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Interpretation of Convolutional Neural Networks for Acid Sulfate Soil Classification

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Convolutional neural networks (CNNs) have been originally used for computer vision tasks, such as image classification. While several digital soil mapping studies have been assessing these deep learning algorithms for the prediction of soil properties, their potential for soil classification has not been explored yet. Moreover, the use of deep learning and neural networks in general has often raised concerns because of their presumed low interpretability (i.e., the black box pitfall). However, a recent and fastdeveloping sub-field of Artificial Intelligence (AI) called explainable AI (XAI) aims to clarify complex models such as CNNs in a systematic and interpretable manner. For example, it is possible to apply model-agnostic interpretation methods to extract interpretations from any machine learning model. In particular, SHAP (SHapley Additive exPlanations) is a method to explain individual predictions: SHAP values represent the contribution of a covariate to the final model predictions. The present study aimed at, first, evaluating the use of CNNs for the classification of potential acid sulfate soils located in the wetland areas of Jutland, Denmark (c. 6,500 km²), and second and most importantly, applying a modelagnostic interpretation method on the resulting CNN model. About 5,900 soil observations and 14 environmental covariates, including a digital elevation model and derived terrain attributes, were utilized as input data. The selected CNN model yielded slightly higher prediction accuracy than the random forest models which were using original or scaled covariates. These results can be explained by the use of a common variable selection method, namely recursive feature elimination, which was based on random forest and thus optimized the selection for this method. Notably, the SHAP method results enabled to clarify the CNN model predictions, in particular through the spatial interpretation of the most important covariates, which constitutes a crucial development for digital soil mapping.

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1 INTRODUCTION

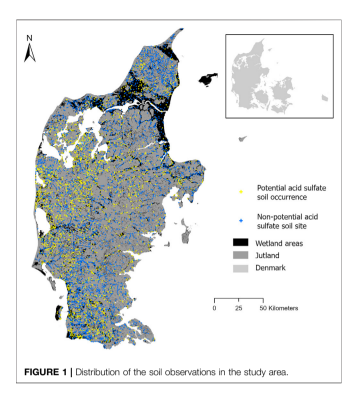
Digital soil mapping (DSM) techniques traditionally relate soil observations with point information extracted from different environmental covariates at corresponding locations. This representation of data as vectors at point location only partially describe a soil property or class. Taking into account spatial contextual information from covariates thus represents a crucial step forward for DSM and

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can be referred to as contextual mapping. During the last 2 decades, various studies attempted to incorporate spatial context through the preprocessing of environmental covariates. These original contextual mapping studies have used: different neighboring size to compute terrain attributes (Smith et al., 2006; Behrens et al., 2010b), adaptative filters applied on elevation data or derived attributes (Behrens et al., 2018b), covariates spatially transformed with wavelet analysis (Lark and Webster, 2001; Sun et al., 2017), hypercovariates created from elevation data (Behrens et al., 2010a; Behrens et al., 2014), and a multi-scale approach using aggregated covariates (Miller et al., 2015). All these studies relied on heavy preprocessing and/or subjective decisions concerning the resolution of covariates. Furthermore, Behrens et al. (2018a) compared different methods for multi-scaling analysis of terrain attributes combined to random forest with a deep neural network (DNN) algorithm (for clay and zinc content mapping). Other DSM studies investigated the use of DNNs for mapping soil moisture content (Song et al., 2016) and soil organic carbon content (Taghizadeh-Mehrjardi et al., 2020b; Emadi et al., 2020; Tao et al., 2020), identifying and delineating soil master horizons (Jiang et al., 2021), or spectral modelling (Singh and Kasana, 2019; Gholizadeh et al., 2020).

Within DNNs, convolutional neural networks (CNNs) represent a powerful and promising method for contextual mapping. CNNs have been originally used for computer vision tasks, such as image classification (Krizhevsky et al., 2012) and object detection. While the DNNs utilized in the aforementioned soil mapping studies use vector data extracted from covariates as input, CNNs use spatial contextual information extracted from covariates around each soil observation as input images (i.e., images or patches as xD arrays made of x covariates). Several DSM studies have been assessing CNNs for the mapping of soil organic carbon content (Padarian et al., 2019b; Wadoux et al., 2019; Wadoux, 2019), particle size fraction (Wadoux, 2019; Taghizadeh-Mehrjardi et al., 2020a), pH and nitrogen content (Wadoux, 2019), or for spectral modelling (Padarian et al., 2019c,a; Ng et al., 2019a,b; Xu et al., 2019; Pyo et al., 2020; Tsakiridis et al., 2020; Yang et al., 2020; Haghi et al., 2021; Zhong et al., 2021), but their potential for soil classification has not been investigated yet.

While DNNs provide high predictive power, their use almost systematically raise concerns because of their presumed low interpretability (i.e. the black box pitfall; Samek et al. (2019); Khaledian and Miller (2020)). However, a recent sub-field of Artificial Intelligence (AI) called explainable AI (XAI) focuses on clarifying complex AI models, such as DNNs, in a systematic and interpretable manner (Samek et al., 2019). For instance, it is possible to apply model-agnostic interpretation methods (Molnar, 2021) to enable extracting interpretations from any machine learning model. In particular, SHapley Additive exPlanations (SHAP) developed by Lundberg and Lee (2017) constitute a method to explain individual predictions, SHAP values representing the contribution of a covariate to the final model predictions. As of today, one DSM study (Padarian et al., 2020) evaluated the use of SHAP for interpreting the predictions of a CNN model, in particular in the geographical space, while



two other DSM studies used SHAP to interpret spectral models (Haghi et al., 2021; Zhong et al., 2021).

During the last decade, a few classical DSM methods were evaluated for mapping acid sulfate (AS) soils at different extents. Clustering through fuzzy k-means was tested to map coastal AS soils on a field extent in Australia (Huang et al., 2014a,b). Fuzzy logic was used to generate a preliminary map for AS soil occurrence at regional extent in Finland (Beucher et al., 2014). Artificial Neural Networks (ANNs) were assessed to map AS soil occurrence and several soil properties characterising the related environmental risks at catchment extent in southwestern Finland (Beucher et al., 2013, 2015). ANNs were also evaluated to map potential AS soils in the wetland areas of Jutland, Denmark (Beucher et al., 2017).

The present study will evaluate the use of CNNs for the classification of potential AS located in the wetland areas of Jutland, Denmark (c. $6,500~\rm km^2$). The two main objectives are, firstly, to assess the use of CNNs in comparison with random forest for AS soil classification, and secondly and most importantly, to apply a model-agnostic interpretation method to clarify the final model predictions.

2 MATERIAL AND METHODS

2.1 Study Area

The wetlands located over Jutland, the continental part of Denmark, constitute the study area (**Figure 1**). They cover about 6,500 km² and represent approximately 20% of the Jutland peninsula (Madsen et al., 1985). Wetlands are saturated soils, such as histosols, fluvisols and gleysols (IUSS

Working Group WRB, 2006). These areas were mainly used for hay production until the second half of the 19th century when tile drainage was introduced (Greve et al., 2014). Most of the wetlands (c. 5,100 km²) have been artificially drained and intensively farmed, the main crop being cereals and grass (Bou Kheir et al., 2010). Low-relief sandy glaciofluvial outwash plains from the Weichselian glaciation (i.e., Last Glacial Maximum) dominate in the western part of the study area (c. 1,200 km²). These plains area surrounded by slightly protruding islands of older and strongly eroded moraine landforms from the Saalian glaciation (c. 700 km²) (Madsen et al., 1985; Madsen and Jensen, 1988). The eastern part of the study area consists of Weichselian moraine landforms (c. 900 km²) while the northern part is composed of late- and post-glacial marine sediments (c. 2,400 km²) (Madsen and Jensen, 1988). Cretaceous limestone dominates in northern Jutland and Djursland. Tertiary mica-rich sand and clay prevail in the rest of Jutland (Madsen and Jensen, 1988). The study area has a temperate climate. The mean temperature is 0°C in the winter and 16°C in the summer. The average annual precipitation is about 800 mm in central Jutland (Danmarks Meteorologiske Institut, 1998).

2.2 Soil Observations

Soil observations were extracted from the Ochre Classification database which resulted from the potential AS soil mapping conducted in the 1980s (Madsen et al., 1985). Soils in wetland areas were targeted and surveyed through conventional mapping. Field work was carried out from May to October over a 3-year period (1981-83). The original selection of about 8,000 sampling sites was based on historical topographic maps, geological maps, soil maps and maps from previous moorland studies, representing an even distribution in wetlands and soil types (Madsen et al., 1985). The carbonate-free samples were considered as potential AS soils if their pH dropped below 3.0 within 16 weeks of incubation at room temperature (Madsen et al., 1985). For samples containing calcium carbonates, the acid-neutralizing capacity (ANC) was calculated from calcium and magnesium concentrations. The samples were classified as potential AS soils if the calculated ANC was smaller than the amount of sulfuric acid potentially produced by the oxidation of the pyrite contained in the sample (Madsen et al., 1985). A subset of 5,885 points was selected to be used in the present study as the original set included points which were incorrectly classified. The main misclassification concerned points displaying a minimum incubation pH between 3.1 and 4.0 and wrongly classified as potential AS soils. For more details concerning the soil observations and their classification, we refer the reader to Beucher et al. (2017). A binary classification was used within the present study: potential AS soil occurrences (n = 2,309) and non-potential AS soil sites (i.e. soils which are not and will not become AS soils; n = 3,576) (Figure 1).

2.3 Environmental Data and Variable Selection

For the present study, the available 23 gridded environmental data can be divided into three groups: topography (21 covariates), soil (1 covariates) and climate (1 covariate). Danish variables

typically have a 30.4-m resolution because the digital elevation model (DEM) was originally derived from the 1.6-m resolution national airborne LiDAR (Light Detection and Ranging) data and resampled to 30.4-m resolution (Adhikari et al., 2014). Variable selection was employed in order to follow the principle of parsimony (i.e., Occam's razor) which indicates that a better model can explain the same phenomena from fewer variables (Batty and Torrens, 2001). Moreover, using fewer covariates may ease result interpretation and enable faster computer processing (Brungard et al., 2015). Recursive feature elimination (RFE) was applied using the R package caret (Kuhn, 2021) to select the optimal set of covariates for modelling based on the random forest (RF) algorithm (R Core Team, 2021; Cai et al., 2018). First, a RF model was trained on all 23 gridded environmental data, yielding the importance of each covariate. Then, the least important covariates were removed. This was recursively done until the optimal number of covariates is reached. The optimal number of covariates is defined through cross-validation. RFE analysis was carried out with the following tuning parameters: the number of tested environmental covariates (m_{trv}) at each split as default and the number of trees (n_{tree}) at 1,000. Finally, 14 environmental covariates were selected and utilized as input data for further modelling (Table 1). Among them, 10 terrain attributes were derived from the DEM within ArcGIS (ESRI, 2020) or SAGA (System of Automated Geoscientific Analyses) GIS (Conrad et al., 2015).

2.4 Random Forest and Scaled Environmental Data

Random forest (RF) is used as a reference modelling technique for comparison with the CNN approach. RF is a tree-based ensemble method which has been widely used in digital soil mapping studies to carry out classification and regression predictions (Hengl et al., 2018). In this study, RF tuning parameters such as n_{tree} (the number of trees) and m_{trv} (the number of input environmental covariates in each random subset), were determined using a Bayesian optimization procedure within R with packages ranger (Wright and Ziegler, 2017) and rBayesianOptimization (Yan, 2021). Bayesian optimization constitutes a sequential model-based optimization procedure that enables the identification of the optimum internal parameters of models in a more efficient way than grid, random or manual (i.e., trial-and-error) search (Hinz et al., 2018). Within Bayesian optimization, the unknown objective function is considered as a random function defined by a prior probability distribution. Using function evaluations as data, the prior is updated to form the objective function posterior distribution. The posterior distribution is in turn used to determines the next evaluation point. Both CNN and RF models utilized this optimization procedure, as well as the same calibration and test sets. Smoothing mean filters with different neighbourhood sizes (i.e., 1, 3, 5, 7, 9, 15, 21, and 29 pixels) were used to provide the RF models with scaled versions of the original environmental covariates, in an attempt to achieve a fairer comparison between CNN and RF (Taghizadeh-Mehrjardi et al., 2020a). RF models were thus trained both with the original

TABLE 1 | Environmental covariates selected for the modelling.

Predictor variables (clarification name)	Range	Description (source)
Soil		
Depth to pre-Quaternary deposits (dpth2prek)	0–294 m	Derived using the DEM and the pre-Quaternary surface topography Binzer and Stockmarr (1994
Climate		
Precipitation (precip61_90)	520–960 mm	Mean precipitation measured from april to October for the years 1961-1990 Scharling (2000)
Topography		
Elevation (elev)	0–161 m	LiDAR-based elevation of land surface (Adhikari et al., 2014)
Detrended DEM (demdetrd)	-58-82 m	Elevation minus the mean elevation within a 4 km radius
Bluespot (bluespot)	0–21 m	Depth of sinks
Direct sunlight insolation (dirinsola)	610-1,611	Direct sunlight insolation Böhner and Antonić (2009)
Mid-slope position (midslppos)	0-1	Relative vertical distance to mid-slope position Bendix (2004)
Multi-resolution valley bottom flatness (mrvbf)	0–10	Calculates depositional areas to identify flat valley bottoms Gallant and Dowling (2003)
Profile curvature (curv_prof)	-2.4-2.3	Curvature parallel to the maximum slope
SAGA wetness index (sagawi)	8.1-19.1	Topographic wetness index with modified catchment area Böhner et al. (2002)
Slope (slope)	0-29°	Slope gradient
Valley depth (valldepth)	0–79 m	Relative height difference to adjacent channel
Vertical distance to channel network (vdtochn)	0–10 m	Vertical distance to nearest waterbody
Distance to former Littorina sea coastline (dist2lit)	0-47.5 km	Distance to former Littorina sea coastline

14 selected environmental covariates and the scaled versions of this selection (i.e., 112 covariates in total). Smoothing mean filters constitute a common technique to derive scaled versions of environmental data in digital soil mapping studies (Grinand et al., 2008; Behrens et al., 2010a; Behrens et al., 2010b). Mean filtering consists in replacing each pixel value in an image with the mean (i.e., average) value of its neighbours, including itself (Shapiro, 1970; Boyle and Thomas, 1988). The R package *raster* (Hijmans, 2021) was used for filtering the original environmental covariates.

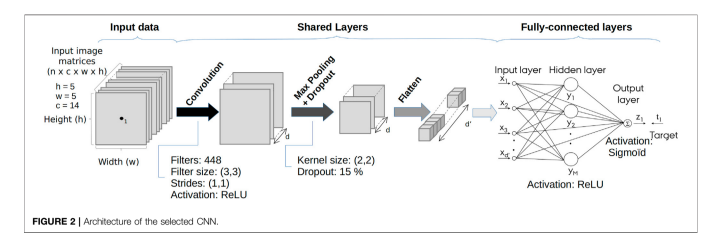
2.5 Convolutional Neural Networks 2.5.1 General

A Convolutional Neural Network (CNN) used for classification or regressions tasks can be divided in two parts: the first part consists in convolutional filtering for hierarchical feature extraction and the second part comprises fully-connected layers of neurons which calculate an output value, finalising the classification or regression task. This second part is equivalent to a classical ANN. Feature extraction by convolutional filtering is carried out through different convolution and pooling steps or layers that can be repeated several times. The objective of this first step is to extract the most relevant features and the multiple representations of the spatial scales from the x environmental covariates fed as an input xD array or patch (equivalent to an image) to the CNN (Taghizadeh-Mehrjardi et al., 2020a). The first convolutional layers extract low-level features or small local patterns (i.e., edges and corners) while the last convolutional layers extract high-level features, such as image structures (Mahdianpari et al., 2018). CNNs can also be more complex and include additional parts for more complex tasks, such as object detection.

A convolution layer consists in a filtering stage and the application of an activation function. This layer is defined by the convolutional filter size and the number of filters which represents the depth of the output feature map produced by the convolutional layer (Mahdianpari et al., 2018). A filter or moving window with typical sizes of 3×3 or 5×5 pixels progresses over the input xD array or patch (from left to right and from top to bottom).

The values of the filter represent the weights for each pixel. These filter weights are randomly initialised and adjusted by backpropagation which is a common algorithm in the training of feed-forward neural networks. Typically, an optimizer, such as stochastic gradient descent or ADAM, supports backpropagation. Specifically, this algorithm computes the gradient of the loss function using the weights of the model for a particular input and output. The filter weights are first multiplied by the overlapping pixel values of the input patch. Then, they are all summed up to a new output value which is assigned to the center location of the filter. The area of input patch which is overlaid by the filter is called receptive field. As the filter progressively moves over the input image, a new feature map is formed, constituting a combination of all environmental covariates. For comparison, the filter weights for all pixels of a feature map are the same in a CNN while each neuron has its own set of weights in a classical ANN. Applying different filter weights enables the identification of distinct specific spatial properties within the input data.

A convolution step can be carried out several times. While the first convolution is applied to the original input patch (i.e., made from the original set of environmental covariates), the following convolution steps are applied to the previously generated feature maps, creating more generalized and spatially coarser levels of information. The size of the convolution filter and the stride control the generation of convoluted feature maps in a CNN. A stride is the step width used by the filter to progress over the input image. For example, a stride of 2 implies that only every second pixel is considered and every other is skipped, which results in a reduced size of the feature maps. A convolution step is followed by a transformation or calculation step through the use of an activation function. This activation function represents a mathematical non-linear transformation function determines the relationship between input values and output value. It calculates the output value of a neuron by non-linearly transforming the weighted sum of the input value, and passes this output to the next layer in the network. The rectified linear unit (i.e.



ReLU) constitutes a commonly used nonlinear activation function for deep learning.

A pooling layer constitutes a resampling or filtering step where kernel sizes of 2×2 pixels and a stride of 2 are generally used. By reducing the number of pixels in feature maps, the spatial dimensions and computation costs are also minimised in order to avoid overfitting and further generalize the information. The maximum value within the kernel is selected when using the max-pooling operator.

Convolution and pooling steps constitute the shared layers in a CNN and are followed by the fully-connected network layers. A flattening step enables the transition from the shared layers into a fully-connected network (i.e., single-pixel feature maps transformed into feature vectors to be used as input within the fully-connected network). Within the second part of a CNN, the extracted spatially local features from the shared layers are basically combined. A main difference between the shared layers and the fully-connected layers is that the neurons in a convolution layer are only connected to the neurons in the receptive field and not to all the neurons in the previous layer. In order to avoid or reduce overfitting, a technique called dropout can be used within the shared layers and the fully-connected network (Krizhevsky et al., 2012). Dropout randomly mutes neurons during the training phase.

2.5.2 Input Data Augmentation

In addition to dropout, data augmentation enables to further avoid overfitting problems (Krizhevsky et al., 2012). Data augmentation generates more training samples from the available training data using a number of possible transformations without changing its meaning (Simard et al., 2003). For example, the original training data can be rotated by 90°, 180° and 270°, increasing the number of training samples. The main goal is to enable the model to explore more aspects of the training samples and, thus to increase the generalization ability of the CNN. This is crucial as the prediction ability of a CNN greatly depends on the amount of the existing training samples (Krizhevsky et al., 2012).

2.5.3 CNN Architecture Selection and Parameter Optimization

The architecture of the CNN model used in the present study and its specifications are represented in **Figure 2**. All environmental

covariates preprocessed through standardization (i.e., centred on 0 and scaled to a standard deviation of 1). An optimal window size for model training was determined by testing various sizes of 3, 5, 7, 9, 15, 21, and 29 pixels for the prediction of potential AS soil occurrence. Importantly, Bayesian optimization was used to identify the optimal network hyperparameters (Hinz et al., 2018). In this study, the number of filters and neurons, filter size, max-pooling kernel size, learning rate (i.e., how many weights are updated during training), dropout rate, activation function and optimizer were determined using Bayesian optimization with 60 iterations. This optimization algorithm employs an objective function (in our case, the overall accuracy) to train a model through a Gaussian process using training samples as input. Different network depths (i.e., number of layers within the CNN model) were also tested. The simple architecture presented in Figure 2 was selected as the increase in accuracy yielded by more complex models was not substantial. The selected model includes a sequence of shared layers: first, a convolutional layer with a filter size of 3 × 3 and a ReLu activation function, second, a maxpooling layer with a size of 2 × 2, third, a dropout layer which randomly disconnects 15% of the connections in order to prevent overfitting, and fourth, a flatten layer. The flatten layer enables the connection between the shared layers and the succeeding fullyconnected layers by converting matrices into vectors. The fullyconnected layers performs the classification part of the CNN, receiving information from the previous layers to generate predictions (Taghizadeh-Mehrjardi et al., 2020b).

CNNs were implemented in *Python* (v3.8) (Python Software Foundation, 2021) using Tensorflow/Keras (v2.6) (Abadi et al., 2016; Keras, 2021) and Tensorboard (v2.6). The experiments were carried out on an Intel CPU i7-8,700 machine with 3.2 GHz 6-core CPU and 32 GB RAM memory with a Nvidia GeForce RTX 2070 GPU under CUDA version 11.4.

2.5.4 Training, Validation and Test Datasets

In order to evaluate the performance of our CNN models to predict potential AS soils, 75 and 25% were randomly selected as training and test dataset, respectively. On the remaining 75%, data augmentation was carried out to increase the amount of the training samples. Additionally, to find the optimal hyperparameters and to avoid overfitting, we used a random

subset (80 and 20%) of the augmented data for training and validation of the CNN. The test dataset (25%), which is independent of the training and validation samples, is used to estimate the generalisation error of the proposed CNN and used for the final model comparison (i.e., between RF and CNN). The relationship between observed and predicted points for the training and validation data sets was evaluated with the overall prediction accuracy (OA) which represents the number of correctly classified points divided by the total number of points.

2.6 Interpretability With SHAP

SHAP (SHapley Additive exPlanations) constitutes a method based on Shapley values that enables explaining individual predictions (Lundberg and Lee, 2017). Within a game theory context, each feature (or covariate) value can be defined as a player in a coalitional game (i.e., a modelling task) where the prediction represents the payout. Shapley values are used to estimate how the payout is distributed among the covariates (Molnar, 2021). In order to compute Shapley values, two models are fitted for a covariate i: one $f_{S \cup \{i\}}$ including the covariate *i* and another f_S withholding it. The difference between the predictions of the two models on the current input x, $f_{S \cup \{i\}}(x_{S \cup \{i\}}) - f_S(x_S)$, constitutes the specific contribution of covariate i (Lundberg and Lee, 2017). Since the specific contribution of a covariate depends on the other utilised covariates, the preceding calculation is computed for all the possible covariate subsets $S \subseteq F$, where *F* is the set of all covariates. The overall covariate contribution ϕ_i represents the weighted average of all specific contributions as follows:

$$\phi_{i} = \sum_{S \subset F \setminus \{i\}} \frac{|S|! (|F| - |S| - 1)!}{|F|!} \left[f_{S \cup \{i\}} \left(x_{S \cup \{i\}} \right) - f_{S} \left(x_{S} \right) \right]$$
(1)

The SHAP method defined by Lundberg and Lee (2017) adapts the procedure described above by applying sampling approximations to Eq. 1 and approximating the effect of removing a covariate by using other samples from the training dataset. In short, a SHAP value indicates the relative contribution, positive or negative, of a covariate in a given sample to the model's output (Zhong et al., 2021). Moreover, Lundberg and Lee (2017) developed a Deep SHAP explainer (DeepExplainer) to compute SHAP values specifically for DNNs. The method applies linear approximations which are aggregated by using a composition rule enabling the approximation of SHAP values for the whole model. Basically, in order to explain a complex original model f, we use an explanation model g simpler than the original. The explanation model is also based on simplified inputs x' which map to the original inputs x through a mapping function $x = h_x(x')$ (Lundberg and Lee, 2017). This local method try to ensure that $g(z') \approx f(h_x(z'))$ when $z' \approx x'$. In order to assess the effect of varying the input x, z' is defined from reference values sampled from observations from the training dataset (Padarian et al., 2020). The contribution ϕ_i of each covariate (Eq. 1) is then included in a linear function of binary variables (i.e., an explainable variable being either present, 1, or absent, 0):

$$g(z') = \phi_0 + \sum_{i=1}^{M} \phi_i z_i'$$
 (2)

where $z' \in \{0,1\}^M$, M is the number of simplified input features, and $\phi_i \in \mathbb{R}$. The explanation model g (i.e., the additive feature attribution explanation model as defined by Lundberg and Lee (2017)) thus represents an approximation of the prediction f(x) by summing all the covariate contributions (or attributions) $\phi_i z_i'$ and including ϕ_0 , the contribution when all the covariates are missing (i.e., mean of the predictions for the complete dataset Padarian et al. (2020)).

The Deep SHAP Explainer is included in the SHAP *Python* library (Lundberg and Lee, 2017) and was used for CNN interpretability purposes. Deep SHAP supports both TensorFlow and Keras models by extending the DeepLIFT algorithm from (Shrikumar et al., 2017) using a distribution of background samples to calculate SHAP values for the selected network. The data utilized to perform the approximations correspond to the CNN training dataset. In order to confirm that the model is capturing sensible relationships between the class target (i.e., potential AS soil occurrence) and the environmental covariates, we applied SHAP to visualize the distribution of the main contributing environmental covariates over the whole study area, both as a map and as a histogram. We also focused on a selection of main contributing environmental covariates and inspected their distribution as a single layer over the study area.

3 RESULTS AND DISCUSSION

3.1 RF Models

The predictive classification abilities of RF were evaluated for the classification of potential AS soil occurrence in the wetlands of Jutland. RF models were tested using firstly, the 14 environmental covariates selected with RFE, and secondly, the 112 scaled covariates for which smoothing mean filters were utilized. Based on the Bayesian optimization procedure, the optimal RF tuning parameters were defined for each RF model. The RF model built with the original 14 covariates (RF_{original}) used a random subset size of 10, a minimal nodesize of 5 and a tree size of 1,600 while the RF model developed with the scaled covariates (RF_{scaled}) used a random subset size of 5, a minimal nodesize of 3 and a tree size of 680. Both models used the Gini splitting rule. RF_{original} and RF_{scaled} models yielded comparable OA values for validation, 61 and 63%, respectively. These results indicate that the RF_{scaled} model did not seem to use the contextual spatial information to classify potential AS soils more accurately than the RF_{original} model. For the RF_{original} model, the most important variables were elevation, depth to pre-Quaternary deposits, detrended elevation model and mid-slope position. For the RF_{scaled} model, they were elevation and detrended elevation model at different scales (fine first, then coarse). AS soils occur in low-lying areas which supports the importance of elevation layers. The depth to pre-Quaternary deposits also represents a useful covariate in particular for central and southern Jutland where potential AS soils can form in non-marine environments (i.e. glaciofluvial outwash plains and moraine landforms) overlaying pyrite-containing Tertiary sediments (Beucher et al., 2017).

3.2 CNN Models

Using Bayesian optimization, the optimal hyperparameter values were selected for our CNN models. Considering the CNN model yielding the highest OA (68%), the hyperparameters were: ReLU

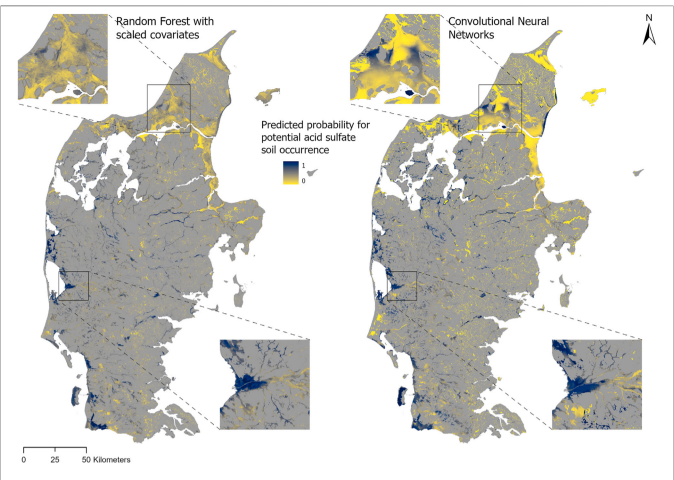


FIGURE 3 | Maps of the predicted probability for potential acid sulfate soil occurrence generated using Random Forest (RF) with scaled covariates (left) and Convolutional Neural Networks (CNN; right).

as activation function, ADAM as optimizer, binary cross entropy as loss function, a learning rate of 0.001, a convolution filter size of 3×3 and 448 filters, a max-pooling kernel size of 2×2 , and a dropout rate of 15%. This model also used augmented data. However, the CNN model with similar architecture and parameters based on data without augmentation already yielded a comparable OA value (63%). The optimal patch size for training our CNN models was determined by assessing different sizes (i.e. 3, 5, 7, 9, 15, 21, and 29) for the classification of potential AS soils. An increase in patch size up to 5 increases OA which then decreases from the patch size of 9. A patch size of 5 was thus found to be optimal. Considering the spatial resolution of covariates of 30.4 m, a patch of 5×5 includes the contextual information of approximately 76 m (i.e., $5/2 \times 30.4$ = 76 m). Therefore, the contextual information of environmental covariates within a 76 m radius was demonstrated as valuable for classifying potential AS soils compared to the local information usually used at a specific point. This result is in line with findings of past studies (Behrens et al., 2010a; Behrens et al., 2010b; Behrens et al., 2018a; Behrens et al., 2018b; Padarian et al., 2019b; Wadoux et al., 2019), which reported that the

prediction accuracy increased by including contextual information within the modelling.

3.3 CNN/RF Model Comparison

The best performing CNN model yielded a slightly higher OA than both RF models, improving the prediction by 5 and 7% compared to RF_{scaled} and RF_{original} models, respectively. This relatively small difference in performance can might be explained by the use of RFE as a common variable selection method. For this study, RFE was based on the RF algorithm and thus rendered a selection of covariates optimized for this method. As indicated by Brungard et al. (2015), the use of RF within RFE most likely explains the consistent higher accuracies rendered by RF models in comparison with other models. Even though ANNs in general, and CNNs in particular, have the ability to extract relevant information from large datasets (Gershenfeld, 1999; Chagas et al., 2013) and generalize from relatively imprecise data (Porwal et al., 2003), variable selection might constitute an important processing step when implementing them. Moreover, a variable selection optimized for RF, a method resistant both to noise (Strobl et al., 2007) and multicollinearity (Hengl et al., 2018), might not be the most sound

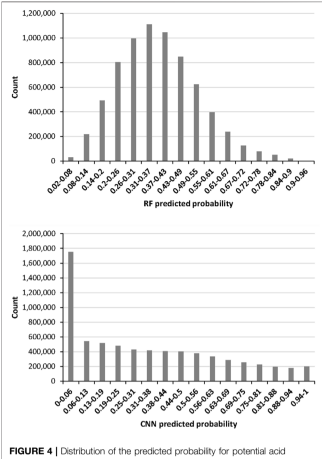


FIGURE 4 | Distribution of the predicted probability for potential acid sulfate soil occurrence generated using Random Forest (RF) with scaled covariates (top) and Convolutional Neural Networks (CNN; bottom).

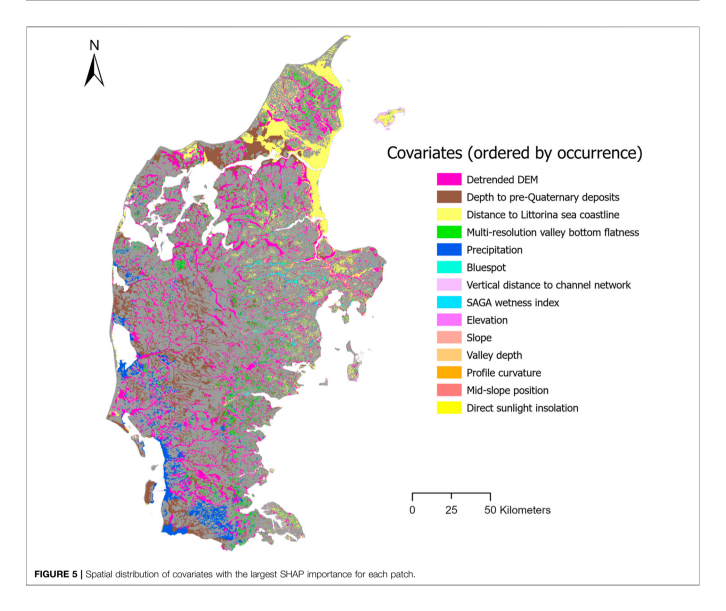
option for CNNs. However, the use of a specific variable selection method for CNNs might also yield a different selection of covariates, which would hinder comparing the two methods. Another possible most probable reason for the comparable performances of RF and CNN lies in is the relatively coarse spatial resolution of the environmental covariates used as input data, which might may have represented a strong limitation for the CNN models. As stated previously using a patch of 5×5 with 30.4-m resolution covariates resulted in the use of contextual information within a 76-m radius around soil observations for the training of CNN models. Employing covariates with a finer resolution, for example 10 m, would enable to use larger size patches and thus feed the CNN models with more details to investigate in each input image. Moreover, this limitation due to the resolution could also explain the fact that CNN models with more complex architectures did not yield notable increases in accuracy compared to the simple CNN selected in this study (Figure 2). While the first convolution layer is applied to the input environmental covariates and extracts low-level features, the following convolution layers are applied to the created feature maps and thus, produce more generalized and spatially coarser level of information. In our case, the additional convolution layers might only have extracted coarser information from coarse input data, which did not enable the fully-connected layers to generate more accurate predictions.

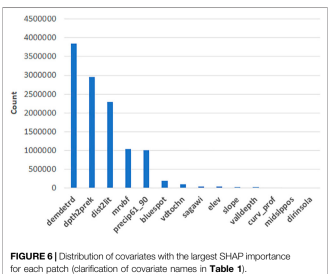
Comparing the prediction maps in **Figure 3**, both models predicted high probability areas located along the western coast of Jutland, for example the delta region of the Skjern River (i.e. the bottom located close-up in **Figure 3**) which constitutes a well-known potential AS soil area (Madsen and Jensen, 1988). However, the RF model tended to over-predict mid-range values (**Figure 4**), which can be observed in the northern part of Jutland, in particular in the top-located close-up in **Figure 3**. This can be interpreted as RF poorly generalizing from input data. In comparison, the CNN model over-predicted low probability values (**Figure 4**), which can be explained by the fact that the set of non-potential AS soils (n = 3,576) is larger than the one of potential AS soils (n = 2,309).

3.4 Interpretability Using SHAP

In order to compute SHAP values, we used 200 training points from each class as background data (i.e., data to perform the approximations). These values were found empirically as it was determined to give a good trade-off between consistency of SHAP values and computation time since the computation of SHAP values from the explainer increases linearly with the number of samples (i.e., training points).

In order to gain more insight as to the spatial interpretation of the model over Jutland, the most important covariates are displayed in Figure 5. The map shows the spatial distribution of the covariates with the largest SHAP value for each 5×5 patch, meaning the largest positive or negative deviation from the baseline SHAP values calculated from the background samples. Moreover, Figure 6 shows the distribution of the covariates with the largest SHAP contribution for each patch. Figure 6 indicates that the detrended DEM, depth to pre-Quaternary deposits and distance to Littorina sea coastline constitute the most important covariates in terms of occurrence, followed by the multi-resolution valley bottom flatness and precipitation. Figure 5 shows that the detrended DEM is the most important covariate in wetland areas all over Jutland, which can be expected as soils typically occur in low-lying areas. The depth to pre-Quaternary deposits represents the second most important covariate, occurring mostly in the western half of Jutland. This can be explained by the fact that potential AS soils can form in non-marine environments (i.e., in glaciofluvial outwash plains and moraine landforms as seen in Figure 7) overlaying pyrite-rich Tertiary sediments (Beucher et al., 2017). The distance to the former Littorina sea coastline constitutes the third most important covariate and almost perfectly concurs with post-glacial marine deposits located in northern Jutland and the easternmost part of central Jutland (Figure 7). This result confirms the suggestion of Beucher et al. (2017) that such a predictor enables targeting potential AS soils formed within post-glacial marine deposits and most probably occurring close to this limit. The multi-resolution valley bottom flatness and precipitation also represent important covariates. The former occurs as a most important covariate in wetlands almost all over Jutland, particularly in the eastern part of Jutland, and most probably emphasizes flat, low-relief areas in which AS soils typically appear. Precipitation mostly occurs as a most important covariate in the southern and westernmost areas of Jutland. These areas are particularly flat and low-lying with a





very high groundwater table. Perhaps, precipitation constitutes an indirect indicator in these specific areas which are all located in or upstream from marsh areas (**Figure 7**).

A summary plot for the selected CNN can be seen in **Figure 8**. Each point on the plot shows the yielded SHAP values for each covariate for all 5×5 patches. It should be noted that if multiple patches for a given covariate have the same value the points are stretched along the *y*-axis. Notably, the plot confirms the previously developed interpretation results concerning the most important covariates in terms of SHAP contribution and enables further interpretation. Focusing on the detrended DEM, **Figure 8** shows that low elevation values (in blue) mostly have a strong positive impact on the model output, which can be explained as soils typically occur in low-lying areas. Shallow depths to pre-Quaternary deposits and small distances to the former *Littorina* sea coastline (in blue) also present a substantial positive impact on the model output. This confirms that potential AS soils can be targeted in the corresponding areas, either in former marine environments

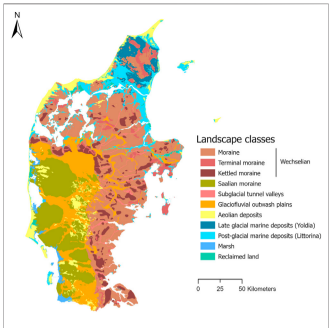


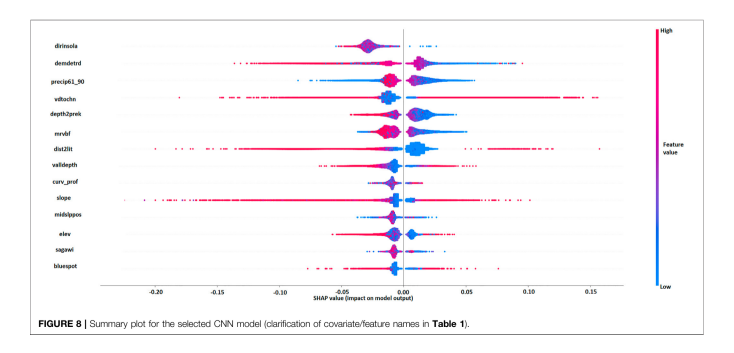
FIGURE 7 | Map of the main landscape types (i.e., spatially homogeneous geomorphological units) in Jutland (derived using the existing digital vector landscape map at 1:100000 scale originally from Madsen et al. (1992)).

close to the former *Littorina* sea coastline or in non-marine environments above shallow pyrite-rich Tertiary deposits.

Next, the distribution of SHAP values for the top three covariates, namely the detrended DEM, depth to pre-Quaternary deposits and distance to *Littorina* sea coastline, is shown in **Figure 9**. While **Figures 5**, **6** display the most important covariates spatially and quantitatively, **Figure 9** shows the distribution of SHAP values, both

positive and negative, for each of the top three covariates. The three maps confirm previous findings. Firstly, the distance to the former Littorina sea coastline (Figure 9C) presents the most positive SHAP values (i.e. strong positive impact on model output) in northern Jutland and the eastern part of central Jutland, which correspond to post-glacial marine deposit areas (Figure 7). The most negative SHAP values (i.e., negative impact on model output) are mostly located in western and southwestern inland areas of Jutland. Secondly, the depth to pre-Quaternary deposits (Figure 9B) show the most positive SHAP values for areas mostly located in the western half of Jutland, where potential AS soils can form in nonmarine environments, such as glacial flood plains and moraine landforms overlaying pyrite-rich Tertiary sediments (Figure 7). The most negative SHAP values are located in coastal areas in northern and southwestern Jutland. Thirdly, the detrended DEM (Figure 9A) presents mostly a positive impact in southern, western and parts of northern Jutland. Negative impact occurs in eastern Jutland and the remaining parts of northern Jutland.

Finally, the Deep SHAP explainer enables assessing covariate importance (as can be done for classical machine learning models such as RF), most importantly in a spatial context, producing different maps of covariate contributions for the CNN predictive model. Concurring with Padarian et al. (2020), we emphasize that the use of SHAP values for model interpretation represents a substantial development within the DSM framework. Samek et al. (2019) states that discovering new patterns within the data through the use of XAI methods such as SHAP (i.e. explaining and interpreting what covariates/features were used for prediction) is often more important than the prediction itself. The results of the present study are consistent with this statement. Furthermore, as the amount of high resolution data will substantially increase in the near future, the use of deep learning models will also most probably expand. Interpretation methods such as SHAP will thus become necessary to 1) improve



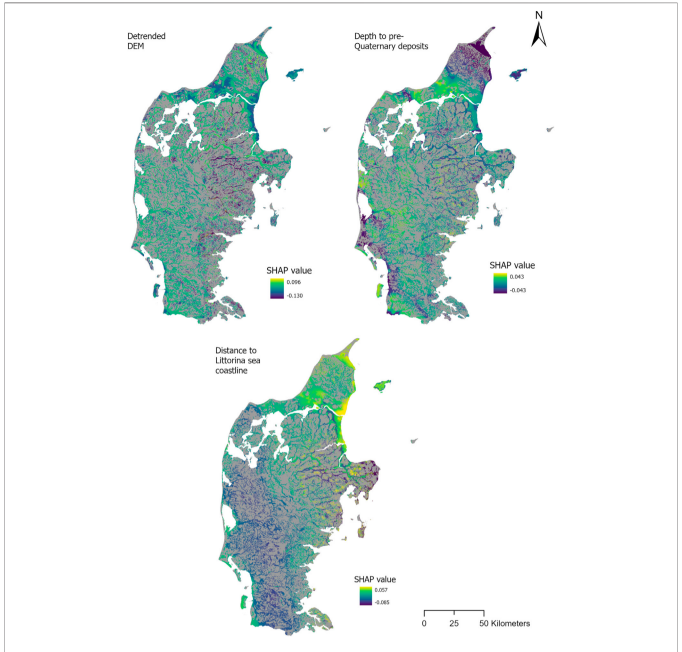


FIGURE 9 | Spatial distribution of SHAP values for the three most important covariates: detrended DEM, depth to pre-Quaternary deposits and distance to Littorina sea coastline.

our understanding of complex model results, 2) possibly develop our knowledge of soil systems and 3) enable us to better explain our findings both to scientific and non-scientific audience.

3.5 Other Challenges Associated With CNNs and Future Possible Development

When employing CNNs, another substantial challenge lies in the selection of a model architecture. Two approaches can be utilized.

With the first approach, pre-existing architectures of well-known CNNs such as DenseNet (Huang et al., 2017), Inception (Szegedy et al., 2015), VGG (Simonyan and Zisserman, 2015), Xception (Chollet, 2017) and ResNet (He et al., 2016) are employed. Primarily defined for computer vision, these CNNs have been used for example in land cover classification studies (e.g., Mahdianpari et al. (2018)). With the second approach, a CNN architecture is fully designed and the model trained from scratch. In this case, the network hyperparameters are determined through a

highly iterative process which can be based on manual (trial-anderror), purely random or grid search, sequential model-based optimization such as Bayesian optimization or a combination of approaches (Bergstra and Bengio, 2012; Hinz et al., 2018). Within the existing DSM studies investigating the use of CNNS, this second approach is employed. However, the selection of the architecture is generally not clarified, with the exception of Taghizadeh-Mehrjardi et al. (2020b) and Wadoux et al. (2019) which both relied on Bayesian optimization. while the present study utilized Bayesian optimization for the selection of several hyperparameters, it still resorted to a trial-and-error process to design the CNN architecture (i.e., number and type of layers), which constitutes a substantial weakness. A future development of the present work would be to test and clarify different existing methods for hyperparameter optimization, such as tree of Parzen estimators, genetic algorithm and sequential model-based algorithm configuration (Hinz et al., 2018) for the specific purpose of soil mapping.

Another crucial challenge consists in the assessment of uncertainty for CNN model predictions. Padarian et al. (2019b) used bootstrapping to derive confidence intervals while Wadoux (2019) employed a method based on bootstrap plus variance estimates (Khosravi et al., 2011). The latter study also referred to two other possible methods for uncertainty quantification of neural networks, namely the Bayesian uncertainty analysis (Mackay, 1992; Gal and Ghahramani, 2016) and the Delta (De Vleaux et al., 1998) methods, which have not been tested yet. Considering AS soils, the use of contextual information may considerably benefit the mapping of their occurrence. A future development of the present study would be the prediction of key soil properties in strategic hot spot areas for AS soil occurrence using CNNs. Accounting for the spatial variability of soil properties, such as incubation pH and titratable acidity, would enable an improved management of environmental risks related to AS soils.

4 CONCLUSION

The present study aimed at investigating first, the use of convolutional neural networks for the classification of potential acid sulfate soils located in the wetlands of Jutland, Denmark, and second and most importantly, the application of a model-agnostic interpretation method called SHAP to the selected convolutional neural networks model. The deep learning model yielded slightly higher prediction accuracy than the random forest models which were using original or scaled covariates. These results could most

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Behrens, T., Schmidt, K., Zhu, A. X., and Scholten, T. (2010a). The ConMap Approach for Terrain-Based Digital Soil Mapping. *Eur. J. Soil Sci.* 61, 133–143. doi:10.1111/j.1365-2389.2009.01205.x likely be explained by the use of a common variable selection method, namely recursive feature elimination. This technique being based on the random forest algorithm most probably rendered a selection of covariates optimized for this method. However, the main focus of the study was to assess the SHAP interpretation method. In particular, the spatial interpretation of the most important covariates enabled to clarify the predictions of the selected convolutional neural networks model. While classical machine learning methods such as random forest or cubist provide variable importance or usage, SHAP allows the visualization of covariate contribution to the convolutional neural network model at different levels in geographical space, such as the general spatial distribution of all most important covariates in one map and the specific spatial distribution of SHAP values for one covariate. The possibility to inspect the contribution of covariates in space constitutes a crucial development for digital soil mapping in general and for contextual mapping in particular. The combination of convolutional neural networks and SHAP represent a powerful and promising progress for contextual mapping. Furthermore, the rapid developments within the field of explainable artificial intelligence will most likely deliver novel tools to visualize, explain and interpret deep learning methods, which may lead to discovering new insights, improving both knowledge and understanding of soil systems.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

AB conceived, executed the research and wrote the paper. CR participated in executing the research and writing the paper. TM reviewed the paper. MG gave suggestions about the approach and reviewed the paper.

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