

Kaggle Challenge: Predicting Rossmann Store Sales

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1 Abstract

Our project attempted to apply linear regression and various machine learning techniques to a real-world problem of predicting drug store sales. Rossmann, Germany's second-largest drug store chain, has provided past sales information of 1115 Rossmann stores located across Germany. We preprocessed, feature-engineered the data, and examined 4 different statistical / machine learning analysis for forecasting sales of each store: multivariate linear regression with various feature selection methods, random forest, and gradient boosting. Then, we compared the methods' predictive power by computing Root Mean Square Percentage Error (RMSPE). We found that Gradient Boosting model performed the best with a RMSPE score of 0.13234 on the test data set.

2 Introduction

The objective of this project is to predict 6 weeks of daily sales for 1115 Rossmann stores located across Germany using the data which was provided by Rossmann through Kaggle. The motivation of this project is the following: by reliably predicting daily sales, store managers may decrease operational costs, and create effective staff schedules (ex. by assigning more staffs during busy days) ("Rossmann Store Sales"). Also, we would like to identify which type of techniques be effective in a real-world sales prediction task. Before building any

regression or machine learning models for the prediction, we attempted to define which features are significant to determine daily sales of stores. By performing exploratory data analysis, we found how some features such as promotion and competitor distance affect the sales.

To confirm our assumption and apply various techniques for the prediction, we first explored several similar efforts that made for sales prediction task in the past in order to set the basis of our study.

Linear Regression: The professor at Western Illinois University suggests multiple linear regression with various feature selection methods and multicollinearity diagnostics to be a popular technique in sales forecasting. This model was applied for predicting the number of subscribers that a cable station can expect, which is closely related to our task (Dehkordi-Vakil). We decided to apply this technique along with partial t-test and stepwise regression to select significant features.

Random Forest: Random Forest is a tree-based ensemble method where all trees depend on a collection of random variables. (Breiman). Breiman, UC Berkeley professor, suggests that Random Forest algorithm is actually from a classification viewpoint but the approach is applicable to regression as well (Breiman). Therefore, we consider the Random Forest is also applicable to our sales prediction regression task. Nils Adriansson and Ingrid Mattsson in their thesis applies the Ran-

dom Forest into a GDP forecast task, showing its power in time-series forecasting (Adriansson). However, for the train dataset, they had only 58 observations compared to our 1,000,000. To make our computation faster, we chose to use “H2O” library which enables parallelized computation although Adriansson used R’s Random Forest. We assume this algorithm would outperform linear regression because Breiman states that the Random Forest does not need to assume any distribution for variables (Breiman).

Gradient Boosting: We also found an useful article by Jay Gu, PhD in Machine Learning from Carnegie Mellon University, on gradient boosting regression. In his article, he asserted that, for predictive analytics, the gradient boosted tree model (GBM) is one of the most effective machine learning models. As he did, we have also witnessed a good number of winning solutions for various kaggle competitions that contain boosted tree models as a crucial component. We got the basic framework from Tianqi Chen’s presentation (Chen). In his “Bike Sharing classification prediction”, we witnessed that Gu reduces his computation time efficiently by dividing out certain features to separately train models. He later combines the predictions of these models to calculate weighted sums for binary classification (Gu). We consider that his separate training of models might force him to also tune hyperparameters separately which would worsen his runtime efficiency. For our regression task, since we have to directly predict the number of sales, we decided to not separately train Gradient Boosting models, but increase our time efficiency by using R’s ‘XGBoost’ library to perform parallelized computation which enables faster cross validations and more hyperparameter tuning.

tary information for each store (41088 lines because there are 41088 stores). “train.csv” and “store.csv” both contain Store id that we can use to join the train and store data sets. Finally, “test.csv” has 41088 lines of daily sales of different stores but the value of Sales column is omitted. We are expected to predict the value of Sales column for the test set (“test.csv”) by using the training set (“train.csv”) and store data (“store.csv”) (“Rossmann Store Sales”).

2.1 Description of Datasets

Data sets are given by Rossmann through Kaggle. There are three files: “test.csv”, “train.csv”, and “store.csv”. “train.csv” contains 1017209 observed daily sales of different stores from 2013 to 2015. “store.csv” contains supplemen-

The following table describes fields in “test.csv” and “training.csv”:

Store	the unique id for each store (positive integer ranging from 1 to 1115)
DayOfWeek	the day of week (positive integer ranging from 1 to 7 where 1, 2, ..., 7 correspond to Monday, Tuesday, ..., Sunday)
Date	the date (string of format YYYYMMDD)
Sales	the total amount of revenue for this given day (positive number between 0 and 41551). This value is what needs to be predicted, so only exists in “training.csv”.
Customers	the number of customers on the given day. This also only exists in “training.csv”.
Open	an indicator of whether the store was open on the given day (0 if closed, 1 if open)
Promo	an indicator of whether the store was running a promotion on the given day (0 if not running, 1 if running)
StateHoliday	an indicator of whether the given day was a state holiday (0 if not a state holiday, a if public holiday, b if eastern holiday, c if christmas)
SchoolHoliday	an indicator of whether the store was affected by a nearby school holiday on the given day (0 if not affected, 1 if affected)

The following table describes fields in “store.csv”:

Store	the unique id for each store (positive integer ranging from 1 to 1115)
StoreType	the store’s model (4 types: a, b, c, d)
Assortment	the store’s assortment level (a = basic, b = extra, c = extended)
Promo2	an indicator of whether the store was participating in a promotion that runs over a period of time (0 if not participating, 1 if participating)
Promo2SinceWeek	the approximate week of year when the store started to participate in promo2 (positive integer ranging from 1 to 50, and NA’s if no promo2)
Promo2SinceYear	the approximate year of time when the store started to participate in promo2 (positive integer ranging from 2009 to 2015, and NA’s if no promo2)
PromoInterval	the months when the promo2 started (comma

3 Methods

We first worked on preprocessing the raw data given by Rossmann.

3.1 Preprocessing

We first merged “train.csv” and “store.csv” by “Store” because we can predict daily sales better with more data related to the sales. We also merged “test.csv” and “store.csv” by “Store”. Then, We removed “Customers” field from the training data because it does not exist in the test data so we cannot use “Customers” to predict “Sales”.

For “CompetitionDistance” with NA values, we replaced NA with 1000000. Since 1000000 is much more bigger than any other distances, replacing NA with the number simulated an effect of having no competitor. In the similar way, we filled missing values for “CompetitionOpenSinceYear” and “CompetitionOpenSinceMonth”.

Similarly, for “Promo2SinceYear” with NA values, we replaced NA with 1990 to simulate the effect of having no promo2.

We splitted “Date” field into three fields: “year”, “month”, and “day” to better account for the effects each components of date have.

We computed average sales for each store to create a new field “Average.Sales”. This variable seemed to be an indicator of how well a store will perform in future. This makes sense because a store with a strong past performance is likely to perform well in future.

We noticed the feature “CompetitionDistance” is highly skewed to right. For the sake of linear regression, we took log transformation on the feature “CompetitionDistance” to create a new field “LogCompetitionDistance” to make sure that the relationship between explanatory variables and dependent variable is approximately linear.

Our R codes for preprocessing are in “Section A: Pre-processing” of the supplementary file “final-project.R”.

3.2 EDA (Exploratory Data Analysis)

To check any distinctive relationships between variables before applications of prediction models, we drew a pearson correlation coefficient plot using R corrplot library. In the plot, we wit-

nessed several interesting behaviors. We decided to explore deeper on the features: “DayOfWeek” and “CompetitionDistance”.

1. “DayOfWeek” and “Sales” are negatively correlated. This implies that, as “DayOfWeek” approaches to Sunday, sales would decrease because drugstores would mostly close on Sunday.
2. “LogCompetitionDistance” and “Sales” are negatively correlated. In other words, lower distance to the competitor implies slightly higher sales. We assume this could occur because stores with a close distance to the competitor are mostly located in crowded regions such as a downtown city with higher sales overall. To check this pattern, we drew a scatterplot of average sales by competitiondistance.

Our R codes for EDA are in “Section B: EDA” of the supplementary file “final-project.R”.

3.3 5-fold Cross Validation

To generalize our predictions to limit problems of overfitting on the training set, we decided to randomly partition the original training observations into 5 equal sized subsamples. The training set is comprised of 4 equal sized subsamples and the rest as the validation set. For the purpose of reproducibility and comparison of the predictive powers of different models, we set the seed (with a value of 42). Our general process would be as the following: 1. Repeat cross-validation process 5 times with each of the 5 subsamples used exactly once as the validation set. 2. Compute the average of 5 different validation error rate. 3. Predict sales on the test set using the model. 4. Submit on Kaggle website to get the test error rate. 5. Compare the average validation error rate and the test error rate among different models.

Our R codes for 5-fold cross validation are in “Section D: Cross Validation” of the supplementary file “final-project.R”.

3.4 Error Metric

As the metrics that Kaggle uses to compute test error is the Root Mean Squared Percentage Error (RMSPE), we used the same metrics to compute our validation error. It's the following:

$$RMSPE = \sqrt{\frac{1}{n} \sum \left(\frac{y_i - \hat{y}_i}{y_i} \right)^2}$$

where y_i is a store's sales on a single day, \hat{y}_{i} is the daily sales prediction for the store on the day, and n is the number of (store, date) pairs in the data set.

Our R codes used to compute error rates are in "Section C: Utility Functions" of the supplementary file "final-project.R".

3.5 Benchmark

We started off by setting a benchmark to obtain a baseline for the prediction. Then, we took 3 different approaches to outperform the benchmark and former model.

Our benchmark is simply the average sales of each store. Precisely, we predict daily sales of a store for a day by the average daily sales the store had in the training data set. This method does not account for many factors (whether the day was holiday, whether the store was running a promotion on the day, etc), so we expect its error rate to have a higher error rate than any other methods we applied.

Our R codes for benchmark are in "Section E: Benchmark" of the supplementary file "final-project.R".

3.6 Linear Regression

Linear regression is an approach for modeling the relationship between dependent variable Y (in our case, "Sales") and explanatory variables X . To apply linear regression on the data, we decided to use R's *lm* function. *lm* is an implementation of multivariate linear regression. If we supply $Y \sim X$ as the input, *lm* builds and returns a model that can be used to predict Y using X . Under the hood, it basically solves the normal equation $\beta = (X^T X)^{-1} X^T Y$.

We first built linear model with all available features (DayOfWeek, Open, Promo, StateHoliday, SchoolHoliday, StoreType, Assortment, CompetitionOpenSinceMonth, CompetitionOpenSinceYear, Promo2, Promo2SinceYear, Average.Sales, LogCompetitionDistance, CompetitionDistance, day, month, year) using the training set. Then, we applied the linear model on validation set.

Our R codes for linear regression are in "Section F: Linear Model" of the supplementary file "final-project.R".

3.6.1 Feature Selection

Then, we did feature selection to reduce complexity of linear model. Before working on feature select, we took a look at the resulting *lm* with R function *summary(lm)* to get some general idea of how important each variable is. All features except CompetitionDistance, CompetitionOpenSinceMonth, CompetitionOpenSinceYear, Promo2, Promo2SinceYear were significant with p-value less than 0.05.

3.6.2 Feature Selection-Backward Elimination

For feature selection, we first applied backward elimination by repeatedly removing features with the highest p-value until no feature has p-value higher than 0.05. As the result, we obtained DayOfWeek, Open, Promo, StateHoliday, SchoolHoliday, StoreType, Assortment, Average.Sales, LogCompetitionDistance, day, month, year. Then, we built linear model with the selected variables using the training set and applied linear regression with the model on the validation set.

Our R codes for linear regression with backward elimination are in "Section G: Variable Selection-Backward Elimination" of the supplementary file "final-project.R".

3.6.3 Feature Selection-Akaike Information Criterion

Another approach we had for feature selection was using Akaike information criterion (AIC). AIC provides a measure of relative quality of models with the equation $AIC = 2k - 2\ln(L)$ where k is the number of estimated parameters in the model and L is the value of likelihood function. We performed AIC by using R function `stepAIC`.

Our R codes for linear regression with AIC are in “Section H: Variable Selection-AIC” of the supplementary file “final-project.R”.

3.7 Random Forest

Random Forest is an ensemble learning method that constructs decision tree using training set. By aggregating many decision trees, Random Forest can make predictions while avoiding overfitting because it uses an ensemble of regression trees over different samples of the data. Here is our Random Forest Regression steps:

1. Construct random forest trees (number of trees is defined by hyper parameter)
2. Classify the incoming data into the decision tree node
3. Calculate the mean value for each node for prediction

In particular, we used H2O's RandomForest package to carry out the training and prediction. Then, we optimized parameters to improve on our prediction model. H2O is the open source math engine for huge dataset (TODO). Since H2O package enables us to compute Random Forest Regression algorithm in parallel distribution, we decided to use the package for faster computation compared to R's Random Forest package.

To find the best hyperparameters, we tried different values of hyperparameter and tested the results on validation set using for loops.

Our R codes for random forest are in “Section I: Random Forest” of the supplementary file “final-project.R”.

3.8 Gradient Boosting

The similarity between Random Forest and Gradient Boosting is that both generate decision trees. However, Random forest uses bootstrapping method for training while Gradient Boosting builds decision tree over the residuals. In other words, Gradient Boosting Tree generates the decision tree as it optimizes on a proper loss function (ρ). This refers that we can establish our own loss function for optimization, while we couldn't do in Random Forest. We set our loss function to be our error metric, RMPSE. Our decision trees here are regression trees. Each regression tree divides the data point at each non-leaf node according to a constraint on a feature. The leaf node value of a data point decides its score, and a data point is predicted by averaging the scores of the leaf node it belongs to.

In particular, we used R's xgboost package. XGBoost also known as Extreme Gradient Boosting is a powerful supervised learning algorithm (ρ). This package enables an automatic parallel computation on a single machine. For our regression task, we put “reg:linear” as a parameter. Our training objective here is :

at most d times, since we might have to try up to d values for X_j before finding a solution

Our R codes for gradient boosting are in “Section J: Gradient Boosting” of the supplementary file “final-project.R”.

4 Supplementary Methods

For anyone who would like to reproduce results in this paper, we provided datasets and R codes we used in tar file “supplementary-methods.tar.gz”.

To untar this tar file, please run the following command: “tar xvfz somefilename.tar.gz”.

This tar file should contain “final-project.R”, “data/store.csv”, “data/test.csv”,

and "data/train.csv".

"final-project.R" contains all R codes used in this project, and it is referenced many time in "Methods" section of this paper. All files under folder "data/" are described in "Description of Datasets" subsection in "Introduction" section of this paper.

5 Results

5.1 Benchmark

Our benchmark is the average sales of each store. The error rate for benchmark was 0.106209 on validation set and 0.25789 on the test set (when submitted to Kaggle website). This is our baseline model. Our goal is to improve our prediction by outperforming the benchmark by large degree at the end.

5.2 Linear Model

The error rates of our different linear models on validation and test sets are the following:

TODO blah blahblahblah features blach-blashblash TODO <http://cseweb.ucsd.edu/jm-cauley/cse255/reports/fa15/022.pdf>

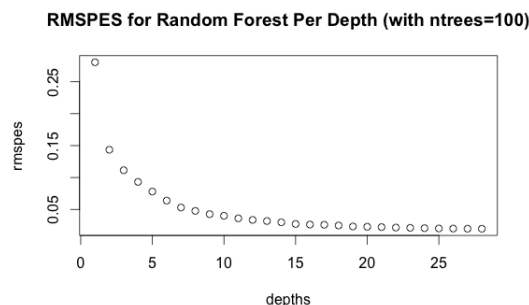
	Validation	Test
Linear Regression (Full Model)	0.09101	0.21036
Linear Regression (Backward Elimination)	0.09108	0.20988
Linear Regression (AIC)	0.091006	0.21039

Through variance selection, the following features CompetitionDistance, CompetitionOpenSinceMonth, CompetitionOpenSinceYear, Promo2, Promo2SinceYear were insignificant so were removed.

5.3 Random Forest

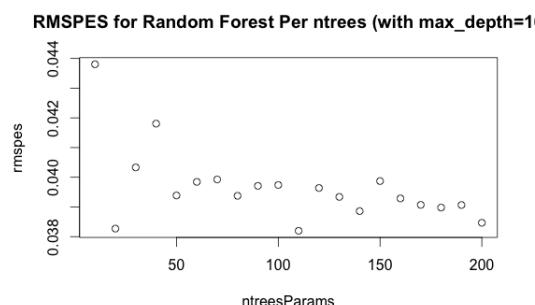
The major parater we tune for Random Forest is the depth and number of trees. We used 5

fold cross validation to compute average RM-SPE while varying the depth and number of trees. The plot for varying depth is shown below:



As the plot illustrates, we see that RMSPE does not change much after depth reaches 20. Also, we confirm that the test error rate

The plot for varying ntrees is shown below:



We see that unlike depth, ntrees does not have much impact on RMSPE (TODO).

As the plot illustrates, we could see that RM-SPE does not change much after the number of tree reaches TODO. Also, we could confirm that the test error rate does not increase even if the number of trees reaches as big as TODO. This is because Random Forest tree utilizes random amount of the data for training. In the algorithm, the training set for a tree is drawn by sampling with replacement.

Hence, we chose 30 as depth, 100 as ntrees to run random forest. The error rate for random forest (depth=30, ntrees=100) was 0.020639 for validation set and 0.14411 on test set.

5.4 Gradient Boosting

Before comparing the results of gradient boosting with other algorithms we performed, let's

explore on the results of gradient boosting itself only. Although we already feature engineered our data set, we decided to also see how our test error rate changes when we exclude stores with 0 sales. As we did in previous methodologies, we tried a range of certain parameters to maximize its prediction accuracy. Since the major issue of all machine learning tasks is to prevent overfitting on training data, we also used 5-fold cross validation to generalize. We computed the average error rates of 5-fold as a parameter changes, and graphed them to choose the best parameter. There are actually more parameters to tune in Gradient Boosting. There are three major parameters subject to be tuned: the step size of each boosting step (learning rate), maximum depth, and the max number of iterations. For our task, we decided to use the default number of trees and max number of iterations, but tune the step size. We might check how the different number of iterations affect the performance, but we checked it did not affect much but exponentially increase the runtime. We decided to set it as 300, and try different step sizes.

The effects of tuning the step size are visualized in the following graph:

As the error rate shows, Gradient Boosting model gives us the most predictive power by far. We were able to achieve test error rate (RMPSE) of 0.13234.

6 Supplementary Methods

7 Discussion

To visualize the overall performance of our different models, we organized the resulting error rates into a simple table.

The error rates of our different models on validation and test sets are the following:

	Validation	Test
Benchmark	0.106209	0.25789
Linear Regression (Full Model)	0.09101	0.21036
Linear Regression (Backward Elimination)	0.09108	0.20988
Linear Regression (AIC)	0.091006	0.21039
Random Forest	0.020639	0.14411.
Gradient Boosting	0.01734	0.13234

Although we readily outperformed our benchmark, as we expected, linear regression model did not perform very well because of the non-linearity of the data. However, we checked the sharp increase from linear regression to Random Forest model. we confirmed that tree based models can approximate functions with any “non-linear shape”, whereas linear models can produce functions with a “linear shape” with respect to a chosen set of features.

There were TODO features removed by feature selection method. We think TODO

As a result, we were able to achieve TODO RMPSE using gradient boosting to predict Rossmann drugstore sales without using customer number information. Although boosting performed very well for this dataset, we witnessed it's harder to tune parameters and took longer time to train the data compared to Random Forest. While the number of tree is only usually tuned to train Random Forest, Gradient Boosting model requires several parameters to be tuned. In addition, tuning such parameters and comparing corresponding performances requires much more time than computing Random Forest algorithm. Also, Random Forest does not easily overfit. Often, many statisticians say Random Forests does not overfit. The testing performance of Random Forests does not decrease (due to overfitting) as the number of trees increases (Breiman). However, Boosting algorithm is more robust to overfitting because it relies on every single data point and tries to find optimal linear combination of regression trees given the train dataset. Thus, it requires proper tuning to avoid it. As shown in the table, it took more than 5 hours

to train Gradient Boosting model, and compute different error rates according to different parameters over 5-fold cross validation. We could have potentially increased the size of the number of tree and get better precision, at the risk of overfitting and longer training time which is a tradeoff between time and accuracy.

In conclusion, we can check that our well-tuned gradient boosting model outperformed Random Forest.

8 References Cited

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9 Author Contributions