Scalable penalized spatiotemporal land-use regression for ground-level nitrogen dioxide

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Abstract

Nitrogen dioxide (NO\textsubscript{2}) is a primary constituent of traffic-related air pollution and has well-established harmful environmental and human-health impacts. Knowledge of the spatiotemporal distribution of NO\textsubscript{2} is critical for exposure and risk assessment. A common approach for assessing air pollution exposure is linear regression involving spatially referenced covariates, known as land-use regression (LUR). We develop a scalable approach for simultaneous variable selection and estimation of LUR models with spatiotemporally correlated errors, by combining a general-Vecchia Gaussian-process approximation with a penalty on the LUR coefficients. In comparisons to existing methods using simulated data, our approach resulted in higher model-selection specificity and sensitivity and in better prediction in terms of calibration and sharpness, for a wide range of relevant settings. In our spatiotemporal analysis of daily, US-wide, ground-level NO\textsubscript{2} data, our approach was more accurate, and produced a sparser and more interpretable model. Our daily predictions elucidate spatiotemporal patterns of NO\textsubscript{2} concentrations across the United States, including significant variations between cities and intra-urban variation. Thus, our predictions will be useful for epidemiological and risk-assessment studies seeking daily, national-scale predictions, and they can be used in acute-outcome health-risk assessments.

Keywords: general Vecchia approximation; spatial statistics; Gaussian process; variable selection; air pollution; Kriging

1 Introduction

Nitrogen dioxide (NO\textsubscript{2}) is a primary constituent of traffic-related air pollution and has well-established harmful environmental and human-health impacts (US Environmental Protection Agency, 2016). For example, exposure to NO\textsubscript{2} is associated with increased all-cause mortality (Hoek et al., 2013), myocardial infarction (Rosenlund et al., 2006, 2009), coronary heart disease (Rosenlund et al., 2008), cardiovascular events (Alexeeff et al., 2018), asthma (Gauderman et al., 2005), autism spectrum disorders (Volk et al., 2013), and impaired neurological...
development and other neurological disorders (Xu et al., 2016). Additionally, atmospheric oxides of nitrogen, including \( \text{NO}_2 \), are precursors to hazardous acid rain (Schindler, 1988), tropospheric ozone (US Environmental Protection Agency, 1999), fine particulate matter (PM\(_{2.5}\)) (US Environmental Protection Agency, 1999), and can result in negative ecological (Schindler, 1988) and economic impacts (Mauzerall et al., 2005).

Knowledge of the spatiotemporal distribution of \( \text{NO}_2 \) is critical for assessing exposure and subsequent risks. A common approach for assessing exposure to outdoor air pollution is linear regression involving spatially referenced covariates, known as land-use regression (LUR). There are many strengths in current implementations of LUR models. First, is the ability to predict a variable of interest in space and time at unmonitored coordinates, including uncertainty quantification. Second, is the use of readily-available, large geospatial datasets such as satellite imagery and census information. Third, is the elucidation and interpretation of coefficients that are possible with linear models, which allows for meaningful policy discussions around factors affecting the distribution of exposure and risk.

Assuming independent and identically distributed (iid) errors, LUR has been implemented for air-quality-exposure modeling of \( \text{NO}_2 \) (Briggs et al., 1997; Hoek et al., 2008; Su et al., 2009; Novotny et al., 2011; Ross et al., 2013; Knibbs et al., 2014; Larkin et al., 2017; de Hoogh et al., 2018) and other air pollutants such as PM\(_{2.5}\) (Henderson et al., 2007; Moore et al., 2007; Ross et al., 2013). Typically, LUR involves model selection or dimension reduction on a large candidate-set of spatially referenced covariates. For example, LUR has been implemented with stepwise model selection for \( \text{NO}_2 \) (Briggs et al., 1997; Su et al., 2009; Novotny et al., 2011; Ross et al., 2013; Knibbs et al., 2014; de Hoogh et al., 2018) and partial-least-squares dimension reduction for \( \text{NO}_2 \) (Young et al., 2016) and PM\(_{2.5}\) (Sampson et al., 2013). \( \text{NO}_2 \) LUR models have also employed penalization-based model-selection methods such as the LASSO (Knibbs et al., 2014; Larkin et al., 2017). Additionally, LUR prediction residuals are often integrated into geostatistical models, such as Kriging (Wu et al., 2013; de Hoogh et al., 2018) and Bayesian maximum entropy (Coulliette et al., 2009; Messier et al., 2012; Beckerman et al., 2013; Reyes and Serre, 2014; Messier et al., 2014, 2015), in a two-stage approach with the goal of improving prediction accuracy.

While LURs have undoubtedly been useful for many exposure and risk assessment studies, the assumption of iid errors is usually violated, because the spatial dependence in the response cannot be captured fully by the covariates, resulting in biased covariate estimates and decreased sensitivity and specificity in the model-selection process. An exception to this case is Holcomb et al. (2018), which implemented backwards model selection in a full Kriging model, but this approach is not feasible for large data sets. Guan et al. (2020) implemented a scalable approach with LUR with spatiotemporal errors, but used principal components instead of model selection to reduce the number of covariates.

In spatial statistics and Gaussian-process modeling, many approaches have been proposed to ensure scalability to large datasets (see, e.g., Heaton et al., 2019; Liu et al., 2020 for recent reviews and comparison), but the focus is often more on prediction based on the (residual) covariance structure, and less on penalized selection from among a large number of spatial or spatiotemporal covariates. Perhaps the most promising approaches for scalable spatial prediction are based on the ordered conditional approximation of Vecchia (1988); here we use and extend the general Vecchia approximation (Katzfuss and Guinness, 2019, Katzfuss et al., 2020a), which is highly accurate, can guarantee linear complexity with respect to the
sample size, and includes many existing Gaussian-process approximations as special cases (e.g., Vecchia, 1988; Snelson and Ghahramani, 2007; Finley et al., 2009; Sang et al., 2011; Datta et al., 2016a; Katzfuss, 2017; Katzfuss and Gong, 2020).

We develop an approach for simultaneous variable selection and estimation of LUR models with spatiotemporally correlated errors, extending the general Vecchia approximation to ensure scalability to large datasets. The resulting dependent-error regression problem can be transformed into standard iid-error regression involving pseudo-data, which can be computed rapidly using Vecchia. This approach can be combined with any existing method for fitting penalized regression models with independent errors, such as least-angle regression (Efron et al., 2004) for LASSO-type L1 penalties (Tibshirani, 1996), and coordinate descent (Breheny and Huang, 2011) for non-convex (e.g., smoothly clipped absolute deviation) penalties (Fan and Li, 2001). The ordering and conditioning-set selection necessary for the Vecchia approximation is carried out based on appropriately scaled spatiotemporal coordinates. All computations necessary for inference scale linearly in the data size for fixed tuning parameters.

The remainder of this article is organized as follows: Section 2 describes the daily, ground-level NO\textsubscript{2} data and the geographic covariates. Section 3 provides a description of LUR with penalization. Section 4 presents our proposed methodology based on the general Vecchia approximation to Kriging models with SCAD penalty. Section 5 compares approaches in simulation studies. In Section 6, we apply our method to the NO\textsubscript{2} concentrations and discuss the results. Section 7 highlights the main conclusions and discusses areas for future research. Further details can be found in Appendices A–B. A separate Supplementary Material document contains Sections S1–S3 with additional plots and details. Code based on the R package GPvecchia (Katzfuss et al., 2020b) for the proposed approach and this study is available at https://github.com/NIEHS/LURK-Vecchia.

2 Ground-level NO\textsubscript{2} data

We consider daily ground-level (i.e., tropospheric) NO\textsubscript{2} concentrations across the conterminous United States, monitored and distributed by the United States Environmental Protection Agency (USEPA) Air Quality System (AQS) (US Environmental Protection Agency, 2019). The date range for our study was July 10, 2018 to May 1, 2019, based on the availability of geographic covariates, primarily the TROPOMI real-time satellite imagery. The final NO\textsubscript{2} dataset contained 76,748 unique spatiotemporal observations distributed across 459 monitoring sites (Figure 1).

2.1 Geographic covariates

For our analysis, we calculated 139 spatial and spatiotemporal geographic covariates representing possible NO\textsubscript{2} sources and attenuation factors. A key characteristic of our and the majority of LUR studies is the presence of highly correlated covariates. In particular, many covariates only differ by their spatial resolution. After all of the covariates are calculated, each covariate is standardized to mean 0 and variance 1. The following subsections explain how each potential covariate was calculated.
2.1.1 TROPOMI

We utilized data from the TROPOspheric Monitoring Instrument (TROPOMI) to calculate many satellite-based spatiotemporal covariates. TROPOMI is the sensor on-board the Copernicus Sentinel-5 Precursor satellite. The TROPOMI-based covariate calculations are performed in Google Earth Engine, a cloud platform for earth observation data analysis that combines a public data catalog with a large-scale computational facility optimized for parallel processing of geospatial data.

TROPOMI provides output (i.e., Level-2 or L2 products) representing atmospheric air pollution and physical properties with a spatial resolution of approximately 3.5 by 7 km. We calculated daily mean values within 1, 10, and 100 km circular buffers for the following TROPOMI L2 products. Note the 1 km buffer is equivalent to the coincident TROPOMI value at the location of the monitor: Total vertical column NO\textsubscript{2} (mol-m\textsuperscript{-2}), tropospheric vertical column NO\textsubscript{2} (mol-m\textsuperscript{-2}), NO\textsubscript{2} slant column density (mol-m\textsuperscript{-2}), tropopause (i.e. boundary between troposphere and stratosphere) pressure (Pa), absorbing aerosol index (AAI; dimensionless), cloud fraction, and the solar azimuth angle (degrees). We used the near real-time TROPOMI product, if available, for the estimation of models. This was the driving factor for the sparsity of the data set within the study range. For prediction, if the real-time data were unavailable, we used the offline data. The near real-time data are available sooner and have small differences with the offline data [Boersma et al., 2007]. Lastly, we used a simple average from 10 nearest-neighbor spatiotemporal coordinates if neither near real-time nor offline were observed at a spatiotemporal coordinate. Potential alternatives for interpolating missing TROPOMI data include longer-time-scale moving-window averages (e.g. monthly) or developing a predictive model [de Hoogh et al., 2019]. The final covariate dataset included 21 TROPOMI-based variables.

2.1.2 Meteorology

Spatial and daily time-resolved meteorological covariates were calculated in Google Earth Engine using the University of Idaho Gridded Surface Meteorological dataset (GRIDMET)
Abatzoglou et al., 2014). GRIDMET provides daily surface fields at approximately 4 km resolution. We calculated average daily values inside 1, 10, and 100 km buffers for the following variables: precipitation (mm), maximum relative humidity (percent), specific humidity (kg-kg$^{-1}$), surface downward shortwave radiation (W-m$^{-2}$), maximum temperature (K), and wind velocity (m-s$^{-1}$). The 1 km buffer is equivalent to choosing the containing grid cell. The final covariate dataset included 18 meteorology-based variables.

### 2.1.3 Vegetative indices

Spatial covariates of vegetative indices were calculated in Google Earth Engine using the MODIS/Terra Vegetative Indices 16-Day L3 Global 500 m SIN Grid (NASA/METI/AIST/Japan SpaceSystems, and U.S./Japan ASTER Science Team, 2019). We calculated spatial averages of the normalized difference vegetative index (NDVI) and the enhanced vegetation index (EVI) in 1, 10, and 100 km circular buffers. The final covariate dataset included 6 vegetative index based variables.

### 2.1.4 Population, traffic, and roads

Population density (people-km$^{-2}$) was calculated in Google Earth Engine from the Gridded Population of World Version 4 (Center for International Earth Science Information Network - CIESIN - Columbia University, 2018). Average population (2015-equivalent) density was calculated in 1, 10, and 100 km circular buffers.

A surrogate for traffic was calculated using the University of Oxford Malaria Atlas Project global travel friction dataset (Weiss et al., 2018). Average travel friction (min-m$^{-1}$), or travel time, was calculated in 1, 10, and 100 km circular buffers.

Road length variables were calculated in ArcMap 10.6.1 and MATLAB R2018a using the ESRI major roads shapefile (Esri, TomTom North America, Inc). The road length in 1, 10, and 100 km circular buffers was calculated for the following road classifications (FRC code in ESRI shapefile): All roads classes, highway (0), major roads (1,2), and secondary roads (3,4,5). The final covariate dataset included 18 population, traffic, or road variables.

### 2.1.5 Land cover

Spatial land-cover attributes were calculated in Google Earth Engine from the National Land Cover Database (Homer et al., 2015). The percent of each land cover class (e.g. water, low developed, deciduous trees, etc.) were calculated in 1, 10, and 100 km circular buffers.

Average elevation was calculated in 1, 10, and 100 km circular buffers using the Japan Aerospace Exploration Agency Advanced Land Observing Satellite global digital surface model with a horizontal resolution of approximately 30 meters (Tadono et al., 2014). The final covariate dataset included 48 land-cover or elevation variables.

### 2.1.6 National Emissions Inventory

Data on point source emissions was downloaded from the USEPA National Emissions Inventory (US Environmental Protection Agency, 2017) for the year 2017. Following Messier
we assume a linear relationship between the response and covariates, pairs $S$ of the $p$ in Section 2.1. We have the values $x_i = x(s_i, t_i) = (x_1(s_i, t_i), \ldots, x_p(s_i, t_i))'$ of the $p = 139$ covariates (described in Section 2.1) at the same (space, time)-coordinate pairs $S = \{(s_1, t_1), \ldots, (s_n, t_n)\}$. Spatial-only covariates are repeated in time as needed. We assume a linear relationship between the response and covariates,

$$z_i = x_i' \beta + \epsilon_i = x_i' \beta + \eta_i + \delta_i = y_i + \delta_i, \quad i = 1, \ldots, n,$$

where $\epsilon_i = \eta_i + \delta_i$ is the regression error consisting of a spatiotemporally dependent component $\eta_i = \eta(s_i, t_i)$ and an independent measurement-noise component $\delta_i \sim N(0, \tau^2)$, $i = 1, \ldots, n$, and $y_i = z_i – \delta_i = x_i' \beta + \eta_i$ is the (noise-free) true log-NO$_2$. We assume that $\eta(\cdot) \sim \mathcal{GP}(0, C_{\theta})$ follows a Gaussian process with covariance function $C_{\theta}$. Throughout, we assume a non-separable spatiotemporal exponential covariance function, $C_{\theta}((s_i, t_i), (s_j, t_j)) = \sigma^2 \exp(-d_{\theta}((s_i, t_i), (s_j, t_j)))$, where

$$d_{\theta}((s_i, t_i), (s_j, t_j)) = \sqrt{\frac{|s_i – s_j|^2}{\gamma_s^2} + \frac{(t_i – t_j)^2}{\gamma_t^2}},$$

and $\theta = (\sigma, \gamma_s, \gamma_t, \tau)$ contains the unknown parameters in the model. We also considered and dismissed a Matérn covariance with estimated smoothness parameter, as this resulted in a smoothness parameter near that of the exponential model (i.e., 0.5), nearly identical negative loglikelihood values, and increased model run time due to evaluation of Bessel functions and larger parameter space. Note that standard LUR models (e.g., Briggs et al. 1997; Hoek et al. 2008; de Hoogh et al. 2018) do not include the dependent component $\eta(\cdot)$ and assume the error terms $\epsilon_1, \ldots, \epsilon_n$ to be iid.

Stacking the quantities in (1), we obtain the regression model

$$z = X\beta + \epsilon, \quad \epsilon \sim \mathcal{N}_n(0, \Sigma_{\theta}),$$

where $\Sigma_{\theta} = C_{\theta} + \tau^2 I_n$, with $C_{\theta} = (C_{\theta}((s_i, t_i), (s_j, t_j)))_{i,j=1,\ldots,n}$. Equivalently, we can write this in terms of a multivariate Gaussian density for the response,

$$f(z; \beta, \theta) = \mathcal{N}_n(z|X\beta, \Sigma_{\theta}).$$

The goal is to estimate the $p$-vector $\beta$ and determine its nonzero elements, which also requires estimation of the covariance parameters $\theta$. Further, given parameter estimates $\hat{\beta}$ and $\hat{\theta}$, we would like to predict the process $y(\cdot)$ at unobserved coordinates.
A standard approach for parameter estimation is to maximize the likelihood in (3) with respect to the parameters $\beta$ and $\theta$. However, we have a large number $p = 139$ of (correlated) covariates, which makes the least-squares or maximum likelihood estimates of $\beta$ unstable.

To alleviate this issue, and to be able to select certain variables and set the coefficients corresponding to the other variables to zero, we instead consider optimizing an objective function consisting of the negative loglikelihood plus a penalization term $p(\beta)$ on $\beta$:

$$Q(\beta, \theta) = -2 \log f(z; \beta, \theta) + \lambda p(\beta) = (z - X\beta)'\Sigma^{-1}_\theta(z - X\beta) + \log |\Sigma_\theta| + \lambda p(\beta),$$  \hspace{1cm} (4)

where $\lambda$ is a shrinkage or tuning parameter, and we have omitted an additive constant in the last equation. In our numerical examples and application, we will use the popular non-convex, smoothly clipped absolute deviation (SCAD) penalty (Fan and Li, 2001):

$$p(\beta) = \begin{cases} 
\lambda|\beta|, & \text{if } |\beta| \leq \lambda, \\
2a\lambda|\beta| - \beta^2 - \lambda^2 \over 2(a-1), & \text{if } \lambda < |\beta| \leq a\lambda, \\
\lambda^2(a+1) \over 2, & \text{otherwise},
\end{cases}$$  \hspace{1cm} (5)

where $a = 3.7$, a popular choice that performs comparably to values based on generalized cross-validation (Fan and Li, 2001). We use the SCAD penalty for its oracle property, but other penalties can be easily be swapped in our framework. Li and Sudjianto (2005) demonstrate that a SCAD-penalized likelihood as in (4) and (5) reduces the variance in the estimates of $\theta$; however, their discussion did not address model selection or large sample sizes.

4 Our methodology

4.1 A general Vecchia approximation of the objective function

Evaluation or optimization of the objective function $Q$ in (4) requires decomposition of the $n \times n$ covariance matrix $\Sigma_\theta$ for many different values of $\theta$, each of which takes $O(n^3)$ time. This is computationally infeasible for the large $n = 76,748$ in our application.

Hence, we will extend the sparse general Vecchia (SGV) approximation (Katzfuss and Guinness, 2019), which we briefly review here, with more details given in Appendix A. SGV applies the approximation of Vecchia (1988) to the vector $u = (y_1, z_1, \ldots, y_n, z_n)'$, which interweaves the latent true-process realizations $y_1, \ldots, y_n$ and the observed noisy data $z_1, \ldots, z_n$. This approximation essentially replaces the conditioning sets in the exact factorization $f(u) = \prod_{j=1}^{2n} f(u_j|u_1, \ldots, u_{j-1})$ by small subsets, resulting in the approximation

$$\hat{f}(u) = \prod_{i=1}^{2n} p(u_i|u_{g(i)}) = N_{2n}(u|(X \otimes 1_2)\beta, (UU')^{-1})$$  \hspace{1cm} (6)

where each $g(i) \subset (1, \ldots, i - 1)$ is a conditioning index set of size $|g(i)| \leq m$, $\otimes$ is the Kronecker product, $1_2$ is a vector consisting of two 1s, and $U = U_\theta$ is a sparse upper triangular matrix whose nonzero entries can be computed easily based on $C_\theta$ and $\tau^2$. Recent results (Schäfer et al., 2020) indicate that the approximation error can be bounded with
the conditioning-set size $m$ increasing logarithmically in $n$ in some settings; in practice, $m \approx 30$ is often sufficient for accurate approximations. We further define $A$ and $B$ as the submatrices of $U$ consisting of the odd- and even-numbered rows of $U$, corresponding to $y$ and $z$, respectively. Then, $W = AA'$ is the implied posterior precision matrix of $y$ given $z$, and we define $V$ as the Cholesky factor based on reverse row-column ordering of $W$.

Our approximation $\hat{f}(u)$ is an extension of the SGV approach for spatial processes described in Katzfuss and Guinness (2019); to approximate the spatiotemporal covariance function $C_\theta$, we modify the ordering and conditioning scheme here to be carried out based on the scaled spatiotemporal distance $\bar{s}$, which depends on unknown parameters and must be updated along with the parameters. Again, more details are given in Appendix A.

The SGV approximation of the density of $u = (y_1, z_1, \ldots, y_n, z_n)'$ in (6) implies an approximation of the distribution for the response:

$$\hat{f}(z) = \int \hat{f}(u) dy,$$

which is also multivariate normal. This concludes our review of Katzfuss and Guinness (2019). Plugging the approximation $\hat{f}(z)$ into (4) results in the Vecchia objective function

$$\hat{Q}(\beta, \theta) = -2 \log \hat{f}_{\beta, \theta}(z) + \lambda p(\beta),$$

where we have now made explicit the dependence of the distribution of $z$ on the parameters $\beta$ and $\theta$. We will use and optimize the Vecchia objective function $\hat{Q}(\beta, \theta)$ in the remainder of the manuscript.

4.2 Inference

The most straightforward way to optimize the objective function is to optimize iteratively with respect to $\theta$ and $\beta$, while holding the respective other parameter vector fixed. In practice, it is usually sufficient to do this just a small number of times, after which there is little change in the parameter values.

As we prove in Appendix B, $\hat{Q}(\beta, \theta)$ in (7) can be written as

$$\hat{Q}(\beta, \theta) = \|\tilde{z}_\theta - \tilde{X}_\theta \beta\|^2_2 + \lambda p(\beta) - 2 \sum_i \log ((U_\theta)_{ii}) + 2 \sum_i \log ((V_\theta)_{ii}),$$

where $\tilde{z}_\theta = B'z + A'W^{-1}AB'z$, $\tilde{X}_\theta = B'X + A'W^{-1}AB'X$, and $V_\theta$ depend on $\theta$ through the matrices $A$, $B$, and $W$ computed from $U = U_\theta$.

4.2.1 Estimation of the regression coefficients

Optimizing $\hat{Q}(\beta, \theta)$ in (8) with respect to $\beta$ for fixed $\theta = \hat{\theta}$ is equivalent to solving a standard penalized regression problem with iid errors, except based on the pseudo-data $\tilde{z}_\theta$ and $\tilde{X}_\theta$.

To develop some intuition, consider briefly the case $m = n - 1$, in which case the approximation $\hat{f}(u)$ in (6) becomes exact. Then, the pseudo-data $\tilde{z}_\theta$ are obtained by first creating an augmented data vector of length $2n$ consisting of $z$ and $E(y | z)$, and then transforming this vector to a vector of iid normal variables based on the joint distribution or covariance
matrix of $z$ and $y$. Interestingly, the resulting inference on $\beta$ is unchanged relative to simply transforming the data $z$ alone, as is often done for general linear models. In the case of small $m$, our sparse general Vecchia approximation allows us to carry out this inference on $\beta$ more accurately based on the first approach.

For example, in the case of the SCAD (Fan and Li 2001) or L1 penalties (Tibshirani 1996), solution paths of optimal $\beta$ values for each value of $\lambda$ can be computed rapidly using coordinate descent (Brehei2011) or least angle regression (Efron et al. 2004), respectively. We select the optimal $\lambda$ and the corresponding $\hat{\beta}$ based on the lowest cross-validated mean square error, which can be performed in many software packages such as ncvreg (Brehei2011) or glmnet (Friedman et al. 2010). Brehei2011 demonstrated for high-dimensional problems that the SCAD penalty estimated with coordinate descent, in combination with cross-validation, leads to the global minimum solution as it likely resides in the locally convex region of $\lambda$.

We considered a simple example to illustrate how quickly the Vecchia solution can converge to the exact solution in the estimation of trend parameters, $\beta$. We simulated $p = 5$ correlated covariates with correlations ranging from .45 to .82, set the true $\beta = 0$, and simulated data with spatially dependent error at $n = 500$ locations. We then computed the exact generalized least-squares (GLS) estimates and the Vecchia GLS estimates implied by the pseudo-data in (8). As shown in Figure 2, the Vecchia solution quickly approached the exact solution going from $m = 0$ (i.e., assuming independent errors) to $m = 10$.

4.2.2 Estimation of the covariance parameters

Defining $\epsilon_\beta = z - X\beta$, (8) can be rearranged to yield

$$\hat{Q}(\beta, \theta) = \|B^T\epsilon_\beta\|^2 - \|V^{-1}AB^T\epsilon_\beta\|^2 - 2 \sum_i \log U_{ii} + 2 \sum_i \log V_{ii} + \lambda p(\beta),$$

(9)

where $V$, $A$, and $B$ implicitly depend on $\theta$ through $U = U_\theta$. Note that this expression is an extension of the Vecchia loglikelihood in Katz2019; we replaced their zero-mean data $z$ with our residuals $\epsilon_\beta$, and we have added the penalization term $\lambda p(\beta)$. This alternative expression of the objective function is important, as it avoids having to compute the pseudo-data for every evaluation as in (8).
For a given $\beta = \hat{\beta}$, (9) can be evaluated cheaply for any given parameter value $\theta$, and hence we can optimize the objective function with respect to $\theta$ using standard numerical optimization algorithms (e.g., Nelder-Mead). As in most Gaussian-process models, there is no guarantee that this procedure will find the global optimum, but we have not observed any negative consequences. Similarly, we have not observed any numerical issues due to nonidentifiability between the variance and range parameters, which theoretically holds under in-fill asymptotics (Zhang, 2004; Tang et al., 2019) but not under the increasing-domain asymptotics that may be more appropriate for our real-data application with small effective ranges relative to the domain size.

We monitor convergence of the overall algorithm by considering the minimum value of (9) achieved at each iteration.

### 4.2.3 Prediction

Often, interest is in prediction of the noise-free process $y_P$ at a set of $n_P$ spatiotemporal coordinates $S_P$, which is equivalent to obtaining the conditional distribution of $y_P$ given the data $z$. To do so, we extend the response-first full-conditioning (RF-full) approach of Katzfuss et al. (2020a), which essentially consists of a general Vecchia approximation $\hat{f}(\tilde{u})$, similar to (6), but now applied to the vector $\tilde{u} = (z', y_{all}')$, where $y_{all} = (y', y_P')$. We have

$$\hat{f}(y_{all}|z) = \frac{\hat{f}(z, y_{all})}{\int \hat{f}(z, y_{all})dy_{all}} = \mathcal{N}_n(\mu_{all}, W_{all}^{-1}).$$

Any quantities of interest can be extracted from this joint distribution, after computing $V_{all}$ as the Cholesky factor based on reverse row-column ordering of $W_{all}$. For example, the prediction mean, also referred to as the kriging predictor, is obtained by subsetting the vector $\mu_{all} = X_{all}\hat{\beta} - (V_{all}')^{-1}V_{all}A_{all}B_{all}'(z - X\hat{\beta}) = (\mu', \mu_P')'$, where $X_{all} = (X', X_P')'$, while the prediction or kriging variances are given by a subset of the diagonal elements of $W_{all}^{-1}$, which can be obtained using selected inversion based on the Takahashi recursions for $V_{all}$. Note that we ignore uncertainty in $\hat{\beta}$ in the predictions, but we conducted experiments that showed this uncertainty is often small relative to the uncertainty in $\eta(\cdot)$. If prediction of $z_P$ is desired, we simply need to add $\tau^2$ to the prediction variances. Our approximation is an extension of the spatial RF-full prediction in Katzfuss et al. (2020a), in that the nonzero mean has to be added and subtracted in the kriging predictor, and we carry out the ordering and conditioning in the scaled spatiotemporal domain. Details are given in Appendix A.

### 4.2.4 Complexity

Our proposed inference procedure is summarized in Algorithm 1. If each conditioning index vector in our Vecchia approximations is at most of size $m$, SGV and RF-full ensure that $U$, $V$, and $V_{all}$ are all highly sparse with at most $m$ nonzero off-diagonal entries per column. As a consequence, for fixed $m$ and $p$, our entire inference procedure requires linear time in the number of observed and prediction coordinates.

More precisely, assuming that $m, p, n_P \leq n$, evaluation of the likelihood and prediction for each parameter value requires $O(nm^3)$ time, coordinate descent for SCAD requires $O(np)$
Algorithm 1: LURK-Vecchia: Land-use regression Kriging with Vecchia approx.

Input: \( z \), \( S \), \( X \), \( S_P \), \( X_P \), \( C_\theta \), \( \text{tol} \)

Result: Parameter estimates \( \hat{\beta} \) and \( \hat{\theta} \); prediction \( \hat{f}(y_{all}|z) \)

1: Initialize \( \hat{\beta}, \hat{\theta}, \text{prev.objective} = \infty \), and \( \text{converged} = \text{FALSE} \)
2: OC: Maxmin ordering and nearest-neighbor conditioning for \( S \) based on \( d_{\hat{\theta}} \) in (2)
3: While \( \text{converged} = \text{FALSE} \) do
4: Compute \( \hat{\theta} = \arg \max_{\theta} \hat{Q}(\hat{\beta}, \theta) \) based on (9) using OC
5: Update OC based on \( d_{\hat{\theta}} \) in (2)
6: Compute pseudo-data \( \tilde{z}_{\hat{\theta}} \) and \( \tilde{X}_{\hat{\theta}} \) as in (8) using OC
7: Estimate \( \beta \) using standard (iid) SCAD based on \( \tilde{z}_{\hat{\theta}} \) and \( \tilde{X}_{\hat{\theta}} \) (see Section 4.2.1)
8: \( \text{new.objective} = \hat{Q}(\hat{\beta}, \hat{\theta}) \)
9: If \( \text{new.objective} > (\text{prev.objective} \times (1-\text{tol})) \) then
10: \( \text{converged} = \text{TRUE} \)
11: Else
12: \( \text{prev.objective} = \text{new.objective} \)
13: End if
14: End while
15: OC_P: Ordering and conditioning for \( S \) and \( S_P \) based on \( d_{\hat{\theta}} \) (see Appendix A)
16: Prediction: Compute relevant summaries of \( \hat{f}(y_{all}|z) \) using \( \text{OC}_P \) (see Section 4.2.3)

Time per iteration, one triangular solve involving \( V \) requires \( O(nm) \) time, and hence computing the pseudo-data \( \tilde{z}_{\hat{\theta}} \) and \( \tilde{X}_{\hat{\theta}} \) requires \( O(nmp) \) time. If we require \( L \) iterations going back and forth between estimating \( \beta \) and \( \theta \), \( L_{\theta} \) iterations to estimate \( \theta \) given \( \hat{\beta} \), and \( L_{\beta} \) iterations in the coordinate descent (including selecting tuning parameters using cross-validation) to estimate \( \beta \) given \( \hat{\theta} \), the overall cost of our algorithm is \( O(nL(L_{\theta}m^3 + L_{\beta}p)) \). We utilize a tolerance \( \text{tol} = 10^{-6} \) for the stopping criterion; a less stringent tolerance may be used, which will likely result in a smaller number of iterations \( L \) and a less accurate approximation of \( \beta \) and \( \theta \).

5 Simulation study

5.1 Simulation scenarios

We sampled 2,000 spatiotemporal coordinates from the possible combinations of 276 unique days and 50 unique spatial locations randomly distributed across the United States. The unique days correspond to the set of dates with complete geographic covariate datasets. The 2,000 coordinates are randomly divided in half for a training set of size \( n = 1,000 \), and a test set of size \( n_P = 1,000 \) that was never used in any model development.

For the spatiotemporal regression errors \( \epsilon_i \) in (11), we specified a baseline (minimum; maximum) scenario of the model, with spatial range parameter \( \gamma_s = 1,000 \) (200; 3,000) km, temporal range \( \theta_t = 30 \) (7; 365) days, total variance (i.e., sill) \( \sigma^2_{\text{total}} = \sigma^2 + \tau^2 = s^2_{\text{trend}} \) \((0.5 \times s^2_{\text{trend}}; 5 \times s^2_{\text{trend}})\), and nugget-to-sill ratio \( \tau^2 / \sigma^2_{\text{total}} = 0.25 \) (0.01; 0.99), where \( s^2_{\text{trend}} \) is the sample variance of the entries of the regression term \( X\beta \), evaluated at the true value of \( \beta \) (see
The nugget-to-sill ratio is the ratio of the noise to total variance. We considered a large number of simulation scenarios in which we varied, in turn, each of these variables, while holding the other variables fixed at their baseline levels. (Results for additional scenarios in which the variables varied jointly, including a spatial range of 30 km, are shown in Section S1.)

For the simulated spatiotemporal coordinates, we created spatiotemporal covariate matrices $X$ and $X_P$ based on the methods described for the NO$_2$ data in Section 2, in order to obtain a realistic simulation setting. To provide a unique set of covariates from the NO$_2$ analysis, we included and removed variables as follows. Ozone TROPOMI satellite data (air mass factor (AMF), total column, and slant) were included with 1, 10, and 100 km buffers. Randomly generated point sources with isotropic exponentially decaying contribution with decay ranges of 1, 10, and 100 km (Messier et al., 2012) and randomly generated spatiotemporal random fields were included in lieu of the NEI and road covariates from the NO$_2$ dataset. The final candidate set for the simulation included 123 potential covariates. The true trend coefficients $\beta$ were assumed to have 8 non-zero coefficients: NO$_2$ Slant 1 km, cloud fraction 10 km, ozone AMF 100 km, precipitation 10 km, NDVI 100 km, developed high intensity 1 km, point sources with 100 km decay range, and a smoothly varying spatiotemporal random field; the corresponding true coefficient values were 5, 5, 3, -3, -5, 10, 3, 5, respectively. The true covariates exhibited low to moderate correlation ($|\rho| \leq 0.41$) with the other true covariates, and low to extremely high correlation ($|\rho| \leq 0.99$) with the extraneous covariates.

5.2 Approaches under comparison

We compared our proposed method to several popular land-use regression approaches:

LUR-iid: An iid land-use regression model, which can be viewed as a special case of (1) with $\eta(\cdot) \equiv 0$. Point predictions are then simply given by $X_P\hat{\beta}$.

LURK-Local: Land-use regression Kriging with a local neighborhood, which is the current state-of-the-art approach in land-use regression (de Hoogh et al., 2018). LURK-local consists of the following steps:

1. Estimate $\hat{\beta}$ as in LUR-iid, and compute residuals $\epsilon_\beta$.
2. Estimate $\theta$ as an average of estimates based on $k = 10$ samples of size $l = (mn_k)^{1/3}$ from $\epsilon_\beta$.
3. Carry out local kriging at each prediction coordinate using the $m$ nearest (in terms of (2)) space-time neighbors among $\epsilon_\beta$.

LURK-Vecchia: Our proposed methodology, summarized in Algorithm 1.

LURK-Full: The full Kriging and SCAD penalized method based on (3). This is equivalent to the proposed LURK-Vecchia approach with $m = n - 1$.

Local-Kriging: Does not use geographic covariates. Similar to LURK-local, we estimate $\theta$ as an average of estimates based on $k = 10$ samples of size $l = (mn_k)^{1/3}$ from $z$, and then make predictions using the $m$ nearest spatiotemporal observations.
LURK-Full can be considered the most accurate approach, but it is computationally infeasible for large \( n \) (in the tens of thousands or more). For the other spatiotemporal approaches, we ensure similar computational complexity by using the same \( m = 25 \).

### 5.3 Prediction scores

For the prediction at unobserved coordinates, we considered three proper scoring rules (e.g., \cite{Gneiting2014}) that all compare the true simulated test data \( y_P^\star \) to the predictive distribution \( f(y_P|z) = N(y_P|\mu_P, \Sigma_P) \) (see Section 4.2.3) as approximated by each method. The mean squared error (MSE) is given by \((1/n_P) \sum_{i=1}^{n_P} (y_{P,i}^\star - \mu_{P,i})^2\), the log-score is given by \(- (1/n_P) \sum_{i=1}^{n_P} N(y_{P,i}^\star | \mu_{P,i}, \Sigma_{P,ii})\), and the continuous ranked probability score is given by \((1/n_P) \sum_{i=1}^{n_P} \int (F_i(x) - 1\{y_{P,i}^\star \leq x\})^2 dx\), where \( F_i \) is the cumulative distribution function of \( N(\mu_{P,i}, \Sigma_{P,ii}) \). Each score is averaged over 20 simulations.

### 5.4 Simulation results

#### 5.4.1 Out-of-sample prediction

Figure 3 shows ridgeline density plots of the prediction scores in Section 5.3 for the methods in Section 5.2 (except LUR-iid, which was not competitive) for the different simulation scenarios described in Section 5.1. The vertically oriented densities are generated as trimmed (inner 98 percent) density functions using the geom_density_ridges function in the ggplot2 and ggridges packages of R. The results for our LURK-Vecchia approach were nearly identical to those for LURK-Full. Across all scenarios, the average LURK-Vecchia scores were consistently between 5 and 60 percent better than those for LURK-Local. Local-Kriging was much worse. (Plots showing percent differences are shown in Section S1.)

#### 5.4.2 Model selection

In terms of model selection, we considered the true negative rate (TNR), true positive rate (TPR), and Cohen’s Kappa (\cite{Banerjee1999}), \( \kappa = (p_o - p_e)/(1 - p_e) \), where \( p_o \) is the observed agreement of coefficient selections and \( p_e \) is the expected agreement based on random chance. Of the methods in Section 5.2, we omitted LURK-Local (because its model selection is identical to LUR-iid) and Local-Kriging (because it does not perform model selection). Figure 4 shows the model-selection statistics for the scenarios in Section 5.1. Similar to the prediction scores, the distributions of the LURK-Vecchia and LURK-Full were very similar, indicating that the LURK-Vecchia approach approximated the full model well in terms of model selection. Compared to LUR-iid, LURK-Vecchia had 5 to 80 percent higher average TPR. The difference in terms of TNR and Kappa was even greater, with LUR-iid selecting a large number of erroneous non-zero coefficients (Figure S3). The results were only comparable for scenarios with negligible spatiotemporal dependence (i.e., high nugget-to-sill ratio, or small ranges). An additional plot showing the percent differences in Kappa is shown in Section S1.
Figure 3: Ridgeline density plots of prediction scores (the lower the better) for different simulation scenarios (Section 5.1), in which the spatiotemporal parameters are singly varied with others held constant at a baseline level. The LURK-Vecchia results were very close to and hence largely cover the results for LURK-Full. Note that each scenario has its own y-axis scale.

Figure 4: Ridgeline density plot comparison of model-selection statistics for simulation scenarios (Section 5.1), in which the spatiotemporal parameters are singly varied with others held constant at a baseline level.
### Table 1: Predictive scores for NO2 10-fold cross-validation

<table>
<thead>
<tr>
<th>Method</th>
<th>MSE (ppb^2)</th>
<th>CRPS</th>
<th>Log-Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local-Kriging</td>
<td>0.30</td>
<td>0.30</td>
<td>0.83</td>
</tr>
<tr>
<td>LUR-iid</td>
<td>0.42</td>
<td>0.38</td>
<td>3.90</td>
</tr>
<tr>
<td>LURK-Local</td>
<td>0.22</td>
<td>0.25</td>
<td>0.68</td>
</tr>
<tr>
<td>LURK-Vecchia</td>
<td>0.20</td>
<td>0.24</td>
<td>0.61</td>
</tr>
</tbody>
</table>

6 Ground-level NO2 analysis

We now return to the daily, US-wide, ground-level NO2 data described in Section 6. The minimum, maximum, mean (standard deviation), and median (interquartile range) observed concentrations were 0.004, 62.9, 8.5 (7.4), and 6.4 (9.0) parts-per-billion (ppb), respectively. Because NO2 is positive and right-skewed, NO2 was natural-log-transformed prior to the analyses.

6.1 Comparison using cross-validation

We compared the predictive accuracy using 10-fold cross-validation for the methods described in Section 5.2; LURK-Full was omitted because it is intractable for the large sample size. We used the same predictive scores as detailed in Section 5.3, except that we considered \( \hat{f}(z_P|z) = \mathcal{N}(z_P|\mu_P, \Sigma_P + \tau^2 \mathbf{I}_{n_P}) \), because the error-free \( y_P \) was unknown.

The results are shown in Table 1. LURK-Vecchia outperformed all other methods in terms of all three scores, resulting in a roughly ten percent decrease in MSE and log-score compared to the next best approach, LURK-Local. Further, LURK-Vecchia resulted in a 20 percent or greater decrease in all of the scores compared to Local-Kriging and LUR-iid.

In terms of model selection, the mean (standard deviation) number of non-zero coefficients was 71 (1.1) and 24 (1.7) for LUR-iid and LURK-Vecchia, respectively. The LUR-iid models were severely affected by multicollinearity; the median (mean) variance inflation factor (VIF) for all 10 cross-validation models was 2.3 (6.7) and 5.7 (18.6) for LURK-Vecchia and LUR-iid, respectively. LURK-Local uses the same model-selection procedure as LUR-iid. Other iid model-selection approaches are likely to result in similarly large numbers of covariates (e.g., Kerckhoffs et al., 2019).

Thus, by appropriately accounting for spatiotemporal dependence, LURK-Vecchia resulted in more accurate, sparser, and more interpretable models than LUR-iid and LURK-Local.

6.2 Prediction maps

Having shown using cross-validation that our proposed LURK-Vecchia approach can outperform the competing methods, we fitted LURK-Vecchia to the entire dataset. The covariance parameters \( \theta = (\sigma^2, \gamma_s, \gamma_t, \tau^2) \) were estimated as (2.2 ppb^2, 1.4 km, 0.63 yr, 0.15 ppb^2), and the trend coefficients are given in Table 1 and discussed in Section 6.3. The entire estimation algorithm required \( L = 4 \) iterations and took approximately 86 minutes on a machine with 16GB RAM and an Intel(R) i7-8665U processor (4 cores, 1.90GHz).
Figure 5 shows the prediction geometric mean, \( \exp(\mu_P) \), for two distinct days, for the entire US domain and for a more detailed 5-county area surrounding Houston, Texas. (Corresponding prediction uncertainties are shown in Figure S4.) For the US domain, predictions were produced on a 200 by 100 grid (10–20 km resolution) across the conterminous United States. For evaluating fine-scale prediction patterns, a 1–2 km grid was produced in the Houston, TX five-county area. Distinct spatiotemporal patterns emerged. Cities and developed areas, such as roadways, showed elevated NO\(_2\) concentrations, as expected for a traffic-related pollutant. However, there was temporal variability in the spatial patterns around the cities and roads. Comparing the the upper, mid-west cities, such as Chicago and Cleveland (blue box, Figure 5), predicted NO\(_2\) was lower on July 11, 2018, than on February 11, 2019. In contrast, in the Houston area sub-figure, predicted NO\(_2\) was higher on July 11, 2018, than on February 11, 2019. Visually inspecting the predictors in the final model can reveal the primary drivers of the spatiotemporal variability, which is easy in linear models with interpretable covariates. Complex machine learning models and dimension-reduction techniques do not allow for such intuitive visual comparisons. Figure S5 shows the predictions and select covariates in the Houston area on July 11, 2018 and February 11, 2019. Visual inspection and correlation of covariate-only predictions with the final predictions show that the TROPOMI NO\(_2\) data are driving the patterns observed on July 11, 2018. Contrarily, on Feb 11, 2019 other factors such as the specific humidity display similar general spatial patterns and have high correlation with the final predictions.

Figure 6 shows spatiotemporal predictions in more detail for a transect through the Houston panel of Figure 5. Moving along the transect, we see a general spatial pattern with modulations in time. For instance, the highway consistently had the highest observed concentrations, but the magnitude of the maximum fluctuated daily, driven by time-varying covariates such as the TROPOMI and meteorological variables (see Section 6.3). July 12, 2018 had consistently higher concentrations than other days across most of the transect locations, including the highway. We also showed prediction uncertainties in terms of geometric standard deviations (SDs), \( \exp(\text{diag}(\Sigma_P)^{1/2}) \). The SD varied over longer time periods than the mean, as evidenced by minor differences within the 2018 and 2019 ranges, but considerable differences between them.

Our predictions will be useful for epidemiological and risk-assessment studies seeking daily, national-scale predictions. For example, [Mills et al. (2015)] provide meta-analysis results for the impacts of 24-hour NO\(_2\) exposure on all-age-group, all-cause mortality, cardiovascular mortality, respiratory mortality, cardiovascular hospital admissions, and respiratory hospital admissions. Our daily NO\(_2\) exposure predictions, combined with population information and with the [Mills et al. (2015)] relative-risk estimates, may be used to develop NO\(_2\) acute-health impact assessments, such as an attributable-fraction of mortality. Please contact the authors to request predictions at the desired spatiotemporal coordinates.

### 6.3 Interpretation of selected covariates

Table 2 shows the 25 variables selected by our LURK-Vecchia procedure, along with their estimated coefficients. We now discuss interpretations and context for each selected variable, grouped by variable category (see Section 6.3):
• **TROPOMI.** Similar to many LUR studies (Novotny et al., 2011; Young et al., 2016; Larkin et al., 2017; de Hoogh et al., 2018), we found satellite observations of NO\textsubscript{2} selected to the LURK-Vecchia model. The two covariates for NO\textsubscript{2} Slant combine for a net positive effect on ground-level NO\textsubscript{2} while providing moderation between local and regional scale effects from the 1 and 100 km circular buffer hyperparameters, respectively. Similarly, the tropospheric NO\textsubscript{2} variables have two variables at different spatial scales that contribute a net positive effect to ground-level NO\textsubscript{2}.

The TROPOMI variables for tropopause pressure at 1 and 100 km were selected with small negative coefficients, indicating areas of reduced ground-level NO\textsubscript{2}. These variables, which have a relatively small impact on the final prediction concentrations, may represent a minor changes in the mixing volume for pollutant molecules.

The dynamic relationship between NO\textsubscript{2} and aerosols is complex and not completely understood. For instance, Grundström et al. (2015) observe weak to moderate correlations between total NO\textsubscript{x} (NO + NO\textsubscript{2}) and particle number concentrations (PNC) depending on meteorological conditions such as wind velocity. Apte et al. (2019) found PNC to have a consistent diurnal pattern of midday new particle formation that is poorly approximated with NO\textsubscript{x}. Without a priori expectations of the AAI coefficient, we observe a 2.6 percent decrease in NO\textsubscript{2} concentrations for every one SD increase.
in AAI. We find a positive coefficient for cloud fraction, which is likely due to the protective effect of clouds on incoming solar radiation.

- **Meteorology.** Two relative humidity and one specific humidity variable contribute a net negative effect on NO₂ concentrations. Similarly to AAI, we expect that water vapor and aerosols to impede solar radiation and breakdown of NO₂ to NO.

  We observe a 55.5 percent increase in NO₂ concentrations for every 1 SD increase in the maximum daily temperature. Hot days are associated with increased solar radiation and O₃ formation. The significant increase in NO₂ concentrations is likely capturing O₃ mediated conversion of NO (NO $\xrightarrow{O_3}$ NO₂) ([Seinfeld and Pandis](2016)).

  We observe a 19.2 percent decrease in NO₂ concentration with a 1 SD increase in wind
<table>
<thead>
<tr>
<th>Variable Category</th>
<th>Variable Name</th>
<th>res. (km)</th>
<th>$\hat{\beta}$</th>
<th>$(e^{\hat{\beta}} - 1) \times 100$</th>
</tr>
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<tr>
<td>Intercept</td>
<td>-</td>
<td>-</td>
<td>0.002</td>
<td>-</td>
</tr>
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<td>TROPOMI (Sec 2.1.1)</td>
<td>NO$_2$ Slant</td>
<td>1</td>
<td>0.086</td>
<td>8.9</td>
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<tr>
<td></td>
<td>NO$_2$ Slant</td>
<td>100</td>
<td>-0.002</td>
<td>-0.21</td>
</tr>
<tr>
<td></td>
<td>NO$_2$ Tropospheric</td>
<td>10</td>
<td>0.036</td>
<td>3.6</td>
</tr>
<tr>
<td></td>
<td>NO$_2$ Tropospheric</td>
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<td>0.44</td>
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<td>-19.2</td>
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<tr>
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<td>NDVI</td>
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<td>Population &amp; Roads (Sec 2.1.4)</td>
<td>Travel Friction</td>
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<td>-0.038</td>
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<td></td>
<td>Total Road</td>
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<td>0.18</td>
<td>20.3</td>
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<td></td>
<td>Mixed Forest</td>
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<td>-18.3</td>
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<td>10</td>
<td>0.017</td>
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<td></td>
<td>Shrub</td>
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</tr>
<tr>
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<td>Herbaceous</td>
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<td>0.92</td>
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</table>

Table 2: Selected covariates for NO$_2$ data: rhmax = maximum relative humidity, vs = wind velocity, sph = specific humidity, tmax = maximum temperature. For every 1-standard-deviation increase in a covariate, we expect an estimated $(e^{\hat{\beta}} - 1) \times 100$ percent increase in NO$_2$.

velocity, which is expected as this increases transport of NO$_2$ and its precursors from the given location.

- **Vegetation.** For every 1 SD increase in NDVI, we observe a 5.1 percent decrease in NO$_2$ concentrations. NDVI represents vegetative greenness, thus this is consistent with the lack of NO$_2$ or NO$_x$ sources.

- **Land cover.** Open water has a small, positive impact on NO$_2$, which is likely due to the concentration of cities and sources near water sources and coastlines or as a proxy variable for ports. Mixed forest (the net sum of short and medium ranges), shrub-land, and herbaceous wetlands have negative contributions to NO$_2$ predictions, which is expected due to the lack of sources. Developed open and low have positive coefficients, while developed low is larger as it represents an increased anthropogenic presence.

For every 1 SD increase in elevation in 1 km buffer, there is a 24.6 percent decrease


in NO₂ concentrations, which is consistent with other LUR models of NO₂ (de Hoogh et al., 2018) and can be due to a combination of atmospheric mixing, fewer sources, decreased average temperature, and increased wind velocity at higher elevations.

- **Population & Roads.** We find travel friction within a 10 km buffer and total road length within a 1 km buffer to result in a 3.7 decrease and 20.3 increase of NO₂ concentrations for every 1 SD increase, respectively. Travel friction is the average travel time, and total road length is a good approximation of vehicle sources, and so they are expected to decrease and increase traffic-related pollutants, respectively.

- **Emissions.** The NEI variable with a 1 km decay range was selected with a small, positive coefficient. Clearly, we expect a covariate representing source emissions of the dependent variable to be contribute positively.

## 7 Conclusions

We analyzed daily ground-level NO₂ concentrations across the United States, using a novel penalized land-use regression approach with spatiotemporally correlated errors that is also scalable to large datasets via a sparse general Vecchia approximation. Our methodological advances can be used in future human health exposure and risk assessment to improve model selection and prediction characteristics. Key results from the NO₂ analysis include: the development of daily NO₂ concentration predictions that can be used for epidemiological analyses of acute health effects such as asthma and increased hospitalizations; the potential to develop annual average concentrations that propagate uncertainty from daily predictions as opposed to those based on direct annual averages; the elucidation of spatiotemporal patterns of NO₂ concentrations across the United States, including significant variations between cities and intra-urban variation; and the resolving of a parsimonious group of geographic covariates describing the spatiotemporal distribution of daily NO₂ concentrations, including satellite imagery, meteorological data, land cover, population distributions, road networks, and point source emissions.

Our methods also offer a scalable way to analyze other large spatiotemporal datasets in environmental and human health risk assessment. For example, in the air-quality research community, mobile monitoring of air pollutants is leading to high-resolution datasets with millions of observations, including campaigns in Zurich, Switzerland (Li et al., 2012), Boston, MA (Padrò-Martínez et al., 2012), Oakland, CA (Apte et al., 2017; Guan et al., 2020), Houston, TX (Miller et al., 2020), and the Netherlands (Kerckhoffs et al., 2019).

Our methods could also be extended to non-Gaussian data (Zilber and Katzfuss, 2020) or on-line spatiotemporal filtering (Jurek and Katzfuss, 2018) using extensions or variations of the general-Vecchia framework.

As mentioned in Section 3, the SCAD penalty used for model selection in Line 7 of Algorithm 1 could be replaced by other penalties, such as LASSO (Tibshirani, 1996), elastic net (Zou and Hastie, 2005), or relaxed LASSO (Hastie et al., 2017), which may result in improvements in prediction accuracy or model selection. While accurate uncertainty quantification and significance assessment is difficult in the context of penalized regression, a potential extension of our approach would be to combine it with existing methods proposed
for this purpose (e.g., [Meinshausen et al. 2009, Chatterjee and Lahiri 2011, Xie et al. 2019]).

This would likely come at an increased computational cost, but it would also allow for the inclusion of covariate uncertainty in predictions. Lastly, another possible avenue is to adjust for spatial confounding as proposed in [Hughes and Haran 2013].

Acknowledgments

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A Review of general Vecchia

We now provide some further details of the general Vecchia approximation [Katzfuss and Guinness 2019, KatzFuss et al. 2020a] that we extended and briefly reviewed in Section 4. Because model (1) implies conditional independence in (6) between \( z_t \) and all other variables in \( u \) given \( y_t \), we assume that \( z_t \) always conditions on only \( y_t \). Hence, we can write the approximation (6) as

\[
\hat{f}(u) = \prod_{i=1}^{n} p(z_i | y_i) p(y_i | \mathbf{y}_{q_y(i)}, \mathbf{z}_{q_z(i)}),
\]

where \( q(i) = q_y(i) \cup q_z(i) \) with \( q(i) \subset \{1, \ldots, i - 1\} \) is the conditioning index vector of size \(|q(i)| \leq m\), and we assume \( q_y(i) \cap q_z(i) = \emptyset \). The ordering of the variables and the choice of conditioning sets can have a strong effect on the approximation accuracy and computational speed.

The ordering of the spatiotemporal coordinates \((s_1, t_1), \ldots, (s_n, t_n)\) implies an ordering of the variables in \( u \). We assume here that the coordinates are ordered and numbered according to a maximum-minimum distance ordering [Guinness 2018], which sequentially picks each coordinate in the ordering to maximize the minimum distance to previous coordinate in the ordering. The conditioning index vectors \( q(i) \) are chosen here as the indices of the nearest \( m \) coordinates previous to \( i \) in this ordering. To determine the ordering and the conditioning sets, we use the scaled spatiotemporal distance (2) as our measure of distance (cf. Datta et al. 2016b). However, this measure of distance depends on the unknown parameters \( \theta \) (specifically, on \( \gamma_s \) and \( \gamma_t \)), and so we update the ordering and conditioning at each iteration (in Line 5) of Algorithm 1 based on the current estimate of \( \theta \).

Different strategies for splitting \( q(i) \) into \( q_y(i) \) and \( q_z(i) \) can also result in vastly different approximation accuracies. In general, conditioning on \( y_j \) is often more accurate but also potentially more computationally expensive than conditioning on \( z_j \). Katzfuss and Guinness (2019) proposed a fast and accurate sparse general Vecchia (SGV) approach that chooses \( q_y(i) \subset q(i) \) such that \( j < k \) can only both be in \( q_y(i) \) if \( j \in q_y(k) \), with the remaining conditioning indices in \( q(i) \) assigned to \( q_z(i) \). Specifically, for \( i = 1, \ldots, n \), SGV finds \( p(i) = \arg \max_{j \in q(i)} |q_y(j) \cap q(i)| \) and \( k_i = \arg \min_{\ell \in p(i)} ||s_i - s_\ell|| \), and then sets \( q_y(i) = (k_i) \cup (q_y(k_i) \cap q(i)) \). We use SGV in all our numerical examples.

The restriction of conditioning only on previous variables in the ordering, \( q_y(i) \subset \{1, \ldots, i-1\} \), ensures that the implied joint distribution is multivariate normal as indicated in (6). To compute the sparse upper-triangular matrix \( \mathbf{U} \), let \( g(i) \) denote the vector of indices of the elements in \( u \) on which \( u_i \) conditions (e.g., if \( u_i = z_k \) then \( g(i) = (i-1) \)). Also define \( K(y_i, y_j) = K(z_i, y_j) = C((s_i, t_i), (s_j, t_j)) \) and \( K(z_i, z_j) =\)
$C((s_i, t_i), (s_j, t_j)) + 1_{i=j}r_i^2$. Then, the $(j, i)$th element of $U$ is

$$U_{ji} = \begin{cases} r_i^{-1/2}, & i = j, \\ -b_i^{(j)} r_i^{-1/2}, & j \in q(i), \\ 0, & \text{otherwise}, \end{cases}$$

where $b_i' = K(u_i, u_{iy(i)})K(u_{iy(i)}, u_{iy(i)})^{-1}r_i = K(u_i, u_i) - b_i'K(u_{iy(i)}, u_i)$, and $b_i^{(j)}$ denotes the $\ell$th element of $b_i$ if $j$ is the $\ell$th element in $q(i)$ (i.e., $b_i^{(j)}$ is the element of $b_i$ corresponding to $u_j$).

For prediction of $y_p$, we employ a spatio-temporal extension of the response-first ordering full-conditioning (RF-full) approach of Katzfuss et al. (2020a), which applies a general Vecchia approximation of the form (6) to $\hat{u} = (z', y_{all}')'$, where $y_{all} = (y', y_p)' = (y_1, \ldots, y_{nall})'$. This results in the approximation

$$\hat{f}(z, y_{all}) = \prod_{i=1}^{n} f(z_i) \times \prod_{i=1}^{n} f(y_{q_y(i)}, z_{q_z(i)})$$

where $y_{q_y(i)}$ and $z_{q_z(i)}$ are chosen as the $m$ variables closest in scaled distance (2) to $y_i$, among those that are previously ordered in $\hat{u}$, where we condition on $y_j$ instead of $z_j$ whenever possible. Specifically, we set $q(i)$ to consist of the indices corresponding to the $m$ nearest spatiotemporal coordinates, including $i$ for $i \leq n$, and not including $i$ for $i > n$. Then, for any $j \in q(i)$, we let $y_j$ condition on $y_j$ if it is ordered previously in $u$, and condition on $z_j$ otherwise. More precisely, we set $q_y(i) = \{j \in q(i) : j < i\}$ and $q_z(i) = \{j \in q(i) : j \geq i\}$.

**B  Alternative expression of the objective function**

We now show that the objective function $\hat{Q}(\beta, \theta)$ can be written as in (8) as:

$$\hat{Q}(\beta, \theta) = \|\hat{z}_0 - \bar{X}_0 \beta\|^2_2 + \lambda \rho(\beta) - 2 \sum_i \log (|U_{ii}|) + 2 \sum_i \log (|V_{ii}|).$$

As in Katzfuss and Guinness (2019) proof of Prop. 2), note that, for any value of $y$, $\hat{f}(z) = \hat{f}(u)/\hat{f}(y|z)$, where $f(u)$ is given in (6), and $f(y|z) = \mathcal{N}_n(\mu, W^{-1})$ with $\mu = X\beta - W^{-1}AB'(z - X\beta)$. Thus, setting $y = \mu$, and denoting by $u_\mu = (\mu_1, z_1, \ldots, \mu_n, z_n)$ the resulting vector $u$, we obtain

$$-2 \log \hat{f}(z) = -2 \log \mathcal{N}_n(u_\mu|(X \otimes 1_2)\beta, \Sigma) + 2 \log \mathcal{N}_n(\mu|\mu, W^{-1})$$

$$= (u_\mu - (X \otimes 1_2)\beta)'UU'(u_\mu - (X \otimes 1_2)\beta) - \log |UU'| + \log |W|$$

$$= \|d\|^2_2 - 2 \sum_i \log U_{ii} + 2 \sum_i \log V_{ii},$$

where

$$d = U'u_\mu - U'(X \otimes 1_2)\beta = B'z + A'\mu - (B'X + A'X)\beta$$

$$B'z - A'W^{-1}AB'z - (B'X + A'X - A'X + A'W^{-1}AB'X)\beta$$

$$= \hat{z} - \hat{X}\beta,$$

where $\hat{z} = B'z + A'W^{-1}AB'z$ and $\hat{X} = B'X + A'W^{-1}AB'X$.

**References**


Supplementary Material

S1 Simulation with jointly varying parameters

Supplementing the simulation study in Section 5, we also evaluated multiple scenarios where the variables are varied jointly. Note we only vary the covariance range in space. We test a 30 scenario set that is all possible combinations of: $\tau^2/\sigma_{total}^2 = \{0.1, 0.33, 0.5, 0.67, 0.9\}$, $\sigma_{total}^2 = \{1, 4\} \times \sigma_{true}^2$, $\theta_s = \{30, 300, 3000\}$ km, and $\theta_t = 30$ days.

Figure S1: Percent difference between in prediction validation statistics for LURK-Local (i) and LURK-Vecchia (ii) approaches as a function of jointly varying spatiotemporal random field parameters. Percent difference is calculated as (i-ii)/i x 100. A positive value indicates that the LURK-Vecchia performance is better. Note that some boxplots are truncated at 40 percent.
Figure S2: Percent difference between in Cohen’s Kappa ($\kappa$) (overall model selection) for LURK-Local (i) and LURK-Vecchia (ii) approaches as a function of jointly varying spatiotemporal random field parameters. Percent difference is calculated as (i-ii)/i x 100. A negative value indicates that the LURK-Vecchia performance is better.

Figure S3: Ridgeline density plot comparison of the number of non-zero coefficients for simulation scenarios [5,2] in which the spatiotemporal random field parameters are singly varied with others held constant. The horizontal black line (y=8) is the number of covariates in the true model.
S2 Prediction uncertainty for NO2 application

Figure S4 shows prediction uncertainties for the noise-free NO$_2$ in terms of geometric standard deviations (SDs), $\exp(\text{diag}(\Sigma)^{1/2})$. The prediction SD varies more smoothly over space than the mean in Figure 5.

![Figure S4: LURK-Vecchia prediction geometric standard deviation (exponentiated log-transform standard deviation) across the conterminous United States (top row) and the five-county subset around Houston, Texas for July 11, 2018 and February 11, 2019.](image)

S3 Prediction and covariate comparison

Figure S5 shows point predictions (log-scale) for select covariates using only the covariate(s) and their estimated coefficients: $X_{\text{select}}\hat{\beta}_{\text{select}}$. The Pearson correlation ($\rho$) and RMSE between the select covariate predictions and the respective full model prediction for that day (top row) are shown (i.e. left-middle vs left-top, left-bottom vs left-top, right-middle vs right-top, and right-bottom vs right-top). The TROPOMI-NO$_2$ covariates are clearly driving the large hot-spot of the final predictions on July 11, 2018, whereas other factors such as specific humidity have more control on February 11, 2019.
Figure S5: Comparison of the LURK-Vecchia predictions in the Houston area with predictions based only on select covariates (i.e., $X_{\text{select}} \hat{\beta}_{\text{select}}$). The rows show the (top) LURK-Vecchia final model predictions, compared to predictions using (middle) TROPOMI-NO$_2$ or (bottom) specific humidity as the only covariate(s). The left and right columns are predictions and comparisons for July 11, 2018 and February 11, 2019, respectively.