

Hyperparameter tuning and performance assessment of statistical and machine-learning algorithms using spatial data

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ABSTRACT

While the application of machine-learning algorithms has been highly simplified in the last years due to their well-documented integration in commonly used statistical programming languages (such as R or Python), there are several practical challenges in the field of ecological modeling related to unbiased performance estimation. One is the influence of spatial autocorrelation in both hyperparameter tuning and performance estimation. Grouped cross-validation strategies have been proposed in recent years in environmental as well as medical contexts to reduce bias in predictive performance. In this study we show the effects of spatial autocorrelation on hyperparameter tuning and performance estimation by comparing several widely used machine-learning algorithms such as boosted regression trees (BRT), k-nearest neighbor (KNN), random forest (RF) and support vector machine (SVM) with traditional parametric algorithms such as logistic regression (GLM) and semi-parametric ones like generalized additive models (GAM) in terms of predictive performance. Spatial and non-spatial cross-validation methods were used to evaluate model performances aiming to obtain bias-reduced performance estimates. A detailed analysis on the sensitivity of hyperparameter tuning when using different resampling methods (spatial/non-spatial) was performed. As a case study the spatial distribution of forest disease (*Diplodia sapinea*) in the Basque Country (Spain) was investigated using common environmental variables such as temperature, precipitation, soil and lithology as predictors. Random Forest (mean Brier score estimate of 0.166) outperformed all other methods with regard to predictive accuracy. Though the sensitivity to hyperparameter tuning differed between the ML algorithms, there were in most cases no substantial differences between spatial and non-spatial partitioning for hyperparameter tuning. However, spatial hyperparameter tuning maintains consistency with spatial estimation of classifier performance and should be favored over non-spatial hyperparameter optimization. High performance differences (up to 47%) between the bias-reduced (spatial cross-validation) and overoptimistic (non-spatial cross-validation) cross-validation settings showed the high need to account for the influence of spatial autocorrelation. Overoptimistic performance estimates may lead to false actions in ecological decision making based on biased model predictions.

1. Introduction

Spatial predictions are of great importance in a wide variety of fields including hydrology (Naghibi et al., 2016), epidemiology (Adler et al., 2017), geomorphology (Brenning et al., 2015), remote sensing (Stelmaszuk-Górska et al., 2017), climatology (Voyant et al., 2017), soil sciences (Hengl et al., 2017) and ecology (Baasch et al., 2010; Muenchow et al., 2013; Murase et al., 2009; Vorpahl et al., 2012). Ecological applications range from species distribution models (Halvorsen et al., 2016; Quillfeldt et al., 2017; Wieland et al., 2017) over plant disease and soil type modeling (Heim et al., 2018; Brungard

et al., 2015) to resource selection (Baasch et al., 2010).

A typical example for a spatial prediction approach in ecology is the detection of fungi infection on Monterey pines (Iturritxa et al., 2014). Fungal species such as *Diplodia sapinea* inflict severe damages to *Pinus radiata* trees which are then subjected to environmental stress (Wingfield et al., 2008). Infected forest stands cause economic as well as ecological damages worldwide (Ganley et al., 2009). In Spain, where timber production is regionally an important economic factor, about 25% of the timber production stems from Monterey pine (*Pinus radiata*) plantations in northern Spain, and here mostly from the Basque Country (Iturritxa et al., 2014). Consequently, the early detection and subsequent containment of

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fungal diseases is of great importance. Statistical and machine-learning models can help in this process by mapping the current infection state and exploring relations between the pathogens and environmental variables. These findings can then be used for spatially predicting the risk of future outbreaks.

1.1. The special role of spatial autocorrelation in predictive modeling

All of the previously mentioned scientific fields have one thing in common: The observations inherit spatial information. One of the main challenges that comes with this information is the accounting for the influence of spatial autocorrelation in the data (Legendre, 1993). Cross-validation and bootstrapping are two widely used performance estimation techniques (Efron, 1983; Gordon et al., 1984; Kohavi et al., 1995). However, in the presence of spatial autocorrelation, estimates obtained using regular (non-spatial) random resampling maybe biased and overoptimistic. This has led to the adoption of spatial resampling in cross-validation and bootstrapping for bias reduction. The mentioned bias inherits from the fact that training and test observations are located close to each other (in a geographical space) if a random sampling is used in cross-validation (CV) (Legendre, 1993). Random sampling in CV leads to the selection of test observations that are spatially close to training observations. According to the first law of geography, close observations are frequently more similar to each other than observations further apart. This violates the fundamental assumption of independence in cross-validation. Hence, algorithms fitted on the training data often achieve very good performance results, simply because the characteristics of the evaluation set are very similar to the training data.

One approach to solve this, which has been applied in various studies in the last decade, builds upon the idea to spatially disjoin training and test set in CV. The naming of this concept varies with the scientific field in which it is applied: Burman et al. (1994); Roberts et al. (2017); Shao (1993) label it “block cross-validation”, Brenning (2005) as “spatial cross-validation”, Pohjankukka et al. (2017) “spatial k-fold cross-validation” and Meyer et al. (2018) “leave-location-out cross-validation”. In this work we use the term “spatial cross-validation” because it is the most generic wording to label this concept and hope that this naming convention will prevail.

Although the importance of bias-reduced spatial resampling methods for performance estimation has been emphasized repeatedly in recent years (Geiß et al., 2017; Meyer et al., 2018; Wenger and Olden, 2012), unfortunately many studies have been published in recent years that did not account for this problem (Bui et al., 2015; Smoliński and Radtke, 2016; Wollan et al., 2008; Youssef et al., 2015).

1.2. Parametric vs. non-parametric algorithms

Supervised learning techniques can be broadly divided into parametric and non-parametric models. Parametric models can be written as mathematical equations involving model coefficients. This enables ecologists to interpret relationships between the response and its predictors. Choosing the best performing algorithm for a specific dataset is an essential step in ecological modeling to maximize predictive accuracy. In this context, model interpretability should certainly be an important criterion in the selection process when the aim is to make inference on the modeled relationship (Johnson and Omland, 2004). While the most commonly used statistical models such as generalized linear mixed models (GLMMs) are parametric, especially machine-learning techniques offer a non-parametric approach to spatial modeling in ecology (De'Ath, 2007). Even though recently a lot of effort has been put into improving the interpretability of machine-learning algorithms, their ability to make inference is still limited compared to parametric ones (Adler et al., 2018; Henelius et al., 2014). The former gained in popularity due to their ability to handle high-dimensional and highly correlated data and their less important model assumptions.

1.3. The importance of hyperparameter optimization

To reach robust performance results with non-parametric models, their respective hyperparameters must be optimized. Default hyperparameter settings cannot guarantee an optimal performance of machine-learning techniques and additional attention should be directed to this critical step. When performance estimation techniques such as cross-validation are used in this step, the adequacy of non-spatial partitioning techniques for spatial datasets can be questioned. Although spatial resampling methods have been used in studies that deal with spatial data for quite some time now (Geiß et al., 2017; Iturritxa et al., 2014; Meyer et al., 2018), there is no analysis of the effect and meaningfulness of using spatial resampling techniques for hyperparameter tuning. This study aims to check if optimizing hyperparameters using a non-spatial sampling may potentially lead to non-optimal performance estimates.

1.4. Main objectives

Overall, the intention of this work is to emphasize the need for spatial CV and corresponding hyperparameter tuning in spatial modeling to receive biased-reduced performance estimates. The following objectives (and hypotheses) are addressed:

- Comparison of the predictive performance of spatial and non-spatial partitioning methods. We expect that non-spatial partitioning methods will yield over-optimistic results in the presence of spatial autocorrelation.
- Exploring the effects of (spatial) hyperparameter tuning for commonly used algorithms in the field of ecological modeling. We propose that optimal hyperparameter tuning has a substantial effect on model performance.
- Comparison of the predictive performance of parametric (GLM, GAM) and non-parametric algorithms (BRT, RF, SVM, KNN). We expect that the predictive performance of non-parametric algorithms is substantially higher.

2. Data and study area

2.1. Summary of the prediction task

This study uses parts of the dataset from Iturritxa et al. (2014). While Iturritxa et al. (2014) focused on the influence of environmental predictors on pathogen probability, the aim of this study is to compare different algorithms with the focus of exploring the influence of spatial autocorrelation on predictive accuracy and hyperparameter tuning. In the present study we also introduced additional predictors (probability of hail damage at trees, soil type, lithology type, pH) to possibly enhance the predictive power of the trained models.

This particular dataset was chosen because it incorporates attributes of common geospatial modeling tasks: An uneven distribution of the binary response variable (25/75), presence of spatial autocorrelation and predictor variables derived from various sources (previous modeling results, remote sensing data, surveyed information). It is representative for many other ecological datasets in terms of sample size ($n = 922$), number of variables ($n = 11$) and predictor types (numeric as well as nominal).

2.2. Variables

The following (environmental) variables were used as predictors: mean temperature (March–September), mean total precipitation (July–September), potential incoming solar radiation (PISR), elevation, slope (degrees), potential hail damage at trees, tree age, pH value of soil, soil type, lithology type, and the year when the tree was surveyed. Temperature, precipitation and PISR are long-term averages (1951–1999) of meteorological stations across the Iberian Peninsula (Ninyerola et al., 2005). Tree infection caused by the fungal pathogen *Diplodia sapinea* represents the

response variable. The ratio of infected and non-infected trees in the sample is roughly 1:3 (223, 703). Precipitation, temperature and PISR were already attached to the dataset. All other variables were extracted to the point data from their raw sources.

[Iturritxa et al. \(2014\)](#) showed in their study that the presence or absence of hail damage observed on trees is an important predictor when modeling pathogen infections of trees in the Basque Country. Because almost every infected tree by *Diplodia sapinea* showed hail damage, it was assumed that the pathogen uses the open wounds caused by the hail damage as an entry point. To make the tree-based hail damage variable spatially available for the whole Basque Country, we spatially predicted hail damage potential (in probabilities from 0 to 1) as a function of climatic variables using a generalized additive model (GAM) ([Schratz, 2016](#)). In the following we shortly describe the source and modifications of the new variables. For the remaining ones, please see [Iturritxa et al. \(2014\)](#).

Soil type was predicted by [Hengl et al. \(2017\)](#) using approximately 150,000 soil profiles at a spatial resolution of 250 m. The age of trees was imputed and trimmed to a value of 40 to reduce the influence of outliers. The pH value was mapped by the [European Commission \(2010\)](#) using a regression-kriging approach based on 12,333 soil pH measurements from 11 different sources. GeoEuskadi provided the lithology types ([GeoEuskadi, 1999](#)). The rock class were aggregated by the respective top level class for magmatic types and sub-classes for sedimentary rocks ([Grotzinger and Jordan, 2016](#)) (Table A.3).

We removed three observations due to missing information in some variables leaving a total of 926 observations (Table A.2). All nominal variables (soil and lithology-type) were dummy-encoded. To avoid introducing collinearity, the following reference levels of the dummy-encoded variables were removed from the data: soil type: “soils with clay enriched sub-soils”. Lithology type: “surface deposits”.

2.3. Study area

The Basque Country in northern Spain represents the study area (Fig. 1). It has a spatial extent of 7355 km². Precipitation decreases toward the south while the duration of summer drought increases. Between 1961 and 1990, mean annual precipitation ranged from 600 to 2000 mm with annual mean temperatures between 8 and 16 °C ([Ganuza and Almendros, 2003](#)). The wooded area covers approximately 54% of the territory (3969.62 km²),

which is one of the highest ratios in the EU. Radiata pine is the most abundant species occupying 33.27% of the total area ([Múgica et al., 2016](#)).

3. Methods

In this study we provide an exemplary analysis combining both tuning of hyperparameters (see Section 1.3) using nested CV (see Section 3.2.1) and the use of spatial CV to assess bias-reduced model performance (see Section 1.1). We compared predictive performance using four settings: non-spatial CV for performance estimation combined with non-spatial hyperparameter tuning (*non-spatial/non-spatial*), spatial CV estimation with spatial hyperparameter tuning (*spatial/spatial*), spatial CV estimation with non-spatial hyperparameter tuning (*spatial/non-spatial*), and spatial CV estimation without hyperparameter tuning (*spatial/no tuning*). We used the open-source statistical programming language R ([R Core Team, 2019](#)). The algorithm implementations of the following packages have been used: *gbm* ([Ridgeway, 2017](#)) (boosted regression trees (BRT), [Elith et al. \(2008\)](#)), *mgcv* ([Wood, 2017](#)) (GAM), *kernlab* ([Karatzoglou et al., 2004](#)) (support vector machine (SVM), [Vapnik \(1998\)](#)), *knnr* ([Schliep and Hechenbichler, 2016](#)) (Weighted k-nearest neighbor (KNN), [Dudani \(1976\)](#)), and *ranger* ([Wright and Ziegler, 2017](#)) (random forest (RF), [Breiman \(2001\)](#)). The spatial partitioning functions of the *sperrrest* package have been integrated into the *mlr* package as part of this work. *mlr* provides a standardized interface for a wide variety of statistical and machine-learning models in R simplifying essential modeling tasks such as hyperparameter tuning, model performance evaluation and parallelization ([Bischl et al., 2016](#)). The complete analysis including data is available as a research compendium at Zenodo (10.5281/zenodo.2582969) ([Schratz et al., 2019](#)).

3.1. Tuning

Determining the optimal (hyperparameter) settings for each model is crucial for the bias-reduced assessment of a model's predictive power. Hyperparameters of machine-learning algorithms need to be tuned to achieve optimal performances ([Bergstra and Bengio, 2012](#); [Duarte and Wainer, 2017](#); [Hutter et al., 2011](#)). Often enough, parametric models do not require tuning to achieve optimal performances. However, some (semi-)parametric algorithms (e.g. GAM, penalized regression methods) can be optimized to possibly increase their performance.

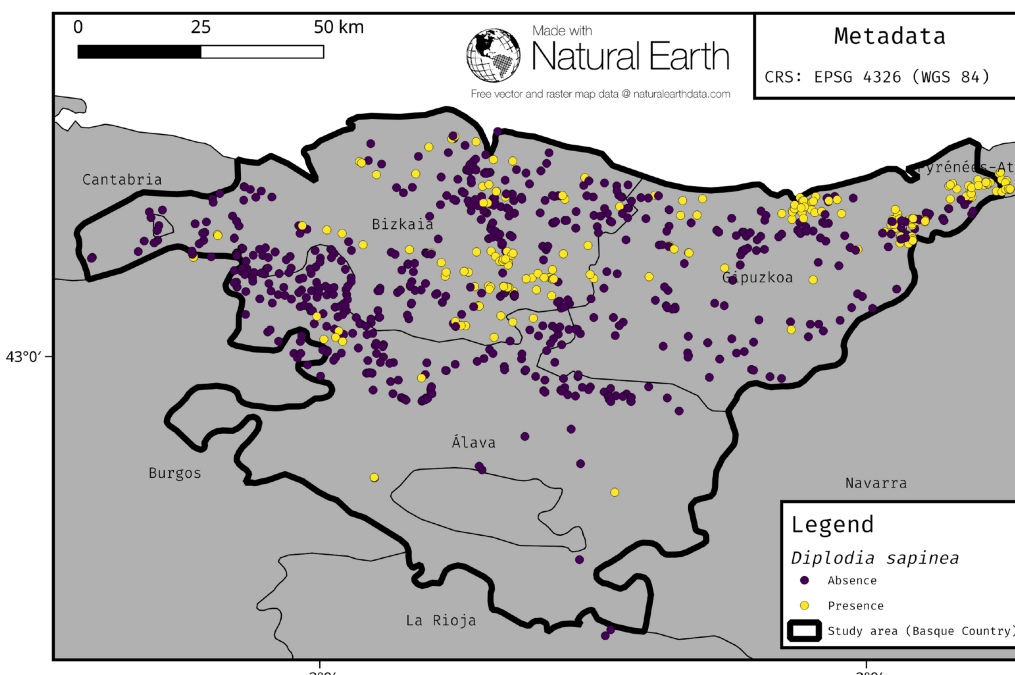


Fig. 1. Spatial distribution of tree observations within the Basque Country, northern Spain, showing infection state by *Diplodia sapinea*.

Table 1

Hyperparameter ranges and types for each model. Notations of hyperparameters from the respective R packages were used. Note that parameter sp of the GAM is a vector with eight entries (one entry for each numeric predictor). p is the number of predictors.

Algorithm (package)	Hyperparameter	Type	Start	End	Default
BRT (gbm)	n.tree	Integer	100	10000	100
	shrinkage	Numeric	0.005	0.2	0.001
	interaction.depth	Integer	1	20	1
KNN (kknn)	k	Integer	1	100	7
	distance	Integer	1	100	2
	kernel	Nominal	*		
GAM (mgcv)	sp	Numeric	0	10^6	–
RF (ranger)	m_{try}	Integer	1	11	\sqrt{p}
	min.node.size	Integer	1	10	1
	sample.fraction	Numeric	0.2	0.9	1
SVM (e1071)	cost	Numeric	2^{-5}	2^{12}	1
	γ	Numeric	2^{-12}	2^3	1

* triangular, Epanechnikov, biweight, triweight, cos, inv, Gaussian, optimal.

3.1.1. Parameter vs. hyperparameter

For parametric models the term “parameter” is often used to refer to the regression coefficients of each predictor of a fitted model. However, for machine-learning algorithms, the terms “parameter” and “hyperparameter” both refer to “hyperparameter” as there are no regression coefficients for these models. In addition, the term “parameter” is often used in programming to refer to an argument of a function. Hyperparameters determine how exactly an algorithm works and they have an influence on the final outcome (Table 1).

Hyperparameters cannot be set manually if the best performance of a model is desired. Automatic optimization is necessary to determine the best setting. This optimization is done via procedures such as *random search* or *Bayesian optimization*. In contrast, parameters of parametric models are estimated when fitting them to the data (Kuhn and Johnson, 2013).

3.1.2. Tuning method

For hyperparameter tuning, we used SequentialModel-Based Optimization (SMBO) as implemented in the *mlrMBO* package (Bischl et al., 2017). At first, n hyperparameter settings are randomly chosen from a user-defined search space. Next, they are evaluated on the chosen resampling strategy. Based on the previous evaluations a regression model is fitted. The regression model estimates the performance of the machine learning method for unknown hyperparameter settings. Using these estimates, a new promising hyperparameter setting is proposed to be evaluated next. This is continued until a termination criterion is reached (Hutter et al., 2011; Jones et al., 1998). In this work we used an initial design of 30 randomly composed hyperparameter settings and a termination criterion of 70 iterations, resulting in a total budget of 100 evaluated settings per fold. This tuning approach substantially reduces the tuning budget that is needed to find a setting that is close to the global minimum compared to methods that do not use information from previous runs such as *random search* or *grid search* (Bergstra and Bengio, 2012).

3.1.3. Hyperparameter search spaces

The boundaries of the hyperparameter search spaces were based on the suggestions of the *mlrHyperopt* package. In cases when the optimal setting of the folds of a model was close to the specified minimum or maximum of the tuning space, we extended the limits. We furthermore checked on the first five inner folds of the first outer fold that the number of tuning iterations set in the SMBO tuning was sufficiently large (Fig. 4). This requirement was met if no new local minimum was found in the last 10 % of the iterations of the selected fold.

In addition, all models were fitted using their respective default hyperparameter settings, i.e. no tuning was performed. For SVM we used $\sigma = 1$ and $C = 1$ to suppress the automatic tuning that is usually applied by the *kernlab* package. These are the default settings set by the package before the automatic tuning is applied. The GAM implementation used in this work performs by default an internal non-spatial generalized cross-validation (GCV) to find the best smoothing parameter λ for each predictor (Wood, 2017). To make the optimization of models comparable, we tuned λ for each covariate using the tuning method that was also applied to the machine-learning algorithms. For the “no tuning” setups, we set $\lambda = 0$ for all predictors. The basis dimension for all GAM setups was set to $k = 15$ for all variables. The search space for λ ($0 - 10^6$) was determined by examining the results of a prior tuning using the internal tuning of the GAM.

3.1.4. Spatial vs. non-spatial hyperparameter tuning

Hyperparameters estimated from a non-spatial tuning lead to fitted models which are more adapted to the training data than models with hyperparameters estimated from a spatial tuning. In a non-spatial tuning setting, hyperparameters that lead to a close fit of the algorithm to the data will be favored in the tuning process due to the presence of spatial autocorrelation.

Models fitted with hyperparameters from a non-spatial tuning can potentially benefit from the remaining spatial autocorrelation in the train/test split (even if a spatial resampling was used) during performance estimation and achieve a better performance than models tuned using a spatial resampling. However, depending on the dataset structure and closeness of the model fit on the data, the reverse effect might occur and models fitted with a spatial tuning setting might yield better results. In the end it depends on whether the training/test difference is more similar to a spatial tuning setting (i.e. more heterogeneous train/test splits) or to a non-spatial tuning setting (i.e. more homogeneous train/test sets).

3.1.5. Practical implementation

Most packages offering CV solutions in R offer only random partitioning methods, assuming independence of the observations. Package *mlr*, which was used as the modeling framework in this work, was missing spatial partitioning functions but provides a unified framework for modeling and simplifies hyperparameter tuning. Within the works of this study we implemented the spatial partitioning methods of package *sperrorest* into *mlr* (Fig. 2).

3.2. Estimation of predictive performance

3.2.1. Nested cross-validation

Cross-validation is a resampling-based technique for the estimation of a model's predictive performance (James et al., 2013). The basic idea behind CV is to split an existing dataset into training and test sets using a user-defined number of partitions (Fig. 3). First, the dataset is divided into k partitions or folds. The training set consists of $k - 1$ partitions and the test set of the remaining partition. The model is trained on the training set and evaluated on the test partition. A repetition consists of k iterations for which every time a model is trained on the training set and evaluated on the test set. Each partition serves as a test set once.

3.2.2. Influence of spatial autocorrelation in cross-validation

In ecology, observations are often spatially dependent (Dormann et al., 2007; Legendre and Fortin, 1989). Subsequently, they are affected by underlying spatial autocorrelation by a varying magnitude (Legendre, 1993; Cliff and Ord, 1970; Telford and Birks, 2005). Model performance estimates are expected to be overoptimistic due to the similarity of training and test data in a non-spatial partitioning setup when using any kind of cross-validation for tuning or validation (Burman et al., 1994; Cliff and Ord, 1970; Racine, 2000). Therefore, cross-validation approaches that adapt to this problem should be used in any kind of performance evaluation when spatial data is involved

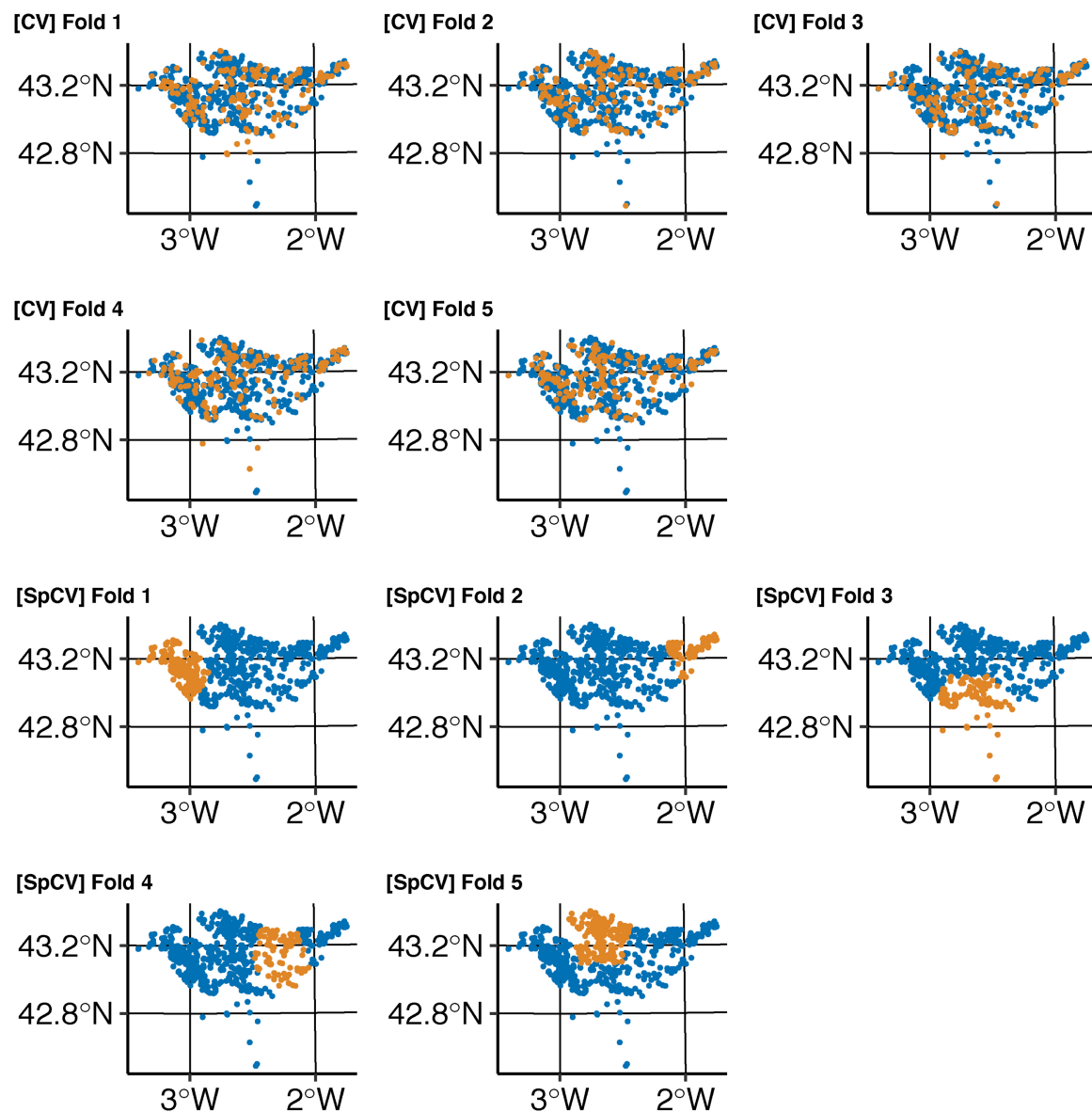


Fig. 2. Comparison of spatial and non-spatial partitioning of the first five folds in spatial and non-spatial cross-validation performance estimation. Blue dots represent the training samples and orange dots the testing sample. “SpCV” stands for spatial cross-validation (spatial sampling of observations) and “CV” for cross-validation (random sampling of observations). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

(Meyer et al., 2018; Telford and Birks, 2009). In this work we use the spatial cross-validation approach after Brenning (2012) which uses k -means clustering to reduce the influence of spatial autocorrelation. In contrast to non-spatial CV, spatial CV reduces the influence of spatial autocorrelation by partitioning the data into spatially disjoint subsets (Fig. 3).

A 100 times repeated (to reduce random variability introduced by partitioning) five-fold partitioning setting was chosen for performance estimation (Fig. 3). For hyperparameter tuning, again five folds were used to partition the training set of each fold. Hyperparameter tuning only applied to the machine-learning algorithms. A sequential model-based optimization approach was used for optimization (see Section 3.1). Model performances of every hyperparameter setting were computed at the tuning level and averaged across folds. The hyperparameter setting with the lowest mean Brier score across all tuning folds was used to train a model on the training set of the respective performance estimation level. This model was then evaluated on the test set of the respective fold (performance estimation level).

3.2.3. Cross-validation settings

To underline the crucial need for spatial CV when assessing a model's performance, and to identify overoptimistic outcomes when neglecting to do so, we used the following CV setups:

- Nested non-spatial CV which uses random partitioning and non-spatial hyperparameter tuning (*non-spatial/non-spatial*)
- Nested spatial CV which uses k -means clustering for partitioning (Brenning, 2005) and results in a spatial grouping of the observations in combination with non-spatial hyperparameter tuning (*spatial/non-spatial*)
- Nested spatial CV including spatial hyperparameter tuning (*spatial/spatial*)
- Spatial CV without hyperparameter tuning (*spatial/no tuning*)
- Non-spatial CV without hyperparameter tuning (*non-spatial/no tuning*)

Setup (*non-spatial/non-spatial*) was only used to show the overoptimistic results when using non-spatial CV with spatial data and

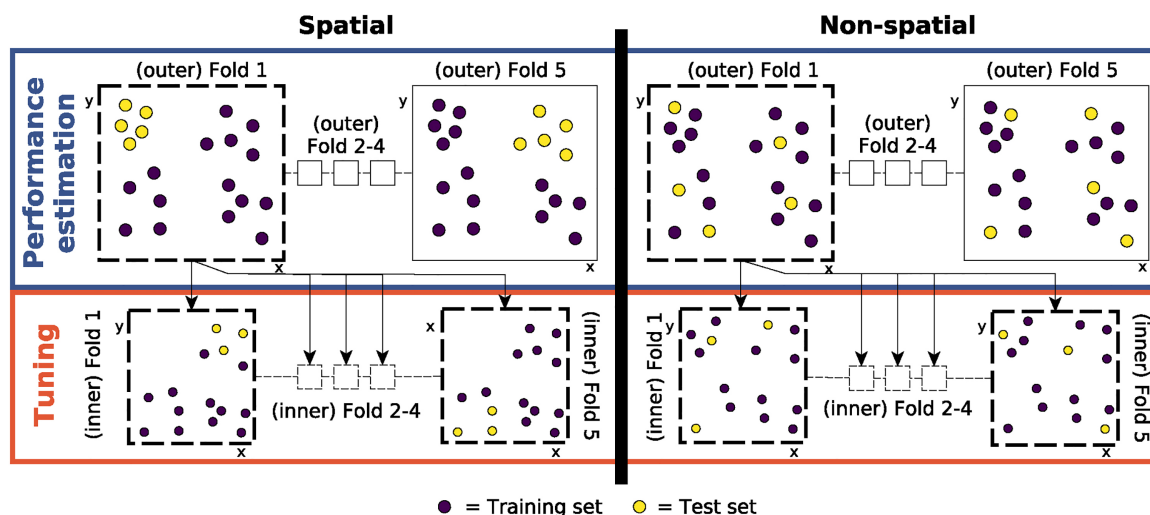


Fig. 3. Theoretical concept of spatial and non-spatial nested cross-validation using five folds for hyperparameter tuning and performance estimation. Yellow/purple dots represent the training and test set for performance estimation, respectively. The tuning sample is based on the respective performance estimation fold sample and consists again of training (orange) and test set (blue). Although the tuning folds of only one fold are shown here, the tuning is performed for every fold of the performance estimation level. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

setups *spatial/non-spatial*, *spatial/spatial* to reveal the differences between spatial and non-spatial hyperparameter tuning. Setup (*spatial/spatial*) should be used when conducting spatial modeling with machine learning algorithms that require hyperparameter tuning.

3.2.4. Performance measure

Brier score was selected as a scoring rule to compare the predictive performances of all algorithms (Brier, 1950). In contrast to other commonly used measures for binary classification (e.g. the area under the receiver operating characteristics curve (AUROC)), Brier score classifies as a proper scoring rule (Byrne, 2016; Gneiting and Raftery, 2007). It is defined as the mean quadratic loss between the predicted and observed probabilities. It ranges from 0 to 1 with low values indicating a good prediction (Brier, 1950).

3.2.5. A note on spatial autocorrelation structures in parametric models

In this work we expect that, on average, the predictive accuracy of parametric models with and without spatial autocorrelation structures incorporated into the model is the same. However, there is little research on this specific topic (Dormann, 2007; Mets et al., 2017) and a detailed analysis goes beyond the scope of this work. In our view, a possible analysis would need to estimate the spatial autocorrelation structure of a model for every fold of a cross-validation using a data-driven approach (i.e. automatically estimate the spatial autocorrelation structure from each training set in the respective CV fold) and compare the results to the same model fitted without a spatial autocorrelation structure. Since we only focused on predictive accuracy in this work, we did not use spatial autocorrelation structures during model fitting for generalized linear model (GLM) and GAM to reduce runtime.

4. Results

4.1. Tuning

4.1.1. Optimization paths

To proof the effectiveness of the tuning, the optimization paths of the first five folds of RF for settings *spatial/spatial* and *spatial/non-spatial* are visualized (Fig. 4). Using 100 SMBO iterations, all shown folds show decrease in Brier score along the optimization path (Fig. 4). Apart from fold 5 of setting *spatial/non-spatial*, all folds show a saturation of at least 15 or more iterations in which no new local minimum was found.

4.1.2. Best hyperparameter settings

There were notable differences in the distribution of the estimated optimal hyperparameters between the spatial (*spatial/spatial*) and non-spatial (*spatial/non-spatial*, *non-spatial/non-spatial*) tuning settings (Fig. 5): in the spatial tuning setting, all models besides BRT show a wide range of optimal hyperparameters across folds. By contrast, the range of optimal settings in the non-spatial tuning case is much smaller and often clusters around a few specific settings (e.g. compare the spatial and non-spatial results of the SVM) (Fig. 5).

For the spatial tuning case of RF, the estimated m_{try} values mainly ranged between 1 and 3 and m_{try} of 1 was most often the optimal value. This stands in strong contrast to the non-spatial tuning setting in which m_{try} mainly ranged between 3 and 5 and m_{try} of 3 was most often the optimal choice. Generally, we observed the tendency that spatially tuned hyperparameters are often close to the limits of the search space especially when compared to their non-spatial counterparts. The GAM results are not included in Fig. 5 as the estimated hyperparameter (smoothing parameter λ) is a vector of length eight (eight being the number of non-linear variables in the model formula) which cannot be visualized in two dimensions.

4.2. Predictive performance

4.2.1. Which models showed the best performance?

For the spatial settings (*spatial/spatial* and *spatial/no tuning*), RF showed the best predictive performance followed by BRT, KNN and GLM (Fig. 6). The absolute difference between the best (RF) and worst (GAM) performing model in our benchmark comparison is 0.039 (mean Brier score (*spatial/spatial*))). The GAM showed a high variance for all spatial settings compared to all other algorithms.

4.2.2. Effect of hyperparameter tuning on predictive performance

The tuning of hyperparameters resulted in a clear increase of predictive performance for BRT (0.183 (*spatial/spatial*) vs. 0.201 (*spatial/no tuning*) mean Brier score), GAM (0.206 (*spatial/spatial*) vs. 0.251 (*spatial/no tuning*) and KNN (0.181 (*spatial/spatial*) vs 0.210 (*spatial/no tuning*) mean Brier score) (Fig. 6). No effect of hyperparameter tuning on predictive performance was visible for RF and SVM. The use of spatial partitioning in hyperparameter tuning (setting (*spatial/spatial*) had a substantial positive impact for BRT and a negative one for GAM and KNN (Fig. 6)).

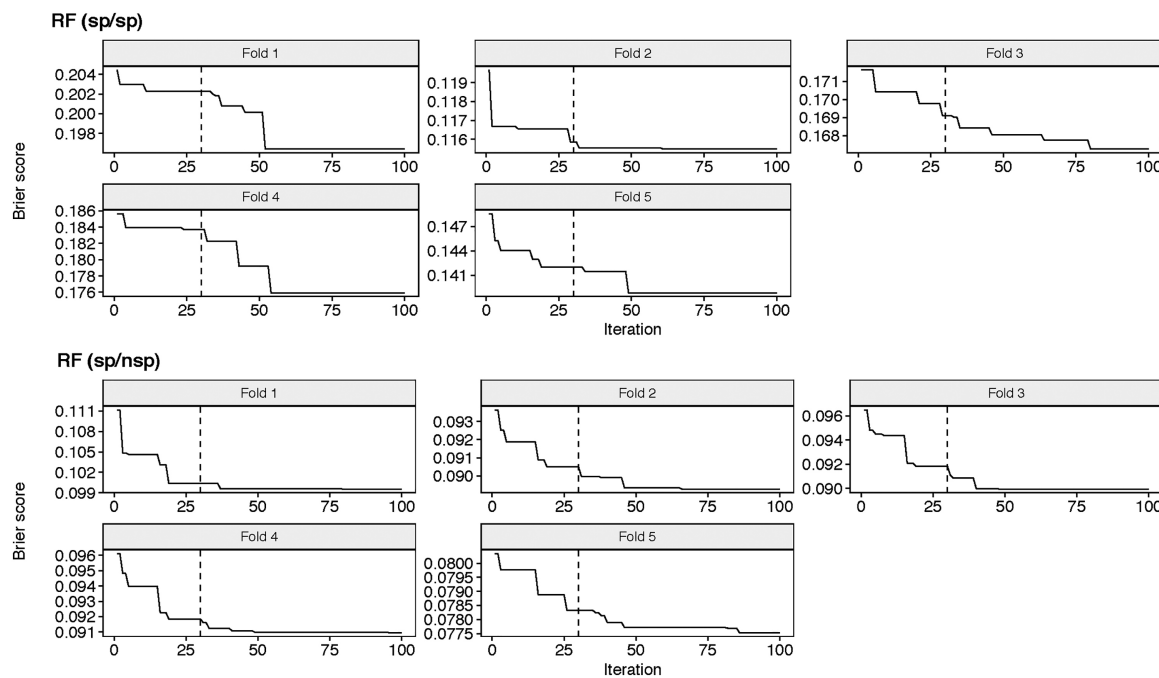


Fig. 4. SMBO optimization paths of the first five folds of the *spatial/spatial* and *spatial/non-spatial* CV setting for RF. The dashed line marks the border between the initial design (30 randomly composed hyperparameter settings) and the sequential optimization part in which each setting was proposed using information from the prior evaluated settings. Optimization paths of the remaining models can be found in the Appendix. Visualizations for other algorithms can be found in the research compendium of this study.

4.2.3. Comparison of spatial vs non-spatial tuning

Predictive performance estimates based on non-spatial partitioning (*non-spatial/non-spatial* or *non-spatial/no tuning*) are around 33–47% higher, i.e. overoptimistic, compared to their spatial equivalents (*spatial/spatial*, *spatial/no tuning*). BRT and RF show the highest differences between these two settings (47% and 46%, respectively) while GLM was the least affected (33%).

5. Discussion

5.1. Tuning

5.1.1. Tuning methods

The question on the most efficient approach of hyperparameter tuning has often been discussed (Bengio, 2000; Probst et al., 2018a; Yang et al., 2017). The goal is to use as few computational resources as possible to find a nearly optimal hyperparameter setting of an algorithm for a specific dataset. In this respect, methods like *random search* are particularly promising

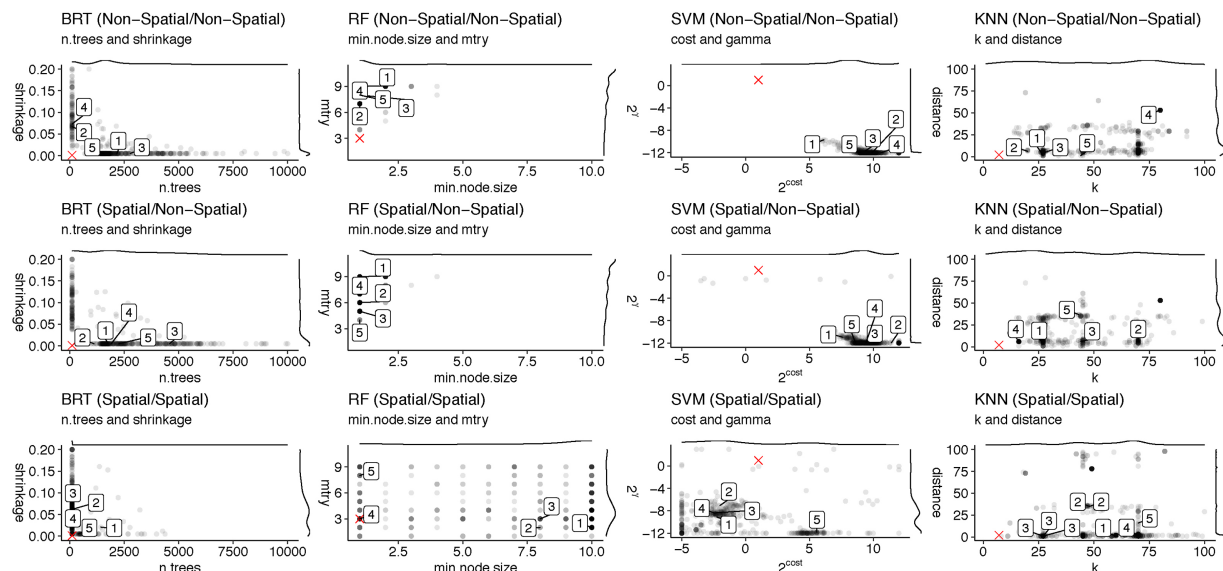


Fig. 5. Best hyperparameter settings by fold (500 total) each estimated from 100 (30/70) SMBO tuning iterations per fold using five-fold cross-validation. Split by spatial and non-spatial partitioning setup and model type. Red crosses indicate the default hyperparameters of the respective model. Black dots represent the winning hyperparameter setting of each fold. The labels ranging from one to five show the winning hyperparameter settings of the first five folds. Density lines on both axes show the density distribution of the respective variable. Visualizations for other algorithms can be found in the research compendium of this study. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

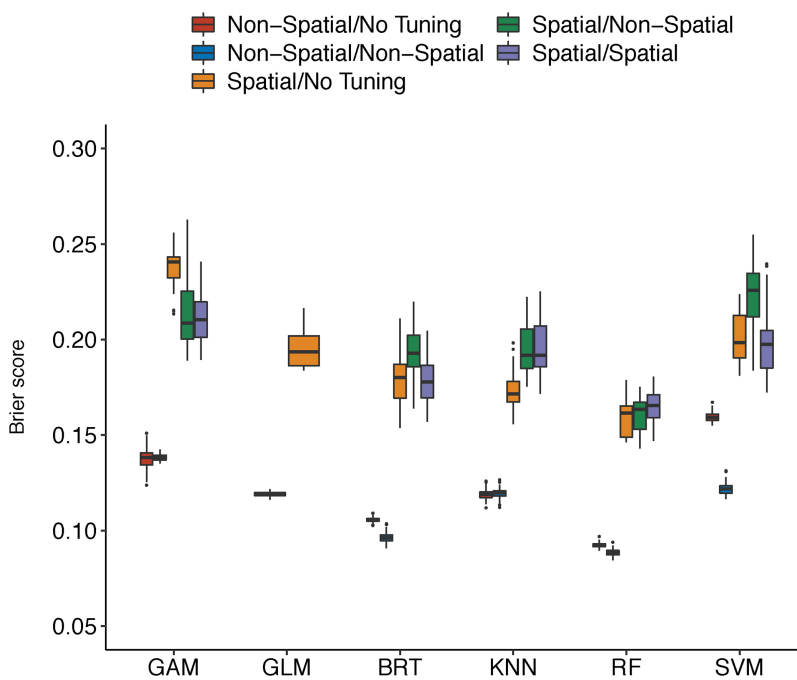


Fig. 6. (Nested) CV estimates of model performance at the repetition level using 100 SMBO iterations for hyperparameter tuning. CV setting refers to performance estimation/hyperparameter tuning of the respective (nested) CV, e.g. “spatial/non-spatial” means that spatial partitioning was used for performance estimation and non-spatial partitioning for hyperparameter tuning.

in multidimensional hyperparameter spaces with possibly redundant or insensitive hyperparameters (low effective dimensionality; (Bergstra and Bengio, 2012)). Adaptive search algorithms offer computationally efficient solutions to these difficult global optimization problems in which little prior knowledge on optimal subspaces is available. Approaches like Bayesian Optimization and F-racing are widely used for optimization of black-box models (Birattari et al., 2002; Bischi et al., 2017; Brochu et al., 2010; Malkomes et al., 2016). In this study, we used a sequential model-based optimization (Bayesian optimization) method. Other tuning methods would be expected to yield almost identical results but at the cost of increased computational efficiency and less robustness in terms of finding the local minimum.

5.1.2. Algorithm sensitivity to tuning

Some models (e.g. RF) are known to be relatively insensitive to hyperparameter tuning (Probst et al., 2018b). However, as the effect of hyperparameter tuning also depends on the dataset, hyperparameters should always be tuned. If no tuning is conducted, it cannot be ensured that the respective model showed its best possible predictive performance on the dataset.

5.1.3. Hyperparameter search spaces

Computational expense, especially when using tuning methods like random search, should focus on plausible parameter settings for each model. It should be ensured by visual inspection that the majority of the obtained optimal hyperparameter settings is not close to the ranges of the tuning space. If the optimal hyperparameter settings are clustered at the borders of the parameter search space, this implies that optimal hyperparameters may actually lie outside the given range. However, extending the tuning space is not always possible nor practical as (1) numerical problems within the algorithm may occur that may prohibit further extension of the tuning space; (2) some algorithms tend to mainly use the limits of the given search space although no substantial increase is achieved (e.g. KNN in the *spatial/spatial* setting).

We encountered exactly these problems in the *spatial/spatial* setting for BRT, KNN and SVM. For example, in the *spatial/spatial* setting, we should have further increased the search space for the mentioned models based on the distribution of the optimal hyperparameters of each fold (Fig. 5). However, using the extended setting, the algorithms did not converge anymore for some folds and at the same time runtime

increased without a substantial increase in predictive performance.

All these points show the need for a thorough specification of parameter search spaces. As the optimal hyperparameter ranges also depend on the dataset characteristics, it is not possible to define a universal search space that works best on every dataset. Nevertheless, the chosen hyperparameter limits of this work can serve as a starting point for future analyses in the spatial modeling field. Within the framework of the *mlr* project a database exists which stores hyperparameter settings of various models from users that can serve as a reference point (Richter, 2017).

5.1.4. Comparison of spatial vs non-spatial tuning

No major differences in model performances were found when using spatial versus non-spatial hyperparameter tuning procedures (e.g. 0.019 for BRT (0.182 vs. 0.201 mean Brier score).

The winning algorithm RF is used to discuss the optimal estimated hyperparameters per fold of the spatial and non-spatial tuning setting in more detail. Although the tuning of RF had no substantial effect on predictive performance (Fig. 6), the estimated optimal hyperparameters of RF differ for the non-spatial and spatial tuning setting (Fig. 5). We split the following discussion into two points: (1) explanation of the differences and (2) the implications of choosing a specific resampling method.

(1) The resampling method has no direct effect on how RF prioritizes variables internally. The Gini Impurity Index which is used to choose the variable that is used for splitting a node is calculated on a bootstrapped sample from the training data of the respective fold (Breiman, 2001). This applies for both spatial and non-spatial tuning. The Gini Index is not affected by spatial autocorrelation in this setting and RF will select the same variables in both resampling settings. Next RF is trained using the specific hyperparameter set which was given in this fold (for example $m_{try} = 3$ and $min.\ node.\ size = 4$). Now the effect of choosing different resampling strategies applies:

- In the spatial setting, RF scored a low performance on the test set. The trained model overfitted on the training set.
- In the non-spatial setting, RF scored a good performance on the test. Here, the test set was highly similar to the training set and fitting the model closely to the training data resulted in good test set results.

The higher m_{try} and the lower $min.\ node.\ size$ are chosen, the more RF will overfit on the given data. This statement is backed up by the

visualization of the chosen hyperparameter settings in each fold (Fig. 5). Ultimately, a spatial resampling in the tuning setting forces the algorithm to create a model that is more regularized than it would be in the case of a non-spatial resampling setting. This applies to all algorithms.

(2) Even though the estimated hyperparameters from a spatial and non-spatial sampling differ, they sometimes achieve the same performance when being evaluated at the performance estimation level of the CV (Fig. 6). This outcome depends on the specific characteristics of the chosen dataset and algorithm. For example, SVM showed substantial differences between the resampling methods chosen during tuning while the effect for KNN was negligibly small. The findings of this work need to be verified by using other spatial datasets (and algorithms). In addition, if a model is going to be evaluated on a specific sampling scheme (here spatial sampling), we see no valid argument why its hyperparameters should be trained on a different sampling scheme if the predictive performances do not differ significantly.

5.2. Predictive performance

5.2.1. Non-spatial vs. spatial CV

Our findings agree with previous studies in that non-spatial performance estimates appear to be substantially “better” than spatial performance estimates (Meyer et al., 2018; Micheletti et al., 2013; Roberts et al., 2017). However, this difference can be attributed to an overoptimistic bias in non-spatial performance estimates in the presence of spatial autocorrelation (Goetz et al., 2015; Meyer et al., 2018; Ruß and Brenning, 2010; Steger et al., 2016). Spatial cross-validation is therefore required for performance estimation in spatial predictive modeling, and similar grouped cross-validation strategies have been proposed elsewhere in environmental as well as medical contexts to reduce bias in predictive performance (Brenning and Lausen, 2008; Meyer et al., 2018; Peña and Brenning, 2015; Pohjankukka et al., 2017; Roberts et al., 2017).

5.2.2. The effect of hyperparameter tuning on predictive accuracy

Although hyperparameter tuning certainly increased the predictive performance for BRT, GAM and KNN in our case, the magnitude always depends on the meaningful/arbitrary defaults of the respective algorithm and the characteristics of the dataset. Naturally, the tuning effect is higher for models without meaningful defaults (such as BRT and KNN) than for models with meaningful defaults such as RF. To underline this statement, there was basically no tuning effect for SVM in this study (Fig. 6) although the SVM usually shows substantial increases when being tuned (Rojas-Dominguez et al., 2018).

5.2.3. Predictive performance across algorithms

Other studies also found that RF showed the best predictive performance (referring to setting *spatial/spatial*) (Bahn and McGill, 2012; Jarnevich et al., 2017; Smoliński and Radtke, 2016; Vorpahl et al., 2012) although this is not always the case (Peña and Brenning, 2015). The fact that the GLM is showing a better performance than the GAM shows the heterogeneous train/test split introduced by spatial partitioning: The GAM was probably not able to generalize enough (i.e. it overfitted on the training set) in the spatial resampling setting. The high variance of the GAM performances in the spatial setting support this assumption: If the training set is similar to the test set, the GAM is able to achieve Brier score results around 0.19. In cases where training and test set are more heterogeneous, the predictive performance showed Brier score estimates up to 0.30. The linear approach of the GLM was able to generalize better in this study and subsequently resulted in a better performance.

It may be surprising at first that the GLM is showing a performance which is similar to that of BRT, KNN and SVM. This is most likely due to the ability of the algorithm to generalize. Highly flexible algorithms have a tendency to overfit when the test set differs substantially from the training set. For instance, a test set located close to the sea might be hard to predict for models trained on data that was almost exclusively located in mountainous regions. In such a setting, a low degree of flexibility will result

in better predictions. This example also shows the importance of traditional parametric approaches in ecological modeling: Often enough ecological datasets show a high degree of diversity and machine-learning models might suffer from overfitting. In this case, the interpretability, speed and generalization capabilities of a GLM make this algorithm a valid choice, especially if the differences in predictive accuracy compared to black-box models are small.

5.2.4. The influence of the performance measure

The choice of the scoring rule for the evaluation of binary classifications is an important decision (Gneiting and Raftery, 2007). Measures that are not classified as “proper” can lead to undetected deviations during scoring that can end up in biased results (Byrne, 2016). One of the most used performance measures in the field of binary classification, the AUROC, is affected by this. In a previous version of this work we used AUROC to rank the algorithms which had the effect of GAM showing a similar performance as RF. So by only changing the measure, GAM went from the best (AUROC) to the worst (Brier score) algorithm. This example highlights the importance of selecting a measure for benchmarking purposes that is classified as a proper scoring rule. However, analyzing the effect of different measures on benchmarking performance across algorithms exceeds the scope of this work. Nevertheless, the use of the AUROC is justifiable in situations where relative indices of susceptibility are sought instead of predicted probabilities (e.g., hazard susceptibility modeling, Goetz et al. (2015)).

5.2.5. A note on spatial autocorrelation structures in parametric models

In this work we expect that, on average, parametric models with and without residual autocorrelation structures are comparable. However, since model comparisons have focused on model behavior in statistical inference there is little research on this specific topic (Dormann, 2007; Mets et al., 2017) and a detailed analysis goes beyond the scope of this work. In our view, a possible analysis would need to re-estimate the spatial autocorrelation structure for every fold of a cross-validation using a data-driven approach (i.e. automatically fit a residual autocorrelation on each in the respective CV fold) and compare the results to the same model fitted without a spatial autocorrelation structure. Since we only focused on predictive accuracy in this work, we did not use spatial autocorrelation structures during model fitting for GLM and GAM to reduce runtime. However, if the aim is statistical inference, it is of utmost importance to include a spatial autocorrelation structure during model fitting.

5.2.6. The effect of overoptimistic performance estimates on ecological decision making

Unbiased model outcomes are the foundation of informed ecological decision-making, biodiversity conservation as well as ecological restoration strategies (Muenchow et al., 2018). In particular, reliable outcomes are indispensable in species distribution (Loehle, 2018), invasive species dispersal (Srivastava et al., 2018), and ecosystem service modeling (Watanabe and Ortega, 2014). Global change makes model predictions uncertain enough (IPCC, 2013). Therefore, it is unnecessary to introduce an additional autocorrelation bias, especially since one can relatively easily account for it. We encourage the use of spatial CV for performance estimation (Ruś and Kruse, 2010; Brenning, 2012), variable importance assessment (Brenning, 2012; Brenning et al., 2012) and hyperparameter tuning (this study).

5.3. Outlook

In this study, we focused on comparing resampling methods (spatial vs. non-spatial strategies) including hyperparameter tuning on a typical ecological dataset. Also we showed how to retrieve a bias-reduced performance estimate in the presence of spatial autocorrelation. Since we only used one dataset, the numeric outcomes are not generalizable. Still, we believe that future studies adapting the approach presented in this work will help with finding general patterns. It would be interesting to see if a spatial

hyperparameter tuning (Fig. 3) shows a more pronounced effect when other datasets are used. Most freely available datasets in the major repositories (Olson et al., 2017; Vanschoren et al., 2014) lack spatial information which obviously is the prerequisite for spatial data analysis.

Finally, ecological observations are often observed repeatedly at the same locations. In this case, the observations are most likely affected by both spatial and temporal autocorrelation. Therefore, one would have to adjust the methodology presented in this manuscript by incorporating the temporal dimension into the spatial resampling strategy.

6. Conclusions

In this study, we compared six statistical and machine-learning models in terms of predictive performance. With the exception of SVM, all machine-learning models outperformed (semi-)parametric models. More importantly, we found that non-spatial partitioning yields largely overoptimistic performance results in the presence of spatial autocorrelation.

By contrast, the effect of hyperparameter tuning on the predictive performance was less obvious, varies by algorithm and was overall smaller than the performance differences between algorithms. Additionally, the performance differences between spatial and non-

spatial hyperparameter tuning were rather small. Still, we would recommend to use spatial CV instead of non-spatial CV for hyperparameter tuning when working with spatial data as only this ensures the assessment of bias-reduced predictive performance results. This is especially important when the corresponding results form the basis of ecological and conservation decision making.

Finally, we recommend to clearly identify the main goal of an analysis from the beginning: If the goal is to disentangle environmental-ecological relationships with the help of statistical inference, (semi-) parametric models should be favored even if they fare less well in terms of predictive accuracy. By contrast, if the intention is to produce highly accurate spatial prediction maps, spatially tuned machine-learning models maybe the better choice.

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Appendix A. Descriptive summary of numerical and nominal predictor variables

Table A.2

Summary of numerical predictor variables. Precipitation (precip) in mm/m², temperature (temp) in °C, solar radiation (pisr) in kW/m², tree age (age) in years. Statistics show sample size (**n**), minimum (**Min**), 25th percentile (**q₁**), median (**̄x**), mean (**̄̄x**), 75th percentile (**q₃**), maximum (**Max**), inner-quartile range (**IQR**) and NA Count (#NA).

Variable	n	Min	q ₁	̄x	̄̄x	q ₃	Max	IQR	#NA
temp	922	12.6	14.6	15.2	15.1	15.7	16.8	1.0	0
precip	922	88.1	181.2	224.6	234.1	252.2	496.6	71.0	0
hail_probability	922	0.0	0.2	0.6	0.5	0.7	1.0	0.5	0
ph	922	4.0	4.4	4.6	4.6	4.8	6.0	0.4	0
slope_degrees	922	0.1	12.3	19.3	19.8	27.0	55.1	14.7	0
pisr	922	-0.1	0.0	0.0	0.0	0.0	0.1	0.1	0
age	922	2.0	13.0	20.0	19.0	24.0	40.0	11.0	0

Table A.3

Summary of nominal predictor variables

Variable	Levels	#	%
Diplo01	0	700	75.9
	1	222	24.1
	All	922	100.0
Soil	soils with clay-enriched subsoil	215	23.3
	Soils with little or no profile differentiation	705	76.5
	Pronounced accumulation of organic matter in the mineral topsoil	1	0.1
	Soils with limitations to root growth	1	0.1
Lithology	All	922	100.0
	surface deposits	31	3.4
	Clastic sedimentary rock	600	65.1
	Biological sedimentary rock	136	14.8
	Chemical sedimentary rock	142	15.4
	Magmatic rock	13	1.4
Year	All	922	100.0
	2009	399	43.3
	2010	260	28.2
	2011	102	11.1
	2012	161	17.5
All	All	922	100.0

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