



Article A Comprehensive Study of Random Forest for Short-Term Load Forecasting

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Abstract: Random forest (RF) is one of the most popular machine learning (ML) models used for both classification and regression problems. As an ensemble model, it demonstrates high predictive accuracy and low variance, while being easy to learn and optimize. In this study, we use RF for short-term load forecasting (STLF), focusing on data representation and training modes. We consider seven methods of defining input patterns and three training modes: local, global and extended global. We also investigate key RF hyperparameters to learn about their optimal settings. The experimental part of the work demonstrates on four STLF problems that our model, in its optimal variant, can outperform both statistical and ML models, providing the most accurate forecasts.

Keywords: random forest; regression tree; pattern representation of time series; short-term load forecasting

1. Introduction

Electricity demand forecasting is extremely important for energy providers to ensure the secure, effective and economic operation of the power system. Short-term load forecasting (STLF) covers a forecast horizon of a few hours to a few days. STLF is necessary for generation resource planning to meet electricity demands and optimize the power flow on the transmission grid to avoid overloads. As electricity demand is a major driver of electricity prices, load forecasting plays a key role in competitive energy markets. The STLF accuracy directly affects the financial performance of energy market participants.

The importance of accurate electricity demand forecasts for the safe, reliable and effective operation of power systems is behind the great interest of researchers in this area. STLF problems are complex because electricity demand time series express a nonlinear trend, multiple seasonality, variable variance, significant random disruptions and changing daily profile. These challenging factors place high demands on STLF models.

1.1. Related Work

Roughly, STLF methods can be divided into statistical and ML methods. The most popular representatives of the first group are: auto-regressive integrated moving average (ARIMA) [1], exponential smoothing (ETS) [2], linear regression [3], and Kalman filtering [4]. The main drawbacks of the statistical methods are their linear character, limited adaptability, limited ability to deal with complex seasonal patterns, and problems with capturing long-term dependencies in time series and introducing exogenous variables into the model [5].

ML models provide more flexibility in modeling nonlinear functions. Unlike statistical methods, they do not require strong assumptions about the mapping function, and they learn relationships between predictors and targets directly from historical data. Among ML methods for forecasting, neural networks (NNs) have gained the most popularity in recent years [6]. The multitude of architectural solutions and mechanisms to improve performance encourage the use of NNs to solve complex forecasting problems such as STLF. Classical NNs were investigated for suitability for STLF in [7]. To deal with triple seasonality in time



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Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). series, patterns of the daily profiles were introduced, which filter out the trend and the weekly and yearly seasonality (a similar approach was used in this study, where different definitions of patterns are examined). Among the considered NN architectures, which included Multilayer Perceptron (MLP), Radial Basis Function NN, General Regression NN (GRNN), Fuzzy Counterpropagation NN, and Self-Organizing Maps, GRNN and MLP stand out as performing with the highest accuracy.

In recent years, a development in the NN field has been the move towards deep and, especially useful in forecasting, recurrent architectures [6]. Deep NNs (DNNs) are especially beneficial in learning the most useful data representation for modeling a given target function, while Recurrent NNs (RNNs) are beneficial in modeling complex, shortand long-term temporal relationships in data. New mechanisms and procedures introduced to RNNs such as delayed connections, attention, hybrid architecture, dynamic training sets, residual connections and flexible loss functions improve their learning capabilities and expressive power to solve forecasting problems [5]. Some examples of using DNNs and RNNs for STLF are [8], where Convolutional NNs (CNNs) are utilized to extract load and temperature features, which are fed as inputs into the bidirectional propagating RNN to perform hourly electrical load forecasting [9], where a recurrent inception CNN is proposed for STLF that combines RNN and 1-dimensional CNN [10], where RNN with attention significantly reduced forecasting errors as compared to the current state-of-the-art results; and [11], where deep residual networks integrate domain knowledge and researchers' understanding of the problem and enables probabilistic load forecasting using Monte Carlo dropout.

An effective way to increase the forecast accuracy and robustness of both statistical and ML models is ensembling. This combines multiple models for a common response to improve both the accuracy and stability of the final solution compared to a single model [12]. The theoretical properties of forecast combination investigated in [13] answer the question why a simple average of forecasts often outperforms forecasts from single models. They also prove that simple averages in many cases perform better than more complicated weighting schemes. The beneficial effects of the forecast aggregation on STLF accuracy are shown in many papers: in [14], several ML methods are aggregated in ensembles for one-day-ahead wind power forecasting; in [15], to forecast an interval-valued load, ensemble of RNNs is applied, which learns on the components of the bivariate empirical mode decomposition; in [5], ensembling of a hybrid model, which combines ETS and RNN, leads to a significant reduction in the forecast error; in [16], an ensemble of randomized DNN combined with a walk-forward decomposition is proposed; in [17], a stacking ensemble approach is used to combine DNNs, and in [18], several methods of aggregating base models (MLPs) are considered. Stacking, used in the last two papers, is a way of combining base models via meta-learning, i.e., a meta-model is trained on the predictions of the base models.

Alternatives to stacking are boosting and bagging. Popular representatives of these are gradient-boosted trees and random forest (RF), respectively. Both, used as forecasting models, are based on regression trees. It was shown in [19] that RF can compete with both classical models and NNs in STLF. It can deal with complex time series using appropriate data preprocessing, which produces normalized patterns of the daily profiles. Based on this research, in this study, to improve RF performance, we extend pattern definitions and introduce additional predictors. To enrich input information, in [20], the input patterns are extracted from electrical, meteorological and calendar data by temporal CNN. Fed with these patterns, a Light Gradient Boosting Machine, a type of gradient-boosted trees algorithm, was able to forecast very volatile industrial customer loads. A novel tree-based ensemble method called Warm-start Gradient Tree Boosting (WGTB) was proposed in [21]. It combines four different inference models and aggregates their outputs by a warm-start, bagging and boosting, which at the same time reduces bias and variance. The result proves the efficiency of the proposed strategy and shows an improvement in STLF accuracy over baseline models. Another type of tree-based ensemble, eXtreme Gradient Boosting (XGBoost), was used in [22] for forecasting electricity consumption by industrial customers. To deal with multiple seasonality, the time series were first decomposed using variational mode decomposition. Then, a linear regression model was applied for the trend series and a XGBoost regression model was applied for each fluctuation sub-series.

To close this section, we note that, according to some studies, tree-based ensembles are not inferior to NNs in terms of forecast accuracy. In [23], the authors have examined and reproduced a number of state-of-the-art DNNs for time series forecasting. DNNs were compared on different datasets to a Gradient Boosting Regression Tree (GBRT). The experimental results show that a conceptually simpler model such as GBRT can compete and sometimes outperform modern DNNs by efficiently feature-engineering the input and output structures of GBRT.

1.2. Motivation and Contribution

Tree-based methods are widely used as prediction models as they have very attractive properties such as a capacity for flexible nonlinear regression, which can capture complex interactions between variables and effectively handle multiple predictors (including exogenous ones) of various types (numeric, binary, and categorical). Moreover, they are robust against over-fitting of the training data, they are relatively simple to tune, and they are easy to implement with the available software. Their effectiveness has been confirmed in many forecasting competitions, for example those carried out on the Kaggle platform [24].

The excellent performance of tree-based approaches was demonstrated in the 2020 M5 forecasting competition. The top places in this competition, in terms of both accuracy and uncertainty, were dominated by entries that used tree-based ML methods such as gradient-boosted trees [25]. Four out of the five winning models used a variant of the tree-based method and most of the other top 50 best-performing models adopted similar approaches to the winning submission by training recursive and non-recursive tree-based models [26]. Thus, tree-based forecasting models appear to be strong competitors to NNs, which in the form of deep learning-based models dominate the recent literature on forecasting methods [6].

In this study, motivated by the excellent results of tree-based models in forecasting competitions, we apply RF to the challenging problem of STLF. RF gives similar results to boosting, but is easier to train and tune [27]. The main contribution of this work is to examine RF models using a variety of time series preprocessing methods and training modes. In the local mode, RF learns on samples similar to the query sample, which enables the model to focus on the local features of the target function around the query pattern and improve accuracy in this region. In the global mode, to achieve the same goal, i.e., focusing on the proper region of the target function, we introduce additional calendar variables. By examining different methods of time series preprocessing, we find the most useful data representation for achieving the highest accuracy of the model. We empirically demonstrate that the proposed approach outperforms in terms of accuracy both standard statistical models as well as more sophisticated ML approaches.

The novelty of this work in relation to our previous work [19] is twofold. First, we extend pattern definition by introducing seven types of patterns based on the historical data. They incorporate daily and/or weekly seasonality, while in [19], the patterns captured only daily seasonality. Second, we introduce a global mode of training with additional predictors representing calendar data. In [19], only a local training was considered without calendar inputs.

The rest of the paper is organized as follows. In Section 2, we propose several data prereprocessing methods for electricity demand times series. Section 3 defines the forecasting problem and RF training modes. Section 4 describes the RF algorithm in application to STLF. The experimental framework used to evaluate the performance of the proposed model and compare it with baseline models is described in Section 5. Finally, Section 6 concludes the work.

2. Data Preprocessing

The power system load or electricity demand time series express a trend, triple seasonality (annual, weekly and daily) and random fluctuations. These components are dependent on the system size, size of the economy served, customer structure, as well as weather and climatic conditions. The daily load profiles that we focus on in STLF vary throughout the year and depend on the day of the week [5].

To forecast future demand with the least possible error, the forecasting model should be fed by the most relevant predictors. In our univariate STLF model, which produces forecasts for the next day, the predictors are selected from recent history and are preprocessed accordingly. The forecasting model based on RF is fed by input patterns x_i and produces encoded forecasts for hour *t* of day *i*, $y_{i,t}$ (MISO model).

Let $\{z_{\tau}\}_{\tau=1}^{M}$ be an electricity demand time series with hourly resolution and vector $\mathbf{z}_{i} = [z_{i,1}, ..., z_{i,24}]$ represents its 24-hour-long sequence for day *i*. To capture the characteristic properties of the series, remove the trend and unify data, we define the input patterns as follows:

r1 The input patterns are defined based on the weekly sequence which precedes forecasted day *i*:

$$\mathbf{x}_i = \frac{\mathbf{s}_i - \bar{s}_i}{\|\mathbf{s}_i - \bar{s}_i\|} \tag{1}$$

where $\mathbf{x}_i \in \mathbb{R}^{168}$ is the input pattern, $\mathbf{s}_i = [\mathbf{z}_{i-7}, ..., \mathbf{z}_{i-1}]$ is the demand sequence of the week preceding the forecasted day *i* and \overline{s}_i is the mean of this sequence.

Input vectors (1), which for successive *i* represent overlapping weekly sequences shifted by one day, are normalized versions of centered vectors \mathbf{s}_i . They all have zero mean, the same variance and the same unity length. However, they differ in shape. Thus, we assume that the weekly shape carries the information about the forecasted demand of the day following this week.

- **r2** The input patterns are defined based on the daily sequence which precedes the forecasted day *i*. The encoding equation is (1), where $\mathbf{x}_i \in \mathbb{R}^{24}$ and $\mathbf{s}_i = \mathbf{z}_{i-1}$ is the demand sequence of the day preceding the forecasted day *i*. In case of r2, a carrier of the information about the forecasted value is the shape of the preceding day.
- **r3** The input patterns are defined based on the sequence composed of the demands at hour *t* of seven consecutive days preceding the forecasted day *i*. In (1), $\mathbf{x}_i \in \mathbb{R}^7$ and $\mathbf{s}_i = [z_{i-7,t}, ..., z_{i-1,t}]$, where *t* is the forecasted hour of day *i*.
- r4 The input patterns are defined based on the sequence composed of the demands at hour *t* of 21 consecutive days preceding the forecasted day *i*. In (1), $\mathbf{x}_i \in \mathbb{R}^{21}$ and $\mathbf{s}_i = [z_{i-21,t}, ..., z_{i-1,t}]$, where *t* is the forecasted hour of day *i*.
- **r5** The input patterns are defined based on the sequence composed of the demands at hour *t* of seven days preceding forecasted day *i* and representing the same day of the week as the forecasted day. For example, when the model predicts demand at hour *t* on Monday, the input pattern is composed of the demands at hour *t* of seven preceding Mondays. In (1), $\mathbf{x}_i \in \mathbb{R}^7$ and $\mathbf{s}_i = [z_{i-49,t}, z_{i-42,t}, ..., z_{i-7,t}]$, where *t* is the forecasted hour of day *i*.
- **r6** Cross-pattern combining r2 and r3—the input patterns are defined based on both: the daily sequence and the sequence composed of the demands at hour *t* of seven consecutive days preceding the forecasted day *i*. In (1), $\mathbf{x}_i \in \mathbb{R}^{30}$ and $\mathbf{s}_i = [\mathbf{z}_{i-1}, z_{i-7,t}, ..., z_{i-2,t}]$, where *t* is the forecasted hour of day *i*.
- r7 Cross-pattern combining r2 and r4—the input patterns are defined based on both: the daily sequence and the sequence composed of the demands at hour *t* of 21 consecutive days preceding the forecasted day *i*. In (1), $\mathbf{x}_i \in \mathbb{R}^{44}$ and $\mathbf{s}_i = [\mathbf{z}_{i-1}, z_{i-21,t}, ..., z_{i-2,t}]$, where *t* is the forecasted hour of day *i*.

Figure 1 shows the sequences which are used for x-patterns construction and Figure 2 shows data used for construction patterns r6 and r7. Depending on the definition, x-patterns introduce different input information to the model. Pattern r1 introduces detailed informa-

tion about the weekly sequence which precedes the forecasted day. Note that r1 expresses both daily and weekly seasonality unlike r2, which carries information only about the daily seasonality. To deal with weekly seasonality when r2-patterns are used, the model can be trained in the local mode, i.e., on the subset of x-patterns corresponding to the forecasting task (see Section 3).



Figure 1. Load time series points used for input patterns construction.

Haur	History (days)												Future	
Hour	 i-22	i-21		i -9	i-8	i-7	i-6	i-5	i-4	i-3	i-2	i-1	i	
1	 14122	12265		14133	14352	14445	14434	14422	13022	12429	14372	14544		
2	 13736	11761		13560	13891	13811	13891	13847	12353	12104	13800	14023		
3	 13640	11592		13343	13581	13527	13637	13483	12025	11961	13587	13778		
4	 13765	11588		13369	13698	13509	13696	13392	11891	11980	13576	13846		
5	 14182	11916		13527	13832	13826	13829	13385	11821	12280	13 <mark>757</mark>	14016		
6	 15500	12882		14147	14531	14458	14580	13433	11953	Inpu	t 505	14675		Forecasted
7	 17251	15218		16140	16469	16383	16546	13981	12047	data	668	16775		value
8	 18424	17033		17661	18033	17814	17949	14824	12380		970	18161		
9	 18961	18111		18515	18814	18632	18593	15876	13352	18575	18720	18968		
10	 19091	18584		18705	19108	18917	18864	16613	14214	18939	18989	19217	X	
11	 19131	18535		18630	19074	18761	18837	16908	14783	18848	18816	19119		
12	 19170	18790		18741	19188	18930	18987	17028	14952	19000	18959	19323		
13	 19135	18708		18745	19151	18835	18851	16812	15066	18974	18907	19279		
14	 18947	18678		18750	19118	18880	18823	16874	15211	19029	19030	19258		
15	 18619	18364		18419	18776	18533	18477	16521	14896	18664	18820	18989		
16	 18257	18077		18225	18461	18350	18251	16155	14658	18468	18452	18727		
17	 18049	17854		17969	18053	18186	18077	15913	14525	18189	18239	18609		
18	 18608	17936		17980	18045	18367	18293	16051	14765	18259	18399	18823		
19	 19860	18939		19145	19170	19639	19601	17451	16132	19663	19961	20302		
20	 20265	20238		20573	20549	20687	20456	18270	17110	20603	20730	20940		
21	 19166	19652		19985	19981	19960	19649	17564	16471	19877	20121	20272		
22	 17549	18164		18504	18351	18562	18201	16239	15451	18318	18627	18850		
23	 16092	16514		16802	16747	16786	16607	15092	14402	16668	16946	17241		
24	 16741	15171		15433	15436	15463	15379	13923	13267	15392	15568	15858		

Figure 2. Cross-patterns: r6 (green + blue) and r7 (green + orange + blue).

Pattern r3 introduces information on the demand in the previous seven days at the same hour as the forecasted one. It expresses only weekly seasonality. Information about the daily seasonality is not included, so the local mode of training, i.e., training on the selected r3-patterns corresponding to the forecasting task (see Section 3), can help with dealing with daily seasonality. Similar information as in r3 is contained in pattern r4 but from a longer 3-week period. Pattern r5 shows neither daily nor weekly seasonality. It carries information about the demand at the same hours as forecasted in previous days of the same type as the forecasted day.

Cross-patterns r6 and r7 express both daily and weekly seasonalities as r1, but in a more sparing form, using respectively 30 or 44 instead of 168 components. In [28], we showed that STLF based on both daily and weekly patterns gives better results than forecasting based on separate daily or weekly patterns. In [28], we aggregated forecasts generated by two neural models: a daily pattern-based model and a weekly pattern-based model, while in this study, we combine daily and weekly patterns into one pattern and use only one model.

Examples of input patterns r1–r7 are depicted in Figure 3. Note different shapes of patterns, carrying different input information.



Figure 3. Examples of input patterns r1-r7.

The output data, i.e., the electricity demand at hour *t* of day *i*, is encoded as follows:

$$y_{i,t} = \frac{z_{i,t} - \bar{s}_i}{\|\mathbf{s}_i - \bar{s}_i\|} \tag{2}$$

where \mathbf{s}_i is the demand sequence preceding forecasted day *i*, defined depending on the x-pattern type r1–r7 (this is the same sequence based on which pattern \mathbf{x}_i was defined) and \bar{s}_i is the mean of this sequence.

The output data are encoded similarly to the input data, using the same coding variables: \bar{s}_i and $\|\mathbf{s}_i - \bar{s}_i\|$. Thus, Equation (2) like Equation (1) filters the data by removing the local trend (\bar{s}_i) and unifying the variance (this is a function of the denominator of these equations, which can be thought of as a measure of diversity of the input sequence). Such filtered and unified data are predicted by the forecasting model (RF). Then, the real forecast is determined from transformed Equation (2):

$$\hat{z}_{i,t} = \hat{y}_{i,t} \| \mathbf{s}_i - \bar{s}_i \| + \bar{s}_i \tag{3}$$

where $\hat{y}_{i,t}$ is the model prediction and $\hat{z}_{i,t}$ is the real forecast.

Note that (3) brings back the local current properties of the time series (level and dispersion), which were removed by (1) and (2) to simplify the relationships between input and output data. We have successfully used this kind of preprocessing of input and output data in our previous load forecasting models to deal with multiple seasonality, simplify the model and speed up training, see, e.g., [7,19,28–32].

3. Forecasting Problem and Training Modes

The forecasting task is defined as follows: predict electricity demand for hour t^* (1, ..., 24) of day i^* based on historical data. Day i^* represents day of the week d^* (*Monday*, ..., *Sunday*). To maximize the forecasting performance and to make the most of all available training data up to day i^* (forecasted day), the forecasting model is trained individually for each forecasting task and it performs only one prediction: \hat{y}_{i^*,t^*} . Note that the "global" generalization property of the model is not important because it is built to make only one prediction. What is important is the "local" performance in the neighborhood of pattern \mathbf{x}_{i^*} . To increase this property, we use two approaches. In the first one, we train the model in the local mode and in the second one, we extend input patterns with calendar variables when we use global learning. For comparison we also train the model in the standard global mode.

The full training set determined on the historical data is $\Psi = \{(\mathbf{x}_i, y_{i,t})\}$, where $i = 1, ..., i^* - 1, t = 1, ..., 24$ and pair $(\mathbf{x}_i, y_{i,t})$ includes the input pattern and target defined according to r1–r7. The three training modes are as follows:

Local The model is trained on the subset of Ψ containing pairs $(\mathbf{x}_i, y_{i,t})$ which correspond to the forecasting task, i.e., the pairs which include targets representing the same day type as forecasted, $d(y_{i,t}) = d^*$, and the same hour as forecasted, $t = t^*$.

Global The model is trained on full training set Ψ .

Global extended The input data are extended with calendar information:

season of the year encoded as follows [29]:

$$\mathbf{p}_i = \left[\sin\frac{2\pi\#i}{366}, \cos\frac{2\pi\#i}{366}\right] \tag{4}$$

where #*i* is the number of day *i* in the year,

- day of the week, d = Monday, ..., Sunday (categorical variable), and
- hour of the day, t = 1, ..., 24 (categorical variable).

The training set in the global extended mode is of the form: $\Psi = \{(\langle \mathbf{x}_i, \mathbf{p}_i, d_i, t \rangle, y_{i,t})\}, i = 1, ..., i^* - 1, t = 1, ..., 24.$

In the local training mode, the model solves the forecasting task by learning on the samples expressing similar properties and relationships between input and output data as those expressed by input pattern \mathbf{x}_{i^*} and forecasted value y_{i^*,t^*} . That is, the training input patterns are limited to those that are similar in shape to pattern \mathbf{x}_{i^*} (further limitations in this regard can be made by selecting training data from the same period of the year as day i^* or by selecting training data based on similarity to pattern \mathbf{x}_{i^*} [7], but we have not employed these approaches in this study). The relationships between input and output data expressed in the local training set are limited to those corresponding to day type d^* and hour t^* , so we expect the model to more accurately approximate the relationship between \mathbf{x}_{i^*} and y_{i^*,t^*} than in case of global training on the full training set expressing the relationship for all day types and hours.

In the global training mode with extended inputs, the model has additional input information to more accurately solve the forecasting task, i.e., the calendar variables. Although the model is global, we expect that the calendar data will help to increase its local accuracy around pattern x_{i^*} . A regression tree as a base forecasting model can more appropriately divide the input space using the calendar variables than without these variables, and therefore approximate locally the target function with greater accuracy. Although this will lead to a more complex model than in the other two training modes.

4. Random Forest for STLF

RF is a ensemble learning algorithm based on decision trees (CART [33]) as the base models [34]. It is suitable for either regression or classification problems. In this study, for forecasting problems, we focus on the regression RF based on regression trees.

RF is devoid of the well-known drawbacks of single trees such as unstable splits and a lack of smoothness [27]. It combines bagging [35] with a random subspace method [36]. The key idea in bagging is to average multiple noisy but approximately unbiased base models and thus reduce the variance. Trees as noisy and low biased models if they have grown sufficiently deep, are great candidates for bagging. The main goal of the random subspace method is to increase diversity between trees by restricting them to work on different random subsets of the full predictor space (more specifically, at each node of the tree, a random predictor subset is selected). Each tree in the forest is built from a bootstrap sample of the original dataset, which is an additional source of diversity. Random predictors selected in the nodes of bagged trees help to decorrelate the trees and improve prediction accuracy as well as reduce the model variance.

The RF algorithm draws a bootstrap sample Ψ_k of size *N* from training set Ψ for each of *K* trees, k = 1, ..., K. For each bootstrapped sample, a tree *T* is grown by recursive partitioning the input space in each node until a minimum leaf size is reached. At each node, data splits based on *p* out of *n* predictors chosen at random are considered. The

best split is determined by maximizing the reduction in mean squared error (MSE) over all splitting candidates and cutpoints. After all *K* trees are grown in this fashion, the RF predictor is [27]:

$$\hat{f}_K(\mathbf{x}) = \frac{1}{K} \sum_{k=1}^{K} T(\mathbf{x}; \Theta_k)$$
(5)

where **x** is the input pattern and Θ_k characterizes the *k*-th tree in terms of split predictors, cutpoints, and terminal-node values.

RF has the following hyperparameters:

- number of trees *K*. Intuitively, the number of trees should be as large as possible because the model variance reduces with *K*. Usually, the forecast error stabilizes with the number of trees, and the most reasonable RF size is selected.
- minimum leaf size m. As deeper trees have low bias and large variance, they are strongly recommended as RF members. Thus, the minimum leaf size should be small. The inventors of the algorithm recommend m = 5 for regression forest.
- number of predictors to select at random for each split p. As p decreases, the correlation between trees reduces, and hence the variance of the average reduces. Typically, a value for p for regression RF is n/3, as the inventors recommend.

In practice the best values for the hyperparameters are dependent on the problem, and they should be treated as tuning parameters.

The standard method of selecting split predictors [33] has two drawbacks. Firstly, it tends to miss important interactions between pairs of predictors and the response. Secondly, it tends to select continuous predictors that have many levels, which masks more important predictors that have fewer levels, such as categorical predictors. To mitigate selection bias and increase detection of important interactions, curvature or interaction tests can be applied [37,38]. Therefore, in this study we consider three methods of selecting the split predictors:

- **s1** Standard CART method. This selects the split predictor that maximizes the splitcriterion gain over all possible splits of all predictors.
- **s2** Curvature test. This selects the split predictor that minimizes the *p*-value of chi-square tests of independence between each predictor and the response.
- **s3** Interaction test. This performs the curvature test extended by the minimization of the *p*-value of a chi-square test of independence between each pair of predictors and the response.

The algorithm of RF construction for STLF is shown in Algorithm 1. It produces a set of *K* trees, $\{T_k\}_{k=1}^{K}$. Based on them, to make a prediction for new point **x**, we use (5). Then, the real forecast is calculated from (3). Note that training set Ψ is prepared for the selected training mode and input pattern type.

Algorithm 1 Random forest construction for STLF

Input: training set Ψ containing *N* samples, number of trees *K*, minimum leaf size *m*, number of predictors to select at random for each split *p*, split predictor selection method *s* **Output:** set of trees $\{T_k\}_{k=1}^K$ **Procedure: for** k = 1 **to** *K* **do** Draw a bootstrap sample Ψ_k of size *N* from Ψ Grow tree T_k to Ψ_k by recursively repeating the following steps for each terminal node, until the minimum node size *m* is reached: – Select *p* predictors from the *n* predictors – Pick the best predictor/cutpoint among the *p* using predictor selection method *s* – Split the node into two daughter nodes **end for** In the experimental study (Section 5), we use RF specified in Algorithm 1 in several variants depending on the data preprocessing method (r1–r7) and training modes, i.e., local, global and global extended. In the global extended mode, the predictor vector is composed of $\langle \mathbf{x}_i, \mathbf{p}_i, d_i, t \rangle$. In the other training modes, the predictor vector is the same as input pattern \mathbf{x} (1).

5. Simulation Study

In this section, we investigate RF variants with different data preprocessing methods and training modes. We compare RF performance with that of other models based on classical statistical methods and ML methods.

STLF for four countries is performed: Poland (PL), Great Britain (GB), France (FR) and Germany (DE). The real-world data was collected from ENTSO-E repository (www.entsoe. eu/data/power-stats; accessed on 6 April 2016). It details the hourly power system load in the period from 2012 to 2015. The last year of the data (2015) is treated as a test period. We predict the daily load profiles for each day of this period, excluding atypical days such as public holidays (between 10 and 20 days a year). RF models were optimized on the data from 2012 to 2014, with validation data composed of 100 patterns selected randomly from 2014 and training data preceding the validation pattern.

5.1. Results for Different Preprocessing Methods and Training Modes

Tables 1 and 2 show mean absolute percentage error (MAPE) and root mean square error (RMSE), respectively, for input patterns r1–r7 and different training modes. Figures 4–6 show the boxplots of MAPE. The results can be summarised as follows:

- It is evident from these tables and figures that the global extended mode yields the lowest errors, when combined with patterns r4 (for PL, FR and DE) or r6 (for GB).
- The local training mode brings lower errors than the global one when patterns r1, r2, r6 and r7 are used, i.e., patterns which are composed of the daily curves. The local mode is usually better than the global extended one when patterns r1 and r2 are used, but it is worse than the global extended mode when cross-patterns are used, which also reflect a weekly seasonality.
- The highest errors for the global mode were observed when patterns r1 and r2 were used. In these cases, the errors were up to nine times greater than in the alternative training modes. Pattern r4 is recommended for the global mode, which for all countries provided the lowest errors. Modifying the global mode by extending the input patterns with calendar variables has always resulted in a reduction of errors.

Data	Training Mode	r1	r2	r3	r4	r5	r6	r7
	Local	1.54	1.55	1.89	2.17	3.66	1.45	1.63
PL	Global	11.55	11.61	2.11	1.30	2.94	1.82	1.82
	Global ext.	1.92	1.79	1.40	1.24	1.96	1.50	1.55
	Local	1.92	1.91	2.29	2.21	3.68	1.76	1.75
GB	Global	14.88	15.01	2.21	1.75	2.92	1.84	1.83
	Global ext.	2.22	2.09	1.79	1.59	2.12	1.47	1.52
	Local	1.88	1.77	2.12	2.43	5.10	1.74	1.86
FR	Global	8.50	8.54	2.09	1.53	2.90	2.11	1.92
	Global ext.	1.98	1.71	1.59	1.37	1.71	1.63	1.64
DE	Local	1.34	1.31	1.67	1.93	3.47	1.21	1.45
	Global	12.33	12.34	1.62	1.07	2.61	1.79	1.54
	Global ext.	1.55	1.57	1.17	0.99	1.65	1.17	1.19

Table 1. Validation MAPE for different input patterns r1–r7 and training modes (lowest errors in bold, second lowest errors in italics).

Data	Training Mode	r1	r2	r3	r4	r5	r6	r7
	Local	436	522	489	540	1014	412	473
PL	Global	2420	2427	577	347	741	454	458
	Global ext.	468	448	364	337	509	406	410
	Local	918	825	1100	1118	1816	811	841
GB	Global	6309	6358	1035	860	1445	811	824
	Global ext.	925	937	784	757	985	669	690
	Local	1630	1576	1851	1939	3893	1406	1626
FR	Global	5602	5635	1714	1144	2697	1538	1437
	Global ext.	1414	1280	1287	1048	1450	1232	1272
DE	Local	1590	1190	1748	2099	3046	1168	1641
	Global	8927	8926	1410	878	2094	1411	1131
	Global ext.	1224	1229	833	713	1261	915	883

Table 2. Validation RMSE for different input patterns r1–r7 and training modes (lowest errors in bold, second lowest errors in italics).



Figure 4. Local training mode: Boxplots of validation MAPE for different input patterns r1-r7.



Figure 5. Global training mode: Boxplots of validation MAPE for different input patterns r1-r7.



Figure 6. Global extended training mode: Boxplots of validation MAPE for different input patterns r1–r7.

Based on the results, the recommended training mode is global extended with r4 patterns for PL, FR and DE, and r6 patterns for GB. These variants of RF were used in the experiments described in the next sections.

5.2. Tuning Hyperparameters

In this experiment, we change the selected hyperparameter in the range shown in Figure 7, keeping the remaining hyperparameters at their constant values as follows: number of trees in the forest—K = 100, minimum number of leaf node observations—m = 1, and number of predictors to select at random for each decision split—p = n/3.

Figure 7 shows the impact of hyperparameters on the forecasting error (MAPE). As expected, the error decreases with the number of trees in the forest. The reduction in MAPE when the RF size changes from 1 do 300 trees was from 38.7% for PL to 50.0% for DE. At the same time, a significant reduction in the forecast variance was also observed from 63.8% for PL to 82.5% for DE. It can be seen from Figure 7b that an increase in the minimum leaf size leads to a deterioration in the results. Small values of *m*, close to 1, are preferred. This means that trees as deep as possible are the most beneficial in RF. The optimal number of predictors selected in the nodes to perform a split varies from country to country (see Figure 7c). For PL and DE it is 15, for GB it is 20, and for FR it is 6. These values differ from the recommended p = n/3, which are 8 for PL, FR and DE, and 11 for GB.



Figure 7. Validation MAPE depending on hyperparameters: number of trees in the forest (**a**), minimum number of leaf node observations (**b**) and number of predictors to select at random for each decision split (**c**).

Using optimal values of the hyperparameters for each country, we investigate the methods of split predictor selection s1–s3. Table 3 shows the results, validation MAPE and RMSE. Both accuracy measures show similar results for all methods of split predictor selection. Therefore, s1 is recommended as a simple, standard method, which does not cause any additional computational burden.

Data	Variant		MAPE		RMSE			
Data	variant	s1	s2	s 3	s1	s2	s 3	
PL	r4, <i>p</i> = 15	1.26	1.30	1.30	339	345	345	
GB	r6, $p = 20$	1.44	1.42	1.42	652	649	649	
FR	r4, $p = 6$	1.35	1.39	1.38	1046	1071	1053	
DE	r4, <i>p</i> = 15	0.98	1.00	1.00	709	714	714	

Table 3. Validation MAPE and RMSE for different methods of split predictor selection (training mode: global extended, #trees: 300, minimum leaf size: 1; lowest errors in bold).

Figure 8 shows the "importance" or "predictive strength" of the predictors estimated on the out-of-bag data (this is discussed further in Section 5.4). As can be seen from this figure, when r4 extended pattern is used (PL, FR and DE), the most important predictor is the last component of the r-pattern, i.e., the predictor expressing electricity demand at forecasted hour *t* of the day preceding the forecasted day, $z_{i-1,t}$. The importance of this predictor reaches 3.5 for FR and over 5 for PL and DE, while the importance of other demand predictors is usually below 2. Among the calendar predictors, the most important for r4 extended pattern are those coding the season of the year, especially for DE. For cross-pattern r6 (GB), the most important predictors are the calendar ones: day of the week and season of the year (*p*1). The next positions are occupied by predictors coding the demand for the last four hours of the day before the forecasted day ($z_{i-1,24}$ is clearly the most important of these) and predictor coding demand at forecasted hour *t* week ago, $z_{i-7,t}$. Note the low importance of the other predictors representing demand at hour *t* of the preceding days, $z_{i-6,t}-z_{i-2,t}$.



Figure 8. Importance of the predictors (x1 - x30—predictors expressing demand pattern r4 or r6, p1 and p2—components of vector **p** expressing season of the year, *d*—day of the week, and *t*—hour of the day).

Table 4 shows the forecasting results for the test set when using RF with the optimal values of hyperparameters. As performance metrics we use: MAPE, MdAPE (median of absolute percentage error), IqrAPE (interquartile range of APE), RMSE, MPE (mean percentage error), and StdPE (standard deviation of PE). MdAPE measures the mean error without the influence of outliers, while RMSE, as a square error, is especially sensitive to outliers.

Table 4. Results for test data (training mode: global extended, #trees: 300, minimum leaf size: 1, splitpredictor selection method: s1).

Data	Variant	MAPE	MdAPE	IqrAPE	RMSE	MPE	StdPE
PL	r4, $p = 15$	1.05	0.78	1.06	259	0.03	1.52
GB	r6, $p = 20$	2.36	1.78	2.39	1058	-0.32	3.36
FR	r4, $p = 6$	1.67	1.18	1.67	1338	-0.22	2.42
DE	r4, <i>p</i> = 15	1.06	0.80	1.07	815	-0.04	1.48

The MAPE and MdAPE values in Table 4 indicate that the most accurate forecasts were obtained for PL and DE, while the least accurate were for GB. MPE allows us to assess the forecast bias. Positive values of MPE indicate underprediction, while its negative values indicate overprediction. Note that for PL and DE the forecast bias was significantly smaller than for GB and FR. The same can be said about the forecasts dispersion measured by IqrAPE and StdPE.

5.3. Results Comparison with Other Models

We compare the performances of RF with other models including statistical models and ML models. The comparative models are outlined below (see [30,39] for further description). Their hyperparameters were selected on the data from 2012–2014 in grid search procedures using a variant of cross-validation or selected by experimentation (this applies to models with a large number of hyperparameters, which are difficult to optimize using standard methods due to the huge search space).

- Naive—naive model: $\hat{\mathbf{z}}_{i^*} = \mathbf{z}_{i^*-7}$;
- ARIMA—Auto-Regressive Integrated Moving Average model;
- ETS—Exponential Smoothing model;
- Prophet—a modular additive regression model with nonlinear trend and seasonal components;
- MLP—perceptron with a single hidden layer and sigmoid nonlinearities;
- SVM—linear epsilon insensitive Support Vector Machine (ε-SVM);
- ANFIS—Adaptive Neuro-Fuzzy Inference System;
- LSTM—Long Short-Term Memory;
- FNM—Fuzzy Neighborhood Model;
- N-WE—Nadaraya–Watson Estimator;
- GRNN—General Regression NN;
- RandNN—ensemble of 100 Randomized NNs;
- MTGNN—Graph NN for multivariate time series forecasting;
- ES-adRNNe—ensemble of five hybrid and hierarchical models combining ETS and dilated RNN with attention mechanism.

We also compare our model with competitive tree-based ensembles: XGBoost [40] and LightGBM [41]. Their predictors include both calendar data (hour of the day, day of the week, quarter, month, year, day of the year, day of the month and week of the year) and historical demands (demands at hour t of 21 consecutive days preceding the forecasted day).

Table 5 compares MAPE for RF and the baseline models. From this table, you can clearly see the better performance of RF compared to the other models. RF outperformed all other models in terms of accuracy for PL, GB and DE. For FR it took third place after RandNN and SVM. To confirm the results, a pairwise one-sided Giacomini-White test was performed (GM test) [42]. Its results, *p*-values, are shown in Figure 9 (we used GW test implementation from [43]). Small *p*-values, below 0.05 (green color), indicate that the model on the X-axis significantly outperforms in terms of accuracy the model on the Y-axis.

Model	PL	GB	FR	DE
RF (proposed)	1.05	2.36	1.67	1.06
Naive	2.96	4.80	5.53	3.13
ARIMA	2.31	3.50	3.00	2.31
ETS	2.14	3.19	2.79	2.10
Prophet	2.63	4.00	4.71	3.23
MLP	1.39	2.84	1.93	1.58
SVM	1.32	2.54	1.63	1.38
ANFIS	1.64	2.80	2.12	2.48
LSTM	1.57	2.92	1.81	1.57
FNM	1.21	3.02	1.84	1.30
N-WE	1.19	3.12	1.86	1.29
GRNN	1.22	3.01	1.81	1.30
RandNN	1.14	2.51	1.57	1.18
MTGNN	1.95	3.44	2.59	2.04
ES-adRNNe	1.22	2.45	1.73	1.15
XGBoost	1.67	3.36	2.51	1.74
LightGBM	1.67	3.39	2.54	1.79

Table 5. MAPE comparison between RF and baseline models (lowest errors in bold).



Figure 9. Results of the Giacomini-White test for the proposed and baseline models (black color is for *p*-values larger than 0.10).

5.4. Discussion

In our previous work [19], we used a local training mode with input patterns r2, which express daily profiles. Our current research revealed that input patterns incorporating weekly seasonality (r4) or both daily and weekly seasonality (r6) combined with global training with extended inputs improve the results (note that in Tables 1 and 2, patterns r4 and r6 provide lower errors than r2 for all countries). Calendar data used as additional input in the global extended training helps the trees to properly partition the input space and thus approximate the target function with greater accuracy. It does not take place without costs: the complexity of the model increases due to learning on all data, not just the selected data as in local training.

Table 5 and Figure 9 show that the proposed RF outperforms classical statistical models (ARIMA and ETS), modern statistical model (Prophet), classical ML models (MLP, SVM, ANFIS, GRNN), modern ML models (LSTM, RandNN), similarity-based models (FNM, N-WE) as well as state-of-the-art ML models (MTGNN, ES-adRNNe) and boosted regression trees (XGBoost, LightGBM). The last two models, as well as the proposed RF model, also used calendar variables, even in larger numbers. However, in contrast to these models, our model uses specific time series preprocessing, which may be a decisive advantage. Our model also outperforms ES-adRNNe, which is a very sophisticated and complex model developed especially for STLF [39]. To increase its predictive power, it is equipped with a new type of RNN cell with delayed connections and inherent attention, it processes time series adaptively, learning their representation and it learns in the cross-learning mode (i.e., it learns from many time series in the same time). It reveals its strength with a large amount of data, numerous and long time series. In our case, this condition was not met-there were only four, relatively short series available for training the models. In this case, the proposed RF model, which learns from individual series, generated more accurate forecasts than ES-adRNNe.

It is worth noting that RF has few hyperparameters to tune, which makes it easy to optimize (compare with DNNs with many hyperparameters). The results of our experiments confirmed that the number of trees in the forest should be as large as possible and the mimimum leaf size can be set to one. Therefore, the key hyperparameter remains the number of predictors to sample in the nodes. Its optimal values significantly differed from the recommended default values. Our attempt to increase the performance of the RF model through alternative methods of selecting predictors for split failed. Neither the curvature test nor the interaction test, which take into account the relationship between predictors and response when splitting data in nodes, improves the results significantly over the default CART method.

In our study, we used both continuous and categorical predictors. Such a mix causes many difficulties for other models such as ARIMA, ETS, NNs, SVM, LSTM and others. Categorical variables cannot be processed by these models directly. Such predictors must be converted into numerical data, so as to maintain the relationship between their values. The method of this conversion can be treated as an additional hyperparameter. RF has no problems with categorical variables, which is its big advantage. Moreover, RF can deal easily with any number of additional exogenous predictors and does not need to unify predictor ranges, which is often necessary for other models. RF can even deal with raw data because the predictors are not processed by the tree in any way, just selected in nodes, to construct a specific decision model (flowchart-like structure).

Regression tree provides fast one-pass training which does not need to repeatedly refer to the data. In contrast, NNs, which use a variant of the gradient descent optimization algorithm with multiple scanning of a dataset, are more time-consuming to train. Additionally due to the number of hyperparameters, they are also much more expensive to optimize in terms of time than RF. The training of a tree does not provide an optimal result because decisions about data split are made in nodes using a local rather than a global criterion, i.e., the split made may not be optimal from the point of view of the final result. However, the NN learning process also does not lead to optimal results due to sensitivity to the starting point and tendency to fall into the traps of the local minimum. Note that non-optimality of the trees is mitigated by their aggregation in the forest. Aggregation also smoothes out functions modeled by individual trees and reduces their variance. The learning process of RF can be easily paralleled because the individual trees learn independently.

One useful feature of RF is that it enables the generalization error to be estimated using out-of-bag (OOB) patterns, i.e., training patterns not selected for the bootstrap sample (approximately one third of the training patterns are left out in each bootstrap sample). Therefore, the time-consuming cross-validation that is widely used in other models for estimating the generalization error is not needed. Using OOB patterns, the generalization error can be estimated during one training session, along the way. Although for forecasting problems, where training patterns should precede validation to prevent data leakage, the OOB approach as well as standard cross-validation may be questionable. For this reason, we did not use the OOB approach in this study. Instead, we applied a different strategy. We chose a set of 100 validation patterns from 2014 and for each of them we trained RF on training patterns preceding the validation patterns.

A valuable feature of RF is its built-in mechanism for predictor selection. In each node, the predictor which improves the split-criterion the most is selected. The splitting criterion favors informative predictors over noisy ones, and can completely disregard irrelevant ones. Thus, in RFs an additional feature selection procedure is unnecessary. Based on the internal mechanism for selecting predictors, the predictor importance or strength can be estimated. The importance measure attributed to the splitting predictor is the accumulated improvement this predictor gives to the split-criterion at each split in each tree. RF also offers another method of estimating the predictor importance based on the OOB patterns [27]. When the tree is grown, the OOB patterns are passed down the tree, and the prediction error is recorded. Then the values for the given predictor are randomly permuted across the OOB samples, and the error is computed again. The importance measure is defined as the increase in error. This measure is computed for every tree, then averaged over the entire forest and divided by the standard deviation over the entire forest. Such a measure is presented in Figure 8. Note that information about the predictor importance is a key factor, which helps to improve the interpretability of the model and can be used for feature selection for other models.

Model interpretability is an emerging area in ML that aims to make the model more transparent and strengthen confidence in its results. This topic is also explored in electricity demand forecasting literature [44]. In [45], it was shown that the predictor importance is related to the model sensitivity to inputs and also to the method of importance estimation. An LSTM-based model, which is proposed in [45], is equipped with a built-in mechanism based on a mixture attention technique for temporal importance estimation of predictors. In the experimental study, this model demonstrated higher sensitivity to inputs than treebased models (RF and XGBoost) which showed very low sensitivity on the predictors except one, which strongly dominated (the authors used built-in functions of scikit-learn to calculate the predictor importance for tree-based models). In our study, the predictor importance is more diverse (see Figure 8), which may result from the fact that our trees are very deep and thus involve a great number of predictors. Note that tree-based models enhance interpretability not only through built-in mechanisms of predictor importance estimation, which show predictive power of individual predictors, but also through their flowchart-like tree structure. They can be interpreted simply by plotting a tree and observing how the splits are made and what is the arrangement of the leaves. It should be noted, however, that while following the path that a single tree takes to make a decision is trivial and self-explanatory, following the paths of hundreds of trees in the ensemble is much more difficult. To facilitate this, in [46], model compression methods were proposed that transform a tree ensemble into a single tree that approximates the same decision function.

In this study we use a standard RF formulation which is a MISO model producing point forecasts. Thus for prediction of 24 values of the daily curve of electricity demand, we need to train 24 RF models. In [32], we proposed a multivariate regression tree for STLF, which produces a vector as an output, representing the 24 predicted values. Using such MIMO trees as ensemble members simplifies and speeds up the forecasting process. A promising extension of the RF in the direction of probabilistic forecasting can be achieved using a quantile regression forest, which can infer the full conditional distribution of the response variable for high-dimensional predictor variables [47].

6. Conclusions

ML ensemble models are state-of-the-art for forecasting problems. They dominate the most recent literature on forecasting. Among them, tree-based ensembles have a solid theoretical basis and have been thoroughly researched in a huge number of papers. Their predictive power has been confirmed in numerous forecasting competitions [24].

In this study, we propose a RF model for a challenging STLF problem with multiple seasonality, nonlinear trend, and varying variance in time series. Unlike DNNs, RF is simple and transparent, it does not require a complex, deep architecture, equipped with additional sophisticated mechanisms to deal with complex time series. The greatest advantages of RF as a forecasting model are: small number of tuning hyperparameters (we show that only one is key), fast training and optimization, ability to deal with multiple exogenous predictors of different types, and built-in mechanism for selecting predictors and estimating their importance.

As with any predictive model, the performance of RF depends significantly on data preprocessing and proper organization of the training process. In the simulation study, we show how the results of RF depend on the training method, definition of input variables and hyperparameters. Based on the results, we recommend the best method of predictor definition (r4 and r6) and training mode (global extended) for STLF. Comparing the performances of RF and baseline models including statistical and ML ones, we showed that RF can successfully compete with them, providing the most accurate forecasts.

In our future work, we plan to extend RF with random data projection (to further smooth the estimator and provide an additional source of diversity) and use RF for probabilistic forecasting. A quantile regression forest [47] is a promising tool for the latter task.

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Abbreviations

The following abbreviations are used in this manuscript:

ANFIS	Adaptive Neuro-Fuzzy Inference System
APE	Absolute Percentage Error
ARIMA	Auto-Regressive Integrated Moving Average
CART	Classification And Regression Trees
CNN	Convolutional Neural Network
DE	Germany
DNN	Deep Neural Network
ES-adRNNe	ensemble of five hybrid and hierarchical models combining ES
	and dilated RNN with attention mechanism
ETS	Exponential Smoothing
FR	France
FNM	Fuzzy Neighborhood Model
GB	Great Britain
GBRT	Gradient Boosting Regression Tree
GRNN	General Regression Neural Network
IqrAPE	Interquartile Range of Absolute Percentage Error
LightGBM	Light Gradient Boosting Machine
LSTM	Long Short Term Memory Neural Network
MAPE	Mean Absolute Percentage Error
MdAPE	Median of Absolute Percentage Error
MIMO	Multiple Input Multiple Output
MISO	Multiple Input Single Output
ML	Machine Learning
MLP	Multilayer Perceptron
MPE	Mean Percentage Error
MTGNN	Graph Neural Network for Multivariate Time series forecasting
N-WE	Nadaraya–Watson Estimator
NN	Neural Network
PE	Percentage Error
PL	Poland
RandNN	Randomized Neural Network
RF	Random Forest
RMSE	Root Mean Square Error
RNN	Recurrent Neural Network
StdPE	Standard Deviation of Percentage Error
SVM	Support Vector Machine
STLF	Short-Term Load Forecasting
WGTB	Warm-start Gradient Tree Boosting
XGBoost	eXtreme Gradient Boosting

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