

A Fast and Exact Simulation Algorithm for General Gaussian Markov Random Fields

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SUMMARY

This paper presents a fast and exact simulation algorithm for a general Gaussian Markov Random Field (GMRF) defined on a $n_r \times n_c$ lattice, $n_r \geq n_c$. For a 5×5 neighborhood, the algorithm has initialization cost of $4n_r n_c^3$ flops and exact samples are then available using $4n_r n_c^2$ flops. Conditioned samples on k general constraints costs only $3n_r n_c k$ extra flops. The algorithm is easy to generalize to non-rectangular lattices, compute the log-likelihood of the sample nearly for free and runs very efficient on the computer.

An exact and fast simulation algorithm for unconditional and conditional GMRF additional to easy access to the log-likelihood, makes GMRF (even more) computational convenient. GMRF has a potential use when modeling non-stationary spatial fields and is commonly used in hierarchical spatial or spatio-temporal Bayesian models analyzed using Markov chain Monte Carlo.

KEYWORDS: Conditional Autoregressive models, Gaussian Markov random field, Markov chain Monte Carlo.

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1 INTRODUCTION

This paper presents a fast and exact algorithm for simulating a general Gaussian random fields with a Markov property (GMRF), also known as conditional auto-regressions. Besag & Kooperberg (1995) give references to GMRF in various applications. The algorithm depends only on the conditional independence structure induced by the Markov property and no assumptions (apart from positive definiteness) are required on the interaction terms. On a $n_r \times n_c$ lattice, $n_r \geq n_c$, and a 5×5 neighborhood, the algorithm has initialization cost of $4n_r n_c^3$ flops and then exact samples are available using only $4n_r n_c^2$ flops. Conditioning on k observed variables or linear combinations of variables, only costs $3n_r n_c k$ extra flops. The simulation algorithm can be extended to GMRF defined on non-rectangular lattices with essentially the same computational cost. Further, the likelihood can be computed for the generated sample nearly for free.

The presented simulation algorithm makes general GMRF even more computational convenient in important applications. Hierarchical spatial and spatio-temporal Bayesian models often require simulation based inference using Markov chain Monte Carlo (MCMC) and a GMRF is a natural choice for one layer in the model due to its Markov properties, see for example Wikle, Berliner & Cressie (1998). A fast and exact sampling algorithm for general GMRF will be useful as it may allow for fast and large block updates of the GMRF in the MCMC algorithm. Further, the inclusion of external covariates into spatial and spatio-temporal Gaussian models through the covariance function is troublesome. Using hierarchical modeling and GMRFs, covariates can be introduced directly into the conditional densities while preserving its Markov property. The presented simulation algorithm can then be used to produce exact unconditional and conditional samples. Since the normalization constant is readily available and the simulation algorithm is general, GMRF is a natural candidate to model non-stationary spatial fields, a topic yet to be explored.

The plan of the paper is as follows. After some preliminary definitions in Section 2, we present our algorithm in Section 3 and discuss generalizations in Section 4. We present an example in Section 5 and conclude in Section 6.

2 PRELIMINARIES

Define the Gaussian random field (GRF) $\boldsymbol{x} = \{x_{ij}, (i, j) \in \Lambda\}$ on the $n_r \times n_c$ lattice Λ , $n_r \geq n_c$. Denote by $\pi(\boldsymbol{x})$ the Gaussian joint density of \boldsymbol{x} and assume \boldsymbol{x} has zero mean. The GRF has a Markov property iff

$$\pi(x_{ij} \mid x_{kl}, (k, l) \in \Lambda \setminus \{ij\}) = \pi(x_{ij} \mid x_{kl}, (k, l) \in \partial_{ij})$$

where ∂_{ij} is the neighborhood of (i, j) . A GRF with a Markov property is a Gaussian Markov random field (GMRF). Typical, ∂_{ij} is a set of neighboring sites, for example

$$\partial_{ij} = \{(k, l) \neq (i, j) : \|(k, l) - (i, j)\| \leq d\}$$

where $\|\cdot\|$ is some norm. Common neighborhoods in the nearest (in Euclidean norm) have 4, 8 and 12 sites, or the nearest 24 using the maximum-norm. We are mainly interested in the neighborhood defined as the 5×5 window centered at each site, but assume a general $(2m_r + 1) \times (2m_c + 1)$ neighborhood throughout the paper.

Let \mathbf{Q} be the $n_r n_c \times n_r n_c$ precision matrix (the inverse covariance matrix) for \mathbf{x} . A Markov property of a GRF induce a band-structure in the precision matrix, as follows:

$$x_{ij} \perp x_{kl} \mid \mathbf{x} \setminus_{ij,kl} \iff Q_{ij,kl} = 0, \quad (1)$$

meaning that x_{ij} and x_{kl} are conditional independent given the others iff the corresponding element in \mathbf{Q} is zero. Thus for a 5×5 neighborhood system, (1) implies that only 25 elements in $Q_{ij,kl}$ are non-zero for each ij , and these corresponds to x_{ij} , $x_{i+1,j}$, $x_{i+2,j}$, $x_{i,j-1}$ and $x_{i,j-2}$ and so on. Appropriate changes at the boundary is needed. Hence, \mathbf{Q} is a sparse block pentadiagonal matrix or a band matrix as shown in Figure 1 for a 15×15 lattice and a 5×5 neighborhood. We will from now treat \mathbf{Q} as a band matrix with bandwidth $b_w = \min\{m_c n_r + m_r, m_r n_c + m_c\}$, where we chose column-wise or row-wise storage scheme to minimize the bandwidth. We will use the band matrix interpretation because the numerical algorithms needed in the simulation algorithms is designed for band matrices.

3 THE ALGORITHM

Based on the preliminaries in Section 2 we can now derive our exact simulation algorithm. We first start by noting that computing the Cholesky factorization of the band matrix \mathbf{Q} (with bandwidth b_w)

$$\mathbf{Q} = \mathbf{L}\mathbf{L}^T \quad (2)$$

is a standard problem in numerics. By Theorem 4.3.1 in Golub & van Loan (1996), \mathbf{L} has lower bandwidth b_w (and is zero above the diagonal) and can be computed in $n_r n_c b_w^2$ flops, see Golub & van Loan (1996, Sec. 4.3.5) for details and algorithm. Note that \mathbf{L} require $n_r n_c b_w U$ bytes of storage where U is the number of bytes needed to store one float. In the next step we let \mathbf{z} be a vector of independent standard Gaussian variates, and note by direct calculation that \mathbf{x} defined by

$$\mathbf{L}^T \mathbf{x} = \mathbf{z} \quad (3)$$

has mean zero and the desired covariance matrix \mathbf{Q}^{-1} . Again, solving (3) is a standard problem in numerics and the solution can be computed in $2n_r n_c b_w$ flops using band back-substitution, see again Golub & van Loan (1996, Sec. 4.3.2) for details and algorithm.

We need to compute the band Cholesky factorization in (2) only once, and repeated samples are then available by solving (3). There are no assumptions of the coefficients in the \mathbf{Q} -matrix (apart from being positive definite), so the algorithm is general.

To study to computational costs, assume for simplicity that $m_r = m_c$, then the cost of computing the band Cholesky decomposition (2) is $n_r n_c^3 m_c^2$, and solving (3) is $2n_r n_c^2 m_c$. The storage needed is

$n_r n_c^2 m_c U$ bytes. Note that these figures are linear in the *largest* dimension n_r , and cubic/quadratic in the *smallest* dimension. Hence, it is far more efficient to sample from rectangular compared to square lattices.

To implement the algorithm, we make use of high-quality routines in the public-domain Lapack-library written in Fortran (Anderson et al. 1995). The appropriate routines are `dpbtf2` (level 2) or `dpbtrf` (level 3) to compute the band Cholesky decomposition (2), and `dtbsv` to do band back-substitution (3). A software-package for these tasks are available from the author.

I agree that the algorithm seems rather trivial, but (unfortunately) sometimes solutions to problems are!

4 GENERALIZATIONS OF THE ALGORITHM

I will now discuss some generalizations of the simulation algorithm to conditional simulation, simulation on a non-rectangular lattice, and the computation of the log-likelihood.

4.1 CONDITIONAL SIMULATION

I consider first consider the general case where we want to generate samples from π conditioned on $\mathbf{Ax} + \epsilon = \mathbf{b}$ where \mathbf{A} is a $k \times n$ matrix with rank k . Then I will discuss conditional sampling of a rectangular window $W \subset \Lambda$ where we condition on the boundary.

4.1.1 GENERAL CONDITIONING

Let the Gaussian noise ϵ has zero mean and known covariance matrix Σ_ϵ , and define $\tilde{\mathbf{x}} = (\tilde{\mathbf{x}}_1^T, \tilde{\mathbf{x}}_2^T)^T = (\mathbf{x}^T, (\mathbf{Ax} + \epsilon)^T)^T$ where $\tilde{\mathbf{x}}$ have dimension $n + k$. Let $\tilde{\Sigma}$ denote the covariance matrix of $\tilde{\mathbf{x}}$. It is easy to simulate $\tilde{\mathbf{x}}$ unconditionally by sampling \mathbf{x} and ϵ , and then use the definition $\tilde{\mathbf{x}} = (\mathbf{x}^T, (\mathbf{Ax} + \epsilon)^T)^T$. A conditional sample of $\tilde{\mathbf{x}}_1 \mid \tilde{\mathbf{x}}_2 = \mathbf{b}$ can be obtained using the following identity

$$\tilde{\mathbf{x}}_1 \mid \tilde{\mathbf{x}}_2 = \mathbf{b} \stackrel{d}{=} \tilde{\mathbf{x}}_1 - \tilde{\Sigma}_{12} \tilde{\Sigma}_{22}^{-1} (\tilde{\mathbf{x}}_2 - \mathbf{b}) \quad (4)$$

where $\tilde{\mathbf{x}}$ on the right hand side is an unconditional sample. A direct calculation shows that (4) is correct. Eq. (4) is commonly referred to as conditioning using Kriging, see Cressie (1993, Sec. 3.6.2). By expanding the right hand side of (4), we obtain

$$\mathbf{x} - \mathbf{Q}^{-1} \mathbf{A}^T (\mathbf{AQ}^{-1} \mathbf{A}^T + \Sigma_\epsilon)^{-1} (\mathbf{Ax} + \epsilon - \mathbf{b}). \quad (5)$$

We need to compute $\mathbf{v} = \mathbf{Q}^{-1} \mathbf{A}^T$, which requires solving $\mathbf{Q}\mathbf{v}_i = (\mathbf{A}^T)_i$ for each of the k columns. Since the band Cholesky decomposition is already available as $\mathbf{Q} = \mathbf{L}\mathbf{L}^T$, we solve $\mathbf{Q}\mathbf{v}_i = (\mathbf{A}^T)_i$ by a forward substitution $\mathbf{L}\mathbf{u}_i = (\mathbf{A}^T)_i$ and a back-substitution $\mathbf{L}^T \mathbf{v}_i = \mathbf{u}_i$. The Lapack routine `dtbsv` do both forward and back-substitutions.

The remaining parts of (5) are now straight forward. To save work, we compute \mathbf{v} and then $\mathbf{A}\mathbf{v}$ (using $4kn_r n_c b_w$ flops) and invert the symmetric $k \times k$ matrix (using k^3 flops) only once. For each sample we need to compute $\mathbf{A}\mathbf{x} + \boldsymbol{\epsilon} - \mathbf{b}$, pre-multiply with the matrix found in the initialization, negate and add \mathbf{x} (in total $3n_r n_c k$ flops).

4.1.2 CONDITIONING ON THE BOUNDARY

Let W be a rectangular window of size $w_r \times w_c$, where $W \subset \Lambda$. The task is to sample \mathbf{x}_W condition on the rest $\mathbf{x}_{\Lambda-W}$. As the GMRF has a Markov property we strictly need to condition only on the sites in the boundary of W , $\mathbf{x}_{\partial W}$, where ∂W is the set of all sites not in W which has a neighbor in W . For notational simplicity, we condition on $B = \Lambda - W$ and it will soon become clear where we can simplify due to ∂W .

Partition as usual the precision matrix into blocks corresponding to W and B ,

$$\mathbf{Q} = \text{Cov}((\mathbf{x}_W^T, \mathbf{x}_B^T)^T)^{-1} = \begin{bmatrix} \mathbf{Q}_{WW} & \mathbf{Q}_{WB} \\ \mathbf{Q}_{BW} & \mathbf{Q}_{BB} \end{bmatrix}. \quad (6)$$

By expressing the conditional mean and precision for $\mathbf{x}_W \mid \mathbf{x}_B$ using the blocks of \mathbf{Q} instead of the similar blocks of the covariance matrix $\boldsymbol{\Sigma}$, we obtain the conditional mean $\boldsymbol{\mu}_{W|B} = \mathbf{Q}_{WW}^{-1} \mathbf{Q}_{WB} \mathbf{x}_B$ and conditional precision $\mathbf{Q}_{W|B} = \mathbf{Q}_{WW}$. Although this is a standard result, it is surprisingly little known.

Since W is a rectangular subset of Λ , \mathbf{Q}_{WW} has the same band structure as \mathbf{Q} hence the algorithm in Section 2 can be used to sample from the conditional density (with zero mean) by computing the band Cholesky decomposition $\mathbf{Q}_{WW} = \mathbf{L}_W \mathbf{L}_W^T$ and solving $\mathbf{L}_W^T \mathbf{y} = \mathbf{z}$. However, we need also to compute the conditional mean, by solving

$$\mathbf{Q}_{WW} \boldsymbol{\mu}_{W|B} = \mathbf{Q}_{WB} \mathbf{x}_B. \quad (7)$$

Since $\mathbf{Q}_{WW} = \mathbf{L}_W \mathbf{L}_W^T$ is already available, we solve first by forward substitution $\mathbf{L}_W \mathbf{v} = \mathbf{Q}_{WB} \mathbf{x}_B$ and then by back substitution $\mathbf{L}_W^T \boldsymbol{\mu}_{W|B} = \mathbf{v}$. Finally, $\mathbf{x}_{W|B} = \boldsymbol{\mu}_{W|B} + \mathbf{y}$.

The dependency of ∂B is most easily incorporated while computing $\mathbf{Q}_{WB} \mathbf{x}_B$, since

$$(\mathbf{Q}_{WB} \mathbf{x}_B)_{ij} = - \sum_{kl \in \partial_{ij} \cap B} Q_{ij,kl} x_{kl}, \quad ij \in W. \quad (8)$$

The computational cost (assuming $m_r = m_c$ and $w_r \geq w_c$) is $w_r w_c^3 m_c^2$ flops for computing the band Cholesky decomposition and $6w_r w_c^2 m_c$ flops for solving the three band linear systems needed. We need to compute the band Cholesky decomposition once only if more than one conditional sample is needed.

4.2 NON-RECTANGULAR LATTICE

I will now relax the assumption that \mathbf{x} is defined on a rectangular lattice. Suppose the region of interest is a non-rectangular lattice Λ . The precision matrix will not have the required band-structure required for the fast simulation algorithm. However, define Λ' as the smallest rectangular lattice covering Λ and extend \mathbf{x} to \mathbf{x}' by adding independent standard Gaussian variates for each site in $\Lambda' - \Lambda$, then the precision matrix for \mathbf{x}' will have the required band structure. Further, \mathbf{x}' restricted to Λ have the desired marginal density. The conditioning discussed in Section 4.1.1 and Section 4.1.2 can both easily be generalized along these lines, also to include the case where W is not a rectangular subset of Λ .

4.3 LIKELIHOOD

Likelihood based inference and testing, computation of acceptance rates in MCMC from GMRF, all depends on the log-likelihood which is easily available for general GMRF. First, if we simulate a sample \mathbf{x} , then note that $\mathbf{x}^T \mathbf{Q} \mathbf{x}$ equals $\mathbf{z}^T \mathbf{z}$, and $|\mathbf{Q}|^{1/2}$ equals $\prod_i L_{ii}$. Hence, the log-likelihood for the sample $L^T \mathbf{x} = \mathbf{z}$ becomes

$$\text{log-likelihood}(\mathbf{x}) = -\frac{n}{2} \log 2\pi + \sum_i \log(L_{ii}) - \frac{1}{2} \mathbf{z}^T \mathbf{z}$$

and the computation of the log-likelihood does not require (nearly) any extra computational costs compared to sampling \mathbf{x} . If \mathbf{x} is given, we need to compute the alternative expression $\mathbf{x}^T \mathbf{Q} \mathbf{x}$. Using the argument as in Section 4.2, we can compute the log-likelihood for a general lattice. The computation of the log-likelihood for the conditional sample in Section 4.1.2 is similar.

5 AN EXAMPLE

To give an idea of the CPU costs for the algorithm, I will present results for my current implementation of the algorithm on a Sun Ultra 2 Model 1296 with a 296 MHz SUNW UltraSPARC-II processor and a 100×100 lattice. Since the sampling algorithm is linear in the largest dimension, the CPU will be doubled for a 200×100 lattice and so on.

With a 3×3 neighborhood the algorithm used 0.92 seconds for the first sample and then produced iid samples using 0.07 seconds each. (All timings include the evaluation of the log-likelihood for the sample.) When I increased the neighborhood to 5×5 , the algorithm used 2.80 and 0.11 seconds, and for a 7×7 neighborhood 6.00 and 0.15 seconds.

Then I added 100 linear conditions requiring the sum of each of the 100 columns is zero. The first evaluation of the conditioning took 14.3, 19.9 and 25.8 seconds for 3×3 , 5×5 and 7×7 neighborhoods, respectively. Then the following samples used only 0.08 seconds in the conditioning, for all three neighborhoods.

Figure 2 shows an unconditioned sample using a 5×5 neighborhood with coefficients chosen to approximate a Gaussian field with correlation function $\exp(-3d^2/20^2)$, where d is the Euclidean distance in pixels. The GMRF fits the Gaussian field surprisingly well and demonstrates that GMRF is indeed appropriate as a computational attractive approximate to Gaussian fields, refer to Rue & Tjelmeland (1999) for details and discussion along these lines.

My current implementation makes about 140 and 37 Megaflops on a 5×5 neighborhood, computing (mainly) the Cholesky-decomposition and solving the band linear system, respectively. To compare the numbers, the plain loop $y_i = y_i + \beta x_i$, makes on the same computer 180 and 195 Megaflops in C and Fortran compiled with SUN compilers using options `-fast -xO5`, respectively.

6 DISCUSSION

We have in this paper demonstrated that Gaussian Markov random fields can be sampled exactly using $\mathcal{O}(n_r n_c^2)$ flops with an initial costs $\mathcal{O}(n_r n_c^3)$, and that a very efficient implementation of the sampler is possible using the `Lapack`-library. Further, the log-likelihood for the sample is easily available, the algorithm is trivial to extend to non-rectangular lattices, we can obtain general conditional samples efficiently using Kriging, and easily compute samples conditioned on the boundary.

The example in Figure 2 shows that a GMRF is indeed capable of approximating a Gaussian fields with (relative) long correlation length, see Rue & Tjelmeland (1999) for further details.

The only negative aspect of the algorithm is the memory-requirement of $\mathcal{O}(n_r n_c^2)$. This prohibits the use of the algorithm for large lattices. For a 256×256 lattice, the algorithm require about 268 MBytes of memory (using 8 Bytes to store a float), still feasible for my computer, but a 512×512 lattice requires about 2.1 GBytes which is too much. Rue, Marthinsen & Husby (1999) solves this problem by developing a split and conquer algorithm which runs with near optimal speedup on parallel computers, where the memory requirement is $\mathcal{O}(n_r n_c)$ and the computational costs is about $\mathcal{O}(n_r^{3.6})$ when $n_c = n_r$. However, their algorithm is intended for large fields and runs much slower for problems where the current algorithm is feasible, see Rue et al. (1999) for details.

I end this paper by pointing out that Lavine (1998) also reports an $\mathcal{O}(n_r n_c^3)$ (unconditional) simulation algorithm for GMRF with 4 nearest neighbors and invariant interactions (the elements in the \mathbf{Q} -matrix) by running a special Bayesian dynamic model. However, for his case we can do even better by making use of the fast block-tridiagonal solver of Swarztrauber (1974) and derive (essentially) an $\mathcal{O}(n_r n_c \log(n_r n_c))$ using the split and conquer algorithm in Rue et al. (1999). However, algorithms for special cases in terms of values of the interactions are usually of less interest compared to a general algorithm.

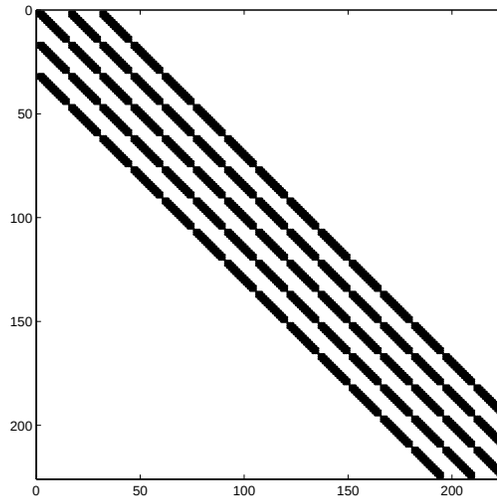


FIGURE 1: The structure in the precision matrix for a 15×15 lattice and a 5×5 neighborhood.

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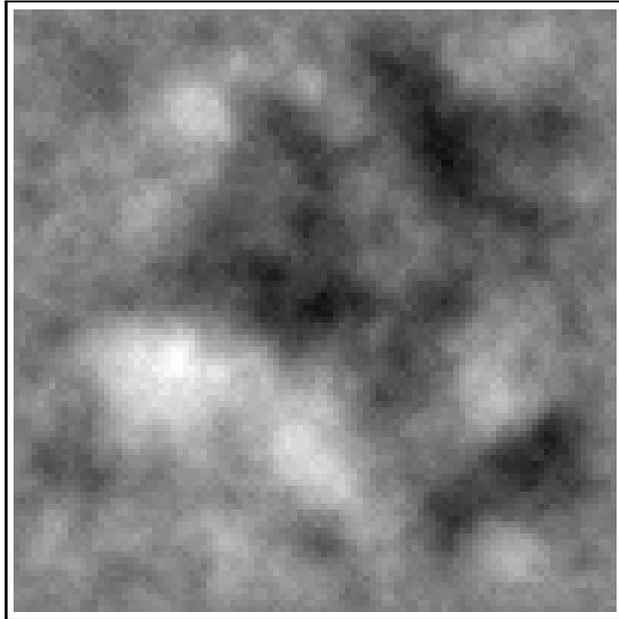


FIGURE 2: An exact sample from a 100×100 GMRF with a 5×5 neighborhood, where the coefficients are chosen to fit a Gaussian field with correlation function $\exp(-3d^2/20^2)$.