Fast Kronecker Inference in Gaussian Processes with non-Gaussian Likelihoods

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Abstract

Gaussian processes (GPs) are a flexible class of methods with state of the art performance on spatial statistics applications. However, GPs require $O(n^3)$ computations and $O(n^2)$ storage, and popular GP kernels are typically limited to smoothing and interpolation. To address these difficulties, Kronecker methods have been used to exploit structure in the GP covariance matrix for scalability, while allowing for expressive kernel learning (Wilson et al., 2014). However, Kronecker methods have been confined to Gaussian likelihoods. We propose new scalable Kronecker methods for Gaussian processes with non-Gaussian likelihoods, using a Laplace approximation which involves linear conjugate gradients for inference, and a lower bound on the GP marginal likelihood for kernel learning. Our approach has near linear scaling, requiring $O(Dn^{D+1})$ operations and $O(Dn^D)$ storage, for $n$ training data-points on a dense $D > 1$ dimensional grid. Moreover, we introduce a log Gaussian Cox process, with highly expressive kernels, for modelling spatiotemporal count processes, and apply it to a point pattern ($n = 233,088$) of a decade of crime events in Chicago. Using our model, we discover spatially varying multiscale seasonal trends and produce highly accurate long-range local area forecasts.

1 Introduction

Gaussian processes were pioneered in geostatistics (Matheron, 1963) where they are commonly known as kriging models Ripley (1981). O’Hagan (1978) instigated their general use, pursuing applications to optimal design, curve fitting, and time series. GPs remain a mainstay of spatial and spatiotemporal statistics Diggle & Ribeiro (2007); Cressie & Wikle (2011) and have gained widespread popularity in machine learning Williams & Rasmussen (2006).

Unfortunately, the $O(n^3)$ computations and $O(n^2)$ storage requirements for GPs has greatly limited their applicability. Kronecker methods have recently been introduced (Saatçi, 2011) to scale up Gaussian processes, with no losses in predictive accuracy. While these methods require that the input space (predictors) are on a multidimensional lattice, this structure is present in many spatiotemporal statistics applications, where predictors are often indexed by a grid of spatial coordinates and time.

A variety of approximate approaches have been proposed for scalable GP inference, including inducing point methods (Quiñonero-Candela & Rasmussen, 2005), and finite basis representations through random projections (Lázaro-Gredilla et al., 2010; Yang et al., 2014). Groot et al. (2014)
use Kronecker inference and low-rank approximations for non-Gaussian likelihoods, focusing on classification only and scaling to moderately sized datasets ($n < 7000$). We perform a detailed comparison and demonstrate that our new approach far outperforms theirs in terms of scalability and accuracy. In this paper, we make the following contributions:

- We extend Kronecker methods for non-Gaussian likelihoods, enabling applications outside of standard regression settings. We use a Laplace approximation on the likelihood, with linear conjugate gradients for inference, and a lower bound on the GP marginal likelihood, for kernel learning. Moreover, our methodology extends to incomplete grids – caused by, for example, water or political boundaries.

- We develop a spatiotemporal log Gaussian Cox process (LGCP), with highly expressive spectral mixture covariance kernels (Wilson & Adams, 2013). Our model is capable of learning intricate structure on large datasets, allowing us to derive new scientific insights from the data, and to perform long range extrapolations. This is the first use of structure learning with expressive kernels, enabling long-range forecasts, for GPs with non-Gaussian likelihoods.

- We apply our model to a challenging public policy problem, that of small area crime rate forecasting. Using a decade of publicly available date-stamped and geocoded crime reports we fit the $n = 233,088$ point pattern of crimes coded as “assault” using the first 8 years of data to train our model, and forecast 2 years into the future. We produce very fine-grained spatiotemporal forecasts, which we evaluate in a fully probabilistic framework. Our forecasts far outperform predictions made using popular alternatives. We interpret the learned structure to gain insights into the fundamental properties of these data.

We begin with a review of Gaussian processes in section 2 and then present the log-Gaussian Cox Process model for spatiotemporal point processes which we wish to fit in section 3. In section 4 we describe the standard Laplace approximation approach to GP inference and hyperparameter learning. In sections 5 and 6 we present our new Kronecker methods for scalable inference, hyperparameter learning, and missing observations. We detail our experiments on synthetic and real data in section 8.

2 Gaussian processes

We assume a basic familiarity with Gaussian processes (GPs) Williams & Rasmussen (2006). We are given a dataset $\mathcal{D} = (\mathbf{y}, \mathbf{X})$ of targets (responses), $\mathbf{y} = \{y_1, \ldots, y_n\}$, indexed by predictors (inputs) $\mathbf{X} = \{x_1, \ldots, x_n\}$. The targets could be real-valued, categorical, counts, etc., and the predictors, for example, could be spatial locations, times, and other covariates. We assume the relationship between the predictors and targets is determined by a latent Gaussian process $f(x) \sim \mathcal{GP}(m, k_\theta)$, and an observation model $p(y(x) | f(x))$. The GP is defined by its mean $m$ and covariance function $k_\theta$ (parametrized by $\theta$), such that any collection of function values $f = f(X) \sim \mathcal{N}(\mu, K)$ has a Gaussian distribution with mean $\mu_i = m(x_i)$ and covariance matrix $K_{ij} = k(x_i, x_j | \theta)$.

Our goal is to infer the predictive distribution $p(f_* | \mathbf{y}, x_*)$, for any test input $x_*$, which allows us to sample from $p(\mathbf{y}_* | \mathbf{y}, x_*)$ via the observation model $p(y(x) | f(x))$:

$$p(f_* | \mathcal{D}, x_*, \theta) = \int p(f_* | \mathbf{X}, x_*, f, \theta) p(f | \mathcal{D}, \theta) df$$  \hspace{1cm} (1)
We also wish to infer the marginal likelihood of the data, conditioned only on kernel hyperparameters \( \theta \),

\[
p(y|\theta) = \int p(y|f) p(f|\theta) \, df,
\]

so that we can optimize this likelihood, or use it to infer \( p(\theta|y) \), for kernel learning. Having an expression for the marginal likelihood is particularly useful for kernel learning, because it allows one to bypass the extremely strong dependencies between \( f \) and \( \theta \) in trying to learn \( \theta \). Unfortunately, for all but the Gaussian likelihood (used for standard GP regression), where \( p(y|f) = N(f, \sigma^2 I) \), equations (1) and (2) are analytically intractable.

3 A motivating example: Cox Processes

In this section, we describe the log-Gaussian Cox Process (LGCP), a particularly important spatial statistics model for point process data (Møller et al., 1998; Diggle et al., 2013). While the LGCP is a general model, its use has been limited to small datasets. We focus on this model because of its importance in spatial statistics and its suitability for the Kronecker methods we propose. Note, however, that our methods are generally applicable to Gaussian process models with non-Gaussian likelihoods, such as Gaussian process classification.

An LGCP is a Cox process (inhomogeneous Poisson process with stochastic intensity) driven by a latent log intensity function \( \log \lambda := f \) with a GP prior:

\[
f(s) \sim GP(\mu(s), k_\theta(\cdot, \cdot)).
\]

Conditional on a realization of the intensity function, the number of points in a given space-time region \( S \) is:

\[
y_S|\lambda(s) \sim \text{Poisson} \left( \int_{s \in S} \lambda(s) \, ds \right).
\]

Following standard practice, we introduce a “computational grid” (Diggle et al., 2013) on the observation window and represent each grid cell with its centroid, \( s_1, \ldots, s_n \). Let the count of points inside grid cell \( i \) be \( y_i \). Thus our model is a Gaussian process with a Poisson observation model and exponential link function:

\[
y_i|f(s_i) \sim \text{Poisson} \left( \exp[f(s_i)] \right).
\]

4 Laplace Approximation

The Laplace approximation models the posterior distribution of the Gaussian process, \( p(f|y, X) \), as a Gaussian distribution, to provide analytic expressions for the predictive distribution and marginal likelihood in Eqs. (1) and (2). We follow the exposition in Williams & Rasmussen (2006).

Laplace’s method uses a second order Taylor expansion to approximate the unnormalized log posterior,

\[
\Psi(f) := \log p(f|D) = \log p(y|f) + \log p(f|X),
\]

centered at the \( \hat{f} \) which maximizes \( \Psi(f) \). We have:
\[ \nabla \Psi(f) = \nabla \log p(y|f) - K^{-1}(f - \mu) \] 
\[ \nabla \nabla \Psi(f) = \nabla \nabla \log p(y|f) - K^{-1} \]

\( W := -\nabla \nabla \log p(y|f) \) is an \( n \times n \) diagonal matrix since the likelihood \( p(y|f) \) factorizes as \( \prod_i p(y_i|f_i) \).

We use Newton’s method to find \( \hat{f} \). The Newton update is

\[ f_{\text{new}} \leftarrow f_{\text{old}} - (\nabla \nabla \Psi)^{-1} \nabla \Psi \] 

Given \( \hat{f} \), the Laplace approximation for \( p(f|y) \) is given by a Gaussian:

\[ p(f|y) \approx N(\hat{f}, (K^{-1} + W)^{-1}) \]

Substituting the approximate posterior of Eq. (9) into Eq. (1), and defining \( A = W^{-1} + K \), we find the approximate predictive distribution is

\[ p(f^*|D, x^*, \theta) \approx N(k^* \nabla \log p(y|\hat{f}), k^* - k^* A^{-1} k^*) \]

where \( k^* = [k(x^*, x_1), \ldots, k(x^*, x_n)]^\top \) and \( k_{ss} = k(x_s, x_s) \).

This completes what we refer to as inference with a Gaussian process. We have so far assumed a fixed set of hyperparameters \( \theta \). For learning, we train these hyperparameters through marginal likelihood optimization. The Laplace approximate marginal likelihood is:

\[ \log p(y|X, \theta) = \log \int \exp[\Psi(f)] df \]
\[ \approx \log p(y|\hat{f}) - \frac{1}{2} \alpha^\top K^{-1} \alpha - \frac{1}{2} \log |I + KW| , \]

where \( \alpha := K^{-1}(\hat{f} - \mu) \). Standard practice is to find the \( \hat{\theta} \) which maximizes the approximate marginal likelihood of Eq. (12), and then condition on \( \hat{\theta} \) in Eq. (10) to perform inference and make predictions.

Both the learning and inference stages require solving linear systems involving matrices of size \( n \times n \). This takes \( O(n^3) \) time and \( O(n^2) \) storage, using e.g. the Cholesky decomposition.

### 5 Kronecker Methods

We provide a brief review of Kronecker methods for efficient Gaussian processes, following Saatci (2011), Gilboa et al. (2013), and Wilson et al. (2014). In the next section we extend these methods to non-Gaussian likelihoods.

The key assumptions enabling the use of Kronecker methods is that the GP kernel is formed by a product of kernels across input dimensions and the inputs are on a Cartesian product grid (multidimensional lattice), \( x \in \mathcal{X} = \mathcal{X}_1 \times \cdots \times \mathcal{X}_D \). (This grid need not be regular and the \( \mathcal{X}_i \) can have different cardinalities.) Given these two assumptions, the covariance matrix \( K \) decomposes as a Kronecker product of covariance matrices \( K = K_1 \otimes \cdots \otimes K_D \).

Saatci (2011) shows that the computationally expensive steps in GP regression can be made much faster assuming Kronecker structure. Inference and learning require solving linear systems \( K^{-1} v \) and
computing log-determinants \( \log |K| \). Typical approaches require \( O(n^3) \) time and \( O(n^2) \) space. Using Kronecker methods, these operations only require \( O(Dn^{D+1}) \) operations and \( O(Dn^{\frac{2D}{D+1}}) \) storage, for \( n \) datapoints and \( D \) input dimensions. In Section A.1 in the Appendix, we review the key Kronecker algebra results that we use, including efficient matrix-vector multiplication and eigendecomposition.

Wilson et al. (2014) extend these efficient methods to partial grids, by augmenting the data with imaginary observations to form a complete grid, and then ignoring the effects of the imaginary observations using a special noise model in combination with linear conjugate gradients. Partial grids are common, and can be caused by, e.g., government boundaries, which interfere with grid structure.

6 Kronecker Methods for Non-Gaussian Likelihoods

We introduce our efficient Kronecker approach for Gaussian processes inference (section 6.2) and learning (section 6.3) with non-Gaussian likelihoods, after introducing some notation and transformations for numerical conditioning.

6.1 Numerical Conditioning

For numerical stability, we use the following transformations: \( B = I + W^{1/2}KW^{1/2}, Q = W^{1/2}B^{-1}W^{1/2}, b = W(f - \mu) + \nabla \log p(y|f), \) and \( a = b - QKb. \) Now \( (K^{-1} + W)^{-1} = K - KQK, \) from the matrix inversion lemma, and the Newton update in Eq. (8) becomes:

\[
\mathbf{f}_{\text{new}} \leftarrow K \mathbf{a} \tag{13}
\]

Using these transformations, the predictive distribution in Eq. (10) becomes:

\[
p(f_*|D, \mathbf{x}_*, \theta) \approx \mathcal{N}(k_*^T \nabla \log p(y|\hat{f}), k_*^T - k_*^T Qk_*) \tag{14}
\]

6.2 Inference

Existing Kronecker methods do not apply to non-Gaussian likelihoods because we are no longer working solely with the covariance matrix \( K. \) We use linear conjugate gradients (LCG), an iterative method for solving linear systems which only involves matrix-vector products, to efficiently calculate the key steps of the inference algorithm in Section 4. Our full algorithm is shown in Algorithm 1. The Newton update step in Eq. (13) requires costly matrix-vector multiplications and inversions of \( B = (I + W^{1/2}KW^{1/2}). \) We replace Eq. (13) with the following two steps:

\[
Bz = W^{-1/2}b \tag{15}
\]

\[
\mathbf{a}_{\text{new}} = W^{1/2}z \tag{16}
\]

For numerical stability, we follow (Williams & Rasmussen, 2006, p. 46) and apply our Newton updates to \( \alpha \) rather than \( f. \) The variable \( b = W(f - \mu) + \nabla \log p(y|f) \) can still be calculated efficiently because \( W \) is diagonal, and Eq. (15) can be solved efficiently for \( z \) using LCG because matrix-vector products with \( B \) are efficient due to the diagonal and Kronecker structure.

The number of iterations required for convergence of LCG to within machine precision is in practice independent of \( n \) (the number of columns in \( B \)), and depends on the conditioning of \( B. \) Solving Eq. (15) requires \( O(Dn^{\frac{D+1}{D}}) \) operations and \( O(Dn^{\frac{2D}{D+1}}) \) storage, which is the cost of matrix
vector products with the Kronecker matrix $K$. No modifications are necessary to calculate the the predictive distribution in Eq. (14). We can therefore efficiently evaluate the approximate predictive distribution in $O(mDn^{D+1})$ where $m \ll n$ is the number of Newton steps. For partial grids, we apply the extensions in Wilson et al. (2014) without modification.

6.3 Hyperparameter learning

To evaluate the marginal likelihood in Eq. (12), we must compute $\log |I + KW|$. Fiedler (1971) showed that for Hermitian positive semidefinite matrices $U$ and $V$:

$$\prod_i (u_i + v_i) \leq |U + V| \leq \prod_i (u_i + v_{n-i+1})$$

(17)

where $u_1 \leq u_2 \leq \ldots \leq u_n$ and $v_1 \leq \ldots \leq v_n$ are the eigenvalues of $U$ and $V$. To apply this bound let $e_1 \leq e_2 \leq \ldots \leq e_n$ be the eigenvalues of $K$ and $w_1 \leq w_2 \leq \ldots \leq w_n$ be the eigenvalues of $W$. Then we use that the eigenvalues of $W^{-1}$ are $w_{n-1}^{-1} \leq w_{n-2}^{-1} \leq \ldots \leq w_1^{-1}$:

$$\log |I + KW| = \log(|K + W^{-1}|W|) \leq \log \prod_i (e_i + w_i^{-1}) \prod_i w_i$$

(18)

$$= \sum_i \log(1 + e_i w_i)$$

Putting this together with Equation (12) we have our bound on the Laplace approximation’s log-marginal likelihood:

$$\log p(y|X, \theta) \geq \log p(y|\hat{f}) - \frac{1}{2}\hat{\alpha}^T K^{-1}\hat{\alpha} - \frac{1}{2} \sum_i \log(1 + e_i w_i)$$

(19)

We chose the lower bound because we use gradient ascent as our learning approach to find the best parameters $\theta$ to maximize the approximate marginal likelihood. We approximately calculate the necessary gradients using finite differences.

6.4 Evaluation of our Learning Approach

The bound we used on the Laplace approximation’s log-marginal likelihood has been shown to be the closest possible bound on Hermitian psd matrices $|U + V|$ in terms of the eigenvalues of $U$ and $V$ (Fiedler, 1971), and has been used for heteroscedastic regression (Gilboa et al., 2014). However, its most appealing quality is computational efficiency. We efficiently find the eigendecomposition of $K$ using standard Kronecker methods, where we calculate the eigenvalues of $K_1, \ldots, K_D$, each in time $O(n^{2\frac{D}{2}})$. We immediately know the eigenvalues of $W$ because it is diagonal. Putting this together, the time complexity of computing this bound is $O(Dn^{\frac{3}{2}})$. The log-determinant is recalculated many times during hyperparameter learning, so its time complexity is quite important to scalable methods.

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1 An alternative would be to try to exactly compute the eigenvalues of $I + KW$ using LCG. But this would require performing at least $n$ matrix-vector products, which could be quite computationally expensive. Note that this was not an issue in computing the Laplace predictive distribution, because LCG solves linear systems to within machine precision for $J \ll n$ iterations. Our approach, with the Fiedler bound, on the other hand, provides an approximation to the Laplace marginal likelihood, and a lower bound which we can optimize, at the cost of a single eigendecomposition of $K$, which is in fact more efficient than a single matrix vector product $Bv$. 

6
Figure 1: We evaluate our lower bound on the Laplace marginal likelihood in Eq. (19), and our upper bound on the log determinant in Eq. (18), compared to exact values and low rank approximations. In a), the approximation ratio is the ratio of the Laplace negative marginal likelihood with a given approximation (our bound or the low rank approximation). In b) and c), the approximation ratio is the ratio of the true log determinant with a given approximation.
As shown in Figure 1(a), as the sample size increases the lower bound on the negative log marginal likelihood approaches the negative log marginal likelihood calculated with the true log determinant. This result makes perfect sense for our Bayesian model, because the log-determinant is a complexity penalty term defined by our prior, which becomes less influential with increasing datasizes compared to the data dependent model fit term, leading to an approximation ratio converging to 1.

Next, we compare the accuracy and run-time of our bound to a recently proposed (Groot et al., 2014) log-det approximation relying on a low-rank decomposition of $K$. In Figure 1(b) we generated synthetic data on an $\sqrt{n} \times \sqrt{n}$ grid and calculated the approximation ratio by dividing the approximate value $\log |I + K W|$ by the true value $\log |I + K W|$ calculated with the full matrix. Our bound always has an approximation ratio between 1 and 2, and it gets slightly worse as the number of observations increases. This contrasts with the low-rank approximation. When the rank $r$ is close to $\sqrt{n}$ the approximation ratio is reasonable, but quickly deteriorates as the sample size increases.

In Figure 1(c) we compare the running times of these methods, switching to a 3-dimensional grid. The exact method quickly becomes impractical. For a million observations, a rank-5 approximation takes 6 seconds, a rank-15 approximation takes 600 seconds, while our bound takes only 0.24 seconds. We do not have the true log-determinant to compare, but our bound is provably a lower bound, so the ratio between the low-rank approximation’s log-determinant and ours is a lower-bound on the true approximation ratio. This ratio is 2.8 for the rank-15 approximation and 30 for the rank-5 approximation (where we have again switched the signs to match the value in the negative log marginal likelihood calculation).

Finally, we know theoretically that Fiedler’s bound is exact when the diagonal matrix $W$ is equal to spherical noise $\sigma^2 I$, which is the case for a Gaussian observation model. Since the Gaussian distribution is a good approximation to the Poisson distribution in the case of a large mean parameter, we evaluated our log-determinant bound while varying the prior mean $\mu$ of $f$ from 0 to 10. As shown in Figure 1(d), for larger values of $\mu$, our bound becomes more accurate. There is no reason to expect the same behavior from a low-rank approximation, and in fact the rank-20 approximation becomes worse as the mean of $\lambda$ increases.

6.5 Algorithm Details and Analysis

For inference, our approach makes no further approximations in computing the Laplace predictive distribution, since LCG converges to within machine precision. Thus, unlike inducing points methods like FITC or approximate methods like Nyström, our approach to inference gives the same answer as if we used standard Cholesky methods.

Pseudocode for our algorithm is shown in Algorithm 1. Given $K_1, \ldots, K_D$ where each matrix is $n^{1/D} \times n^{1/D}$, line 2 takes $O(D n^{2/D})$. Line 5 repeatedly applies Equation (A21), and matrix-vector multiplication $(K_d) v$ reduces to $D$ matrix-matrix multiplications $V K_j$ where $V$ is a matrix with $n$ entries total, reshaped to be $n^{D+1} \times n^{1/D}$. This matrix-matrix multiplication is $O(n^{D+1} n^{1/D} n^{1/D}) = O(n^{D+1})$ so the total run-time is $O(D n^{D+1})$. Line 7 is elementwise vector multiplication which is $O(n)$. Line 8 is calculated with LCG as discussed in Section 6 and takes $O(D n^{D+1})$. Lines 4 through 12 comprise the Newton update. Newton’s method typically takes a very small number of iterations $m \ll n$ to converge, so the overall run-time is $O(m D n^{D+1})$. Line 13 requires $D$
eigendecompositions of matrices $K_1, \ldots, K_D$ which takes time $O(Dn^{3\frac{D}{D+1}})$ as discussed in Section 6.4. Line 14 is elementwise vector multiplication and addition so it is $O(n)$. 

Algorithm 1 Kronecker GP Inference and Learning

1: **Input:** $\theta, \mu, K, p(y|f), y$
2: Construct $K_1, \ldots, K_D$
3: $\alpha \leftarrow 0$
4: repeat
5: $f \leftarrow K\alpha + \mu$  
   # Eq. (A21)
6: $W \leftarrow -\nabla \nabla \log p(y|f)$
7: Solve $Bz = W^{-\frac{1}{2}}b$ with CG  
   # Eq. (15)
8: $\Delta \alpha \leftarrow W^{-\frac{1}{2}}z - \alpha$  
   # Eq. (16)
9: $\hat{\xi} \leftarrow \arg \min_{\xi} \Psi(\alpha + \xi \Delta \alpha)$
10: $\alpha \leftarrow \alpha + \hat{\xi} \Delta \alpha$  
    # Line Search
11: until convergence of $\Psi$
12: $e = \text{eig}(K)$  
    # exploit Kronecker structure
13: $Z \leftarrow \alpha^T(f - \mu)/2 + \sum_i \log(1 + e_i W_i)/2 - \log p(y|f)$
14: **Output:** $f, \alpha, Z$

Overall, the runtime is $O(Dn^{\frac{D+1}{D}})$. There is no speedup for $D = 1$, and for $D > 1$ this is nearly linear time. This is much faster than the standard Cholesky approach which requires $O(n^3)$ time. The memory requirements are given by the total number of entries in $K_1, \ldots, K_D$: $O(Dn^{\frac{D}{D+1}})$. This is smaller than the storage required for the $n$ observations, so it is not a major factor. But it is worth noting because it is much less memory than required by the standard Cholesky approach of $O(n^2)$ space.

7 Model Specification

We propose to combine our fast Kronecker methods for non-Gaussian likelihoods, discussed in section 6, with Cox processes, which we introduced in section 3. We will use this model for crime rate forecasting in section 8.

With large sample sizes but little prior information to guide the choice of appropriate covariance functions, we turn to a class of recently proposed expressive covariance functions called Spectral Mixture (SM) kernels Wilson & Adams (2013). These kernels model the spectral density given by the Fourier transform of a stationary kernel ($k = k(\tau) = k(x - x')$) as a scale-location mixture of Gaussians. Since mixtures of Gaussians are dense in the set of all distribution functions and Bochner’s theorem shows a deterministic relationship between spectral densities and stationary covariances, SM kernels can approximate any stationary covariance function to arbitrary precision. For 1D inputs $z$, and $\tau = z - z'$, an SM kernel with $Q$ components has the form

$$k(\tau) = \sum_{q=1}^{Q} w_q \exp(-2\pi^2 \tau^2 v_q) \cos(2\pi \tau \mu_q).$$  \hspace{1cm} (20)

$w_q$ is the weight, $1/\mu_q$ is the period, and $1/\sqrt{v_q}$ is the length-scale associated with component $q$. In the spectral domain, $\mu_q$ and $v_q$ are the mean and variance of the Gaussian for component $q$. Wilson
et al. (2014) showed that a combination of Kronecker methods and spectral mixture kernels distinctly enables structure discovery on large multidimensional datasets – structure discovery that is not possible using other popular scalable approaches, due to the limiting approximations in these alternatives.

For our space-time data, in which locations $s$ are labeled with coordinates $(x, y, t)$, we specify the following separable form for our covariance function $k_{\theta}$:

$$k_{\theta}((x, y, t), (x', y', t')) = k_x(x, x')k_y(y, y')k_t(t, t')$$

where $k_x$ and $k_y$ are Matérn-5/2 kernels for space and $k_t$ is a spectral mixture kernel with $Q = 20$ components for time. We used Matérn-5/2 kernels because the spatial dimensions in this application vary smoothly, and the Matérn kernel is a popular choice for spatial data Stein (1999).

We also consider the negative binomial likelihood as an alternative to the Poisson likelihood. This is a common alternative choice for count data Hilbe (2011), especially in cases of overdispersion and we find that it has computational benefits. The GLM formulation of the negative binomial distribution has mean $m$ and variance $m + \frac{m^2}{r}$. It approaches the Poisson distribution as $r \to \infty$.

8 Experiments

We evaluate our methods on synthetic and real data, focusing on runtime and accuracy for inference and hyperparameter learning. We implemented our methods as an extension to the GPML framework Rasmussen & Nickisch (2010). We apply our methods to spatiotemporal crime rate forecasting, comparing with FITC, SSGPR Lázaro-Gredilla et al. (2010) and Kronecker methods with a Gaussian observation model.

8.1 Synthetic Data

To demonstrate the vast improvements in scalability offered by our method we simulated a realization from a GP on a grid of size $n \times n \times n$ with covariance function given by the product of three SM kernels. For each realization $f(s_i)$, we then drew $y_i \sim \text{NegativeBinomial}(\exp(f(s_i)) + 1)$. Using this as training data, we ran non-linear conjugate gradients to learn the hyperparameters that maximized the lower bound on the approximate marginal likelihood in equation (19), using the same product of SM kernels. We initialized our hyperparameters by taking the true hyperparameter values and adding random noise. We compared our new Kronecker methods to standard methods and FITC with varying numbers of inducing points. In each case, we used the Laplace approximation. We used 5-fold crossvalidation, relearning the hyperparameters for each fold and making predictions for the latent function values $f_i$ on the 20% of data that was held out. The average MSE and running times for each method on each dataset are shown in Figure 2. We also calculated the log-likelihood of our posterior predictions for varying numbers of observations $n$ for FITC-100, as shown in Table A3 in the Appendix. Our method achieved significantly higher predictive log-likelihood than FITC-100 for $n \geq 1000$.

In our final synthetic test, we simulated 100 million observations from a GP on an 8 dimensional grid, possibly the largest dataset that has ever been modeled with a Gaussian process. This is particularly exceptional given the non-Gaussian likelihood. In this case, we had a simple covariance structure given by a squared exponential (RBF) kernel with different length-scales per dimension. We successfully evaluated the marginal likelihood in 27 minutes.
Figure 2: Run-time and accuracy (mean squared error) of optimizing the hyperparameters of a GP with the Laplace approximation, comparing our new Kronecker inference methods to standard GP inference and FITC. The standard method approach has cubic running time. Each experiment was run with 5-fold crossvalidation but error bars are not shown for legibility. There is no significant difference between the standard and Kronecker methods in terms of accuracy. For grids of size $10 \times 10 \times 10$ observations and greater, FITC has significantly lower accuracy than Kronecker and standard methods.
8.2 Crime Rate Forecasting in Chicago

The City of Chicago makes geocoded, date-stamped crime report data publicly available through its data portal\(^3\). For our application, we chose crimes coded as “assault” which includes all “unlawful attacks” with a weapon or otherwise. Assault has a marked seasonal pattern, peaking in the summer. We used a decade of data from January 1, 2004 to December 31, 2013, consisting of 233,088 reported incidents of assault. We trained our model on data from the first 8 years of the dataset (2004-2011), and made forecasts for each week of 2012 and 2013. Forecasting this far into the future goes well beyond what is currently believed to be possible by practitioners.

LGCPs have been most widely applied in the 2-dimensional case, and we fit spatial LGCPs to the training data, discretizing our data into a \(288 \times 446\) grid for a total of 128,448 observations. Posterior inference and learned hyperparameter are shown in Section A.3 of the Appendix.

For our spatiotemporal forecasting, we used Spectral Mixture (SM) kernels for the time dimension, as discussed in section 7. Specifically, we consider \(Q = 20\) mixture components. For hyperparameter learning, our spatial grid was \(17 \times 26\), corresponding to 1 mile by 1 mile grid cells, and our temporal grid was one cell per week, for a total of 416 weeks. Thus, our dataset of 233,088 assaults was discretized to a grid of size 183,872. Both of these sample sizes far exceed the state-of-the-art in fitting LGCPs, and indeed in fitting most GP regression problems without extreme simplifying assumptions or approximations.

To find a good starting set of hyperparameters, we used the hyperparameter initialization procedure in Wilson et al. (2014) with a Gaussian observation model. We also rescaled counts by the maximum count at that location, log-transformed, and then centered so that they would have mean 0. We ran non-linear conjugate gradient descent for 200 iterations. Using the hyperparameters learned from this stage, we switched to the count data and a negative binomial likelihood. We then ran non-linear conjugate gradient descent for another 200 iterations to relearn the hyperparameters and also the variance of the negative binomial.

The spatial hyperparameters that we learned are \(\sigma^2 = 0.2231\), \(\lambda_1 = 0.11\) and \(\lambda_2 = 0.02\). This means that at this high resolution, with so much temporal data, there was little smoothing in space, with nearby locations allowed to be very different. Yet due to the multiplicative structure of our covariance function, our posterior inference is able to “borrow strength” such that locations with few observations follow a globally-learned time trend. We learned 60 temporal hyperparameters, and the spectral mixture components with the highest weights are shown in Figure 4, visualized in the covariance and frequency domains. We also show what posterior time series predictions would be if only a particular spectral component had been used, roughly giving an idea of the “explanatory” power of separate spectral components. We interpret the components, by decreasing weight, as follows: component 1 has a period and length-scale larger than the observation window thus picking up a decreasing trend over time. Components 2 (with period 1 month) and 4 pick up very-short-scale time variation, enabling the model to fit the observed data well. Component 3 picks up the yearly periodic trend (the spike in the spectral domain is at \(0.02 = \frac{1}{52.1}\)). Component 5 picks up a periodic trend with length longer than a year – 97 weeks, a feature for which we do not have any explanation. The exact hyperparameters are in Table A2 in the Appendix.

After learning the hyperparameters, we made predictions for the entire 8 years of training data and 2 years of forecasts. In Figure A6 in the Appendix we show the time series of assaults for 9 neighborhoods with our predictions, forecasts, and uncertainty intervals. Next, we redischertiz\(^3\)ed

\(^{3}\)http://data.cityofchicago.org
Figure 3: Local area posterior forecasts of assault one year into the future with the actual locations of assaults shown as black dots. The model was fit to data from January 2004 to December 2011, and the forecasts were made for the first week of June 2012 (left) and December 2012 (right).

Our original point pattern to a grid of size $51 \times 78$ ($n = 1.6$ million observations) and made spatial predictions 6 months and 1 year into the future, as shown in Figure 3, which also includes the observed point pattern of crimes. Visually, our forecasts are quite accurate. The accuracy and runtime of our method and competitors is shown in Table 1. The near 0 RMSE for predictions at the training data locations (i.e. the training error) for Kronecker Gaussian SM-20 indicates overfitting, while our model, Kronecker NegBinom SM-20, has a more reasonable RMSE of 0.79, out-performing the other models. The forecasting RMSE of our model was not significantly different than SSGPR or Kronecker Gaussian, while it outperformed FITC. But RMSE does not take forecasting intervals (posterior uncertainty) into account. Kronecker Gaussian and SSGPR had overly precise posterior estimates. Forecast log-likelihood is the probability of the out-of-sample data (marginalizing out the model parameters), so we can use it to directly compare the models, where higher likelihoods are better. The Kronecker Gaussian approach has the lowest forecast log-likelihood, and SSGPR has
the next lowest forecast log-likelihood. FITC was not overconfident, but its posterior forecasts were essentially constant. Our model has the highest forecast log-likelihood, showing a balance between a good fit and correct forecasting intervals. Kronecker Gaussian methods showed the fastest run-times due to the availability of a closed form posterior. FITC was very slow, even though we only used 100 inducing points.

Table 1: Our method, Kronecker NegBinom SM-20, uses Kronecker inference, a negative binomial observation model, and an SM kernel with 20 components. Kronecker Normal SM-20 uses a Gaussian observation model. FITC 100 uses the same observation model and kernel as Kronecker NegBinom SM-20 but uses FITC with 100 inducing points for inference. SSGPR-200 uses a Gaussian observation model and 200 spectral points.

<table>
<thead>
<tr>
<th></th>
<th>Kronecker NegBinom SM-20</th>
<th>Kronecker Gaussian SM-20</th>
<th>FITC-100 NegBinom SM-20</th>
<th>SSGPR-200</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training RMSE</td>
<td>0.79</td>
<td>10^{-11}</td>
<td>2.14</td>
<td>1.45</td>
</tr>
<tr>
<td>Forecast RMSE</td>
<td>1.26</td>
<td>1.28</td>
<td>1.77</td>
<td>1.26</td>
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<tr>
<td>Forecast log-likelihood</td>
<td>-33916</td>
<td>-352320</td>
<td>-42897</td>
<td>-82781</td>
</tr>
<tr>
<td>Run-time</td>
<td>2.8 hours</td>
<td>22 minutes</td>
<td>4.5 hours</td>
<td>2.8 hours</td>
</tr>
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</table>

8.3 Conclusion

We proposed a new scalable Kronecker method for Gaussian processes with non-Gaussian likelihoods, achieving near linear run-times for inference and hyperparameter learning. We evaluated our method on synthetic data, where it outperformed competing alternatives, and demonstrated its real-world applicability to the challenging problem of small area crime rate forecasting. Our kernel learning automatically discovered multiscale seasonal trends and our inference generated highly accurate long-range forecasts, with correct uncertainty intervals.
Figure 4: The five spectral mixture components with highest weights learned by our model are shown as a covariance (top) and spectral density (middle). In the bottom row, time series predictions were made on the dataset (ignoring space) using only that component. Red indicates out-of-sample forecasts.
References


A Appendix

A.1 Kronecker Algebra

We exploit the identity Steeb & Hardy (2011):

\[(B^T \otimes A)v = \text{vec}(AVB)\]  \hspace{1cm} (A21)

where \(v = \text{vec}(V)\) and the vec operator turns a matrix into a vector by stacking columns vertically. Since a full \(n \times n\) matrix is never formed, this approach is very efficient in terms of space and time complexity, relying only on operations with the smaller matrices \(K_i\) and the matrix \(V\) which only has \(n\) entries. We analyzed the complexity in Section 6.5. Another result we use is that given the eigendecompositions of \(K_d = Q_d \Lambda_d Q_d^T\), we have:

\[K = (\bigotimes Q_d)(\bigotimes \Lambda_d)(\bigotimes Q_d^T)\]  \hspace{1cm} (A22)

A.2 Supplementary Results

<table>
<thead>
<tr>
<th>(q)</th>
<th>Weight</th>
<th>Period</th>
<th>Length-scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>52.72</td>
<td>10813.9</td>
<td>133280.2</td>
</tr>
<tr>
<td>2</td>
<td>5.48</td>
<td>4.0</td>
<td>1.1</td>
</tr>
<tr>
<td>3</td>
<td>0.33</td>
<td>52.1</td>
<td>27700.8</td>
</tr>
<tr>
<td>4</td>
<td>0.05</td>
<td>22.0</td>
<td>1.6</td>
</tr>
<tr>
<td>5</td>
<td>0.02</td>
<td>97.4</td>
<td>7359.1</td>
</tr>
</tbody>
</table>

Table A2: The top five spectral mixture components learned for the temporal kernel in the LGCP fit to 8 years of assault data. The components are visualized in Figure 4 where component \(q\) corresponds to the row of the table.

<table>
<thead>
<tr>
<th>(N)</th>
<th>Standard</th>
<th>Kronecker</th>
<th>FITC-100</th>
</tr>
</thead>
<tbody>
<tr>
<td>125</td>
<td>-62.12</td>
<td>-61.52</td>
<td>-61.20</td>
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<tr>
<td>343</td>
<td>-157.47</td>
<td>-157.80</td>
<td>-159.21</td>
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<tr>
<td>1000</td>
<td>-445.48</td>
<td>-443.87</td>
<td>-455.84</td>
</tr>
<tr>
<td>1728</td>
<td>-739.56</td>
<td>-740.31</td>
<td>-756.95</td>
</tr>
<tr>
<td>8000</td>
<td>-3333.10</td>
<td>-3333.66</td>
<td>-3486.20</td>
</tr>
</tbody>
</table>

Table A3: Predictive log-likelihoods are shown corresponding to the experiment in Figure 2. A higher log-likelihood indicates a better fit. The differences between the standard and Kronecker results were not significant but the difference between FITC-100 and the others was significant (two-sample paired t-test, \(p \leq .05\)) for \(n \geq 1000\).

A.3 A two-dimensional LGCP

We used a product of Matérn-5/2 kernels: \(k_x(d)\) with length-scale \(\lambda_x\) and variance \(\sigma^2\) and \(k_y(d)\) with length-scale \(\lambda_y\) and variance fixed at 1: \(k((x, y), (x', y')) = k_x(|x - x'|)k_y(|y - y'|)\).
We discretized our data into a $288 \times 446$ grid for a total of 128,448 observations. Locations outside of the boundaries of Chicago – about 56% of the full grid—were treated as missing. In Figure A5 we show the location of assaults represented by dots, along with a map of our posterior intensity, log-intensity, and variance of the number of assaults. It is clear that our approach is smoothing the data. The hyperparameters that we learn are $\sigma^2 = 5.34$, $\lambda_x = 2.23$, and $\lambda_y = 2.24$, i.e., length-scales for moving north-south and east-west were found to be nearly identical for these data; by assuming Kronecker structure our learning happens in a fashion analogous to Automatic Relevance Determination Neal (1996).
Figure A5: We fit a log Gaussian Cox Process to the point pattern of reported incidents of assault in Chicago (a) and made posterior estimates of the intensity surface (b). The latent log-intensity surface is visualized in (c) and the posterior variance is visualized in (d).
Figure A6: We show the time series of weekly assaults in the nine neighborhoods with the most assaults in Chicago. The blue line shows our posterior prediction (training data, first 8 years of data) and forecast (out-of-sample, last 2 years of data, to the right of the vertical bar). Observed counts are shown as dots. 95% posterior intervals are shown in gray.