

Statistica Sinica Preprint No: SS-2021-0277

Title	DeepKriging: Spatially Dependent Deep Neural Networks for Spatial Prediction
Manuscript ID	SS-2021-0277
URL	http://www.stat.sinica.edu.tw/statistica/
DOI	10.5705/ss.202021.0277
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DeepKriging: Spatially Dependent Deep Neural Networks for Spatial Prediction

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Abstract: In spatial statistics, a common objective is to predict values of a spatial process at unobserved locations by exploiting spatial dependence. Kriging provides the best linear unbiased predictor using covariance functions, and is often associated with Gaussian processes. However, for nonlinear predictions for nonGaussian and categorical data, the Kriging prediction is no longer optimal, and the associated variance is often overly optimistic. Although deep neural networks (DNNs) are widely used for general classification and prediction, they have not been studied thoroughly for data with spatial dependence. In this work, we propose a novel DNN structure for spatial prediction, where we capture the spatial dependence by adding an embedding layer of spatial coordinates with basis functions. We show in theory and simulation studies that the proposed DeepKriging method has a direct link to Kriging in the Gaussian case, and has multiple advantages over Kriging for nonGaussian and nonstationary data. That is, it provides nonlinear predictions, and thus has smaller approximation errors.

Furthermore, it does not require operations on covariance matrices, and thus is scalable for large data sets. With sufficiently many hidden neurons, the proposed method provides an optimal prediction in terms of model capacity. In addition, we quantify prediction uncertainties based on density prediction, without assuming a data distribution. Finally, we apply the method to $PM_{2.5}$ concentrations across the continental United States.

Key words and phrases: Basis function, Deep learning, Feature embedding, Gaussian process, Spatial Prediction.

1. Introduction

Spatial prediction is at the heart of spatial and spatio-temporal statistics. It is aimed at predicting values of a spatial process at unobserved locations by accounting for the spatial dependence in the region of interest. Originally, spatial prediction was applied in the fields of geological and environmental science (Cressie, 2015), but has been extended to other fields, such as the biological sciences, computer vision, economics, and public health (Anselin, 2001; Austin, 2002; Waller and Gotway, 2004; Franchi et al., 2018).

The main spatial prediction methods are based on the best linear unbiased prediction (BLUP), also referred to as Kriging (Matheron, 1963). Kriging prediction is a weighted average of observed data, where the weights are determined by the spatial covariance function or variogram of the random

process. Under the Gaussian assumption, Kriging also provides the full predictive distribution. Applying Kriging requires estimating the spatial covariance function, which is commonly assumed to be stationary. However, physical processes tend to be nonGaussian and nonstationary. For instance, data on wind speed and fine particle ($PM_{2.5}$) exposures are positive, right-skewed, and sometimes heavy-tailed (Hennessey Jr, 1977; Adgate et al., 2002), and the spatial covariance typically varies across space, for example, in urban versus rural areas (Sampson et al., 2013). It is possible to derive the best linear prediction for certain parametric nonGaussian processes (Xu and Genton, 2017; Rimstad and Omre, 2014) and certain nonstationary covariance structures (Fuentes, 2002; Paciorek and Schervish, 2004; Li and Sun, 2019), but Kriging for more general spatial processes remains an open problem. Another drawback of Kriging is that it is computationally prohibitive for large spatial data sets, because it involves inverting an $N \times N$ covariance matrix, where N is the number of observed locations (Heaton et al., 2019), and the computation requires $O(N^3)$ time and $O(N^2)$ memory complexity, based on the typical Cholesky decomposition approach.

Recently, deep learning and deep neural networks (DNNs) have become powerful prediction tools for a wide range of applications, especially in computer vision and natural language processing (LeCun et al., 2015).

DNNs are effective for predictions with complex features such as nonlinearity and nonstationarity, and are computationally efficient when analyzing massive data sets using GPUs (Najafabadi et al., 2015). Although it appears promising to apply DNNs to spatial predictions, classical DNNs cannot incorporate spatial dependence appropriately. Spatial prediction applications with neural networks usually simply include spatial coordinates as features (Cracknell and Reading, 2014), which may not be sufficient. Recently, convolutional neural networks (CNNs, Krizhevsky et al. 2012) have claimed to successfully capture the spatial and temporal dependencies in image processing using relevant filters. However, the framework is designed for applications with a large feature space, and often requires large numbers of training labels as the ground truth. In many spatial prediction problems, only in-situ and sparse observations are available.

To address the above-mentioned problems, we develop an effective DNN for spatial prediction that

- 1) builds a direct link between DNNs and Kriging in spatial prediction;
- 2) models spatial dependence using a set of basis functions;
- 3) does not require matrix operations and is scalable for large data sets;
- 4) provides a nonlinear predictor in covariates or generally in observations;
- 5) has a Gaussian process (GP) representation, and provides more flexible

spatial covariance structures than simply using the coordinates as features;
6) suits different data types, for example, nonGaussian or nonstationary data; and
7) potentially measures uncertainty using predictive density functions, without assuming a data distribution.

We call our method “DeepKriging”, which achieves optimal spatial prediction, similar to the original use of Kriging (Cressie, 1990), but using DNNs. We also conduct simulation studies and apply our approach to $PM_{2.5}$ concentration data from across the continental United States to show how DeepKriging performs relative to Kriging and other naive DNN methods. The rest of our paper is organized as follows. In Section 2, we construct our DeepKriging method, and in Section 3, provide its theoretical properties. Section 4 presents simulation studies that demonstrate the performance of the DeepKriging method. In Section 5, we apply DeepKriging to predict the $PM_{2.5}$ concentration in the United States. Section 6 summarizes the main results and suggests directions for future work.

2. Methodology

2.1 Deep learning in spatial prediction

Suppose $\mathbf{z} = \{z(\mathbf{s}_1), \dots, z(\mathbf{s}_N)\}^T$ are measurements observed at N spatial locations from a real-valued spatial process $\{Y(\mathbf{s}) : \mathbf{s} \in D\}$, where $D \subseteq \mathbb{R}^d$. The goal of spatial prediction is to find the optimal predictor $\hat{Y}^{\text{opt}}(\mathbf{s}_0)$ of the true process at an unobserved location \mathbf{s}_0 , as a function of \mathbf{z} . In decision theory, $\hat{Y}^{\text{opt}}(\mathbf{s}_0)$ is the minimizer of an expected loss function or risk function (DeGroot, 2005). That is,

$$\hat{Y}^{\text{opt}}(\mathbf{s}_0) = \underset{\hat{Y}}{\operatorname{argmin}} \mathbb{E}\{L(\hat{Y}(\mathbf{s}_0), Y(\mathbf{s}_0))\} = \underset{\hat{Y}}{\operatorname{argmin}} R(\hat{Y}(\mathbf{s}_0), Y(\mathbf{s}_0)), \quad (2.1)$$

where $L(\cdot, \cdot)$ is a loss function and $R(\cdot, \cdot)$ is a risk function. Under the mean squared error (MSE) loss, the optimal predictor is $\hat{Y}^{\text{opt}}(\mathbf{s}_0) = \mathbb{E}\{Y(\mathbf{s}_0)|\mathbf{z}\}$, if it is finite. This predictor has multiple good properties, such as unbiasedness and asymptotic normality under regularity assumptions (Lehmann and Casella, 2006). In particular, if $Y(\mathbf{s}_0)$ and \mathbf{z} are jointly Gaussian, the conditional mean is a linear combination of \mathbf{z} ; if $Y(\mathbf{s}_0)$ and \mathbf{z} are not jointly Gaussian, the conditional mean obtained based on a Gaussian assumption remains the BLUP, which is called Kriging. However, as mentioned before, the Kriging predictor is sub-optimal for nonGaussian data, and is not scalable for large data sizes.

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In this work, we use deep learning to approximate the optimal predictor $\widehat{Y}^{\text{opt}}(\mathbf{s}_0)$ in (2.1) using the output of the neural network. The optimal neural network predictor is given by $f_{\text{NN}}^{\text{opt}}(\mathbf{s}_0) = \operatorname{argmin}_{f_{\text{NN}}} R\{f_{\text{NN}}(\mathbf{s}_0), Y(\mathbf{s}_0)\}$, where $f_{\text{NN}}(\cdot) \in \mathcal{F}$ can be any function in the function space \mathcal{F} expressible by a family of neural networks, and $f_{\text{NN}}^{\text{opt}}(\cdot)$ is the best function in \mathcal{F} in terms of minimizing a certain risk $R(\cdot, \cdot)$. The inputs of the neural network can be relevant covariates $\mathbf{x}(\mathbf{s}_0)$ and other features at \mathbf{s}_0 . Typically, we write $f_{\text{NN}}(\mathbf{s}; \boldsymbol{\theta})$ as a parametric model with unknown parameters $\boldsymbol{\theta}$, which include the weights and biases in the neural network. Note that the optimal neural network predictor $f_{\text{NN}}^{\text{opt}}(\mathbf{s}_0)$ is practically unreachable, because $Y(\mathbf{s}_0)$ is unknown. In practice, we approximate the predictor by minimizing the empirical loss function over the training set \mathbf{z} (Goodfellow et al., 2016); that is, the final predictor is $\widehat{Y}_{\text{NN}}(\mathbf{s}_0) = f_{\text{NN}}(\mathbf{s}_0; \widehat{\boldsymbol{\theta}})$, with

$$\widehat{\boldsymbol{\theta}} = \operatorname{argmin}_{\boldsymbol{\theta}} \frac{1}{N} \sum_{n=1}^N L\{f_{\text{NN}}(\mathbf{s}_n; \boldsymbol{\theta}), z(\mathbf{s}_n)\}. \quad (2.2)$$

Applying this classical neural network framework directly to spatial prediction is problematic, for at least two reasons. First, classical DNNs do not account for spatial dependence and, second, spatial prediction typically has limited observed features, rather than excessive features in common applications of neural networks. In particular, assume that the spatial process $Y(\mathbf{s})$ is modeled by $Y(\mathbf{s}) = \mathbf{x}(\mathbf{s})^T \boldsymbol{\beta} + \nu(\mathbf{s})$, where $\mathbf{x}(\mathbf{s}) \in \mathbb{R}^P$ is a

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vector process of P known covariates, β is a vector of coefficients, and $\nu(\mathbf{s})$ is a spatially dependent and zero-mean random process with a generally nonstationary covariance function: $\text{Cov}(\nu(\mathbf{s}), \nu(\mathbf{s}')) = C(\mathbf{s}, \mathbf{s}')$. In neural networks, we usually assume that $Y(\mathbf{s})$ are mutually independent, conditional on the features $\mathbf{x}(\mathbf{s})$. However, this assumption is not reasonable in spatial prediction, because the covariates $\mathbf{x}(\mathbf{s})$ contribute only to the mean structure of $Y(\mathbf{s})$, and $\nu(\mathbf{s})$ remains a spatially correlated process. Hence, we need features in addition to $\mathbf{x}(\mathbf{s})$ to model spatial dependence using neural networks.

To account for spatial information, the most natural way is to add d coordinates (e.g., longitude and latitude) to the features, in the hope that the neural networks can learn the dependent term $\nu(\mathbf{s})$ as a function of \mathbf{s} (Cracknell and Reading, 2014). By doing that, the adjusted features become $\mathbf{x}^{adj}(\mathbf{s}) = (\mathbf{x}(\mathbf{s})^T, \mathbf{s}^T)^T$. However, this does not help to enlarge the feature space, because the dimension of the coordinates is usually $d \leq 3$. Moreover, the associated neural network may not be efficient, because if the true function is far from linear, it may require a significant effort for the neural network to achieve a good approximation. For instance, the optimal predictor under the Gaussian assumption and MSE loss is the Kriging predictor, which is linear in $\mathbf{x}(\mathbf{s})$, but obviously nonlinear in the

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coordinates \mathbf{s} ; this is a special case in which the natural structure of neural networks may not work.

Delving deeper into the form of a Kriging prediction may help us to find an appropriate way to incorporate spatial dependence in a DNN. Suppose \mathbf{z} is observed from a generalized additive model: $Z(\mathbf{s}) = Y(\mathbf{s}) + \varepsilon(\mathbf{s})$, where $Y(\mathbf{s}) = \mathbf{x}(\mathbf{s})^T \boldsymbol{\beta} + \nu(\mathbf{s})$, as defined above, and $\varepsilon(\mathbf{s})$ is a white noise process, called the nugget effect, with zero mean and variance $\sigma^2(\mathbf{s})$, caused by measurement inaccuracy and fine-scale variability. The (universal) Kriging prediction is

$$\hat{Y}_{\text{UK}}(\mathbf{s}_0) = \mathbf{x}(\mathbf{s}_0)^T \hat{\boldsymbol{\beta}} + \mathbf{c}(\mathbf{s}_0)^T \boldsymbol{\Sigma}^{-1}(\mathbf{z} - \mathbf{X}\hat{\boldsymbol{\beta}}), \quad (2.3)$$

where $\mathbf{X} = (\mathbf{x}(\mathbf{s}_1), \dots, \mathbf{x}(\mathbf{s}_N))^T$ is an $N \times P$ matrix, $\mathbf{c}(\mathbf{s}_0) = \text{Cov}(\mathbf{Z}, Z(\mathbf{s}_0))$, $\boldsymbol{\Sigma} = \text{Cov}(\mathbf{Z}, \mathbf{Z}^T)$, and $\hat{\boldsymbol{\beta}} = (\mathbf{X}^T \boldsymbol{\Sigma}^{-1} \mathbf{X})^{-1} \mathbf{X}^T \boldsymbol{\Sigma}^{-1} \mathbf{z}$. The spatial dependence is incorporated in $\hat{Y}_{\text{UK}}(\mathbf{s}_0)$ using a linear function of the covariance vector $\mathbf{c}(\mathbf{s}_0)$, but its coefficient $\boldsymbol{\Sigma}^{-1}(\mathbf{z} - \mathbf{X}\hat{\boldsymbol{\beta}})$ is unknown. This motivates us to use a set of known nonlinear functions as the embedding of \mathbf{s} in the features to characterize the spatial process $\nu(\mathbf{s})$ in the neural network. This can be done using the Karhunen–Loève (KL) theorem (Adler, 2010), which establishes that $\nu(\mathbf{s})$ admits a decomposition $\nu(\mathbf{s}) = \sum_{k=1}^{\infty} w_k \phi_k(\mathbf{s})$, where w_k are pairwise uncorrelated random variables and $\phi_k(\mathbf{s})$ are pairwise orthogonal basis functions in the domain of $\nu(\mathbf{s})$. Hence, $\nu(\mathbf{s})$ can be linearly quantified

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using nonlinear basis functions of \mathbf{s} .

In practice, the prediction of $\nu(\mathbf{s})$ is typically the truncated KL expansion, based on the property that given any orthonormal basis functions $\phi_k(\mathbf{s})$, we can find some large integer K , so that $\nu(\mathbf{s})$ can be approximated by the finite weighted sum of the basis functions, that is, $\hat{\nu}(\mathbf{s}) = \sum_{k=1}^K w_k \phi_k(\mathbf{s})$. Based on the KL theorem, the form of the basis functions is not as important as the number of basis functions when approximating the spatial random effect $\nu(\mathbf{s})$. This result is supported by the additional simulations we conduct in Section S4.1 of the Supplementary Material. Multiple types of basis functions can be used, such as smoothing spline basis functions (Wahba, 1990), wavelet basis functions (Vidakovic, 2009), and radial basis functions (Friedman et al., 2001). By adding an embedding layer with sufficiently large K , the width of the neural network increases greatly, enabling the network to incorporate more spatial information than when using the coordinates alone. A similar idea is used in the recommendation systems of Cheng et al. (2016).

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In this section, we use a simple DNN to illustrate our DeepKriging framework. Our model can potentially be used in other deep learning frameworks,

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such as CNNs and recurrent neural networks (RNNs).

First, choose the value for K and the basis functions to approximate the spatial process $\nu(\mathbf{s})$. We adopt the idea of Nychka et al. (2015), who developed a multi-resolution model for spatial prediction for large data sets. The radial basis functions at each level of resolution are constructed using a Wendland compactly supported correlation function, with the nodes arranged on a rectangular grid. In particular, at a certain level of resolution, let $\{\mathbf{u}_j\}$, for $j = 1, \dots, m$, be a rectangular grid of points (or node points in radial basis function terminology), and let θ be a scale parameter. The basis functions are given by $\phi_j^*(\mathbf{s}) = \phi(\|\mathbf{s} - \mathbf{u}_j\|/\theta)$, where

$$\phi(d) = \begin{cases} (1-d)^6(35d^2 + 18d + 3)/3, & d \in [0, 1] \\ 0, & \text{otherwise.} \end{cases}$$

Hence, the embedding layer uses the mutual distance locally to each knot location, implying that the spatial patterns are location invariant locally. As a result, the proposed DeepKriging method models the spatial nonstationarity; as shown in Section 3.3, the induced covariance functions of an infinitely wide DeepKriging network are in general nonstationary. The scale parameter θ is set to 2.5 times the associated knots spacing, following Nychka et al. (2015). The grid at each finer level increases by a factor of two, and the basis functions are scaled to have a constant overlap. In particular,

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in the h th level, the number of knots is chosen as $K_h = (9 \times 2^{h-1} + 1)^d$, where d is the spatial dimension. For a massive data set and to obtain $K \geq N$, we need $H = 1 + \lceil \log_2(\sqrt[d]{N}/10) \rceil$ levels. Therefore, for a four-level model, for instance, we need $K = 10 + 19 + 37 + 73 = 139$ basis functions in a one-dimensional space, and $K = 10^2 + 19^2 + 37^2 + 73^2 = 7159$ basis functions in a two-dimensional space. This scheme gives a good approximation for standard covariance functions, and is flexible enough to fit more complicated shapes. [Other works use multi-resolution approximation for massive spatial data sets; see Katzfuss \(2017\), and the references therein.](#)

Then, for any coordinate \mathbf{s} , we compute the K basis functions to obtain the embedded vectors $\boldsymbol{\phi}(\mathbf{s}) = (\phi_1(\mathbf{s}), \dots, \phi_K(\mathbf{s}))^T$. The basis functions are recommended to be orthogonal, based on the KL expansion. Then, let $\mathbf{x}_\phi(\mathbf{s}) = (\mathbf{x}(\mathbf{s})^T, \boldsymbol{\phi}(\mathbf{s})^T)^T$ be the embedded input of length $P + K$, and specify an L -layer DNN as

$$\begin{aligned} \mathbf{u}_1(\mathbf{s}) &= \mathbf{W}_1 \mathbf{x}_\phi(\mathbf{s}) + \mathbf{b}_1, \quad \mathbf{a}_1(\mathbf{s}) = \psi_1(\mathbf{u}_1(\mathbf{s})); \\ \mathbf{u}_2(\mathbf{s}) &= \mathbf{W}_2 \mathbf{a}_1(\mathbf{s}) + \mathbf{b}_2, \quad \mathbf{a}_2(\mathbf{s}) = \psi_2(\mathbf{u}_2(\mathbf{s})); \\ &\dots \\ \mathbf{u}_L(\mathbf{s}) &= \mathbf{W}_L \mathbf{a}_{L-1}(\mathbf{s}) + \mathbf{b}_L, \quad f_{\text{DK}}(\mathbf{s}) = \psi_L(\mathbf{u}_L(\mathbf{s})). \end{aligned} \tag{2.4}$$

For the l th layer with N_l neurons, \mathbf{W}_l is an $N_l \times N_{l-1}$ weight matrix, \mathbf{b}_l is a bias vector of length N_l , \mathbf{a}_l is a neuron vector of length N_l , and $\psi_l(\cdot)$ is

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the activation function. The output of this neural network is $f_{\text{DK}}(\mathbf{s})$, which is a function of the weights and the biases. Let $\boldsymbol{\theta}$ be the vector of unknown weights and biases, and $\hat{\boldsymbol{\theta}}$ be the estimate from Equation (2.2) based on the training sample. The final DeepKriging prediction at an unobserved location \mathbf{s}_0 is defined as $\hat{Y}_{\text{DK}}(\mathbf{s}_0) = f_{\text{DK}}(\mathbf{s}_0; \hat{\boldsymbol{\theta}})$.

One major advantage of our DeepKriging method is that we can adjust the number of neurons, activation functions, and loss functions to cater to different data types and model interpretations. For example, for predicting continuous variables, as in a regression problem, we choose $N_L = 1$, $\psi_L(\cdot)$ as an identity function, and the loss function as the MSE. Figure 1 provides a visualization of a DeepKriging structure in a two-dimensional prediction for continuous data. For predicting categorical variables, as in a classification problem, we choose N_L as the number of categories, $\psi_L(\cdot)$ as a softmax function, and the loss function as the cross-entropy loss. For the activation functions in the hidden layers, we choose the rectified linear unit (ReLU) as a default, which allows us to keep the linear relationship in the KL expansion, but add some deactivated neurons to select the best number of basis functions. The DeepKriging structure also allows the covariate effects to be spatially varying.

Regularizing the DeepKriging network structure includes adding dropout

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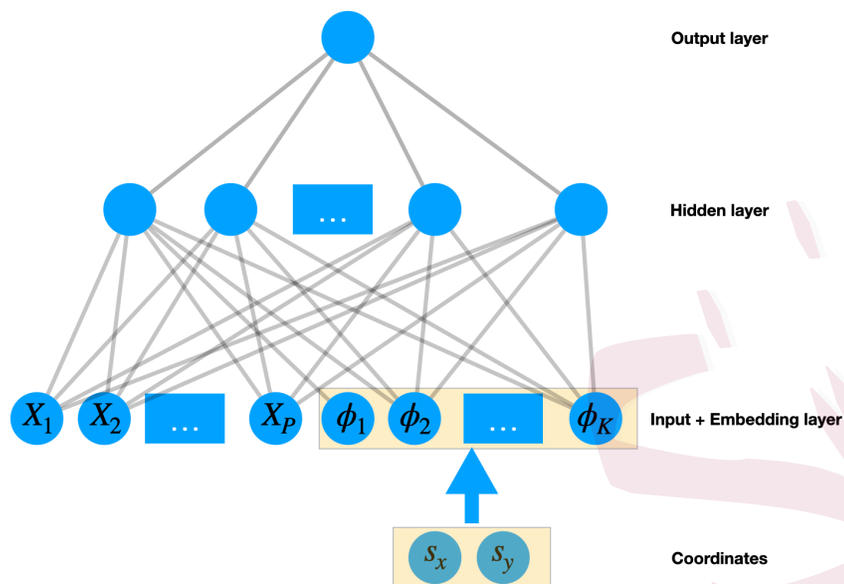


Figure 1: Visualization of the DeepKriging structure in a 2D spatial prediction based on a three-layer DNN

layers to mitigate overfitting, adding batch-normalization layers to regularize the covariates and the basis functions to the same scale, and removing all-zero columns in the basis matrix, owing to the compactly supported structure of the basis function. Details of the default setting of our DeepKriging network structure are included in Section S2 of the Supplementary Material. The time complexity of our DeepKriging method is about $O(N_{\text{neuron}})$, where N_{neuron} is the number of neurons in the network. The computation cost depends on the width and depth of the network. However, the computation is highly parallelizable, and can be accelerated significantly using CPUs and GPUs.

3. Theoretical Properties of DeepKriging

DeepKriging provides a novel spatial prediction framework using deep learning. It differs from classical Kriging methods in several aspects. First, Kriging prediction is a linear combination of observations. In contrast, DeepKriging prediction is linked to the observations by the weights and biases through model training, and is typically nonlinear in the observations (see Section S3.1). Second, DeepKriging does not assume a GP with a certain covariance function, instead modeling the spatial dependence using basis functions. Third, Kriging predicts the random process $Y(\mathbf{s})$ at an unobserved location; in contrast, DeepKriging approximates the process using a deterministic continuous function.

In this section, we provide important theoretical properties of DeepKriging, including 1) the underlying relationship between DeepKriging and Kriging, 2) the accuracy of DeepKriging in terms of the prediction error compared with that of Kriging, and 3) how the spatial dependence is measured in the DeepKriging framework. These three aspects are critical to understanding our DeepKriging method.

3.1 The link between DeepKriging and Kriging-based methods

3.1 The link between DeepKriging and Kriging-based methods

DeepKriging is closely related to Kriging and its associated variants, which can be classified as multi-resolution processes (Nychka et al., 2015; Kleiber and Nychka, 2015; Katzfuss, 2017) and Gaussian predictive processes (Banerjee et al., 2008, 2010). These processes lead to spatial predictions that can be treated as linear functions of embedded features $\mathbf{x}_\phi(\mathbf{s}_0)$, and thus can be approximated using DeepKriging.

One example is the fixed-rank Kriging (FRK) proposed by Cressie and Johannesson (2008), who use one of the low-rank approximations of the covariance matrix to speed up the computation of universal Kriging. Similar to DeepKriging, they represent the spatial random effects $\nu(\mathbf{s})$ by K basis functions, that is, $\nu(\mathbf{s}) = \boldsymbol{\phi}(\mathbf{s})^T \boldsymbol{\eta}$, where $\boldsymbol{\eta}$ is a K -dimensional Gaussian random vector, with $\text{Cov}(\boldsymbol{\eta}) = \boldsymbol{\Sigma}_K$. They assume that the model for $Y(\mathbf{s})$ is $Y(\mathbf{s}) = \mathbf{x}(\mathbf{s})^T \boldsymbol{\beta} + \nu(\mathbf{s}) = \mathbf{x}(\mathbf{s})^T \boldsymbol{\beta} + \boldsymbol{\phi}(\mathbf{s})^T \boldsymbol{\eta}$. The covariance matrix of $Z(\mathbf{s}) = Y(\mathbf{s}) + \varepsilon(\mathbf{s})$, where $\varepsilon(\mathbf{s})$ is white noise with variance $\sigma^2(\mathbf{s})$, is given by $\boldsymbol{\Sigma} = \boldsymbol{\Phi} \boldsymbol{\Sigma}_K \boldsymbol{\Phi}^T + \mathbf{V}$, where $\boldsymbol{\Phi} = \{\boldsymbol{\phi}(\mathbf{s}_1), \dots, \boldsymbol{\phi}(\mathbf{s}_N)\}^T$ is an $N \times K$ basis matrix, and $\mathbf{V} = \text{diag}\{\sigma^2(\mathbf{s}_1), \dots, \sigma^2(\mathbf{s}_N)\}$ is an $N \times N$ diagonal matrix. The FRK prediction as a linear function of \mathbf{z} is given by

$$\hat{Y}_{\text{FRK}}(\mathbf{s}_0) = \mathbf{x}(\mathbf{s}_0)^T \hat{\boldsymbol{\beta}} + \boldsymbol{\phi}(\mathbf{s}_0)^T \boldsymbol{\Sigma}_K \boldsymbol{\Phi}^T \boldsymbol{\Sigma}^{-1} (\mathbf{z} - \mathbf{X} \hat{\boldsymbol{\beta}}), \quad (3.1)$$

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where $\mathbf{X} = (\mathbf{x}(\mathbf{s}_1), \dots, \mathbf{x}(\mathbf{s}_N))^T$ is an $N \times P$ matrix, $\hat{\boldsymbol{\beta}} = (\mathbf{X}^T \boldsymbol{\Sigma}^{-1} \mathbf{X})^{-1} \mathbf{X}^T \boldsymbol{\Sigma}^{-1} \mathbf{z}$, and $\boldsymbol{\Sigma}^{-1}$ has a computationally simple form that involves inverting the fixed-rank $K \times K$ positive-definite matrix $\boldsymbol{\Sigma}_K$ and the $N \times N$ diagonal matrix \mathbf{V} . Writing Equation (3.1) as $\hat{Y}_{\text{FRK}}(\mathbf{s}_0) = \mathbf{x}(\mathbf{s}_0)^T \hat{\boldsymbol{\beta}} + \boldsymbol{\phi}(\mathbf{s}_0)^T \hat{\boldsymbol{\alpha}}$, where $\hat{\boldsymbol{\alpha}} = \boldsymbol{\Sigma}_K \boldsymbol{\Phi}^T \boldsymbol{\Sigma}^{-1} (\mathbf{z} - \mathbf{X} \hat{\boldsymbol{\beta}})$, implies that the FRK prediction $\hat{Y}_{\text{FRK}}(\mathbf{s}_0)$ is linear in the P covariates $\mathbf{x}(\mathbf{s}_0)$ and K basis functions $\boldsymbol{\phi}(\mathbf{s}_0)$. This is a special case of DeepKriging in which we set all activation functions to be linear.

FRK usually chooses K to be much smaller than N in order to speed up the computation for large data sets. Because the covariance $\boldsymbol{\Phi} \boldsymbol{\Sigma}_K \boldsymbol{\Phi}^T$ has at most rank K , such a low-rank approximation of the covariance matrix may fail to capture the high-frequency variation or small-scale spatial dependence in the spatial process (Stein, 2014). In contrast, for DeepKriging, K needs to be sufficiently large ($K > N$) in order to have a good approximation of the spatial random effect $\nu(\mathbf{s})$. Thus our method captures more spatial information in the prediction.

By setting $K = N$ in the FRK, we can see that the (universal) Kriging prediction in Equation (2.3) is also a linear function of $\mathbf{x}_\phi(\mathbf{s}_0) = (\mathbf{x}(\mathbf{s}_0)^T, \boldsymbol{\phi}(\mathbf{s}_0)^T)^T$.

A detailed proof is provided in Section S1.1 of the Supplementary Material. This result implies that a Kriging prediction with any covariance function can be expressed linearly by the embedding features $\mathbf{x}_\phi(\mathbf{s}_0)$. In this

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sense, DeepKriging generalizes Kriging by allowing for nonlinear functions of $\mathbf{x}_\phi(\mathbf{s}_0)$ in the prediction.

3.2 DeepKriging in decision theory

Our DeepKriging prediction procedure conventionally follows an approximation-estimation decomposition described in Fan et al. (2019). Let \mathcal{F} be the function space expressible by a particular DNN model, and $\hat{Y}_N(\mathbf{s}_0)$ be the final prediction from the model based on N observed locations. The following decomposition of the total risk between the true value $Y(\mathbf{s}_0)$ and the prediction $\hat{Y}_N(\mathbf{s}_0)$ implies three sources of errors:

$$R\{Y(\mathbf{s}_0), \hat{Y}_N(\mathbf{s}_0)\} = \underbrace{R\{Y(\mathbf{s}_0), \hat{Y}_{\mathcal{F}}^{\text{opt}}(\mathbf{s}_0)\}}_{\text{approximation error}} + \underbrace{R\{\hat{Y}_{\mathcal{F}}^{\text{opt}}(\mathbf{s}_0), \hat{Y}_N^{\text{opt}}(\mathbf{s}_0)\}}_{\text{estimation error}} + \underbrace{R\{\hat{Y}_N^{\text{opt}}(\mathbf{s}_0), \hat{Y}_N(\mathbf{s}_0)\}}_{\text{optimization error}}.$$

The approximation error relates to the model capacity, and is defined as the risk between the true process $Y(\mathbf{s}_0)$ and the optimal predictor $\hat{Y}_{\mathcal{F}}^{\text{opt}}(\mathbf{s}_0) = \operatorname{argmin}_{\hat{Y} \in \mathcal{F}} R(\hat{Y}(\mathbf{s}_0), Y(\mathbf{s}_0))$ as a function in \mathcal{F} . The estimation error is defined as the risk between $\hat{Y}_N^{\text{opt}}(\mathbf{s}_0)$ and $\hat{Y}_{\mathcal{F}}^{\text{opt}}(\mathbf{s}_0)$, where $\hat{Y}_N^{\text{opt}}(\mathbf{s}_0) = \hat{Y}_N(\mathbf{s}_0; \hat{\boldsymbol{\theta}})$, with $\hat{\boldsymbol{\theta}} = \operatorname{argmin}_{\boldsymbol{\theta}} \frac{1}{N} \sum_{n=1}^N L\{\hat{Y}_N(\mathbf{s}_n; \boldsymbol{\theta}), z(\mathbf{s}_n)\}$; this type of error is affected by the complexity of \mathcal{F} , and relates to the generalization power of the model. The optimization error is the empirical risk between $\hat{Y}_N^{\text{opt}}(\mathbf{s}_0)$ and $\hat{Y}_N(\mathbf{s}_0)$.

The function class of the Kriging prediction in Equation (2.3), \mathcal{F}_{UK} , can be viewed as the space of linear functions of $\mathbf{x}(\mathbf{s}_0)$ and \mathbf{z} taking the

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form $\mathbf{x}(\mathbf{s}_0)^T \boldsymbol{\beta} + \mathbf{z}^T \boldsymbol{\gamma}$, whereas the function class of DeepKriging, \mathcal{F}_{DK} , is the function space generated by the DNN described in (2.4). The universal approximation theorem (Theorem 2.3.1 of Csáji (2001)) claims that every continuous function of the features $\mathbf{x}_\phi(\mathbf{s})$, denoted as $\mathbb{C}(\mathbf{x}_\phi)$, can be arbitrarily well approximated by a feed-forward neural network with a single hidden layer that contains a finite number of hidden neurons and with an arbitrary activation function. This indicates that the optimal DeepKriging prediction with a single hidden layer and a finite loss function has the largest model capacity in $\mathbb{C}(\mathbf{x}_\phi)$, that is, $\hat{Y}_{\mathcal{F}_{\text{DK}}}^{\text{opt}}(\mathbf{s}_0) = \hat{Y}_{\mathbb{C}(\mathbf{x}_\phi)}^{\text{opt}}(\mathbf{s}_0)$. This result holds for any type of data (i.e., continuous or discrete) and for any type of task (i.e., regression or classification). Therefore, the optimal DeepKriging prediction has larger capacity than the Kriging prediction in terms of minimizing the approximation error, that is, $\mathbb{E}\{L(\hat{Y}_{\mathcal{F}_{\text{DK}}}^{\text{opt}}(\mathbf{s}_0), Y(\mathbf{s}_0))\} \leq \mathbb{E}\{L(\hat{Y}_{\mathcal{F}_{\text{UK}}}^{\text{opt}}(\mathbf{s}_0), Y(\mathbf{s}_0))\}$. A detailed proof is provided in Section S1.2 of the Supplementary Material. Similarly, the optimal DeepKriging prediction has larger model capacity than that of the FRK prediction. FRK can be viewed as DeepKriging, with a single hidden layer containing a finite number of neurons and a linear activation function. By allowing for a large number of basis functions, multiple layers, more flexible activation functions, and a wide network, DeepKriging yields nonlinear predictions that

can appropriately capture the spatial dependence in a spatial process.

3.3 DeepKriging as a GP

Neal (1994) showed that a single-layer fully connected neural network with an independent and identically distributed (i.i.d.) prior over its parameters (i.e., weights and biases) is equivalent to a GP with the limit of an infinite network width (i.e., an infinite number of hidden neurons). Later, Lee et al. (2018) derived the exact equivalence between infinitely wide deep networks and GPs. Consequently, a similar correspondence to GPs also holds for our DeepKriging network.

We start with a regression-type DeepKriging model with a single hidden layer containing N_1 neurons. The input features are $\mathbf{x}_\phi(\mathbf{s}) = (\mathbf{x}(\mathbf{s})^T, \boldsymbol{\phi}(\mathbf{s})^T)^T \in \mathbb{R}^{P+K}$, and the output is $\hat{Y}_{\text{DK}}(\mathbf{s}) = b^1 + \sum_{j=1}^{N_1} w_j^1 a_j^1(\mathbf{s})$, where $a_j^1(\mathbf{s}) = \psi_1(b_j^0 + \sum_{i=1}^{P+K} w_{ji}^0 \mathbf{x}_\phi^{(i)}(\mathbf{s}))$, with $\mathbf{x}_\phi^{(i)}(\mathbf{s})$ being the i th component of $\mathbf{x}_\phi(\mathbf{s})$. The weights (w_j^1, w_{ji}^0) and biases (b^1, b_j^0) are independent and drawn randomly to have a zero mean and variances σ_w^2/N_1 and σ_b^2 , respectively. Consequently, the post-activations a_j^1 and $a_{j'}^1$ are independent for $j \neq j'$. Moreover, because $\hat{Y}_{\text{DK}}(\mathbf{s})$ is a sum of i.i.d terms, it follows from the central limit theorem that in the limit of infinite width $N_1 \rightarrow \infty$, $\hat{Y}_{\text{DK}}(\mathbf{s})$ follows a Gaussian distribution from the multi-dimensional central limit theorem, any

3.3 DeepKriging as a GP

finite collection of $\{\hat{Y}_{\text{DK}}(\mathbf{s}_1), \hat{Y}_{\text{DK}}(\mathbf{s}_2), \dots, \hat{Y}_{\text{DK}}(\mathbf{s}_n)\}$ has a joint multivariate Gaussian distribution, which is exactly the definition of a GP. Therefore, we conclude that with sufficiently large N_1 , \hat{Y}_{DK} is a GP with zero mean and covariance function

$$C^1(\mathbf{s}, \mathbf{s}') = E\{\hat{Y}_{\text{DK}}(\mathbf{s})\hat{Y}_{\text{DK}}(\mathbf{s}')\} = \sigma_b^2 + \sigma_w^2 E\{a_j^1(\mathbf{s})a_j^1(\mathbf{s}')\} = \sigma_b^2 + \sigma_w^2 C(\mathbf{s}, \mathbf{s}'),$$

where $C(\mathbf{s}, \mathbf{s}')$ is obtained by integrating against the distribution of w^0 , b^0 , as in Neal (1994).

For DeepKriging with deeper layers, the induced covariance function can be obtained in a recursive way, following Lee et al. (2018):

$$C^l(\mathbf{s}, \mathbf{s}') = \sigma_b^2 + \sigma_w^2 F_\psi(C^{l-1}(\mathbf{s}, \mathbf{s}'), C^{l-1}(\mathbf{s}, \mathbf{s}), C^{l-1}(\mathbf{s}', \mathbf{s}')), \quad (3.2)$$

where $F_\psi(\cdot)$ is a deterministic function that depends only on the activation function ψ . An iterative series of computations yields the covariance C^L for the GP describing the network's final output, $\hat{Y}_{\text{DK}}(\mathbf{s})$. For the base case, $C^0(\mathbf{s}, \mathbf{s}') = \sigma_b^2 + \sigma_w^2 \{\mathbf{x}_\phi(\mathbf{s})^T \mathbf{x}_\phi(\mathbf{s}') / (P + K)\}$. The aforementioned results require the assumption of infinitely many hidden neurons in each layer. However, when the prior distributions of the weights and biases are Gaussian, this condition is not needed.

For certain activation functions, Equation (3.2) can be computed analytically. The simplest case occurs when the activation function is an iden-

3.3 DeepKriging as a GP

tivity function $\psi_l(x) = x$ and no covariate effect exists. Then, $\hat{Y}_{\text{DK}}(\mathbf{s})$ is a linear function of the basis functions $\boldsymbol{\phi}(\mathbf{s})$, that is, $\hat{Y}_{\text{DK}}(\mathbf{s}) = b + \mathbf{w}^T \boldsymbol{\phi}(\mathbf{s})$, where b and \mathbf{w} are combined biases and weights, respectively. In this case, the induced covariance function of \hat{Y}_{DK} is given by $C^L(\mathbf{s}, \mathbf{s}') = \sigma_b^2 + \sigma_w^2 \boldsymbol{\phi}(\mathbf{s})^T \boldsymbol{\phi}(\mathbf{s}')$, which is the basis approximation of a spatial covariance function.

In the case of ReLU nonlinearity, Equation (3.2) has a closed form of the well-known arc-cosine kernel (Cho and Saul, 2009):

$$C^l(\mathbf{s}, \mathbf{s}') = \sigma_b^2 + \frac{\sigma_w^2}{2\pi} \sqrt{C^{l-1}(\mathbf{s}, \mathbf{s})C^{l-1}(\mathbf{s}', \mathbf{s}')} \left\{ \sin(\theta_{\mathbf{s}, \mathbf{s}'}^{l-1}) + (\pi - \theta_{\mathbf{s}, \mathbf{s}'}^{l-1}) \cos(\theta_{\mathbf{s}, \mathbf{s}'}^{l-1}) \right\},$$

where $\theta_{\mathbf{s}, \mathbf{s}'}^l = \cos^{-1}(C^l(\mathbf{s}, \mathbf{s}') / \sqrt{C^l(\mathbf{s}, \mathbf{s})C^l(\mathbf{s}', \mathbf{s}')}))$. When no analytic form of the resulted covariance function exists, it can be computed numerically, as described in Lee et al. (2018).

Consider a regression-type DeepKriging model, with a single hidden layer and no covariate effects. It can be shown that with infinitely many hidden neurons, the covariance function of the output $\hat{Y}_{\text{DK}}(\mathbf{s})$ for any two nearby locations has the form

$$C(\mathbf{s}, \mathbf{s}') = v(\mathbf{s}) + v(\mathbf{s}') - c \|\boldsymbol{\phi}(\mathbf{s}) - \boldsymbol{\phi}(\mathbf{s}')\|^2, \quad (3.3)$$

where $\boldsymbol{\phi}(\mathbf{s})$ is the basis vector at location \mathbf{s} , $v(\mathbf{s}) > 0$ is related to the variance when $\mathbf{s} = \mathbf{s}'$, and c is a scaling parameter. The proof is provided in Section S1.3 of the Supplementary Material. As a special case, if only

3.3 DeepKriging as a GP

the coordinates are used in the features, then $\|\phi(\mathbf{s}) - \phi(\mathbf{s}')\|^2 = \|\mathbf{s} - \mathbf{s}'\|^2$, $v(\mathbf{s}) = v(\mathbf{s}') = v$, and thus $C(\mathbf{s}, \mathbf{s}') = v - c\|\mathbf{s} - \mathbf{s}'\|^2$, which contains less information than in Equation (3.3). Therefore, the embedding layer in DeepKriging yields spatial covariance structures that are more flexible than when simply using the coordinates.

Next, we show how the DeepKriging-induced covariance function can approximate the common stationary covariance functions in spatial statistics. Let the basis functions be $\phi_l(\mathbf{s}) = k(\mathbf{s}, \mathbf{u}_l)$ based on a certain kernel function $k(\cdot, \cdot)$ and knot \mathbf{u}_l , for $l = 1, \dots, K$. If the \mathbf{u}_l form a fine grid of knots covering the spatial domain, then

$$\begin{aligned} \|\phi(\mathbf{s}) - \phi(\mathbf{s}')\|^2 &= \sum_{l=1}^K \{k(\mathbf{s}, \mathbf{u}_l) - k(\mathbf{s}', \mathbf{u}_l)\}^2 \approx \int \{k(\mathbf{s}, \mathbf{u}) - k(\mathbf{s}', \mathbf{u})\}^2 d\mathbf{u} \\ &= \int k(\mathbf{s}, \mathbf{u})^2 + k(\mathbf{s}', \mathbf{u})^2 - 2k(\mathbf{s}, \mathbf{u})k(\mathbf{s}', \mathbf{u}) d\mathbf{u}. \end{aligned}$$

Note that the last term is the kernel convolution approximation to a covariance function. Higdon (2002) shows that by selecting an appropriate kernel function, we can approximate any stationary covariance function based on the kernel convolution. Furthermore, the induced covariance function of DeepKriging possesses favorable physical interpretations. For example, DeepKriging can yield the Matérn covariance function, also commonly used in Kriging, because it is related to a stochastic partial differential equa-

tion (SPDE) of Laplace type (Whittle, 1954). In addition, DeepKriging can induce a GP that approximates a fractional Brownian motion, based on the example of a DNN provided in Neal (1996).

4. Simulation Studies

4.1 DeepKriging on a one-dimensional GP

We first consider the performance of DeepKriging when data are simulated from a one-dimensional stationary GP, where the Kriging prediction is optimal. We also compare DeepKriging with two naive DNNs: a DNN with the intercept $x(s) = 1$ as its only input, and a DNN with $x(s) = 1$ and coordinate s as the inputs. We also consider a Kriging prediction with the true covariance function and with an estimated Matérn covariance function. The simulation design is illustrated in Section S3.1 of the Supplementary Material.

Figure S1 in the Supplementary Material shows the prediction for one of the sample data sets using each of the five prediction methods. The DNN with only the intercept predicts the mean of the process. Although including the coordinate s in the DNN improves the prediction, it fails to capture the high-frequency variability, and cannot reflect the spatial correlations of the true process. Moreover, the DeepKriging prediction and

4.2 DeepKriging on two-dimensional nonstationary data

the optimal Kriging prediction almost overlap.

To further validate the performance, we calculate the root MSE (RMSE) and mean absolute percentage error (MAPE) on the testing data over 100 replicated samples in Table S1 in the Supplementary Material, where MAPE is defined as $\frac{1}{N_{\text{test}}} \sum_{n=1}^{N_{\text{test}}} \frac{Y_n^{\text{pred}} - Y_n^{\text{true}}}{Y_n^{\text{true}}}$, N_{test} is the number of testing samples, Y_n^{pred} is the predicted value, and Y_n^{true} is the true value. As the minimum-MSE predictor, the Kriging prediction with the true covariance function has the smallest RMSE, as expected. DeepKriging performs similarly to the two Kriging predictions, and significantly outperforms the two naive DNN models. We also provide the results on the training set in Table S1 in the Supplementary Material. Again, the Kriging prediction with the true covariance function performs best. The DeepKriging prediction is comparable with the optimal Kriging prediction, and outperforms the Kriging prediction with an estimated covariance function and the two naive DNN models in terms of both the RMSE and the MAPE.

4.2 DeepKriging on two-dimensional nonstationary data

In this section, we evaluate the performance of DeepKriging on two-dimensional nonstationary data, so that the procedure is designed to resemble the real-data application in Section 5. The simulation details are included in Section

4.2 DeepKriging on two-dimensional nonstationary data

S3.2 of the Supplementary Material.

We use 10-fold cross-validation to show the performance of DeepKriging, Kriging with an estimated stationary covariance function, and the baseline DNN with only the coordinates s in the features. We calculate the RMSEs and MAPEs on the testing data set, and show the results in Figure S2(b) and Table S2. DeepKriging significantly outperforms Kriging in terms of the RMSEs and MAPEs, because Kriging assumes a stationary covariance function, whereas DeepKriging captures the nonstationarity in the data. In addition, the baseline DNN is better than Kriging in this example, because [the data are nonGaussian, and Kriging is no longer optimal](#). Moreover, the baseline DNN performs worse than DeepKriging, as expected. The MAPE from DeepKriging is lower than that of the baseline DNN, but higher than that of Kriging. This can happen because we are using the MSE as the loss function in DeepKriging, so it does not necessarily possess the lowest MAPE. We also calculate the RMSEs and MAPEs on the training data set (see Table S2). Kriging outperforms the other two models in terms of both metrics. This is because the errors for the training data set can be viewed as the variance estimates of the assumed model, much as in a regression model. Kriging tends to underestimate this variance, leading to a worse prediction on the testing data set.

4.2 DeepKriging on two-dimensional nonstationary data

Additional simulations (see Section S3 of the Supplementary Material) show that DeepKriging is nonlinear in observation, whereas Kriging is linear. Furthermore, computation time comparison based on the same simulation study shows that Kriging is faster for small sample sizes ($N < 1,500$), but that DeepKriging is much more scalable when the sample size increases. This is because when the sample size is small, the computation time is still under control for Kriging, but for DeepKriging, the number of parameters is large, owing to the large width and depth of the network, making the computation time longer than that of Kriging. When the sample size increases, the computational burden of both methods increases, but for DeepKriging, we can use parallel computing with CPUs or GPUs to accelerate the computation. Therefore, our DeepKriging method is much more scalable to large data sizes. For example, when $N = 12,800$, it takes more than 1.5 hours (5,663 seconds) to implement a Kriging model, whereas DeepKriging takes only 3.5 minutes (214 seconds) without GPU acceleration, and 1.5 minutes (94 seconds) with a Tesla P100 GPU.

5. Application

5.1 Challenges of predicting $PM_{2.5}$ concentration

$PM_{2.5}$, fine particulate matter of less than $2.5 \mu m$, is a harmful air pollutant. Its adverse effects are associated with many diseases, such as respiratory disease (Peng et al., 2009) and myocardial infarction (Peters et al., 2001); see the review by the World Health Organization (2013). Therefore, it is essential to obtain a high-resolution map of $PM_{2.5}$ exposure in order to assess its impact. The measurements from monitoring networks are the best characterization of $PM_{2.5}$ concentration at a given time and location. However, data from monitoring locations are often sparsely distributed, so are out of spatial and temporal alignment with health outcomes. At the same time, it is known that $PM_{2.5}$ concentration is associated with meteorological conditions such as temperature and relative humidity (Jacob and Winner, 2009), where the meteorological data or data products are often easy to access with good spatial coverage and resolutions. Hence, interpolations of $PM_{2.5}$ concentration using data from monitoring networks and other meteorological data is a promising field of research (Di et al., 2016), where spatial prediction plays a central role.

The modeling and prediction of $PM_{2.5}$ concentration are challenging.

5.2 Data and preprocessing

First, $\text{PM}_{2.5}$ concentration data are obviously nonGaussian, and thus classical Kriging methods are inappropriate here. Second, $\text{PM}_{2.5}$ data from monitoring stations are irregular and sparse, but many interpolation methods require lattice data. Third, it is more important, but challenging, to understand the risk of high pollution and to predict pollution levels, such as being low, medium, or high; statistically, these two questions are related to estimating the probability over a threshold and a classification problem, respectively. Quantile regression and CNNs have been used to overcome some of the above issues (Reich et al., 2011; Porter et al., 2015; Di et al., 2016). However, there is not yet a unified method that can handle all of the aforementioned tasks.

5.2 Data and preprocessing

To tackle the above-mentioned problems, we apply the proposed DeepKriging method to the spatial prediction of $\text{PM}_{2.5}$ concentrations based on meteorological variables. Meteorological data are obtained from the NCEP North American Regional Reanalysis (NARR) product. Reanalysis is a gridded data set that represents the state of the atmosphere, incorporating observations and outputs of numerical weather prediction models from past to present-day. Reanalysis data are often used to represent the “true state”

5.2 Data and preprocessing

of the atmosphere according to observations, and thus we use these data as the “observed data” for the covariates. Six meteorological variables are used in this study: 1) air temperature at 2 m; 2) relative humidity at 2 m; 3) accumulated total precipitation; 4) surface pressure; 5) the u-component of wind; and 6) the v-component of wind at 10 m. The covariates from the NARR product are gridded data on June 05, 2019, with a spatial resolution of about 32×32 km covering the continental United States, containing 7,706 gridded cells in total. Because the units of the meteorological variables vary, we use min-max normalization to re-scale the data before implementing the models. Daily averaged data of $\text{PM}_{2.5}$ concentrations are observed from 841 monitoring stations. Because the coordinates from NARR and those from stations are not identical, and because some stations are too close to each other, we keep the spatial resolution of NARR and average the $\text{PM}_{2.5}$ measurements of nearby monitoring stations in the same grid cell. After the matching, 604 grid cells remain for the model training, with the $\text{PM}_{2.5}$ concentration value at each location shown in Figure 2(a). Our goal is to predict the $\text{PM}_{2.5}$ concentrations at any s_0 of the other $7,706 - 604 = 7,102$ locations, where the $\text{PM}_{2.5}$ concentrations are not observed, but the covariates are provided by the reanalysis data.

5.3 Model fitting and results

Our aim is to predict the $PM_{2.5}$ concentration values at unobserved grid cells where the six meteorological variables are provided. We use 10-fold cross-validation to verify the performance of DeepKriging. For comparison purposes, we also show the results from Kriging and the baseline DNN with the six covariates and coordinates. We calculate the MSEs and MAEs as the validation criteria, shown in the first two rows of Table 1, both of which show that DeepKriging outperforms the baseline DNN and Kriging.

To assess the risk of high $PM_{2.5}$ pollution, we can use DeepKriging for spatial data classification. Specifically, we threshold the $PM_{2.5}$ concentrations by $12.0 \mu g/m^3$, which is the threshold between “good” and “moderate” levels for the daily mean of EPA national ambient air quality standards (NAAQS) (EPA, 2012). Based on the classified data, we can implement a binary classification using DeepKriging by assuming that the actual values of $PM_{2.5}$ concentration are unknown. A direct comparison with Kriging is not feasible, because Kriging is not suitable for binary classification. Instead, we predict the continuous $PM_{2.5}$ concentrations using Kriging, and classify the predictions by thresholding them at $12.0 \mu g/m^3$. We then use 10-fold cross-validation to show the classification accuracy, presented in the last row of Table 1. As shown, DeepKriging significantly outperforms

5.3 Model fitting and results

Kriging and the baseline DNN in terms of classification accuracy.

Table 1: Model performance based on 10-fold cross-validation. The MSEs and MAEs of the predictions and the classification accuracy (ACC) for predicting $\text{PM}_{2.5}$ concentrations above $12.0 \mu\text{g}/\text{m}^3$ are used as validation criteria. The mean and standard deviation (SD) of the 10 sets of validation errors or accuracy are provided in the table.

Parameters	DeepKriging		Baseline DNN		Kriging	
	Mean	SD	Mean	SD	Mean	SD
MSE	1.632	.572	3.632	.925	3.361	.773
MAE	.892	.103	1.448	.162	1.365	.178
ACC	95.2%	2.6%	89.6%	4.8%	88.5%	4.6%

Based on the model fitting, we can predict the $\text{PM}_{2.5}$ concentration, level of pollution, and risk of a high pollution level over the threshold of $12 \mu\text{g}/\text{m}^3$ at unobserved locations using the NARR data. Figure 2(a) shows the raw $\text{PM}_{2.5}$ station data from the AQS database. Figure 2(b) shows a smooth map of the predicted $\text{PM}_{2.5}$ concentration from DeepKriging. We also provide the distribution prediction (details and algorithms are included in Section S5 of the Supplementary Material) to obtain the predicted risk defined as $\mathbb{P}\{\text{PM}_{2.5} > 12 \mu\text{g}/\text{m}^3\}$, shown in Figure 2(c). This map implies that high $\text{PM}_{2.5}$ pollution risks exist over much of the eastern United

States. We further compare the results to the Kriging prediction in Figure 2(d), showing that DeepKriging provides more local features/patterns than Kriging does.

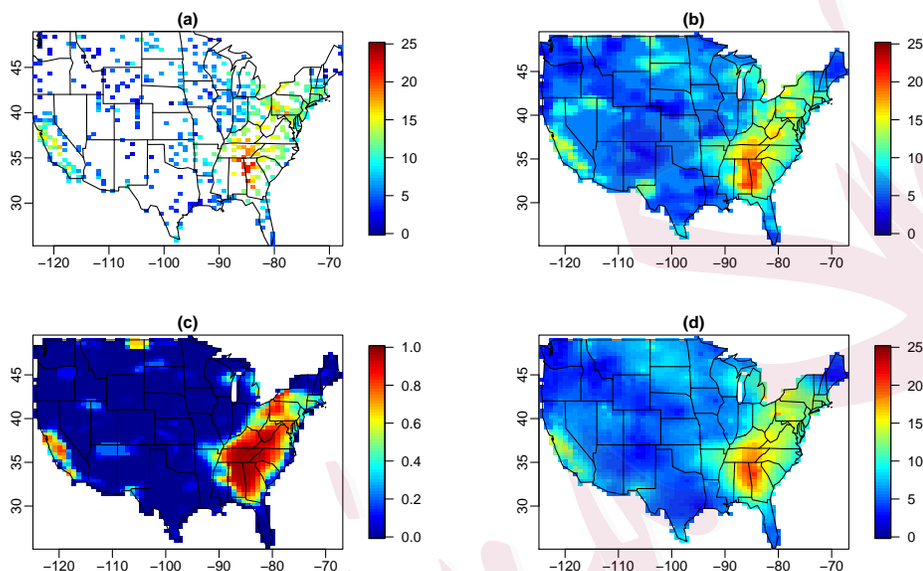


Figure 2: (a) $\text{PM}_{2.5}$ concentration ($\mu\text{g}/\text{m}^3$) collected from monitoring stations. (b) Predicted $\text{PM}_{2.5}$ concentration using DeepKriging. (c) Predicted risk of high pollution $\mathbb{P}\{\text{PM}_{2.5} > 12 \mu\text{g}/\text{m}^3\}$ based on a distribution prediction using DeepKriging. (d) Predicted $\text{PM}_{2.5}$ concentration using Kriging.

6. Discussion

We have proposed a new spatial prediction model using a DNN that incorporates spatial dependence by using a set of basis functions. Our method

does not assume parametric forms of the covariance functions or data distributions and, in general, is compatible with nonstationarity, nonlinear relationships, and nonGaussian data. Our DeepKriging framework can provide uncertainty quantification using the distribution prediction method described in Section S5 of the Supplementary Material.

Classical Kriging methods consider predictions as linear combinations of observations, which impedes their interaction with several machine learning frameworks. Some evidence of the equivalence between Kriging and radial basis function interpolation has been provided by Matheron (1981). However, without modern machine learning tools, only a linear combination and a limited number of radial basis functions have been investigated, and are viewed as a less favorable choice to Kriging (Dubrule, 1983, 1984). This work provides a new perspective on deep learning in spatial prediction with a large number of basis functions. We have shown that the proposed method is superior to Kriging in many aspects, both theoretically and numerically, in our simulation and real application. For instance, DeepKriging is more scalable large data sets and suits for more data types than Kriging does. DeepKriging also has a GP representation with flexible spatial covariance structures, which enables Bayesian inference on regression tasks by evaluating the corresponding GP. More importantly, the proposed DeepKriging

framework connects regression-based prediction and spatial prediction so that many other machine learning algorithms can be applied.

In general applications, it is possible that the covariates at the new location \mathbf{s}_0 are not observed. One promising approach for coping with this problem is to find the true values of the missing covariates for a subset of the observations, and then, to train a machine learning algorithm to predict the values of those covariates for the rest (see, e.g., Imai and Khanna (2016)). However, Fong and Tyler (2021) showed that plugging in these predictions without regard for the prediction error renders regression analyses biased, inconsistent, and overconfident. They describe a procedure to avoid these inconsistencies that combines a new sample-splitting scheme and a general method of moments (GMM) estimator to form an efficient and consistent estimator. Overall, it is nontrivial to address the problem of missing covariates: intuitive strategies such as plugging in machine learning predictions lead to bias and inconsistency, while the implementation of a more complicated method, such as that in Fong and Tyler (2021), requires extra assumptions (e.g., the exclusion restriction condition) and increases the computational burden. If the goal is to predict both the response and the covariates, a multivariate version of DeepKriging could be developed. These are left to future work.

Supplementary Material

The online Supplementary Material contains proofs of the lemmas and theorems (Section S1), the settings for the DeepKriging network structure (Section S2), details of the simulation studies (Section S3), additional simulation studies (Section S4), distribution prediction and uncertainty quantification (Section S5), and the source code and data required to reproduce the research (Section S6).

Acknowledgments

This research was supported by the National Key Research and Development Program (2021YFA1000101), Zhejiang Provincial Natural Science Foundation of China (LZJWY22E090009), Natural Science Foundation of Shanghai (22ZR1420500), Open Research Fund of Key Laboratory of Advanced Theory and Application in Statistics and Data Science-MOE, ECNU, and King Abdullah University of Science and Technology (KAUST), Office of Sponsored Research (OSR), under Award No: OSR-2019-CRG7-3800.

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