Sampling Strategies for Fast Updating of Gaussian Markov Random Fields

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Abstract

Gaussian Markov random fields (GMRFs) are popular for modeling dependence in large areal datasets due to their ease of interpretation and computational convenience afforded by the sparse precision matrices needed for random variable generation. Typically in Bayesian computation, GMRFs are updated jointly in a block Gibbs sampler or componentwise in a single-site sampler via the full conditional distributions. The former approach can speed convergence by updating correlated variables all at once, while the latter avoids solving large matrices. We consider a sampling approach in which the underlying graph can be cut so that conditionally independent sites are updated simultaneously. This algorithm allows a practitioner to parallelize updates of subsets of locations or to take advantage of ‘vectorized’ calculations in a high-level language such as R. Through both simulated and real data, we demonstrate computational savings that can be achieved versus both single-site and block updating, regardless of whether the data are on a regular or an irregular lattice. The approach provides a good compromise between statistical and computational efficiency and is
accessible to statisticians without expertise in numerical analysis or advanced computing.

*Keywords:* Bayesian computation, Cholesky factorization, chromatic Gibbs sampling, conditional autoregressive model, graph coloring, Markov chain Monte Carlo
1 INTRODUCTION

Suppose we have observed data $\mathbf{y} = (y_1, \ldots, y_n)^T$ in which each $y_i$ summarizes information over an area $i$, $i = 1, \ldots, n$, such as a sum or average of individuals in the area. For instance, Self et al. (2018) investigate regional trends of occurrence of Lyme disease, where the data are the number of positive disease cases observed in each county in the United States. Other examples include Brown et al. (2014), who consider functional magnetic resonance imaging data in which each $y_i$ quantifies the neuronal changes associated with an experiment observed in the $i^{th}$ three-dimensional pixel in a brain image, where the goal is to identify those areas exhibiting statistically significant changes. Waller et al. (1997) estimate spatially-varying risks of developing lung cancer using reported deaths in each county of the state of Ohio. In the examples we consider in this work, $y_i$ is either the observed intensity at pixel $i$ in an image or the number of votes cast for a particular candidate in voting precinct $i$ in the state of New York. The task in the former is to reconstruct an underlying true image that has been corrupted with noise; in the latter we aim to estimate spatially-varying trends in voter preference throughout the state.

What these examples, and countless others, have in common is that the data are correlated so that the value at one location is influenced by the values at nearby locations. While this dependence can be directly modeled in the likelihood of $\mathbf{y}$, it is often reasonable to assume that it can be explained by an unobservable process $\mathbf{x} = (x_1, \ldots, x_n)^T$, where $x_i$ is the realization of the process at node (location) $i$. Then a typical Bayesian analysis of this problem takes the $y_i$’s to be conditionally independent given $\mathbf{x}$; $y_i \mid \mathbf{x} \sim f_i(\cdot \mid \mathbf{x})$, $i = 1, \ldots, n$. In other words, the correlation is assumed to be completely explained by $\mathbf{x}$. For more flexibility and to more fully account for sources of uncertainty, one might assume that the distribution of $\mathbf{x}$ is determined by an unknown parameter vector $\mathbf{\theta}$ (usually of much smaller dimension than $\mathbf{x}$) which is itself assigned a hyper-prior. Thus, the Bayesian model is

$$y \mid \mathbf{x} \sim f(\cdot \mid \mathbf{x})$$
$$\mathbf{x} \mid \mathbf{\theta} \sim \pi_x(\cdot \mid \mathbf{\theta})$$
$$\mathbf{\theta} \sim \pi_\theta(\cdot).$$

Inference proceeds by evaluating (or estimating) characteristics of the posterior distribution,
determined via Bayes’ rule as \( \pi(x, \theta \mid y) \propto f(y \mid x) \pi_x(x \mid \theta) \pi_\theta(\theta) \).

A widely adopted approach for modeling the dependence structure in this problem is to assume \( x \) satisfies a Markov property. In the simplest case, this means that if \( x_j \) is in between \( x_i \) and \( x_k \), then \( x_i \) and \( x_k \) are conditionally independent, given \( x_j \). (Higher-order neighborhoods are also sometimes used where conditioning on more values is necessary.) If \( x \) satisfies this property, then \( x \) is said to be a Markov random field (MRF). MRFs are useful tools in a variety of challenging applications, including disease mapping (Waller et al., 1997; Self et al., 2018), medical imaging (Higdon 1998; Brown et al., 2014), and gene microarray analysis (Xiao et al., 2009; Brown et al., 2017a). Even autoregressive time series models are instances of Markov random fields; though this work is primarily motivated by models for spatially-indexed data in which there is no clear direction of influence. Awareness of such models was raised after the seminal work of Besag (1974), after which they came to be known in the statistics literature as conditional autoregressive (CAR; Banerjee et al., 2015) models. Since then, they have become popular for modeling temporally- or spatially-dependent areal data due to their interpretability and computational tractability afforded by the conditional independence induced by the Markov property. This property is particularly important for modern Markov chain Monte Carlo (MCMC; Gelfand and Smith, 1990) methods. Indeed, the ease with which Markov random fields can be incorporated into a Gibbs sampling algorithm (Geman and Geman, 1984) has contributed to their popularity in Bayesian statistics.

We are concerned in this work with models in which \( x \mid \theta \) is a Gaussian Markov random field. Gaussian Markov random fields (GMRFs; Rue and Held, 2005) are simply MRFs in which the conditional distribution of each (scalar) random variable is Gaussian. GMRFs typically are specified either implicitly by providing the complete set of full conditional distributions \( p(x_i \mid x_1, \ldots, x_{i-1}, x_{i+1}, \ldots, x_n) \), \( i = 1, \ldots, n \), or explicitly by defining the precision (inverse covariance) matrix instead of the covariance function as would be done in Gaussian process modeling (Schabenberger and Gotway, 2005). Further, GMRFs do not usually yield stationary processes due to a so-called “edge effect” in which the marginal variances vary by location. Corrections can be made to yield a stationary process such as a periodic boundary assumption (Fox and Norton, 2016) or algorithmic specification of
the precision matrix \cite{Dempster1972}. Sometimes the effect can simply be ignored with little effect on inference \cite{Besag1995}. Efforts have been made to use GMRFs to approximate Gaussian processes with specified covariance functions \cite{Rue2002,Song2008,Lindgren2011}, but much work still remains.

A particularly intuitive instance of a GMRF is one that centers the distribution of each $x_i$ at the average of its neighbors; i.e., $x_i \mid \mathbf{x}_{(-i)} \sim N(\bar{x}_i, \sigma_i)$, where $\mathbf{x}_{(-i)} = (x_1, \ldots, x_{i-1}, x_{i+1}, \ldots, x_n)^T$, $\bar{x}_i$ is the average of the values adjacent to $x_i$, and $\sigma_i^2$ is obtained by scaling a common variance term by the number of neighbors at site $i$. The precision matrix determined by this model is only positive semi-definite and thus not invertible, meaning that the joint distribution is improper. Such models are called intrinsic autoregressive (IAR; \cite{Besag1995}) models and are popular as Bayesian prior distributions, due in part to their interpretability.

Belonging to the Gaussian class of distributions, GMRFs are the most widely studied Markov random fields. See \cite{Rue2005} for an overview of relevant work. The literature includes techniques for efficiently sampling from GMRFs. As we discuss in Section 2, the two most common methods for sampling both have caveats when working with extremely high-dimensional data. So-called block sampling involves Cholesky factorizations of large precision matrices and thus carries high computational and memory costs. While a GMRF prior induces sparsity which can be exploited to economize such calculations, conditional posterior precision matrices arising in Bayesian models such as \cite{1} typically depend on parameters that change in each iteration of an MCMC algorithm and the required repeated factorizations can be extremely time consuming. On the other hand, so-called single-site samplers work by only considering scalar random variable updates. In addition to being more loop-intensive than block samplers, single-site samplers are known to exhibit slow convergence when the variables are highly correlated \cite{Carlin2009}. The competing goals of statistical efficiency and computational efficiency have led to recent innovations in alternative sampling approaches for GMRFs. Some of these approaches require considerable expertise in numerical analysis or message passing interface (MPI) protocol, but others are relatively easy to implement and hence can be quite useful for statisticians. Specifically, the recently proposed chromatic Gibbs sampler \cite{Gonzalez2015}.  

is easy to implement and is competitive with or even able to improve upon other existing strategies. It allows a practitioner to parallelize sampling or to take advantage of ‘vectorized’ calculations in a high-level language such as R \cite{R Core Team,2018} without requiring extensive expertise in numerical analysis or MPI.

The chromatic sampler appearing in Gonzalez et al. \cite{Gonzalez et al.,2011} was motivated by and demonstrated on binary MRFs. However, it is straightforward to carry over the same idea to the Gaussian case. In this paper, we discuss block updating and single-site updating of GMRFs and compare them to chromatic sampling. Rather than focusing on theoretical convergence rates or an otherwise overall “best” approach, we view these techniques through the lens of a practitioner looking for easily implemented yet efficient algorithms. To the best of our knowledge, this work is the first time chromatic Gibbs sampling has been directly compared to the standard approaches for sampling of GMRFs.

There exist fast approximation methods for estimating features of a posterior distribution without resorting to Markov chain Monte Carlo. One of the most popular of these is integrated nested Laplace approximation \cite{INLA; Rue et al.,2009}, the R implementation of which is the \texttt{R-INLA} package \cite{Lindgren and Rue,2015}. Such approximation methods are useful when certain quantities need to be estimated quickly, but they are only approximations and thus are not interchangeable with Markov chain Monte Carlo algorithms that converge to the exact target distribution and allow for the approximation of virtually any posterior expected value with the same Monte Carlo sample. Indeed, INLA provides the most accurate approximations around the posterior median and can disagree with MCMC in tail probability approximations \cite{Gerber and Furrer,2015}. These disagreements are more pronounced in cases where the full conditional distribution of the random field is non-Gaussian (for which INLA uses Laplace approximations) and a GMRF is used as a proposal in a Metropolis-Hastings algorithm. Further, the \texttt{R-INLA} package is a “black box” that works well for a set of pre-defined models. For more flexibility to manipulate non-standard models, there is the need to break open the black box to customize an algorithm to suit one’s needs. In the context of GMRFs, this requires more direct interaction with the random fields, motivating this work. Efficient strategies such as those considered here are not intended to be substitutes for INLA or other approximation methods. Rather, they
are complementary procedures that are useful when one is interested in direct MCMC on challenging posterior distributions.

In Section 2, we briefly motivate our sampling problem and review GMRFs. We then compare chromatic sampling to block updating and single-site sampling of GMRFs. In Section 3 we compare the performance of single-site sampling, block updating, and the chromatic approach in a numerical study using a simple Bayesian model with spatial random effects on simulated, high-dimensional imaging data, as well as a real application involving non-Gaussian polling data. We conclude in Section 4 with a discussion.

2 MCMC SAMPLING FOR GAUSSIAN MARKOV RANDOM FIELDS

In modern Bayesian analysis, it is common for the posterior distribution to have no known closed form. Hence, expectations with respect to this distribution cannot be evaluated directly. If one can obtain a sample from this distribution, though, laws of large numbers allow us to approximate quantities of interest via Monte Carlo methods. A common approach to obtaining a sample from a posterior distribution is Markov chain Monte Carlo (MCMC), particularly Gibbs sampling.

One reason for the popularity of Gibbs sampling is the ease with which the algorithm can be constructed. For an estimand \( \mathbf{\mu} = (\mu_1, \ldots, \mu_p)^T \), it proceeds simply by initializing a chain at \((\mu^{(0)}_1, \ldots, \mu^{(0)}_p)^T\) and, at iteration \(t\), sampling \(\mu^{(t)}_m \sim \pi(\mu_m | \mu^{(t)}_1, \ldots, \mu^{(t)}_{m-1}, \mu^{(t-1)}_{m+1}, \ldots, \mu^{(t-1)}_p)\), \(m = 1, \ldots, p\). Under suitable conditions, ergodic theory (e.g., Robert and Casella [2004]) establishes that the resulting Markov chain \(\{(\mu^{(t)}_1, \ldots, \mu^{(t)}_p)^T : t = 0, 1, \ldots\}\) has \(\pi(\mathbf{\mu})\) as its limiting distribution. In practice, for GMRFs with target distribution \(\pi(\mathbf{x}, \mathbf{\theta} | \mathbf{y})\), implementing this algorithm requires the ability to draw \(\mathbf{x} | \mathbf{\theta}, \mathbf{y}\) thousands of times. This is computationally expensive and thus quite challenging when \(\mathbf{x}\) is high dimensional, as we discuss in this Section.
2.1 Gaussian Markov Random Fields

Consider a GMRF $\mathbf{x} = (x_1, \ldots, x_n)^T$, where $x_i$ is the realization of the field at node $i$, $i = 1, \ldots, n$. The density of $\mathbf{x}$ is given by

$$
\pi(\mathbf{x} | \boldsymbol{\mu}) \propto \exp \left( -\frac{1}{2} \mathbf{x}^T \mathbf{Q} \mathbf{x} + b^T \mathbf{x} \right),
$$

(2)

where $\boldsymbol{\mu} \in \mathbb{R}^n$ and $b = \mathbf{Q} \boldsymbol{\mu}$. If $\mathbf{Q}$ is nonsingular, then this distribution is proper (i.e., $\int \pi(\mathbf{x} | \boldsymbol{\mu}) d\mathbf{x} < \infty$, for all $\boldsymbol{\mu}$) and the normalizing constant is $(2\pi)^{-n/2} \det(\mathbf{Q})^{1/2}$. Intrinsic GMRFs are such that $\mathbf{Q}$ is rank deficient and only positive semidefinite. In this case, we may define the density with proportionality constant $(2\pi)^{-\left(\frac{n-k}{2}\right)} \det^* (\mathbf{Q})^{1/2}$, where $n - k$ is the rank of $\mathbf{Q}$ and $\det^*(\cdot)$ is the product of the $n - k$ non-zero eigenvalues of $\mathbf{Q}$ (Hodges et al., 2003; Rue and Held, 2005). Such improper GMRF models are common in Bayesian disease mapping (Waller et al., 1997) and linear inverse problems (Bardsley, 2012), as they are easily interpretable and usually yield proper posterior distributions.

An appealing feature of GMRFs is the ability to specify the distribution of $\mathbf{x}$ through a complete set of full conditional distributions, $\{p(x_i | x_{(\neq i)}): i = 1, \ldots, n\}$. For instance, we can assume each $x_i | x_{(\neq i)} \sim N(\eta_i, \sigma_i^2)$, with $\eta_i = \mu_i + \sum_{j \sim i} c_{ij} (x_j - \mu_j)$ and $\sigma_i^2 > 0$, where $i \sim j$ if and only if node $i$ is connected to (i.e., a neighbor of) node $j \neq i$ and $c_{ij}$ are specified weights such that $c_{ij} \neq 0$ if and only if $i \sim j$. Specification of a Markov random field through these so-called local characteristics was pioneered by Besag (1974), after which such models came to be known as conditional autoregressive (CAR) models. Besag (1974) uses Brook’s Lemma and the Hammersley-Clifford Theorem to establish that the set of full conditionals collectively determine a joint density, provided a positivity condition holds among $\mathbf{x}$. In this case, we have that $Q_{ij} = (I(i = j) - c_{ij} I(i \neq j))/\sigma_i^2$, where $I(\cdot)$ is the indicator function. The condition $\sigma_j^2 c_{ij} = \sigma_i^2 c_{ji}$, for all $i, j$, is necessary to ensure symmetry of $\mathbf{Q}$. The ease with which these full conditional distributions can be incorporated into a Gibbs sampling algorithm has led to a dramatic increase in the popularity of CAR models over the past twenty years or so (Lee, 2013; Banerjee et al., 2015). Indeed, there exists user-friendly software that facilitates incorporating CAR models into Bayesian spatial models without detailed knowledge of their construction. Examples include the GeoBUGS package in WinBUGS (Lunn et al., 2000) and the R package CARBayes (Lee, 2013).
GMRFs may be specified according to an undirected graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$, where $\mathcal{V}$ indicates nodes (the vertices) and $\mathcal{E} = \{ (i, j) : i \sim j \}$ is the edge set. The precision matrix $Q$ is determined by $(Q)_{ij} \neq 0$ if and only if $(i, j) \in \mathcal{E}$. Specifying the density through the precision matrix $Q$ instead of a covariance matrix induces a Markov property in the random field $\text{[Rue and Held 2005 Theorem 2.2]}$. For any node $i$, $x_i \mid x_{(-i)} \overset{d}{=} x_i \mid x_{\mathcal{N}(i)}$, where $\mathcal{N}(i) = \{ j : (i, j) \in \mathcal{E} \}$ is the neighborhood of node $i$ and $x_A := (x_i : i \in A)^T$ for some index set $A$. That is, $x_i$ is conditionally independent of the rest of the field given its neighbors. Most GMRFs assume that each node has relatively few neighbors, resulting in $Q$ being sparse. The typical sparsity of the precision matrix is another reason GMRFs are widely used to model dependence in areal data.

With the need to model extremely large datasets with nontrivial correlation has come the need for efficient sampling techniques whereby posterior distributions arising from fully Bayesian models can be simulated. When periodic boundary conditions on $x \in \mathbb{R}^n$ can be assumed (i.e., each $x_i$ has the same number of neighbors, including the edge nodes, as in a pixelized image with zero-padded boundaries), $\text{[Fox and Norton 2016]}$ note that the sampling problem can be diagonalized via the Fast Fourier Transform (with complexity $O(n \log n)$), whence a sample can be drawn by solving a system in $O(n)$ operations. They propose reducing the total number of draws from the conditional distribution of $x$ by using a “marginal-then-conditional” sampler in which the MCMC algorithm operates by completely collapsing over $x$ and subsequently sampling $x$ using only the approximately independent draws of the hyperparameters obtained from a full MCMC run on their marginal distribution. In many applications, though, the periodic boundary assumption may not be realistic (e.g., administrative data indexed by irregular geographic regions or pathways in microarray analysis consisting of different numbers of genes), and sampling from the marginal distribution of hyperparameters can itself be challenging. To avoid the computational difficulties associated with full GMRFs, $\text{[Cai et al. 2013]}$ propose using a pairwise graphical model as an approximate GMRF for high-dimensional data imputation without specifying the precision matrix directly. The authors admit, however, that this procedure is very hard to implement $\text{[Cai 2014 p. 7]}$. In cases where we are given $Q$ and $b$ in (2) with the goal of estimating $\mu$, $\text{[Johnson et al. 2013]}$ express the Gibbs sampler as a Gauss-Seidel
iterative solution to $Q\mu = b$, facilitating the “Hogwild” parallel algorithm of Niu et al. (2011) in which multiple nodes are updated simultaneously without locking the remaining nodes. In the Gaussian case, Johnson et al. (2013) prove convergence to the correct solution when the precision matrix $Q$ is symmetric diagonally dominant. Motivated by Johnson et al. (2013), Cheng et al. (2015) use results from spectral graph theory to propose a parallel algorithm for approximating a set of sparse factors of $Q^l, -1 \leq l \leq 1$, in nearly linear time. They show that it can be used to construct independent and identically distributed realizations from an approximate distribution. This is opposed to a Gibbs sampler, which produces approximately independent samples from the correct distribution (possibly after thinning). Similar to the Gauss-Seidel splitting considered by Johnson et al. (2013), Liu et al. (2015) propose an iterative approach to approximating a draw from a GMRF in which the corresponding graph is separated into a spanning tree and the missing edges, whence the spanning tree is randomly perturbed and used as the basis for an iterative linear solve.

The aforementioned algorithms can be difficult to implement and require substantial knowledge of graph theory, numerical analysis, and MPI programming. This makes such approaches inaccessible to many statisticians who nevertheless need to work with large random fields. Further, they are iterative routines for producing a single draw from an approximation to the target distribution. This feature makes them less appealing for users who work in R or MATLAB (The MathWorks, Natick, MA). It is well-known that loops should be avoided in these languages to avoid repeated data type interpretation and memory overhead issues. In response to these difficulties while still faced with the problem of efficient updating of GMRFs inside a larger MCMC algorithm, additional R packages have been made available which are beneficial for manipulating the sparse matrices associated with GMRFs, including Matrix (Bates and Maechler, 2016), SparseM (Koenker and Ng, 2016), and spam (Furrer and Sain, 2010; Gerber and Furrer, 2015).

2.2 Block and Single-Site Gibbs Sampling

In this Section, it is helpful to distinguish between sampling $x$ directly from a prior GMRF and from the full conditional distribution of $x$ derived from an hierarchical Bayesian model with a GMRF prior on $x$. For an unconditional (and proper) GMRF, the distribution
is of the form \( x \sim N(\mu, Q^{-1}) \), where \( \mu \) and \( Q \) are generally unrelated. When drawing from the full conditional distribution as in a Gibbs sampler, the distribution is of the form \( x \sim N(Q_p^{-1}b, Q_p^{-1}) \), where \( Q_p \neq Q \) is an updated precision matrix. For example, in a typical linear model \( y \mid x, \Sigma \sim N(Ax, \Sigma) \) with \( A \) fixed and \( x \sim N(\mu, Q^{-1}) \), standard multivariate normal theory yields \( x \mid y, \Sigma \sim N(Q_p^{-1}b, Q_p^{-1}) \), where \( Q_p = A^T \Sigma^{-1} A + Q \) and \( b = A^T \Sigma^{-1} y + Q \mu \).

Two approaches to updating GMRFs inside a Markov chain Monte Carlo algorithm are so-called single site sampling in which individual sites are updated one at a time using the available full conditional distributions, and block Gibbs sampling in which the entire random field is updated all at once via sampling from a known multivariate Gaussian distribution induced by the GMRF. Block sampling improves the convergence of Gibbs samplers in the presence of a posteriori correlated variables by allowing the chain to move more quickly through its support (Liu et al., 1994). The drawback is in the manipulation and solution of large covariance matrices necessary for both random variable generation and evaluation of the likelihood in a Metropolis-Hastings algorithm (Metropolis et al., 1953; Hastings, 1970). Single site updating uses the conditional distributions of each scalar random variable, thus avoiding large matrix computations. In single-site sampling, though, statistical efficiency may be sacrificed as updating a group of possibly highly correlated parameters one at a time can result in slow exploration of the support, slowing convergence of the Markov chain.

In single-site Gibbs sampling, we sequentially draw from each univariate conditional distribution with density \( p(x_i \mid x_{(-i)}) \), \( i = 1, \ldots, n \). Broadly speaking, this requires alternating \( n \) times between using \( x_{(-i)} \in \mathbb{R}^{n-1} \) to calculate the conditional mean and drawing from \( p(x_i \mid x_{(-i)}) \), meaning that single-site updating essentially is an \( O(n^2) \) operation. However, by exploiting the fact that the conditional mean (usually) only depends on a relatively few neighbors of \( x_i \) (i.e., \( \mathcal{N}(i) \ll n \)), updating it after each draw becomes negligible, reducing the complexity to \( O(n) \). This algorithm has little regard for the ordering of the nodes, making such sampling strategies very easy to implement. Compared to block updating, though, many more Gibbs scans may be required to sufficiently explore the support of the distribution. This approach is the most iteration-intensive of any of the approaches considered here. As such, its implementation in R can result in a large amount
of overhead associated with loops, considerably slowing the entire routine.

Efficient block sampling schemes for GMRFs are discussed in [Rue (2001) and Knorr-Held and Rue (2002)]. What most of these schemes have in common is the use of a Cholesky factorization to solve a system of equations. For the case typically encountered in a Gibbs sampler, [Rue and Held (2005)] provide an algorithm for simulating from $N(Q_p^{-1}b, Q_p^{-1})$. This algorithm, presented in Algorithm 1 in the Supplementary Material, requires one Cholesky factorization and three linear solves via forward or backward substitution.

In general, for a matrix of dimension $n \times n$, the Cholesky factorization is an $O(n^3/3)$ operation and each linear solve costs $O(n^2)$ flops ([Golub and Van Loan] 1996). This can be particularly onerous in a fully Bayesian approach in which hyperpriors are assigned to hyperparameters $\theta$ that appear in the precision matrix $Q_p \equiv Q_p(\theta)$. However, the key to making block updating feasible on high-dimensional data lies in the computational savings that can be achieved when $Q_p$ is sparse. Sparse matrix algebra is itself a non-trivial problem requiring specialized knowledge beyond the expertise of many statisticians. Indeed, concerning this point, [Rue and Held (2005), p. 52] recommend “leaving the issue of constructing and implementing algorithms for factorizing sparse matrices to the numerical and computer science experts.” In practice, most statisticians rely on special functions for sparse matrices such as those found in the Matrix, SparseM, or spam packages in R. Of these three, spam is the most specifically tailored for repeatedly sampling GMRFs in MCMC.

For simulating posterior distributions via block Gibbs sampling, we are interested in drawing from full conditional distributions. In this case, sparsity of the entire precision matrix is contingent upon the sparsity of $A^T \Sigma^{-1}A$. This is often the case in practice. For instance, in disease mapping and related applications, it is common to place a spatially correlated random effect at each location to encourage smoothing of the incidence rate over space (e.g., [Waller et al., 1997] [Banerjee et al., 2015]). In terms of the linear model, this can be expressed as $y - X\beta = Z\gamma + \varepsilon$, where $X\beta$ corresponds to fixed effects and $\gamma$ contains the spatially-varying effects. With site-specific random effects, $Z$ is diagonal or block diagonal. The diagonal case (e.g., $Z = I$) is especially amenable to efficient block Gibbs sampling as well as chromatic sampling (see Subsection 2.3), since the underlying graph $G$ for the full conditional distribution is exactly the same as the prior graph.
For a fully Bayesian model, the full conditional precision matrix associated with the GMRF will generally depend on parameters that are updated in each iteration of an MCMC routine, meaning that the Cholesky factorization has to be recomputed on each iteration. Often, though, the neighborhood structure and thus the sparsity pattern of the Cholesky factor remain fixed. The sparse matrix implementation in the spam package exploits this fact to accelerate repeated block GMRF updates. After finding the initial Cholesky factorization using so-called supernodal elimination trees (Ng and Peyton [1993]), spam stores the symbolic factorization and only performs numeric factorizations on subsequent iterations. Even with sparse matrix algebra, though, block sampling $x$ from a GMRF in very high dimensions can be problematic due to the computational cost of even an initial factorization as well as the associated memory overhead (Rue [2001], p. 331).

2.3 Chromatic Gibbs Sampling

Consider the graph representation of the GMRF, $\mathcal{G} = (\mathcal{V}, \mathcal{E})$. The local Markov property says that $x_i \perp x_{-i,N(i)} | x_{N(i)}$, where $x_{-i,N(i)}$ denotes all $x$ except $x_i$ and the neighborhood of $x_i$, and $\perp$ denotes (statistical) independence. An extension of the local Markov property is to let $C \subset \mathcal{V}$ denote a separating set, or cut, of $\mathcal{G}$ such that nodes in a set $A \subset \mathcal{V}$ are disconnected from nodes in $B \subset \mathcal{V}$ after removing the nodes in $C$ from the graph. Then the global Markov property states that $x_A \perp x_B | x_C$. Chromatic sampling exploits this property by partitioning the nodes according to a graph coloring whereby the nodes in each subset can be updated simultaneously.

A coloring $f : \mathcal{V} \rightarrow \{1, \ldots, k\}$, $k \in \mathbb{N}$, is a collection of labels assigned to nodes on a graph so that no two nodes that share an edge have the same label. A $k$-coloring induces a partition of the nodes $\{A_1, \ldots, A_k\}$, where $A_j = f^{-1}(\{j\}) \subset \mathcal{V}$. For example, Figure 1 displays a 4-coloring that could be used for data that lie on a regular two-dimensional lattice; e.g., imaging data. Given a $k$-coloring of the MRF graph, we can determine a cut $C_j$ corresponding to each color $j$ by assigning all nodes that are not of that color to be in the cut; i.e., $C_j = A_j^c$, $j = 1, \ldots, k$. Defining cuts in this way for $j = 1, \ldots, k$, we have that $x_i | x_{C_j} \overset{\text{indep.}}{\sim} N(\eta_i, \sigma_i^2)$, for all $i \in A_j$, where each $\eta_i$ and $\sigma_i^2$ depend on $x_{C_j}$. That is, all nodes of the same color are conditionally independent and can be sampled in parallel,
given the rest of the field. The use of graph colorings in this way lead to term the approach *chromatic Gibbs sampling*.

Algorithm 1 presents a general chromatic Gibbs sampler for GMRFs. An advantage of using this approach is in step 3 of the algorithm. When updates of the random variables indexed by $A_j$ are distributed across several processors, the computational effort of updating the entire field can potentially be dramatically reduced, even compared to the approximate linear complexity obtained from sparse matrix factorization. Given $c$ processors and a $k$-coloring of a Markov random field over $n$ nodes, and assuming calculating conditional means is $O(1)$, the chromatic Gibbs sampler generates a new sample in approximately $O(n/c + k)$ operations (Gonzalez et al., 2011, p. 326). Observe that single-site Gibbs sampling can be obtained as a case of chromatic Gibbs sampling with $k = n$ colors.

The best computational savings under chromatic sampling will be achieved by using the *chromatic index* for the coloring, defined as the minimum $k$ so that a $k$-coloring of $G$ exists. The minimal coloring problem for a graph is NP-hard and thus very challenging except in simple situations. On regular lattices with commonly assumed neighborhood structures (e.g., Figure 1), such colorings can be found by inspection without complicated algorithms. Coloring more general graphs is more involved. However, it is important to observe that for fixed sparsity patterns (and hence fixed Markov graphs) such as those we consider here, graph coloring is a pre-computation. It is only required to run the algorithm once prior to

![Figure 1: An example of a $k$-coloring ($k = 4$) for nodes on a regular two-dimensional lattice.](image)
**Input:** Current state of GMRF, $x^{(t)}$, a $k$-coloring of the MRF graph, $\{A_j : j = 1, \ldots, k\}$.

**Output:** New draw $x^{(t+1)}$ from the GMRF.

```
1 for $j = 1$ to $k$ do
2    For $i \in A_j$, calculate conditional means and standard deviations $\eta_i, \sigma_i^2$ using $x^{(t+1)}_{A_1}, \ldots, x^{(t+1)}_{A_{j-1}}, x^{(t)}_{A_{j+1}}, \ldots, x^{(t)}_{A_k}$
3    Draw $x_{A_j} \sim N(\eta_1, \ldots, \eta_{|A_j|})^T, \text{diag}(\sigma_i^2, i = 1, \ldots, |A_j|)$
4 end
5 Return $x^{(t+1)}$
```

**Algorithm 1:** Chromatic Gibbs step updating of a GMRF.

A straightforward approach to graph coloring is the greedy algorithm, but it is known to generally produce suboptimal colorings. In fact, for random graphs in which any two vertices have probability 1/2 of sharing an edge, the greedy algorithm is known to asymptotically produce, on average, twice as many colors as necessary (Grimmett and McDiarmid, 1975). We illustrate this with an example in the Supplementary Material in which the greedy algorithm produces the optimal coloring under one permutation of the vertices, and over twice as many colors with another permutation. The sensitivity of the greedy algorithm to the ordering of the vertices was recognized by Culberson (1992), who proposes an iterative approach in which the greedy algorithm is repeatedly applied to permutations of the vertices so that the optimal coloring can be better approximated. Beyond greedy algorithms, Krager et al. (1998) cast the $k$-coloring problem as a semidefinite program and propose a randomized polynomial time algorithm for its solution. A full exposition of coloring algorithms is well beyond the scope of this work. However, our experience is that even the suboptimal colorings produced by the simple greedy algorithm are still able to facilitate vast computational improvements over block or single-site updating. As such, we provide in the Supplementary Material Algorithm 2, an easily-implemented greedy algorithm that is accessible to most statisticians looking for a quick way to color their MRF graph. We also remark that there exist R packages containing functions for coloring graphs; e.g., the
MapColoring package (Hunziker, 2017) which implements the DSATUR algorithm (Brélaz, 1979).

Most computers today have parallel processing capabilities, and any distributed processing over $c$ processors can reduce the computational burden by an approximate factor of $1/c$. Regardless of the number of processors available to the user, though, savings can still be realized when working in a high-level language such as R by ‘vectorizing’ the updating of the conditionally independent sets. Vectorizing still ultimately uses a for loop on each set of nodes, but the loops are performed in a faster language such as C or Fortran. It also minimizes the overhead associated with interpreting data types; i.e., vectorizing allows R to interpret the data type only once for the entire vector instead of repeatedly for each element of the vector.

3 NUMERICAL ILLUSTRATIONS

In this Section we compare chromatic sampling to block Gibbs and single-site sampling with both simulated data on large regular arrays and real, non-Gaussian (binomial) data on an irregular lattice. Our emphasis here is on ease of implementation for statisticians who may not be as comfortable with low-level programming as they are in R. Thus, most of our comparisons are done by examining the computational effort associated with programming entirely in R. We emphasize that computational improvements may be realized without direct parallel processing. We simply vectorize the simultaneous updating steps, thereby avoiding direct for loops in R. It is important to note that our implementation of chromatic sampling involves updating the means after each simultaneous draw via matrix-vector multiplication. The necessary matrices are stored in sparse format. Without sparse representations, the computational effort would be dramatically increased. Near the end of Subsection 3.1 we consider also a parallel implementation of the chromatic sampler in R, as well as what happens when the single-site updating step is done in C++ rather than R. To implement the block Gibbs sampler, we use the spam package (Furrer and Sain, 2010), since it is specifically tailored for GMRFs inside MCMC routines by storing the sparsity structure for repeated use. To make the spam functions as efficient as possible, we follow the authors’ suggestion and turn off the symmetry check and safe mode op-
tions (options(spam.cholsymmetrycheck= FALSE, spam.safemodevalidity= FALSE)). The computer code is available as supplementary material.

3.1 Simulated Imaging Problem

Image analysis involves attempting to reconstruct a true latent image, where the ‘image’ may mean a true physical structure as in clinical medical imaging, or an activation pattern or signal as in, e.g., functional magnetic resonance imaging (Lazar, 2008). The available data consist of pixel values, often corresponding to color on the grayscale taking integer values from 0 to 255. The true values are assumed to have been contaminated with error due to the image acquisition process. This area is one of the original motivating applications for Markov random fields (Besag, 1986; Besag et al., 1991).

There is growing interest in the statistical analysis of ultra-high dimensional imaging data. For example, structural magnetic resonance images of the human brain may consist of 20-40 two dimensional slices, each of which has 256 × 256 resolution or higher. Spatial Bayesian models for even a single slice of such data can involve GMRFs over lattices of dimension $n = 256^2$ (Brown et al., 2017b) and thus are very computationally challenging when drawing inference via Markov chain Monte Carlo. Motivated by such applications, we consider images consisting of $p \times p$ pixels, each of which has an observed value $y_{ij} = x_{ij} + \varepsilon_{ij}$, where $x_{ij}$ is the true value of the $(i, j)^{th}$ pixel in the latent image and $\varepsilon_{ij}$ represents the corresponding contamination. To simulate the data, we take the error terms to be independent, identically distributed $N(0, 1)$ random variables. The true image in this case is a rescaled bivariate Gaussian density with $x_{ij} = 5 \exp\left(-\|v_{ij}\|^2/2\right)/\pi$, where $v_{ij} = (v_i, v_j) \in [-3, 3] \times [-3, 3]$ denotes the center of the $(i, j)^{th}$ pixel, evenly spaced over the grid, and $\| \cdot \|$ denotes the usual Euclidean norm. Figure 2 depicts the true generated image (in 50 × 50 resolution) and its corrupted counterpart. To study each of the three sampling algorithms, we consider first an image with dimension $n = p \times p = 50^2$.

The assumed model for the observed image is given by $y = 1\beta_0 + \gamma + \varepsilon$, where $y \in \mathbb{R}^n$ is the vector of the observed pixel values, $1 = (1, \ldots, 1)^T \in \mathbb{R}^n$, $\beta_0 \in \mathbb{R}$ is a constant intercept parameter, $\gamma$ is the vector of spatial effects, and $\varepsilon$ is the vector of errors assumed to follow $N(0, \sigma^2 I)$. To capture local homogeneity of the image, we assume
Figure 2: True image (left panel) and corrupted image (right panel) for the simulated image reconstruction example. (These particular images have resolution 50 × 50.)

the spatial random effects obey an IAR model with mean zero; i.e., the density of $\gamma$ is $f(\gamma) \propto (\tau^2)^{-(n-1)/2} \exp\{(2\tau^2)^{-1} \gamma^T(D - W)\gamma\}$, $\gamma \in \mathbb{R}^n$, where $W = \{w_{ij} := I(i \sim j)\}_{i,j=1}^n$ is the incidence matrix of the underlying graph and $D = \text{diag} \left( \sum_{j=1}^n w_{ij} : i = 1, \ldots, n \right)$. Here we assume a first-order neighborhood structure in which each interior pixel has eight neighbors. We ignore edge effects induced by the perimeter pixels of the image. We specify inverse gamma priors for the variance components and a flat prior for the intercept; i.e., $\sigma^2 \sim \text{InvGam}(\alpha, \alpha)$, $\tau^2 \sim \text{InvGam}(\alpha, \alpha)$, $\alpha > 0$, and $\pi(\beta_0) \propto 1$, $\beta_0 \in \mathbb{R}$. To approximate vague priors for the variance components, we take $\alpha = 0.001$. It has been observed that an inverse gamma prior on $\tau^2$ sometimes can yield undesirable behavior in the posterior (Gelman, 2006); but our focus is on sampling the random field and thus we use this prior simply for convenience. For posterior sampling, our modeling assumptions lead to a Gibbs sampler having the following full conditional distributions: $\beta_0|\mathbf{y}, \gamma, \sigma^2 \sim N \left( \frac{1^T(\mathbf{y} - \gamma)}{n}, \frac{\sigma^2}{n} \right)$, $\sigma^2|\mathbf{y}, \gamma, \beta_0 \sim \text{InvGam}\left( \alpha + n/2, \alpha + \|\mathbf{y} - 1\beta_0 - \gamma\|^2/2 \right)$, $\tau^2|\gamma \sim \text{InvGam}\left( \alpha + (n - 1)/2, \alpha + \gamma^T(D - W)\gamma/2 \right)$, and $\gamma|\mathbf{y}, \sigma^2, \tau^2 \sim N \left( Q_p^{-1}b, Q_p^{-1} \right)$, where $Q_p = \sigma^{-2}I + \tau^{-2}(D - W)$ and $b = (\mathbf{y} - 1\beta_0)/\sigma^2$. We remark that it is not unusual for spatial prior distributions to have zero or constant mean functions (Bayarri et al., 2007), since the a posteriori updated spatial model will still usually capture the salient features. In the presence of noisy data, however, identifiability is limited, meaning that the parameters will more closely follow the assumed correlation structure in the prior. In terms of statistical efficiency, this situation favors block sampling over componentwise updating, as
To implement the Gibbs sampler, three strategies are employed, with the only difference being how we sample the full conditional distribution of $\gamma$. First, since $Q_p$ is sparse, we consider full block Gibbs sampling based on Algorithm 1 in the Supplementary Material to sample $\gamma$ in a single block. The second strategy is to obviate the large matrix manipulation by employing single-site Gibbs sampling using the local characteristics, $\gamma_i|\gamma_{(-i)}, y, \sigma^2, \tau^2 \sim N(\mu_i, \sigma_i^2)$, $i = 1, \ldots, n$, where $\mu_i = \sigma_i^2(\sigma^{-2}y_i + \tau^{-2}\sum_{j \in \mathcal{N}(i)} w_{ij}\gamma_j)$ and $\sigma_i^2 = \tau^2\sigma^2\{\sigma^2(D)_{ii} + \tau^2\}^{-1}$. The final sampling strategy we implement is chromatic Gibbs sampling discussed in Subsection 2.3. This approach uses the coloring depicted in Figure 1 as a 4-coloring of the pixels in the image. Following the notation in Subsection 2.3, we have that $\gamma_i | \gamma_{C_j}, y, \sigma^2, \tau^2 \overset{\text{iid}}{\sim} N(\mu_i, \sigma_i^2)$, $i \in \mathcal{A}_j$, $j = 1, \ldots, 4$. The most important feature of the chromatic sampler is that $\gamma_i | \gamma_{C_j}, y, \sigma^2, \tau^2$, $i \in \mathcal{A}_j$, can be drawn simultaneously. All of the necessary conditional means and variances for a given color can be computed through matrix-vector multiplication. In general, multiplication of an $n \times n$ matrix with an $n \times 1$ vector has $\mathcal{O}(n^2)$ complexity. Similar to the block sampler, though, we use sparse representations of the necessary matrices for the chromatic sampler, which reduces the complexity to $\mathcal{O}(n)$ since each pixel has relatively few Markov neighbors.

We implement the three sampling strategies so that each procedure performs 10,000 iterations of the Monte Carlo Markov chain to approximate the posterior distribution of the model parameters. For each approach, three chains are run using dispersed initial values. We assess convergence of the chains via trace plots, Gelman plots (Brooks and Gelman, 1998), and plots of cumulative ergodic averages of scalar hyperparameters. We discard the first 8000 iterations as a burn-in period and assess convergence using the last 2000 realizations of each Markov chain. The simulations, coded entirely in R, are carried out on a Dell Precision T3620 desktop running Windows 10 with an Intel Xeon 4.10 GHz CPU and 64 GB of RAM.

Figure 3 displays the trace plots and empirical autocorrelation functions for the hyperparameters $\sigma^2$ and $\tau^2$ for chromatic, block, and single-site sampling. We see very similar behavior in terms of autocorrelation across all three sampling approaches. From Supplementary Figure 1, we glean that each sampling approach has approximately converged in

previously mentioned.
Figure 3: MCMC Trace plots (two left columns) and empirical ACF plots (two right columns) of single chains each for $\sigma^2$ and $\tau^2$ for the $50 \times 50$ regular array example. The top, middle, and bottom rows are from the chromatic, block, and single-site chains, respectively.

the $\sigma^2$ and $\tau^2$ chains after 2,000 iterations, although the block sampler evidently has the longest convergence time according to the Gelman plots. While each approach produces estimates of $\sigma^2$ and $\tau^2$ that tend to the same value, the block sampler exhibits slightly larger Monte Carlo standard error than the other two approaches. This partly explains the slight difference in empirical distribution from the block sampler versus that of the chromatic and single-site samplers, as depicted in Figure 4. Regardless, the joint and marginal density estimates largely agree. This agreement is also evident in Figure 5, which displays the posterior mean estimates of the true image $\beta_0 \mathbf{1} + \gamma$, the primary quantity of interest.

To assess exploration of the posterior distributions, the Figure also depicts point-wise ratios of sample standard deviations for each pair of algorithms. All three samplers produce distributional estimates that are virtually indistinguishable.

The primary advantage of the chromatic approach versus the other two is in the computational cost incurred to obtain each sample. Of course, the same number of samples from two different algorithms is not guaranteed to provide the same quality of approximation to the target distribution. To accommodate the different convergence characteristics of the
Figure 4: Left panel: Scatterplot and estimated marginal posterior densities (left) and empirical CDFs (right) from the three sampling approaches in the $50 \times 50$ array example. The left panel was created using code available at https://github.com/ChrKoenig/R_marginal_plot.

Figure 5: Posterior mean estimates of the true image obtained from each sampling approach (top row) along with pairwise standard deviation ratios (bottom row) in the $50 \times 50$ array example.

three algorithms while still considering total computation time (including the burn-in period), we measure the cost per effective sample (Fox and Norton, 2016), $CES := N^{-1} \kappa T$, where $T$ is the total computation time, $N$ is the size of the retained sample from the Markov
Table 1: CPU times to draw 2,000 realizations (including 8,000 burn-in iterations) from one $\tau^2$ Markov chain under each sampling approach in the 50 $\times$ 50 array example. Also reported are the effective sample sizes (ESS), integrated autocorrelation times (IAT), and costs per effective sample (CES).

| Sampler   | CPU Time (s) | ESS  | IAT  | CES  |
|-----------|--------------|------|------|------|
| Chromatic | 10.99        | 65.74| 30.42| 0.17 |
| Block     | 49.63        | 32.71| 61.15| 1.52 |
| Single-Site | 3331.68    | 53.54| 37.36| 62.23|

chain (after burn-in), and $\kappa$ is the integrated autocorrelation time (IAT; Kass et al., 1998; Carlin and Louis, 2009). CES measures the total computational effort required to generate an effectively independent sample from the target distribution. Table 1 displays the total CPU times, effective sample sizes, integrated autocorrelation times, and CES for the $\tau^2$ chains under each sampling approach. Here we see an approximately 89% improvement in computational effort between independent samples compared to block Gibbs sampling. Single-site sampling (when coded in R) is by far the worst performer, as expected. It is interesting to note that in this case, the chromatic sampler has the shortest IAT of the three methods considered.

To further study the performance of block sampling versus chromatic sampling, particularly how they scale with regular arrays of increasing dimension, we repeat the model fitting procedure using data simulated as before, but with images of size $p \times p$, for $p = 80, 128, 256$, and 512. To create a more challenging situation, we add considerably more noise to the images by assuming $Var(\varepsilon) = 50^2I$. This makes the underlying spatial field much more weakly identified by the data and thus more strongly determined by the prior. Hence the GMRF parameters ($\beta_0, \gamma, \tau^2$) will be more strongly correlated in the posterior, creating a more challenging situation for any MCMC algorithm. For each $p$, we run the same model with the same prior specifications as in the first example. We again run each MCMC algorithm for 10,000 iterations, treating the first 8,000 as burn-in periods.

Supplementary Figures 2 through 8 display diagnostics and posterior mean estimates produced by the different sampling procedures under $p = 50, 80, 128$ with noisy data. The R-coded single-site sampler was not computationally feasible for images with resolution
Figure 6: Simulated data and posterior mean estimates of the true image from the chromatic and block sampling approaches in the noisy $128 \times 128$ array example.

$p \geq 80$ and so was not considered. As expected, we see the autocorrelation in the $\tau^2$ chains increased with the noisy data, regardless of the sampling approach, whereas the data-level variance $\sigma^2$ remains well identified. The three approaches still produce parameter estimates that agree with each other. The Gelman plots indicate that the chromatic sampler takes longer to converge than the other two approaches, but still becomes an acceptable approximation to a posterior sample after about 7,000 iterations. The joint and marginal densities of $(\sigma^2, \tau^2)$ are more diffuse than the low-noise case, again agreeing with intuition. Despite the deterioration of the $\tau^2$ chain, the quantity of interest as in many imaging problems is a function of the model parameters. In this case, we are mainly interested in $\varphi := \beta_0 1 + \gamma$, meaning that we want to explore the so-called *embedded posterior distribution* of $\varphi$. The parameter $\varphi$ (i.e., the underlying image) converges well under chromatic and block sampling. Hence we are able to recover a reasonable approximation of the target image, as evident in Figure 6 and Supplementary Figure 7. This phenomenon echoes the observation of Gelfand and Sahu (1999) that even when a Gibbs sampler is run over a posterior distribution that includes poorly identified parameters ($\tau^2$ in this case), inferences can still be drawn for certain estimands living in lower dimensional space than the full posterior.
As noted at the beginning of this Section, the preceding results are obtained by only coding in R and without any parallel processing. However, if one is interested in accelerating the conditional GMRF updating step, they might choose to simply code the single-site sampler in C or Fortran and incorporate it into a larger MCMC algorithm. On the other hand, a researcher might want to fully exploit the ability to do chromatic updates in parallel, as opposed to vectorized updates. To examine the computational gains obtained by a simpler algorithm in a faster language, we run the single-site algorithm with the simulated imaging data, but where the GMRF updating step is passed to C++ via the Rcpp package (Eddelbuettel and François, 2011). Further, we implement the chromatic sampler with truly parallelized updates in R by distributing the independent updates (those corresponding to the same color in the graph) over eight processors (five in the 50×50 case) via the parallel package (R Core Team, 2018). In terms of the generated Markov chains, the Rcpp single site sampler and parallel chromatic sampler are algorithmically identical to the R-coded single-site and vectorized chromatic, respectively, so we do not look at their convergence characteristics separately. The code for implementing the C++ and parallel approaches is also available as supplementary material.

Figure 7 displays the total CPU time required to complete 10,000 iterations for \( p = 50, 80, 128, 256, 512 \). As previously mentioned, the R-coded single-site sampler is only feasible in the \( p = 50 \) case, as it is by far the most inefficient implementation due to the nested loops. Chromatic sampling requires much less computing time than block sampling, and scales at a lower rate. This is due in part to the fact that no Cholesky factorizations are required for chromatic sampling. Such factorizations with even sparse matrices can be expensive, and repeated multivariate Gaussian draws are still required even when the symbolic factorization is stored throughout the MCMC routine. The parallel implementation of the chromatic sampler requires more CPU time at lower resolutions, but is more scalable than the vectorized version. The cost of parallelization becomes comparable to the vectorized implementation at 256 × 256 and is slightly faster than the vectorization at \( p = 512 \). This illustrates how the overhead associated with distributing data across processors cancels out any computational gain at smaller scales. Parallelizing becomes worthwhile for extremely large datasets in which splitting up a huge number of pixels is worth spending the overhead.
There is also overhead associated with passing the data and parameter values to a function written in C++, as we see in the Rcpp implementation of the single-site sampler. At small to moderate resolutions, the Rcpp version is much faster than the chromatic sampler and the block sampler. However, the computational cost of Rcpp scales at a much faster rate than the chromatic versions, so much so that both chromatic versions are an order of magnitude faster than the C++ single-site sampler at $p = 512$. Similar to how a parallel implementation depends on the size of the data, we see that the benefit of coding a chromatic sampler in R versus passing to C++ is more pronounced when the dataset is extremely large.

There is also considerable memory overhead associated with both sparse Cholesky block updating and chromatic sampling. The total required memory for Cholesky-based block updating depends on the storage scheme used by the sparse matrix implementation. The spam implementation in our example uses a variant of the so-called compressed sparse row format [Sherman 1975]. Chromatic sampling, on the other hand, requires no matrix storage at all, but only lists of identifiers associated with each graph color. The right panel of Figure 7 illustrates the consequent savings in total memory allocations and how they scale with arrays of increasing dimension. In terms of total memory allocations, both the vectorized and parallel chromatic samplers require considerably less than block updating. In fact, block updating for $p = 256$ and $p = 512$ was not possible due to memory limitations. The Cholesky factorization failed, returning Cholmod error: ‘problem too large’.

3.2 Binomial Election Data on an Irregular Lattice

Here we examine the performance of the block Gibbs and chromatic sampling strategies on an irregular lattice, since both the structure of $Q$ and the possible colorings of the underlying graph are more complicated. Moreover, we illustrate the performance of these procedures when applied to non-Gaussian data. In particular, we examine geographical trends in voter preference using binomial outcomes. The data were obtained from the Harvard Election Data Archive (https://dataverse.harvard.edu) and are depicted in Supplementary Figure 10.

Our data consist of polling results from the 2010 New York Governor’s race in which Democratic candidate Andrew Cuomo defeated Republican candidate Carl Paladino and
Green Party candidate Howie Hawkins. During this election, the state of New York had 14,926 precincts, with polling data being available on 14,597 precincts. The 329 precincts for which data are unavailable is attributable to improper reporting or lack of voter turnout.

Let $Y_i$ be the number of votes cast for the Democratic candidate out of $m_i$ total votes in precinct $i$, $i = 1, \ldots, n$. Then we assume that $Y_i | \pi_i, m_i \overset{\text{indep.}}{\sim} \text{Bin}(m_i, \pi_i)$, where $g^{-1}(\pi_i) = \beta_0 + \gamma_i$, $g(\cdot)$ is the usual logistic link function, and $\gamma = (\gamma_1, \ldots, \gamma_n)^T$ is a vector of random effects inducing spatial homogeneity. We suppose that $\gamma$ follows a proper IAR model; i.e., $\gamma \sim N(0, \tau^2(D - \rho W)^{-1})$, where $D$ and $W$ are as defined in Section 3.1. Here, the “propriety parameter” $\rho \in (\lambda_1^{-1}, \lambda_n^{-1})$ ensures that the precision matrix is non-singular, where $\lambda_1 < 0$ and $\lambda_N > 0$ are the smallest and largest eigenvalues of $D^{-1/2}WD^{-1/2}$, respectively (Banerjee et al., 2015). Proper IARs are sometimes used as approximations to the standard IAR when a proper prior distribution is desired. For simplicity, we fix $\rho = 0.995$. The model is completed with the prior assumptions that $\beta_0 \sim N(0, 1000)$ and $\tau^2 \sim \text{InvGam}(1, 1)$.

Under the logistic link, we can simplify posterior sampling via data augmentation. This
technique exploits the fact that \( \exp(\eta) a (1 + \exp(\eta)) - b = 2 - b \exp(\kappa \eta) \int_0^\infty \exp(-\psi \eta^2 / 2) p(\psi | b, 0) d\psi \), where \( \eta \in \mathbb{R} \), \( a \in \mathbb{R} \), \( b \in \mathbb{R}^+ \), \( \kappa = a - b / 2 \), and \( p(\cdot | b, 0) \) is the probability density function of a Pólya-Gamma random variable with parameters \( b \) and \( 0 \) \cite{Polson:2013}. Using this identity, the observed data likelihood can be written as \( \pi(Y | \beta_0, \gamma) \propto \prod_{i=1}^n \exp\{\kappa_i \eta_i\} \times \int_0^\infty \exp(-\psi_i \eta_i^2 / 2) p(\psi_i | m_i, 0) d\psi_i \), where \( Y = (Y_1, \ldots, Y_n)^T \), \( \eta_i = \beta_0 + \gamma_i \) and \( \kappa_i = Y_i - m_i / 2 \). Thus, by introducing \( \psi_i \) as latent random variables, we have that \( \pi(Y, \psi | \beta_0, \gamma) \propto \exp\{-((\beta_0 1 + \gamma)^T D_\psi (\beta_0 1 + \gamma) \} / 2 \} \prod_{i=1}^n p(\psi_i | m_i, 0) \), where \( \psi = (\psi_1, \ldots, \psi_n)^T \), \( \kappa = (\kappa_1, \ldots, \kappa_n)^T \), and \( D_\psi = \text{diag}(\psi) \). By including \( \psi \) in the MCMC algorithm, we induce a Gaussian full conditional distribution on \( \gamma \), facilitating GMRF updates without having to tune a Metropolis-Hastings algorithm. Additional implementation details are provided in the Supplementary Material.

In order to implement chromatic sampling, a coloring of the underlying Markov graph has to be found. Using the greedy algorithm given in Algorithm 2 in the Supplementary Material, we obtain a 7-coloring, so that the chromatic sampler can update the entire \( n = 14,926 \)-dimensional field in seven steps. The coloring is depicted in Supplementary Figure 11.

We implement Gibbs sampling with both the block Gibbs and chromatic updates for 10,000 iterations, discarding the first 5,000 as a burn-in period. The code is run on a desktop using Windows 10 with an Intel Core i5-3570 3.40GHz CPU with 16GB of RAM. The trace plots and empirical ACF plots for \( \beta_0 \) and \( \tau^2 \) are depicted in Supplementary Figure 12, along with the Gelman plots of these two parameters in Supplementary Figure 13. We see adequate convergence in the same number of iterations under both sampling approaches. Table 2 summarizes the results for both samplers in terms of CPU time and cost per effective sample of the intercept and variance terms. We again see a savings in CPU time under chromatic sampling, so much so that it offsets the slightly larger autocorrelation time. Thus we are able to obtain effectively independent samples with less computational effort. The posterior mean maps of the voter Democratic preference \( (\pi_i) \) obtained under each sampling strategy are displayed in Figure 8. We see essentially identical results under both strategies.
Table 2: CPU times to draw 5,000 realizations (after 5,000 burn-in iterations) from one Markov chain under each sampling approach in the New York election example. Also reported are the effective sample sizes (ESS), autocorrelation times (ACT), and costs per effective sample (CES).

| Sampler            | CPU Time (s) | ESS      | ACT  | CES   |
|--------------------|--------------|----------|------|-------|
| $\beta_0$ Chromatic| 222.06       | 2445.74  | 2.04 | 0.0935|
| $\beta_0$ Block   | 294.16       | 2732.82  | 1.83 | 0.1018|
| $\tau^2$ Chromatic| 222.06       | 1916.63  | 2.61 | 0.1155|
| $\tau^2$ Block    | 294.16       | 2186.021 | 2.29 | 0.1332|

3.3 Summary

These numerical experiments illustrate potential improvements that chromatic Gibbs sampling can offer versus the two most common strategies of block sampling and single-site sampling. In simulated image reconstruction, we find that for every considered resolution, the chromatic sampler is computationally much cheaper than the full block Gibbs and single-site samplers coded entirely in R. We observe in both chromatic and block sampling the deterioration in Monte Carlo Markov chains that is known to occur as the dimension of a GMRF increases (Rue and Held, 2005; Agapiou et al., 2014). Even in this case, however, any of the approaches considered are able to estimate the posterior mean of the latent field and obtain equivalent recovery of the quantity of interest. The chromatic sampler is able to do so much more quickly and with much less memory overhead, the latter of which allows the chromatic sampler to scale to images of extremely large dimension beyond the capability of standard Cholesky factorization routines available in R. The potential advantages extend to irregular arrays and non-Gaussian data, as demonstrated in the election data example.

4 DISCUSSION

Over the last twenty years, Gaussian Markov random fields have seen a dramatic increase in popularity in the applied Bayesian community. In this work, we discussed approaches for simulating from Gaussian Markov random fields that are commonly used in practice.
We compared the two dominant approaches in the statistics literature, single-site and block updating, to chromatic Gibbs sampling. Each procedure has theoretical guarantees, but our criteria have been pragmatic; i.e., how can statisticians effectively lower the computational cost of sampling from the target distribution without resorting to esoteric knowledge from graph theory, numerical analysis, or parallel programming? Taking this view, we have shown that chromatic sampling is competitive with and often able to improve upon single-site and full block Gibbs. In a large-scale scenario, we demonstrated improvements afforded by chromatic sampling even when compared to passing the single-site updating step to C++. This shows that computational efficiency gains are achievable even when a researcher desires (or needs) to do as much programming as possible in a high-level interpreted language.

Motivated by large-scale clinical imaging data, we illustrated potential advantages on a regular array with Gaussian response, finding that chromatic sampling scales to settings where memory limitations prevent direct sparse matrix manipulations. We also considered
a real example with binomial election data on an irregular lattice with almost 15,000 areal units, showing that chromatic sampling is useful even without a provably optimal coloring of the MRF graph. Both block sampling and chromatic sampling tend to be far superior to single-site sampling when working in R.

While facilitating parallel or vectorized simultaneous updates, each individual draw under chromatic sampling is still at the level of a single site. Thus, for variables that are highly correlated in the target distribution, convergence can be slow. To handle this, Gonzalez et al. (2011) propose also a “splash sampler” to combine the blocking principle of updating sets of correlated variables together with the parallelizability afforded by graph colorings. Splash sampling is more involved than simple chromatic sampling. It requires careful construction of undirected acyclic graphs subject to a known tree width determined by individual processor limitations, and hence much more computing effort and more familiarity with graph theory. In the Gaussian case, splash sampling would require repeated Cholesky factorizations, each on matrices of smaller dimension, but without being able to save the sparsity structure. In the presence of highly correlated variables in the target distribution with GMRF updates, it might be preferable to use ordinary block updates with sparse matrix algebra and the algorithms suggested by Rue (2001). However, our numerical experiments demonstrate that the gain in computational efficiency from the simple chromatic sampler can still outweigh the loss of statistical efficiency. This leads to an overall improvement in a variety of situations without resorting to more sophisticated approaches that might be inaccessible to most statisticians.

In both the C++ and parallel implementations, we assume that the researcher would prefer to keep as much of the code in R as possible due to its user-friendly functionality and their desire to avoid low-level programming. Since R itself is a software suite written primarily in C, even matrix-vector calculations and vectorized functions are ultimately executed via loops in C (or a another low-level language such as Fortran). Thus, if one were to code our entire MCMC algorithm with ‘vectorized’ chromatic updates in C or C++, the result would be an algorithm that is essentially identical to a single-site sampler up to the order in which the sites are updated. In other words, if a researcher is working purely in C or Fortran, then the computational advantages of chromatic sampling can only be
fully realized with a distributed parallel approach. However, parallel programming in such low-level languages is much more difficult and nuanced than it is in R and thus beyond the expertise of many statisticians and data scientists. Indeed, one of the main reasons for the popularity of R is the ease with which extremely complicated tasks (e.g., MCMC on exotic hierarchical Bayesian models or sparse matrix factorizations) can be executed. There is nothing R can do that cannot be done in a low-level language if one has the time and patience to write the code for it. Our purpose in this work is to discuss and demonstrate computational savings that may be realized without having to leave the more comfortable environment of a high-level, interpreted programming language.

The parallel implementation used in this work used only eight processors, but this still showed modest acceleration over vectorized chromatic sampling with the largest dataset we considered. The difference would no doubt be much more pronounced by distributing the effort over more processors. With the advent of modern computing clusters and GPU computing, it is becoming more common for researchers to have available thousands of cores for use simultaneously. In fact, it is not difficult to envision scenarios where the number of processors available is $O(n)$, in which case the complexity of parallel chromatic sampling reduces to $O(k)$, where $k$ is often fixed as $n$ increases due to the structure of the data (Gonzalez et al., 2011). Thus the scaling potential of parallel chromatic sampling is enormous and worthy of further investigation. We defer such an exploration to future work.

In this paper, we examined the performance of chromatic sampling versus single-site and block Gibbs on high density data in which the entire study region is sufficiently sampled and in which a first-order Markov neighborhood can capture the salient features of the data. This situation is applicable to many, but not all, analyses of areal data. There remains the issue of how chromatic sampling would perform in the presence of sparse observations from an underlying smooth process, where the autocorrelation of the Markov chain would be expected to be higher than in the high-density case. Related to this point is that, to the best our knowledge, the convergence rates associated with chromatic sampling Markov transition kernels remain unknown. We leave these questions to be explored at a later date.

Given the current trajectory of modern data analysis, the utility of GMRFs is not likely to diminish anytime soon. However, with their use comes the need for efficient yet acces-
sible sampling strategies to facilitate Bayesian posterior inference along with appropriate measures of uncertainty. This area remains an active area of research among statisticians, computer scientists, and applied mathematicians. Fortunately, the increasingly interdisciplinary environment within which researchers are operating today makes it more likely that significant advancements will be widely disseminated and understood by researchers from a wide variety of backgrounds. This is no doubt a promising trend which will ultimately benefit the broader scientific community.

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