'More Bikes': Experiments in Univariate Regression

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1 Task description

The assignment is to predict the number of available bikes at 75 rental stations in three hours' time for a period of three months from November 2014, i.e., a supervised univariate regression problem. It is divided into three sub-tasks, which differ in the information that is available. For sub-task 1, the number of available bikes at each of the 75 stations for the month of October 2014 is provided (section 4). This sub-task may be approached by building a separate model for each station or a single model for all 75 stations. For sub-task 2, a set of linear models that were trained on the number of available bikes at each of a separate set of 200 stations are provided (section 5). Finally, for sub-task 3, both sources of information may be used. Additionally, the number of available bikes at the first ten stations between June 2012 and October 2014 is provided for analysis (see fig. 3).

The predictions are evaluated by the mean absolute error (MAE) between the predicted and true numbers of available bikes. Hereafter, the MAE is referred to as the 'score'.

$$MAE = \frac{1}{n} \sum_{i=1}^{n} |y_i - \hat{y}_i|$$
 (1)

The evaluation data was not available to participants, but the score achieved on a held-out test set is reported on the task leaderboard. I give the score achieved on this test set by the best estimators for each model class and sub-task alongside the mean scores achieved on cross-validation folds of the data provided for sub-task 1 in TODO.

This report begins with a description of the general approach taken to the assignment in section 2. Section 3 presents a preliminary analysis of the data. Finally, sections 4 and 5 detail the results of the methods applied to each of the sub-tasks.²

2 Approach

I used the scikit-learn Python package (Pedregosa et al. 2011) throughout this report. In each case, preprocessing and feature selection were performed by *estimators* that implemented the *transformer* interface, prediction was performed by estimators that implemented the *predictor* interface, and estimators were composed into Pipeline objects over which hyperparameter search was performed (Buitinck et al. 2013, pp. 4–9).

Generally, standard k-fold cross-validation is disfavoured for time-series data due to the inherent correlation between successive folds (Bergmeir et al. 2018). Instead, I used nested time-series cross-validation³ with ten folds to evaluate the models, which is illustrated in fig. 1. I determined the best

¹https://www.kaggle.com/c/morebikes2023/leaderboard

²The code that produced these results is available at \mathbf{Q} tslwn/more-bikes.

³See sklearn.model_selection.TimeSeriesSplit.

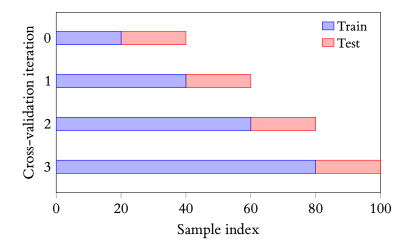


Figure 1: A visualization of the nested time-series cross-validation behaviour that I used, after the documentation for sklearn.model_selection.TimeSeriesSplit.

estimator for each model class by grid search⁴ with the mean absolute error as the scoring function, following the task description (section 1). Finally, I assessed the statistical significances of the differences between the mean scores of the best estimators by paired t-tests of the scores on the cross-validation folds, I describe these methods in more detail in sections 4 and 5.

3 Data analysis

The data provided for sub-task 1 is recorded at hourly intervals with n = 54385 instances across the 75 stations. A summary of its features and the distributional characteristics of the non-temporal quantitative features are given in tables 1 and 2. This analysis revealed that many of the features, including the target variable bikes, are missing for n = 73 instances, which were henceforth excluded. Additionally, the 'profile' features, i.e., the features derived from the numbers of available bikes at preceding times, are not defined for the first week of instances at each station. Hence, they are missing for approximately $\frac{1}{4}$ of the instances. The meteorological features are constant for all stations at a given timestamp. Finally, some features have a natural range – in particular, the number of available bikes at a given station is bounded by zero and the number of docks at that station (section 3.2).

3.1 Feature selection

Intuitively, features that have zero variance are not informative for regression analysis, so I automatically excluded them.⁵ In the case of sub-task 1, the available data is limited to the month of October 2014, so these included the month and year. The 'station' features (table 1) are constant for all instances at a given station, so were likewise excluded in this case. Finally, the precipitation feature is zero for all instances. Correlations between features are undesirable in regression analysis (Alin 2010), so I also sought to identify and exclude redundant features. To determine these, I computed the Pearson correlation coefficients between pairs of quantitative features (fig. 2), which yielded the following:

- bikes_avg_full and bikes_avg_short are perfectly correlated (r = 1.00).
- bikes_3h_diff_avg_full and bikes_3h_diff_avg_short are perfectly correlated (r = 1.00).
- wind_speed_max and wind_speed_avg are highly correlated (r = 0.96).

 $^{^4}$ See sklearn.model_selection.GridSearchCV and sklearn.model_selection.HalvingGridSearchCV.

 $^{^5\}mathrm{See}$ sklearn.feature_selection.VarianceThreshold.

Category	Feature	Data type	Kind	
	station	int	ordinal	
C	latitude	float		
Station	longitude	float		
	docks	int		
	timestamp	int		
	year	int		
	month	int		
Tomporel	day	int		
Temporal	hour	int		
	weekday	str	ordinal	
	weekhour	int		
	is_holiday	bool	categorical	
	wind_speed_max	float		
	wind_speed_avg	float		
	wind_direction	float		
Meteorological	temperature	float		
	humidity	float		
	pressure	float		
	precipitation	float		
Bikes	bikes	int		
	bikes_avg_full	float		
	bikes_avg_short	float		
	bikes_3h	int		
	bikes_3h_diff_avg_full	float		
	<pre>bikes_3h_diff_avg_short</pre>	float		

Table 1: A summary of the features and the target variable (bikes). Except where indicated, the features are quantitative. The features have been renamed to be easier to read.

Feature	n/a	μ	σ^2
wind_speed_avg	73	4.69	21
wind_direction	365	170.23	7,553.8
temperature	73	21.71	10.7
humidity	73	65.94	279.7
pressure	73	1,002.26	1,808.27
bikes	73	7.35	43.15
bikes_avg_full	12,264	7.31	35.82
bikes_avg_short	12,264	7.31	35.82
bikes_3h	292	7.34	43.17
bikes_3h_diff_avg_full	12,483	$4.1 \cdot 10^{-3}$	22.4
bikes_3h_diff_avg_short	12,483	$4.1 \cdot 10^{-3}$	22.4

Table 2: The numbers of missing values (n/a) and distributional characteristics of the non-temporal quantitative features and the target variable (bikes). Features that have zero variance for the first case of sub-task 1 are excluded (section 3.1).

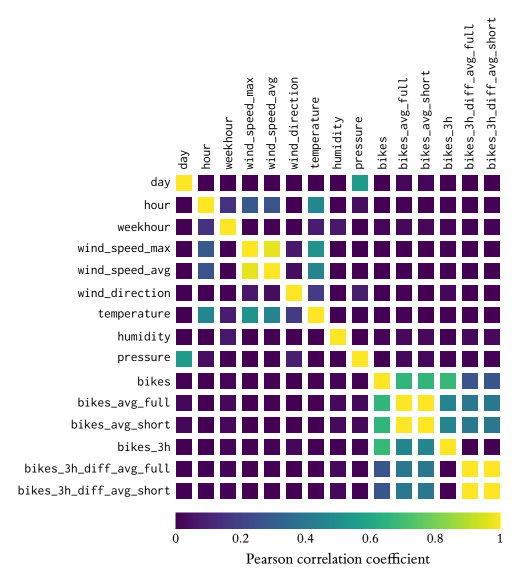


Figure 2: The Pearson correlation coefficients between pairs of the quantitative features and the target variable (bikes). The ordering follows table 1. Features that have zero variance for the first case of sub-task 1 are excluded (section 3.1). The timestamp feature is also excluded because it is naturally correlated with the other temporal features.

Hence, for sub-task 1, the second of each of these pairs of features was also excluded. For sub-task 2, the 'profile' features were retained because they are inputs to the pre-trained linear models (table 6).

3.2 Feature engineering

For sub-task 1, the bounds on the number of available bikes at a given station were enforced by predicting the *fraction* of bikes, i.e., the number of bikes divided by the number of docks at the station. I implemented this by extending the TransformedTargetRegressor meta-estimator to permit data-dependent transforms.⁶ Another possible transformation of the target variable would have been to predict the *change* in the number of available bikes, but I did not explore this option.

Initially, I investigated introducing derived temporal features, e.g., by discretizing the hour of the day into 'day' and 'night' intervals. However, as shown in fig. 3, the average fraction of available bikes throughout the day differs substantially between the training data and the data from the first ten

⁶See sklearn.compose.TransformedTargetRegressor.

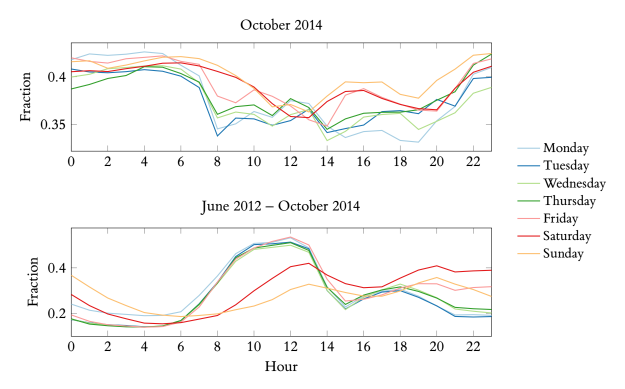


Figure 3: The average fraction of available bikes at each hour of the day, separated by the day of the week. In the top chart, the data is limited to the month of October 2014 for the 75 stations to predict. In the bottom chart, the data from the first ten stations is also included (section 1).

stations (section 1). Hence, I considered that a feature of this kind would be unlikely to generalize well to the test data. Additionally, having elected to investigate tree models (section 4), which partition the spaces of quantitative features (Flach 2012, p. 155), discretization may be irrelevant.

4 Sub-task 1

4.1 Model classes

Gradient-boosted decision trees are a popular choice of model class for time-series forecasting problems (Bojer and Meldgaard 2021). In particular, scikit-learn provides an optimized implementation of gradient-boosted decision trees inspired by LightGBM (Ke et al. 2017). An advantage of this model class is that it supports missing values, which are evident in the data (table 2). For both cases of sub-task 1, I elected to compare the performance of individual decision trees and gradient-boosted decision trees. As a baseline, I predicted the arithmetic mean of the fraction of available bikes.

The best estimator for each model class was determined by grid search with ten-fold cross-validation. With a separate model for each of the 75 stations, I found that it was too computationally expensive to perform grid search over a wide range of hyperparameters. Hence, I performed grid search for the case of a single model for all stations, then chose the values of the best estimator for the case of a separate model for each station. I fixed the scoring criteria to the mean absolute error, as per section 2. The resultant parameter values are listed in table 3.

4.2 Results

The mean scores achieved by the best estimator for each model class over the cross-validation folds and stations are listed in table 4. With a separate model for each station, there is greater variance in the mean score because there are 75 times as many cross-validation folds with commensurately fewer

Parameter	Values
criterion	absolute_error
<pre>max_depth max_leaf_nodes min_samples_leaf</pre>	None, 1, 2, 5, 10, 20, 50 None, 7, 15, 31, 63 1, 2, 5, 10, 20, 50, 100

(a) Decision tree

Parameter	Values
loss	absolute_error
scoring	neg_mean_absolute_error
l2_regularization	0.1, <u>0.2</u> , 0.5, 1.0
learning_rate	0.02, 0.05, <u>0.1</u> , 0.2, 0.5
max_depth	None, 1, 2, 5, 10, 20, <u>50</u> , 100
max_iter	20, 50, <u>100</u> , 200, 500
max_leaf_nodes	None, 7, <u>15</u> , 31, 63
min_samples_leaf	1, <u>2</u> , 5, 10, 20, 50, 100

(b) Gradient-boosted decision tree

Table 3: The parameter grids over which I performed hyperparameter search for sub-task 1. Except where stated, the default values were used, and the parameters of the best estimator are underlined. For a description of the parameters and their default values, see sklearn.tree.DecisionTreeRegressor and sklearn.ensemble.HistGradientBoostingRegressor.

instances per fold. Intuitively, the baseline achieved a lower score with a separate model for each station. For both cases of sub-task 1, both model classes achieved a lower score than the baseline.

To determine whether the differences between the mean scores achieved by the best estimators were statistically significant, I performed paired t-tests of the scores on the cross-validation folds. The null hypothesis was that the mean scores of the best estimators were equal, i.e., that the differences between them were due to chance. I did not compare the scores between estimators for the different cases of this sub-task because the samples are not paired. With a separate model for each station, both decision trees and gradient-boosted decision trees performed significantly better than the baseline for more than half of the stations (table 5a). However, gradient-boosted decision trees only performed significantly better than decision trees for seven stations. With a single model for all stations, the results were more decisive – the best gradient-boosted decision tree performed significantly better than the best decision tree (table 5b).

Model	μ	σ^2	Model	μ	σ^2
Baseline	4.43	3.82	Baseline	5.45	0.06
Decision tree	3.53	3.52	Decision tree	3.51	0.08
Gradient-boosted decision tree	3.37	2.61	Gradient-boosted decision tree	2.60	0.08

(a) Separate model for each station

(b) Single model for all stations

Table 4: The mean scores achieved by the model classes for each case of sub-task 1. In table 4a, μ is the mean of the scores for the 75 stations and ten cross-validation folds. In table 4b, μ is the mean of the scores for the ten cross-validation folds only.

Model 1	Model 2	<i>t</i> > 0	<i>p</i> < 0.05
Baseline	Decision tree	70	39
Baseline	Gradient-boosted decision tree	70	44
Decision tree	Gradient-boosted decision tree	44	7

(a) Separate models for each station. The numbers of positive t-statistics and significant p-values of the mean scores of the best estimators for each model class on the ten cross-validation folds for each of the 75 stations.

Model 1	Model 2	t-statistic	<i>p</i> -value
Baseline	Decision tree		1.70×10^{-9}
Baseline	Gradient-boosted decision tree	31.6	1.55×10^{-10}
Decision tree	Gradient-boosted decision tree	8.2	1.83×10^{-5}

⁽b) Single model for all stations. The t-statistics and p-values for paired t-tests of the mean scores of the best estimators for each model class on the ten cross-validation folds for all 75 stations.

Table 5: The results of paired *t*-tests of the scores achieved by the best estimators for each model class for the cases of sub-task 1. A positive *t*-statistic indicates that 'Model 2' achieved a lower mean score than 'Model 1'.

5 Sub-task 2

5.1 Model classes

The available information for sub-task 2 is a set of linear models that were trained on the number of available bikes at each of a separate set of 200 stations (section 1). For each station, there are six models generated by the rlm function from the R package MASS.⁷ Each of the models uses a different set of features, which are listed in table 6. Unlike the gradient-boosted decision tree implementation in scikit-learn, the pre-trained models do not support missing values, so I imputed these⁸ with the mean value of the feature over the training data. I used stacked generalization⁹ to combine the predictions of the models (Wolpert 1992). This model class is a meta-estimator that trains a second-level regressor on the predictions of a set of first-level regressors (the pre-trained models).

For this sub-task, I also excluded zero-variance and highly correlated features but retained the 'profile' features (section 3.1). However, the full and short features are perfectly correlated, and the latter were excluded for sub-task 1. Therefore, I first assessed the performance of the different kinds of pre-trained models. A box plot of the mean scores achieved on the data for all 75 stations is shown in fig. 4. This shows that individual rlm_short and rlm_short_temp models generally achieved lower scores. However, with the default second-level regressor (ridge regression), the results of stacked generalization with each kind of model were very similar, so I elected to include all the models in the following analysis. For the second-level regressor, I opted to compare ridge regression, decision trees, and gradient-boosted decision trees, as in section 4.1. The best second-level regressor was likewise determined by grid search with ten-fold cross-validation (section 2).

5.2 Results

References

Alin, Aylin (2010). "Multicollinearity". In: WIREs Computational Statistics 2.3, pp. 370-374.

 $^{^7\}mathrm{See}$ https://www.rdocumentation.org/packages/MASS/versions/7.3-54/topics/rlm.

⁸See sklearn.impute.SimpleImputer.

⁹See sklearn.ensemble.StackingRegressor.

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rlm_short_full_temp	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	
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Table 6: The features used by the different kinds of linear models for sub-task 2. The features are listed in table 1 and follow the same ordering.

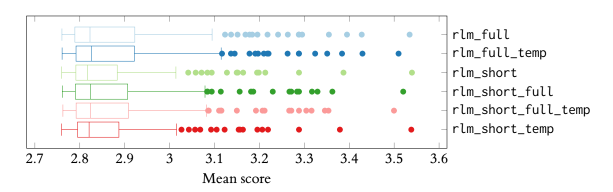


Figure 4: A box plot of the mean scores achieved by the different kinds of models for sub-task 2 over the ten cross-validation folds of the data from the 75 stations for the month of October 2014, i.e., the same data as for the second case of sub-task 1. The kinds of models are listed in table 6.

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