Diabetes 130 US Hospitals for Years 1999-2008: a Machine Learning Approach to Predict Readmission

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Introduction

In this project, we aim to improve the prediction on whether a patient will be readmitted to the hospital within 30 days or not by using a diabetes dataset. Previously the diabetes readmission problem was partially investigated by simple mean comparisons (Ostiling et al., 2017) or multivariate logistic regression (Strack et al., 2014), which had a hidden assumption of no multicollinearity among the independent variables, and therefore requires rigorous feature selections. For the current project, we incorporated 10 machine learning models following feature exploration and data preprocessing. After hypertuning, we were able to better predict the possibility of 30-day readmission rate, and to determine the driving features. Critically, we systematically compared undersampling versus undersampling methods to address the problem of imbalanced class. Our results suggest 1) Advanced algorithms performed better than the simple method in predicting the 30-day diabetes readmission rates. Among them Cat Boosting Classifier had the best scores, outperforming the existing models using a similar approach. 2) Oversampling and undersampling do not produce the same effects for all models. Our results show that oversampling improves the performance of Gradient Boosting Classifier, but is harmful to tree-based models like Random Forest. 3) Hypertuning improves the performances of all learners except for the fully-connected neural network. 4) The novel features we engineered became the dominant features for prediction. These results suggest that advanced machine learning classifiers, relative to simple linear models, are more useful for predicting 30-day readmission rate in diabetic patients based on their profiles. Importantly, we should use proper feature engineering and preprocessing skills, as well as appropriate sampling methods for different models.

Problem Statement

Several challenges are worth mentioning. First, some patients occurred more than once in the dataset, leading to a concern of the i.i.d. assumption. We quantitatively tried to address why we decided to keep the repeats. Second, some features contain many missing values, so we evaluated each of them to decide whether we would create

a new category or to drop them. Some important features contain up to hundreds of distinct values, and to avoid too many binary features when using one-hot encoding, we either used domain knowledge to reduce the categories or engineered some features that would incorporate more information. Finally, we tried to tackle the issues regarding the imbalance between two classes. Besides different sampling methods, it is also critical to find good metrics for model evaluation other than accuracy. As we have more information on the negative group and more negative group samples in the testing set, it is likely that the models would produce lots of negative predictions and few positive predictions, so high accuracy doesn't indicate the model is learning. Therefore, we chose AUC score to capture the trade-off between specificity and sensitivity. However, it is still possible that the AUC is high due to low positive predictions in general, so we further included the Harmonic Mean F1 score to get a measure of the trade-off between precision and recall, where precision is calculated by the F1 score of probability of being in negative group, and the recall is calculated by the probability of being in positive group.

$$F_1 = 2 * \frac{precision * recall}{precision + recall}$$

Methods

We first did feature exploration to remove redundant features and deal with missing data. More than 75% of the patients have only one encounter record (**Figure 1**), we decided not to remove the repeated records, as we are unsure how the repeated visits were sampled. The feature "Medical_specialty" has 50% of the missing data, but we did not remove it because this feature is very important in predicting the readmission rate. We removed non-related and redundant features with too many missing data (**Table 2**).

We then did feature engineering such as integer encoding and one-hot encoding. First, we regrouped some features with too many distinct values into several categories, such as the primary, secondary and third diagnosis (Strack et al., 2014). Second, we introduced a novel method of reducing the 3x9 one-hot features by creating only the binary feature per disease group, and encoded the values based on whether this disease has occurred in any of the diagnosis (**Table 3**). Finally, we splitted our data into a training set (70%), a validation set (15%), and a test set (15%), followed by rescaling some numerical features and combining them with binary features. We also preprocessed the dataset before training the models. As the target class is unbalanced (about 15% of our training data readmitted within 30 days), we thus experimented both oversampling and undersampling to balance our dataset. We then trained 10 different machine learning algorithms with both oversampling and undersampling to preprocess the dataset and hyperparameter tuning to find the best hypertuned model and dominant

features to predict the 30-day readmission rate: Neural Network with "lbfgs" optimizer, Neural Network with "adam" in Pytorch, Stochastic Gradient Descent, Random Forest, Gradient Boosting, AdaBoosted, CatBoosted, Naïve Bayes, Logistic Regression and Decision Tree.

Experiments

Experimental setup

We conducted experiments for feature selection, preprocessing, different learning algorithms and hyperparameter tuning for each learning algorithm. Given the imbalance dataset, we implemented both undersampling and oversampling for the training data using all the models. We have 88 features in total after one-hot encoding, feature selection and feature engineering. After initial model exploration, hyperparameter tuning was conducted for the best six models (Stochastic Gradient Descent, Random Forest, Gradient Boosting, Adaboost Classifier, Cat boosting Classifier and Neural Network with "Ibfgs" optimizer) using their default setting.

Experimental results

We first compared undersampling and oversampling results using each model. We identified that the tree-based models, especially the random forest, performed very poorly in the validation data when using oversampling method, and hyperparameter tuning couldn't improve its performance. This is because oversampling the positive group data (readmitted <30 days) led to overfitting in the tree models (**left bottom panel in Figure 2**, f1 harmonics score closes to 1.0 in the training data but closes to 0.0 in the validation data). When using undersampling, the tree-based models showed tremendous improvement, increasing the h1 harmonics score around 0.4 (**right bottom panel in Figure 2**). The undersampling did not indicate lower performances in other models by comparing the AUC and the harmonic f1 score with the oversampling, even though 90% of the data were dropped (**Figure 2**). The Cat Boosting classifier performed the best in the validation data, and the neural network using "lbfgs" optimizer, logistic regression and Random Forest classifier fell closely behind (**right panel in Figure 2**).

We then conducted hyperparameter tuning for the six best models, as it was too computationally expensive to run all ten models. All of the models showed improvements after hyperparameter tuning, except for the Neural Network with "lbfgs" optimizer, where the scores stay the same. The Cat Boosting Classifier performed the best among the four models (**Figure 3**). **Figure 4** showed the learning curve of the Cat Boosting Classifier, where the training score continued to decrease and the validation score continued to increase with more samples. This indicated that the Cat Boosting Classifier is robust and did not encounter overfitting problems.

Figure 5 showed the features sorted by dominance used to predict the 30-day readmission rate in the Cat Boosting Classifier. The top five dominant features are number of inpatients, discharge disposition id, discharge disposition id, age and number of diagnostics, which provided more information compared to previous studies without considering all of the features (Strack et al., 2014). Finally, we evaluated the final model fits among the best four hypertuned models Catboosting, Gradient Boosting Classifier, Random Forest and Neural Network with "lbfgs" optimizer using the test datasets (**Figure 6**). The ROC curve and the performance score (i.e. AUC and f1 harmonics score) confirmed that the Cat Boosting Classifier performed best in the testing sets, even though these four models did not show very obvious differences.

Discussion

In this project, our best performance model trained to predict the patients 30-day readmission rate was the Cat Boosting Classifier. We were also able to determine the driving features used to achieve the performance. We can conclude that feature exploration and data preprocessing are very important before applying machine learning algorithms. However, we did not compare our model with a reduced model with fewer essential features, as we included all the features we considered to be important. In future work, we could conduct more experiments after feature selection, such as re-train our model and compare the results using the group with and without the feature "medical_specialty", which has over 50% missing value. Moreover, as we already identified the value of novel feature engineering on the 9 disease groups, we should compare models using the raw features (27 one-hot features) versus models only using the 9 features. If the results are promising, it could shed new light on clinical practice, as well as health care data mining.

All the code could be found on Github following https://github.com/91104311/Predict-Hospital-Readmission-for-Patients-with-Diabetes-

References

Strack B, DeShazo J, et al. Impact of HbA1c Measurement on Hospital Readmission Rates: Analysis of 70,000 Clinical Database Patient Records. *BioMed Research International*. 2014. doi: 10.1155/2014/781670

Ostling S, Wyckoff J, Ciarkowski SL, et al. The relationship between diabetes mellitus and 30-day readmission rates. *Clin Diabetes Endocrinol*. 2017;3:3. Published 2017 Mar 22. doi:10.1186/s40842-016-0040-x

Table 1: Missing Values

| Feature | Action | Feature | Action |
|------------------------|--|--------------------|-------------------------------------|
| Weight, Payer_code | Remove whole features (96%, 40% missing) | Race | Group Nan into a new category (2%) |
| Diag_1, Diag_2, Diag_3 | Remove Rows with Nan (Less than 1%) | Medical_Speciality | Group Nan into a new category (50%) |

Table2: Remove Uninformative Features¶

| Feature | Discarding Reasons | |
|--|---|--|
| Encounter_id, Patient_nbr | Not Related to our target (readmitted) | |
| Glipizide-metformin, Glimepiride-pioglitazone, Metformin-rosiglitazone, Metformin-pioglitazone, Acetohexamide, Tolbutamide, Troglitazone, Tolazamide | Few sample (less than 20 patients) use these drugs | |
| Repaglinide, Nateglinide, Chlorpropamide, Acarbose, Miglitol, Glyburide-metformin | Over 98% percent of patients didn't use these drugs | |

Table3: Feature Engineering

| Feature | Changes | Feature | Changes |
|---|--|--------------------------|--|
| Gender | Remove 3 rows with Invalid gender, convert to 1/0 for male and female | Age | Convert each range to the average (like [50,60) to 55) |
| Race | One-hot Encoding | Change, DiabetesMed | 0/1 Encoding |
| Max_glu_serum | 'None' => 0, 'Norm' => 1, '>200' => 2, '>300' => 3 | A1Cresult | 'None' => 0, 'Norm' => 1, '>7' => 2, '>8' => 3 |
| Metformin, Glimepiride, Glipizide, Glyburide, Pioglitazone, Rosiglitazone, Insulin | 'No' => 0, 'Down' => 1, 'Steady' => 2, 'Up' => 3 | Diag_1, Diag_2, Diag_3 | Group 800 distinct values to 9 diagnosis and then One-hot Encoding (Circulatory, Respiratory, Digestive, Diabetes, Injury, Genitourinary, Neoplasms, Musculoskeletal, Others |
| Ever diagnosed | Create a new feature to check whether a patient is diagnosed with any specific diagnosis during diag_1, diag_2 or diag_3 | Discharge_disposition_id | Removing all encounters discharge to a hospice or patient death. Decreasing distinct values from 28 to 8 by grouping similar type, then one-hot encoding |
| Admission_type_id | Just pick 3 more frequent types and group others. Then one-hot encoding | Admission_source_id | Decreasing distinct values from 17 to 6 by grouping similar type, then one-hot |
| Readmitted | 'No' & '>30' => 0, '<30' => 1 | Medical_specialty | Reduce 20 distinct values to 7 by grouping low frequency values, then one-hot encoding |

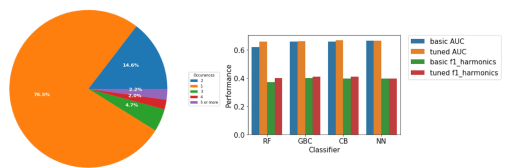


Figure 3: The effect of hyperparameter tuning on AUC and f1 harmonics scores for different models

Figure 1: Percentage of different numbers of occurrence per patient ID

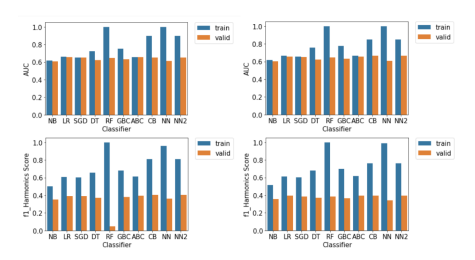


Figure 2: The AUC and f1 harmonics score for each model in the oversampling (left panel) and undersampling (right panel) on training and validation set. NN and NN2 represents Neural Network with "adam" in Pytorch and "lbfgs" optimizer, respectively.

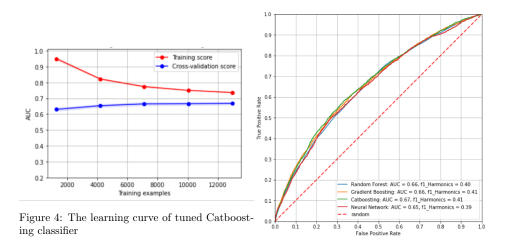


Figure 6: ROC curve for different tuned models based on testing dataset

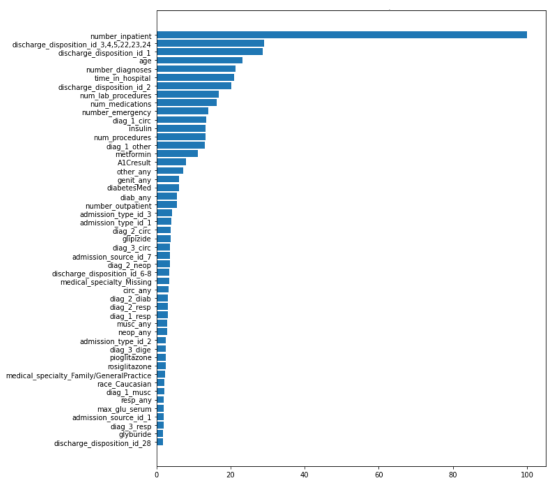


Figure 5. The relative feature importance in Cat Boosting Classifier compared to the most important feature (unit: %).