. Then we present **ngspatial** 1.0, which permits composite likelihood and Bayesian inference for the centered autologistic model, and Bayesian inference for the sparse SGLMM. We conclude with a summary.

The models supported by ngspatial 1.0

Areal models

The autologistic model and the sparse SGLMM are areal models, i.e., models for data observed at the vertices of a graph G = (V, E), where $V = \{1, 2, \ldots, n\}$ are the vertices, and $E \subset V \times V$ are the edges. Each vertex of G represents an areal unit, i.e., an area over which measurements have been aggregated. For example, areal units could be counties, census tracts, voxels, pixels, or provinces. An edge (i,j) of G represents the spatial adjacency of areal units i and j. Typically, two areal units are considered adjacent if they share a boundary, but other definitions are possible. We will assume that G is undirected and free of loops and parallel edges.

The centered autologistic model

The traditional autologistic model was proposed by Besag (1972). The model is a Markov random field (MRF) model (Kindermann and Snell, 1980), which is to say that G describes conditional independencies among the random variables Z_i (i = 1, ..., n) associated with V. For the autologistic model, the ith observation is Bernoulli distributed conditional on its neighbors:

$$\log \frac{\mathbb{P}(Z_i = 1 \mid \{Z_j : (i, j) \in E\})}{\mathbb{P}(Z_i = 0 \mid \{Z_j : (i, j) \in E\})} = \boldsymbol{x}_i' \boldsymbol{\beta} + \eta \sum_{j: (i, j) \in E} Z_j,$$

where x_i is a p-vector of spatial predictors associated with the ith areal unit, β is a p-vector of spatial regression coefficients, η is a spatial dependence parameter, and $\sum Z_i$ is the so called autocovariate.

We see that η is a measure of Z_i 's reactivity to its neighbors. If $\eta=0$, the model reduces to the ordinary Bernoulli GLM, while $\eta>0$ (< 0) corresponds to positive (negative) spatial dependence. We will assume that $\eta>0$ since the model is usually applied to phenomena that exhibit spatial attraction rather than repulsion.

Caragea and Kaiser (2009) showed that the traditional autologistic model is confounded. This is because the traditional autocovariate is not well suited to the task of fitting small-scale structure in the data, i.e., clustering induced by spatial dependence and residual to the large-scale structure $x'\beta$. Instead, the traditional autocovariate and the spatial predictors "compete" to explain the data, which prevents the model from isolating the role of the spatial predictors.

Caragea and Kaiser (2009) reparameterized the model by centering the autocovariate. The resulting conditional log odds are

$$\log \frac{\mathbb{P}(Z_i = 1 \mid \{Z_j : (i,j) \in E\})}{\mathbb{P}(Z_i = 0 \mid \{Z_j : (i,j) \in E\})} = x_i' \beta + \eta \sum_{j:(i,j) \in E} (Z_j - \mu_j),$$

where $\mu_j = \{1 + \exp(-x_j'\beta)\}^{-1}$ is the independence expectation of Z_j . Centering allows the autocovariate to fit only residual structure so that fitting the large-scale structure is left to the regression term. Thus the centered model restores to β and η their desired interpretations as regression coefficients and dependence parameter, respectively.

Maximum likelihood and Bayesian inference for the autologistic model are complicated by an intractable normalizing function. To see this, first let $\mathbf{Z} = (Z_1, \dots, Z_n)'$; let \mathbf{X} be the design matrix; let $\mathbf{A} = [1\{(i,j) \in E\}]$ be the adjacency matrix of G, where $1\{\cdot\}$ is the indicator function; let $\mathbf{\theta} = (\mathbf{\beta}', \mathbf{\eta})'$ be the full parameter vector; and let $\mathbf{\mu} = (\mu_1, \dots, \mu_n)'$ be the vector of independence expectations. Then, assuming G has clique number 2, the joint pmf of the centered model is

$$\pi(\mathbf{Z} \mid \boldsymbol{\theta}) = c(\boldsymbol{\theta})^{-1} \exp\left(\mathbf{Z}' \mathbf{X} \boldsymbol{\beta} - \eta \mathbf{Z}' \mathbf{A} \boldsymbol{\mu} + \frac{\eta}{2} \mathbf{Z}' \mathbf{A} \mathbf{Z}\right), \tag{1}$$

where

$$c(\boldsymbol{\theta}) = \sum_{\boldsymbol{Y} \in \{0,1\}^n} \exp\left(\boldsymbol{Y}' \boldsymbol{X} \boldsymbol{\beta} - \eta \boldsymbol{Y}' \boldsymbol{A} \boldsymbol{\mu} + \frac{\eta}{2} \boldsymbol{Y}' \boldsymbol{A} \boldsymbol{Y}\right)$$

is the normalizing function (Hughes et al., 2011).

The normalizing function is intractable for all but the smallest datasets because the sample space $\{0,1\}^n$ contains 2^n points. Our package offers two ways to solve this problem: (1) composite likelihood inference, which sidesteps $c(\theta)$, and (2) auxiliary-variable MCMC for Bayesian inference, which allows $c(\theta)$ to cancel from the Metropolis-Hastings acceptance probability. See below for details.

The sparse spatial generalized linear mixed model

The traditional SGLMM for areal data—sometimes referred to as the BYM model—was proposed by Besag, York, and Mollié (1991). The BYM model is hierarchical, inducing spatial dependence by way of a latent autonormal random vector. Conditional on these spatial random effects, the observations are independent and follow an ordinary GLM. Specifically, the transformed conditional means are

$$g(\mu_i) = \mathbf{x}_i' \mathbf{\beta} + S_i, \tag{2}$$

where g is a link function, $\mu_i = \mathbb{E}(Z_i \mid S_i)$, and S_i is the spatial random effect for the ith areal unit.

The most common specification for $S = (S_1, ..., S_n)'$ is the so called intrinsic conditional autoregression (ICAR), a zero-mean Gaussian Markov random field (GMRF; Rue and Held, 2005) with a

singular precision matrix that corresponds to the intuitively appealing full conditional distributions

$$S_i \mid \{S_j : (i,j) \in E\} \sim \mathcal{N}\left(\frac{1}{d_i} \sum_{j:(i,j) \in E} S_j, \frac{1}{\tau d_i}\right),$$

where d_i is the degree of vertex i and $\tau > 0$ is a smoothing parameter. These conditionals correspond to the joint distribution

$$S \sim \mathcal{N}\{\mathbf{0}, (\tau \mathbf{Q})^{-1}\},$$

where $\mathbf{Q} = \mathbf{D} - \mathbf{A}$, with $\mathbf{D} = \operatorname{diag}(d_i)$. Since \mathbf{Q} is singular, the BYM model is employed from the Bayesian point of view, with the ICAR a prior distribution on \mathbf{S} . Packages CARBayes (Lee, 2013) and spdep (Bivand, 2015) provide tools for fitting the ICAR and other CAR models.

Reich et al. (2006) showed that the BYM model is spatially confounded in the sense that the random effects can "pollute" the regression manifold $C(\mathbf{X})$, which can lead to a biased and variance-inflated posterior for $\boldsymbol{\beta}$. To see this, first let \mathbf{P} be the orthogonal projection onto $C(\mathbf{X})$, so that $\mathbf{I} - \mathbf{P}$ is the orthogonal projection onto $C(\mathbf{X})^{\perp}$. Now eigendecompose \mathbf{P} and $\mathbf{I} - \mathbf{P}$ to obtain orthogonal bases $(\mathbf{K}_{n \times p} \text{ and } \mathbf{L}_{n \times (n-p)}, \text{ say})$ for $C(\mathbf{X})$ and $C(\mathbf{X})^{\perp}$. Then (2) can be rewritten as

$$g(\mu_i) = x_i'\beta + k_i'\gamma + l_i'\delta$$
,

where $\gamma_{p\times 1}$ and $\delta_{(n-p)\times 1}$ are random coefficients. This form shows that **K** is the source of the confounding, for **K** and **X** have the same column space.

Since the columns of K are merely synthetic predictors (i.e., they have no scientific meaning), Reich et al. (2006) recommend removing them from the model. The resulting model (henceforth the RHZ model) has

$$g(\mu_i) = x_i' \beta + l_i' \delta,$$

so that spatial smoothing is restricted to the orthogonal complement of $C(\mathbf{X})$. In a subsequent paper, Hodges and Reich (2010) referred to this technique as restricted spatial regression.

Restricted spatial regression is not only an effective remedy for confounding but also speeds computing. Because the columns of **L** are orthogonal, the RHZ model's random effects are approximately *a posteriori* uncorrelated. This yields a fast mixing Markov chain, and the cost per iteration is reduced because a simple spherical normal proposal is sufficient for updating the random effects. But fitting the RHZ model to large areal datasets is still quite burdensome computationally because the random effects remain high dimensional.

By taking full advantage of G, Hughes and Haran (2013) were able to greatly reduce the number of random effects while also improving regression inference. Hughes and Haran (2013) begin by defining the so called Moran operator for X with respect to G: (I - P)A(I - P). This operator appears in the numerator of a generalized form of Moran's I, a popular nonparametric measure of spatial dependence (Moran, 1950):

$$l_{\boldsymbol{X}}(\mathbf{A}) = \frac{n}{\mathbf{1}'\mathbf{A}\mathbf{1}} \frac{\boldsymbol{Z}'(\mathbf{I} - \mathbf{P})\mathbf{A}(\mathbf{I} - \mathbf{P})\boldsymbol{Z}}{\boldsymbol{Z}'(\mathbf{I} - \mathbf{P})\boldsymbol{Z}}.$$

Boots and Tiefelsdorf (2000) showed that (1) the (standardized) spectrum of a Moran operator comprises the possible values for the corresponding $I_X(\mathbf{A})$, and (2) the eigenvectors comprise all possible mutually distinct patterns of clustering residual to $C(\mathbf{X})$ and accounting for G. The positive (negative) eigenvalues correspond to varying degrees of positive (negative) spatial dependence, and the eigenvectors associated with a given eigenvalue (λ_i , say) are the patterns of spatial clustering that data exhibit when the dependence among them is of degree λ_i .

In other words, the eigenvectors of the Moran operator form a multiresolutional spatial basis for $C(\mathbf{X})^{\perp}$ that exhausts all possible patterns that can arise on G. Since we do not expect to observe repulsion in the phenomena to which these models are usually applied, we can use the spectrum of the operator to discard all repulsive patterns, retaining only attractive patterns for our analysis (although it can be advantageous to accommodate repulsion (Griffith, 2006). The nature of the attractive Moran eigenvectors is illustrated in Figure 1.

By retaining only eigenvectors that exhibit positive spatial dependence, we can usually reduce the model dimension by at least half *a priori*. And Hughes and Haran (2013) showed that a much greater reduction is possible in practice, with 50–100 eigenvectors being sufficient for most datasets.

Let $\mathbf{M}_{n \times q}$ contain the first $q \ll n$ eigenvectors of the Moran operator. Then the sparse SGLMM has first stage

$$g(\mu_i) = \boldsymbol{x}_i'\boldsymbol{\beta} + \boldsymbol{m}_i'\boldsymbol{\delta}_s,$$

where m_i is the *i*th row of **M** and δ_s is a *q*-vector of random coefficients. This implies p + q + 1 model

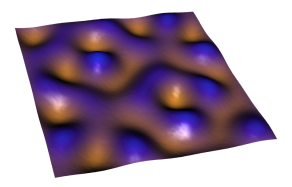


Figure 1: A Moran eigenvector exhibiting strong positive spatial dependence. The graph is the square 100×100 lattice. The coordinates of the vertices ($\mathbf{x}_i = (x_i, y_i)'$ for $i = 1, \dots, 10\,000$) were restricted to the unit square, and those coordinates were used as spatial covariates, i.e., $\mathbf{X}' = (\mathbf{x}_1 \cdots \mathbf{x}_{10\,000})$.

parameters, compared to p + n + 1 for the traditional model and p + (n - p) + 1 = n + 1 for the RHZ model. This dramatic reduction in dimension speeds computation considerably, allowing even the largest areal datasets to be analyzed quickly.

We note that the sparse SGLMM was partly inspired by spatial filtering, also known as principle coordinates of neighbor matrices. See, e.g., Griffith (2003); Dray et al. (2006); Borcard et al. (2011) for more information. Package **spdep** provides tools for spatial filtering.

Package ngspatial

Package **ngspatial** supports composite likelihood and Bayesian inference for the centered autologistic model, and Bayesian inference for the sparse SGLMM. Usage of the package's two main functions, autologistic and sparse.sglmm, is simple and intuitive while offering considerable flexibility.

Fitting the centered autologistic model

The fitting function for the centered autologistic model is called autologistic. This function returns an object of class "autologistic". Auxiliary functions residuals, summary, and vcov accept an object of type "autologistic" and return various kinds of residuals, summarize the fit, and return an estimated covariance matrix, respectively.

A fast perfect sampler for the centered autologistic model

A key component of our autologistic framework is function rautologistic, which simulates a draw from a centered autologistic model. The function has the following signature.

rautologistic(X, A, theta)

The three arguments are as described above. The functions of **ngspatial** require that an adjacency matrix be binary and symmetric (see function isSymmetric from the **base** package) and of type matrix. Note that package **spdep** provides function nb2mat, which constructs an adjacency matrix from a neighbors list. **spdep** also provides many other functions that are useful for handling adjacency structures.

This function employs coupling from the past (Propp and Wilson, 1996) to generate a vector distributed exactly according to (1). We use perfect sampling, rather than ordinary MCMC, for two reasons. First, the MCMC algorithm we use to do Bayesian inference requires that we draw a perfect sample during each iteration (Møller et al., 2006). Were we to use an approximation, the resulting Markov chain would not necessarily have the true posterior as its stationary distribution (Murray et al., 2006). Second, although perfect sampling can be computationally burdensome, a carefully implemented perfect sampler is fast enough to permit composite likelihood analysis of even very large datasets, while obviating convergence diagnosis.

Our perfect sampler was implemented in C++. More specifically, we used the Armadillo linear algebra library (Sanderson, 2010), which provides two benefits: (1) intuitive syntax for linear algebra (e.g., $\mathbf{Z}'\mathbf{X}\boldsymbol{\beta}$ can be computed as Z.t() * X * beta), and (2) speed (Armadillo uses delayed evaluation to combine multiple operations, thus reducing or eliminating the need for temporaries). We integrated the C++ and R code using the Rcpp and RcppArmadillo packages (Eddelbuettel and François, 2011).

We tested our Armadillo-based sampler in a number of circumstances and found it to be over three times as fast as an optimal pure R version.

Composite likelihood inference for the centered autologistic model

One way to overcome the intractability of the normalizing function $c(\theta)$ is to avoid it. This can be accomplished by considering the so called pseudolikelihood (PL), which is a composite likelihood (Lindsay, 1988) of the conditional type. Each of the n factors in the pseudolikelihood is the likelihood of a single observation, conditional on said observation's neighbors:

$$p_i(\boldsymbol{\theta})^{z_i} \{1 - p_i(\boldsymbol{\theta})\}^{1 - z_i} = \mathbb{P}(Z_i = z_i \mid \{Z_j : (i, j) \in E\}) = \frac{\exp[z_i \{\boldsymbol{x}_i' \boldsymbol{\beta} + \eta \mathbf{A}_i (\boldsymbol{Z} - \boldsymbol{\mu})\}]}{1 + \exp\{\boldsymbol{x}_i' \boldsymbol{\beta} + \eta \mathbf{A}_i (\boldsymbol{Z} - \boldsymbol{\mu})\}},$$

where z_i is the observed value of Z_i , and A_i is the ith row of A. Since the p_i are free of the normalizing function, so is the log pseudolikelihood, which is given by

$$\ell_{\text{PL}}(\boldsymbol{\theta}) = \boldsymbol{Z}' \{ \boldsymbol{X}\boldsymbol{\beta} + \eta \boldsymbol{A}(\boldsymbol{Z} - \boldsymbol{\mu}) \} - \sum_{i} \log[1 + \exp\{\boldsymbol{x}_{i}'\boldsymbol{\beta} + \eta \boldsymbol{A}_{i}(\boldsymbol{Z} - \boldsymbol{\mu}) \}].$$
(3)

Although (3) is not the true log likelihood unless $\eta=0$, Besag (1975) showed that the MPLE converges almost surely to the maximum likelihood estimator (MLE) as the lattice size goes to ∞ (under an infill regime). For small samples the MPLE is less precise than the MLE (and the Bayes estimator), but point estimation of β is generally so poor for small samples that precision is unimportant. When the sample size is large enough to permit accurate estimation of β , the MPLE is nearly as precise as the MLE (Hughes et al., 2011).

We find the MPLE $\tilde{\theta} = \arg \max \ell_{PL}(\theta)$ by using the optim function to minimize $-\ell_{PL}(\theta)$. Although the Nelder-Mead simplex algorithm finds the minimum quickly, we opted for the much faster BFGS method. To speed computation even further, we supply the score function

$$\nabla \ell_{\text{PL}}(\boldsymbol{\theta}) = ((\boldsymbol{Z} - \boldsymbol{p})'(\mathbf{I} - \eta \mathbf{A}\mathbf{D})\mathbf{X}, (\boldsymbol{Z} - \boldsymbol{p})'\mathbf{A}(\boldsymbol{Z} - \boldsymbol{\mu}))',$$

where
$$p = (p_1, ..., p_n)'$$
 and $D = \text{diag}\{\mu_i(1 - \mu_i)\}.$

Confidence intervals can be obtained using a parametric bootstrap or sandwich estimation. For the former we generate b samples from $\pi(\mathbf{Z} \mid \tilde{\boldsymbol{\theta}})$ and compute the MPLE for each sample, thus obtaining the bootstrap sample $\tilde{\boldsymbol{\theta}}^{(1)}, \ldots, \tilde{\boldsymbol{\theta}}^{(b)}$. Appropriate quantiles of the bootstrap sample are then used to construct approximate confidence intervals for the elements of $\boldsymbol{\theta}$.

The second approach for computing confidence intervals is based on (Varin et al., 2011)

$$\sqrt{n}(\tilde{\boldsymbol{\theta}} - \boldsymbol{\theta}) \Rightarrow \mathcal{N}\{\mathbf{0}, \boldsymbol{\mathcal{I}}^{-1}(\boldsymbol{\theta})\boldsymbol{\mathcal{J}}(\boldsymbol{\theta})\boldsymbol{\mathcal{I}}^{-1}(\boldsymbol{\theta})\},$$
 (4)

where $\mathcal{I}^{-1}(\theta)\mathcal{J}(\theta)\mathcal{I}^{-1}(\theta)$ is the so called Godambe sandwich matrix (Godambe, 1960). The "bread" in this sandwich is the inverse of the information matrix $\mathcal{I}(\theta) = -\mathbb{E}\nabla^2\ell_{\text{PL}}(\theta)$, and the "filling" is the variance of the score: $\mathcal{J}(\theta) = \mathbb{E}\nabla\nabla'\ell_{\text{PL}}(\theta)$. We use the observed information (computed by optim) in place of \mathcal{I} and estimate \mathcal{J} using a parametric bootstrap. For the bootstrap we simulate b samples $\mathbf{Z}^{(1)},\ldots,\mathbf{Z}^{(b)}$ from $\pi(\mathbf{Z}\mid\tilde{\theta})$ and estimate \mathcal{J} as

$$\hat{\boldsymbol{\mathcal{J}}}(\tilde{oldsymbol{ heta}}) = rac{1}{b} \sum_{k=1}^b
abla
abla' \ell_{ ext{PL}}(ilde{oldsymbol{ heta}} \mid oldsymbol{Z}^{(k)}).$$

Because the bootstrap sample can be generated in parallel (using the **parallel** package) and little subsequent processing is required, these approaches to inference are very efficient computationally, even for large datasets. We note that sandwich estimation is over twice as fast as the full bootstrap. Moreover, asymptotic inference and bootstrap inference yield comparable results for practically all sample sizes because (4) is not, in fact, an asymptotic result. This is because the log pseudolikelihood is approximately quadratic with Hessian approximately invariant in law, which implies that the MPLE is approximately normally distributed irrespective of sample size (Geyer, 2013).

Bayesian inference for the centered autologistic model

For Bayesian inference we use the auxiliary-variable MCMC algorithm of Møller et al. (2006), which allows us to construct a proposal distribution so that the normalizing constant cancels from the Metropolis-Hastings ratio. The method requires that we can draw independent realizations from the unnormalized density for any value of θ , which we can do using our perfect sampler.

Let $h(Z \mid \theta) = \exp\{Q(Z \mid \theta)\}$ denote the unnormalized density, where

$$Q(Z \mid \theta) = Z'X\beta - \eta Z'A\mu + \frac{\eta}{2}Z'AZ$$

is the so called negpotential function; let $Y \in \Omega$ denote the auxiliary variable; and let $f(\cdot)$ denote a prior distribution. Then the Metropolis-Hastings random walk acceptance probability for the algorithm of Møller et al. is given by

$$\alpha = \frac{h(\mathbf{Y}^* \mid \tilde{\boldsymbol{\theta}})h(\mathbf{Z} \mid \boldsymbol{\theta}^*)h(\mathbf{Y} \mid \boldsymbol{\theta})f(\boldsymbol{\theta}^*)}{h(\mathbf{Y} \mid \tilde{\boldsymbol{\theta}})h(\mathbf{Z} \mid \boldsymbol{\theta})h(\mathbf{Y}^* \mid \boldsymbol{\theta}^*)f(\boldsymbol{\theta})} \frac{c(\tilde{\boldsymbol{\theta}})}{c(\tilde{\boldsymbol{\theta}})} \frac{c(\boldsymbol{\theta})}{c(\boldsymbol{\theta}^*)} \frac{c(\boldsymbol{\theta}^*)}{c(\boldsymbol{\theta})}$$
$$= \frac{h(\mathbf{Y}^* \mid \tilde{\boldsymbol{\theta}})h(\mathbf{Z} \mid \boldsymbol{\theta}^*)h(\mathbf{Y} \mid \boldsymbol{\theta})f(\boldsymbol{\theta}^*)}{h(\mathbf{Y} \mid \tilde{\boldsymbol{\theta}})h(\mathbf{Z} \mid \boldsymbol{\theta})h(\mathbf{Y}^* \mid \boldsymbol{\theta}^*)f(\boldsymbol{\theta})},$$

where Y^* is the proposed auxiliary variable, $\theta^* = (\beta^{*\prime}, \eta^*)'$ is the proposed θ , and $\tilde{\theta}$ is the maximum pseudolikelihood estimate of θ . Thus $\log \alpha$ is

$$\log \alpha = Q(Y^* \mid \tilde{\boldsymbol{\theta}}) - Q(Y^* \mid \boldsymbol{\theta}^*)$$

$$+ Q(Z \mid \boldsymbol{\theta}^*) - Q(Z \mid \boldsymbol{\theta})$$

$$+ Q(Y \mid \boldsymbol{\theta}) - Q(Y \mid \tilde{\boldsymbol{\theta}})$$

$$+ \log f(\boldsymbol{\theta}^*) - \log f(\boldsymbol{\theta})$$

$$= Y^{*'}X(\tilde{\boldsymbol{\beta}} - \boldsymbol{\beta}^*) + \frac{\tilde{\eta} - \eta^*}{2}Y^{*'}AY^* + \eta^*Y^{*'}A\mu^* - \tilde{\eta}Y^{*'}A\tilde{\mu}$$

$$+ Z'X(\boldsymbol{\beta}^* - \boldsymbol{\beta}) + \frac{\eta^* - \eta}{2}Z'AZ + \eta Z'A\mu - \eta^*Z'A\mu^*$$

$$+ Y'X(\boldsymbol{\beta} - \tilde{\boldsymbol{\beta}}) + \frac{\eta - \tilde{\eta}}{2}Y'AY + \tilde{\eta}Y'A\tilde{\mu} - \eta Y'A\mu$$

$$+ \log f(\boldsymbol{\theta}^*) - \log f(\boldsymbol{\theta}).$$
(5)

Because the auxiliary proposals cannot be generated in parallel, this approach to Bayesian analysis is computationally expensive. Our optimized perfect sampler eases the burden somewhat, as does our Armadillo implementation of (5). We achieve an additional gain in efficiency as follows.

We use a normal random walk Metropolis-Hastings algorithm, and so our proposal for θ is (p+1)-variate normal, i.e., $\theta^{*(k+1)} \mid \theta^{*(k)} \sim \mathcal{N}_{p+1}(\theta^{*(k)}, \Sigma)$, for a suitably chosen covariance matrix Σ . Taking $\Sigma = \tau^2 \mathbf{I}$ tends to result in a low acceptance rate unless τ^2 is quite small, in which case a long run is required to adequately explore the posterior. Instead, we begin by using

$$\mathbf{\Sigma}_0 = egin{pmatrix} \mathbf{\hat{\Sigma}}_{GLM} & \mathbf{0} \\ \mathbf{0} & 0.01 \end{pmatrix}$$
 ,

where $\hat{\Sigma}_{GLM}$ is the estimated asymptotic covariance matrix for $\hat{\beta}$ obtained from fitting the ordinary logistic model to the data. We use Σ_0 for a training run (the number of iterations can be chosen by the user), and then we use the posterior sample covariance matrix from the training run as the proposal covariance matrix for a subsequent run. We use the latter sample to do inference. The training run usually results in a much better mixing chain, which reduces the total number of iterations. Still, rigorous Bayesian inference (following Møller et al.) is impractical for large lattices because perfect sampling does not scale well.

As for the prior $f(\theta)$, we assume that β and η are a priori independent, i.e., $f(\theta) = f_1(\beta)f_2(\eta)$. The prior for β is $\mathcal{N}(\mathbf{0}, \sigma^2\mathbf{I})$. The common standard deviation defaults to 1,000 but can be specified by the user via control parameter sigma. The prior for η is $\mathcal{U}(0, \eta^{\max})$. The right endpoint defaults to 2 but can be chosen by the user via control parameter eta.max.

Usage examples for the centered autologistic model

Now we present some usage examples for the autologistic model. The fitting function is called autologistic and has the following signature.

autologistic(formula, data, A, method =
$$c("PL", "Bayes")$$
, model = TRUE, x = FALSE, y = FALSE, verbose = FALSE, control = list())

Arguments formula, data, model, x, and y are analogous to those that appear in the signatures of other commonly used R model fitting functions, e.g., 1m and g1m. A is of course the binary adjacency matrix for G. The method argument is used to select pseudolikelihood inference or Bayesian inference.

If verbose is equal to TRUE, various messages may be sent to the standard output stream. Most of those messages have to do with the default values for control parameters that were not set by the user via the control argument.

For the following example the underlying graph is the 50×50 square lattice. First the corresponding adjacency matrix is constructed. Then the vertices are assigned coordinates restricted to the unit square centered at the origin. The vertex locations are used as spatial covariates, with regression coefficients $\boldsymbol{\beta}=(2,2)'$. The resulting independence expectations $\boldsymbol{\mu}=\mathbf{X}\boldsymbol{\beta}$, which range from approximately 0.1 to approximately 0.9, are level plotted in grayscale to show that the large-scale structure corresponds to a probability gradient that increases as one moves from southwest to northeast. Then, for $\eta=0.6$, a dataset is simulated and plotted. The dataset is then fitted using the PL method, and no confidence intervals are computed. Finally, the fit is summarized.

```
R> library("ngspatial")
R> library("lattice")
R > n < -50
R> A <- adjacency.matrix(n)</pre>
R > x < -rep(0:(n-1) / (n-1), times = n) - 0.5
R > y < -rep(0:(n-1) / (n-1), each = n) - 0.5
R > X < - cbind(x, y)
R > beta <- c(2, 2)
R> mu <- exp(X %*% beta)
R> mu <- mu / (1 + mu)
R> col1 <- "white"
R> col2 <- "black"
R> colfunc <- colorRampPalette(c(col1, col2))</pre>
R> dev.new()
R> levelplot(mu ~ x * y, aspect = "iso", col.regions = colfunc(n^2))
R> theta <- c(beta, eta = 0.6)
R> set.seed(123456)
R> Z <- rautologistic(X, A, theta)
R> dev.new()
R > levelplot(Z \sim x * y, aspect = "iso", col.regions = c("white", "black"),
             colorkey = FALSE)
R> fit <- autologistic(Z ~ X - 1, A = A, control = list(confint = "none"))
R> summary(fit)
Call:
autologistic(formula = Z ~ X - 1, A = A, control = list(confint = "none"))
Control parameters:
confint none
Coefficients:
    Estimate Lower Upper MCSE
Хx
      1.846 NA NA NA
       1.920
Χv
               NA
                      NA NA
       0.483 NA
eta
                    NA
                         NA
```

The next two examples compute confidence intervals for the same dataset. The first example uses the normal approximation. The "filling" in the sandwich matrix is estimated using a parallel parametric bootstrap, where the computation is distributed across six cores on the host workstation. The second example computes bootstrap confidence intervals, where the size of the bootstrap sample is 500. The computation is once again parallelized.

Control parameter 'bootit' must be a positive whole number. Setting it to the default

Number of iterations: 0

```
value of 1,000.
Control parameter 'parallel' must be a logical value. Setting it to the default value
Control parameter 'type' must be "SOCK", "PVM", "MPI", or "NWS". Setting it to "SOCK".
Warning: Bootstrapping may be time consuming.
Call:
autologistic(formula = Z ~ X - 1, A = A, verbose = TRUE,
   control = list(confint = "sandwich", nodes = 6))
Control parameters:
confint sandwich
bootit
        1000
parallel TRUE
type
        SOCK
nodes
Coefficients:
   Estimate Lower Upper MCSE
     1.846 1.4630 2.2300 NA
Хx
      1.920 1.5410 2.2990 NA
Xγ
      0.483 0.3619 0.6042 NA
eta
Number of iterations: 1000
R> set.seed(123456)
R> fit <- autologistic(Z \sim X - 1, A = A, verbose = TRUE,
                      control = list(confint = "bootstrap", bootit = 500, nodes = 6))
R> summary(fit)
Control parameter 'parallel' must be a logical value. Setting it to the default value
of TRUE.
Control parameter 'type' must be "SOCK", "PVM", "MPI", or "NWS". Setting it to "SOCK".
Warning: Bootstrapping may be time consuming.
Call:
autologistic(formula = Z \sim X - 1, A = A, verbose = TRUE,
   control = list(confint = "bootstrap", bootit = 500, nodes = 6))
Control parameters:
confint bootstrap
bootit
        500
parallel TRUE
         SOCK
type
nodes
Coefficients:
   Estimate Lower Upper
     1.846 1.4980 2.2510 0.007607
Хx
      1.920 1.5400 2.3270 0.007098
Xγ
      0.483 0.3595 0.6063 0.003406
eta
Number of iterations: 500
```

It took several minutes to run each of the two preceding examples on a Mac Pro with two quad-core 2.8 GHz Intel Xeon CPUs. Since Bayesian inference would be prohibitively expensive for a dataset so large, the next example was run on a dataset for which G was the 20×20 square lattice.

We use, and recommend, fixed-width output analysis (Flegal et al., 2008). In fixed-width analysis, one chooses a tolerance and stops sampling when all Monte Carlo standard errors fall below the tolerance. The output shown below indicates that the Monte Carlo standard errors fell below the default tolerance of 0.01 after 714,025 draws were made from the posterior, which is why the analysis was terminated before having reached the (default) maximum number of iterations (maxit = 1e6). We use the batchmeans package (Haran and Hughes, 2012) to compute Monte Carlo standard errors (denoted MCSE in the output).

In this example we use a training run of length 10,000, and the length of the subsequent inferential run will be at least 10,000.

```
R> n <- 20
R> A <- adjacency.matrix(n)</pre>
R> x \leftarrow rep(0:(n-1) / (n-1), times = n) - 0.5
R> y \leftarrow rep(0:(n-1) / (n-1), each = n) - 0.5
R> X <- cbind(x, y)
R> set.seed(123456)
R> Z <- rautologistic(X, A, theta)
R> fit <- autologistic(Z ~ X - 1, A = A, verbose = TRUE, method = "Bayes",
                        control = list(trainit = 10000, minit = 10000))
R> summary(fit)
Control parameter 'tol' must be a positive number. Setting it to the default value
of 0.01.
Control parameter 'maxit' must be a positive whole number greater than or equal to
'minit'. Setting it to 1e+06.
Control parameter 'sigma' must be a positive number. Setting it to the default value
of 1,000.
Control parameter 'eta.max' must be a positive number. Setting it to the default
value of 2.
Warning: MCMC may be time consuming.
Progress => 5%
Progress => 10%
Progress => 15%
Call:
autologistic(formula = Z ~ X - 1, A = A, method = "Bayes", verbose = TRUE,
    control = list(trainit = 10000, minit = 10000))
Control parameters:
trainit 1e+04
       1e-02
tol
        1e+04
minit
       1e+06
maxit
        1e+03
sigma
eta.max 2e+00
Coefficients:
    Estimate Lower Upper
      2.4450 1.2120 3.667 0.009917
Xx
      1.6800 0.4767 2.832 0.009800
Ху
      0.7363 0.4805 1.028 0.002631
Number of iterations: 714025
```

The object returned by function autologistic is a list with many useful elements. The following output resulted from applying the names function to the object returned in the second example above.

R> names(fit)

```
[1] "coefficients" "fitted.values" "linear.predictors" "residuals" "convergence"
[6] "message" "value" "iter" "sample" "mcse"
[11] "xlevels" "call" "terms" "method" "control"
[16] "model"
```

The standard coef and fitted functions can be applied to an object of type 'autologisticThe residuals method for 'autologistic' objects can be used to extract deviance (default), Pearson, or response residuals.

Other fields of interest are value, which contains $-\ell_{PL}(\tilde{\theta})$, and sample, which contains the bootstrap or posterior sample.

Fitting the sparse SGLMM

The fitting function for the sparse SGLMM is called sparse.sglmm. The function returns an object of class "sparse.sglmm". Auxiliary functions residuals, summary, and vcov accept an object of type "sparse.sglmm" and return various kinds of residuals, summarize the fit, and return an estimated covariance matrix, respectively.

Bayesian inference for the sparse SGLMM

The second stage of the sparse SGLMM, i.e., the prior for δ_s , is

$$f(\boldsymbol{\delta}_{s} \mid \tau_{s}) \propto \tau_{s}^{q/2} \exp\left(-\frac{\tau_{s}}{2}\boldsymbol{\delta}_{s}'\mathbf{M}'\mathbf{Q}\mathbf{M}\boldsymbol{\delta}_{s}\right)$$
,

where τ_s is a smoothing parameter. The prior for β is spherical p-variate normal with mean zero and common variance sigma.b, which defaults to 1,000 but can be controlled by the user via argument hyper. The prior for τ_s is gamma with parameters 0.5 and 2,000 (Kelsall and Wakefield, 1999). This prior is attractive because it corresponds to the prior belief that the fixed effects are sufficient to explain the data (since a large value for τ_s implies small variances for the random effects) and because it discourages artifactual spatial structure in the posterior.

When the response is normally distributed, the identity link is assumed, in which case the first stage is

$$\mu = \mathbf{X}\boldsymbol{\beta} + \mathbf{M}\boldsymbol{\delta}_{\scriptscriptstyle S} + \mathbf{M}\boldsymbol{\delta}_{\scriptscriptstyle h},$$

where δ_h are heterogeneity random effects. When the response is Poisson distributed, heterogeneity random effects are optional. In any case, the prior on δ_h is spherical q-variate normal with mean zero and common variance $1/\tau_h$. The prior for τ_h is gamma with parameters a.h and b.h. The default values for these parameters are 0.01 and 100, respectively, or their values can be set by the user through argument hyper.

If the response is Bernoulli or Poisson, β and δ_s are updated using Metropolis-Hastings random walks with normal proposals. The proposal covariance matrix for β is the estimated asymptotic covariance matrix from a glm fit to the data. The proposal for δ_s is spherical normal with common standard deviation sigma.s. If the response is Poisson distributed and heterogeneity random effects are included, those random effects are updated using a Metropolis-Hastings random walk with a spherical normal proposal. The common standard deviation is sigma.h. Both sigma.s and sigma.h default to 0.01, but they can be set via argument tune.

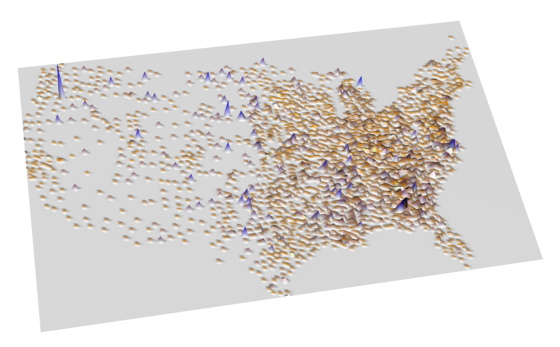


Figure 2: The empirical infant mortality rates.

The updates for τ_s and τ_h are Gibbs updates irrespective of the response distribution. If the response is normally distributed, all updates are Gibbs updates.

Usage examples for the sparse SGLMM

Function sparse. sglmm has the following signature.

```
sparse.sglmm(formula, family = gaussian, data, offset, A, attractive = 50,
repulsive = 0, tol = 0.01, minit = 10000, maxit = 1e+06, tune = list(),
hyper = list(), model = TRUE, x = FALSE, y = FALSE, verbose = FALSE)
```

Most of the arguments are analogous to those already described above. Arguments attractive and repulsive are used to select the number of Moran eigenvectors. Both attractive and repulsive eigenvectors are permitted, although repulsive defaults to 0, which corresponds to pure spatial smoothing. Function sparse.sglmm checks the validity of attractive and repulsive by eigendecomposing the Moran operator and examining the spectrum. Execution is terminated (with an informative error message) if either value is invalid. Arguments tune and hyper can be used to supply values for tuning parameters and hyperparameters, respectively, as described above.

We will illustrate the usage of our sparse SGLMM function by analyzing the data shown in Figure 2. The plot shows infant mortality data for 3,071 US counties. Each shaded peak represents a ratio of deaths to births, i.e., an empirical infant mortality rate, for a given county. The data were obtained from the 2008 Area Resource File (ARF), a county-level database maintained by the Bureau of Health Professions, Health Resources and Services Administration, US Department of Health and Human Services. Specifically, three variables were extracted from the ARF: the three-year (2002–2004) average number of infant deaths before the first birthday, the three-year average number of live births, and the three-year average number of low birth weight infants.

To these data we fit the sparse Poisson SGLMM with

```
\log \mathbb{E}(\text{DEATHS}_i \mid \boldsymbol{\beta}, \boldsymbol{\delta}_s) = \log \text{BIRTHS}_i + \beta_0 + \beta_1 \text{LOW}_i + \beta_2 \text{BLACK}_i + \beta_3 \text{HISP}_i + \beta_4 \text{GINI}_i + \beta_5 \text{AFF}_i + \beta_6 \text{STAB}_i + \boldsymbol{m}_i' \boldsymbol{\delta}_s,
```

where DEATHS is the number of infant deaths; BIRTHS is the number of live births; LOW is the rate of low birth weight; BLACK is the percentage of black residents (according to the 2000 US Census); HISP is the percentage of hispanic residents (2000 US Census); GINI is the Gini coefficient, a measure of income inequality (Gini, 1921); AFF is a composite score of social affluence (Yang et al., 2009); and STAB is residential stability, an average *z*-score of two variables from the 2000 US Census.

These data were analyzed previously in Hughes and Haran (2013), where it was found that the model with 50 attractive eigenvectors yields the smallest value of DIC among the sequence of models

with 25, 50, 100, and 200 attractive eigenvectors. The following code duplicates the analysis of Hughes and Haran (2013).

```
R> data(infant)
R> infant$low_weight <- infant$low_weight / infant$births
R> attach(infant)
R> Z <- deaths
R> X <- cbind(1, low_weight, black, hispanic, gini, affluence, stability)
R> data(A)
R> set.seed(123456)
R> fit <- sparse.sglmm(Z \sim X - 1 + offset(log(births)), family = poisson, A = A,
                       tune = list(sigma.s = 0.02), verbose = TRUE)
R> summary(fit)
Hyperparameter 'sigma.b' must be a positive number. Setting it to the default
value of 1,000.
The Moran operator is being eigendecomposed. This operation may be time consuming.
Warning: MCMC may be time consuming.
Progress => 5%
Progress => 100%
Call:
sparse.sglmm(formula = Z \sim X - 1 + offset(log(births)), family = poisson,
   A = A, tune = list(sigma.s = 0.02), verbose = TRUE)
Tuning parameters:
sigma.s 0.02
Hyperparameters:
sigma.b 1000
Coefficients:
            Estimate
                        Lower
                                   Upper
            -5.435000 -5.621000 -5.250000 9.466e-04
Xlow_weight 8.821000 7.552000 10.050000 5.673e-03
Xblack
           0.004184 0.002864 0.005545 6.568e-06
Xhispanic -0.003792 -0.004870 -0.002678 6.342e-06
           -0.553200 -0.984300 -0.126000 2.070e-03
Xgini
Xaffluence -0.075600 -0.087760 -0.063740 5.148e-05
Xstability -0.028670 -0.043350 -0.013980 6.360e-05
DIC: 10110
Number of iterations: 1000000
```

The analysis took just under two hours. Fitting the RHZ model to these data took approximately two weeks and required over 40 GB of secondary storage.

We also used **CARBayes** to apply the ICAR model to these data. The results for the two fits are shown together in Table 1. We see that the ICAR fit suffers from spatial confounding. Specifically, some of the ICAR point estimates differ markedly from the sparse SGLMM estimates, the ICAR intervals are wider, and the Gini coefficient is not a significant predictor in the ICAR fit.

The object returned by function sparse.sglmm is a list. The following output resulted from applying the names function to the object produced in the example above.

	Sparse SGLMM		ICAR	
Covariate	Estimate	95% HPD Interval	Estimate	95% HPD Interval
Intercept	-5.435	(-5.621, -5.250)	-5.558	(-5.766, -5.354)
Low Weight	8.821	(7.552, 10.050)	7.754	(6.396, 9.112)
Black	0.004	(0.003, 0.006)	0.004	(0.002, 0.006)
Hispanic	-0.004	(-0.005, -0.003)	-0.003	(-0.005, -0.002)
Gini	-0.553	(-0.984, -0.126)	-0.079	(-0.554, 0.406)
Affluence	-0.076	(-0.088, -0.064)	-0.077	(-0.091, -0.063)
Stability	-0.029	(-0.043, -0.014)	-0.042	(-0.060 - 0.024)

Table 1: Results for sparse SGLMM and ICAR fits to the infant mortality data.

R> names(fit)

```
[1] "coefficients" "fitted.values" "linear.predictors" "residuals"
                                                                      "beta.sample"
[6] "gamma.sample"
                    "tau.s.sample"
                                    "beta.mcse"
                                                         "gamma.mcse"
                                                                      "tau.s.mcse"
                                                         "D.bar"
[11] "gamma.est"
                    "tau.s.est"
                                    "iter"
                                                                      "DD"
[16] "dic"
                                                                      "call"
                    "beta.accept"
                                     "gamma.accept"
                                                         "xlevels"
[21] "terms"
                    "formula"
                                    "family"
                                                         "offset"
                                                                      "model"
[26] "tune"
                    "hyper"
```

The standard coef and fitted functions can be applied to an object of type 'sparse.sglmm'. The residuals method for 'sparse.sglmm' objects can be used to extract deviance (default), Pearson, or response residuals.

Other particularly useful fields include beta.sample, gamma.sample, and tau.s.sample, which contain the posterior samples for β , δ_s , and τ_s , respectively. (The package uses γ and δ , respectively, in place of δ_s and δ_h . This paper has used δ_s and δ_h because here it was necessary to present the RHZ model, which is not discussed in the package documentation.) Monte Carlo standard errors are also returned, as are acceptance rates.

Conclusion

This article introduced version 1.0 of R package **ngspatial**, which supports two promising new models for areal data, namely, the centered autologistic model and the sparse SGLMM. The package is user friendly because its model-fitting functions and auxiliary functions were designed to behave like the analogous functions (e.g., 1m, g1m, summary) in the **stats** package. The package is also efficient because the code uses vector and matrix operations where possible, and because key portions of the code were written in C++.

Acknowledgments

This work was supported by a University of Minnesota Grant-in-Aid of Research, Artistry, and Scholarship.

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