September 19, 2022

1. INTRODUCTION

Diabetes is a metabolic disease impacting 37.3 million Americans. Those affected by the disease have complications producing insulin, a chemical messenger that our body uses to store energy. Although it is uncommon, diabetics can be hospitalized for having critically low or high blood glucose levels. These hospitalizations can be life threatening and should be minimized at all costs.

In this case study, we will use a diabetes data set procured by Dr. Slater, to identify what factors most significantly result in diabetics getting readmitted to hospitals. To accomplish this, we will build a Logistic Regression model and extract its respective feature importances. It is our hope that this research can be leveraged by medical professionals to help treat hospitalized diabetics and to ensure that these patients are not readmitted in the future.

DATA UNDERSTANDING:

Data used in this case study was a diabetes.csv provided by Dr. Slater. Our diabetes.csv contained data related to hospitalized diabetic patients including columns such as: readmitted, patient_nbr, insulin, and time in hospital, Upon reviewing the contents of our data set, we saved the data into a data frame named "diabetes_data" and began pre-processing.

2. METHODS

DATA PREPROCESSING:

The first step we performed in pre-processing was reviewing our full data set. Immediately, we recognized that missing values existed in the columns of:

- 1. race 2. weight
- 3. payer_code
- 4. medical_specialty

5. diag_1 6. diag_2

7. diag_3

Given that machine learning models do not handle missing values well, we imputed them using appropriate statistical methods. Full details on how these columns were imputed can be found in the sub-header of this case study titled, "Data Imputation."

After identifying that missing values existed in our data set, we ran the command, diabetes_data_info() and noted the following details of our data frame:

The last step performed in pre-processing was calculating the percentage of missing values in our categorical columns. Exhibit 1.1 details our findings:

Our data frame contained 101,766 rows. Our data frame contained 50 columns. No null values existed in our data frame contained 13 numeric columns. Our data frame contained 37 categorical columns.

From this output we recognized that one hot encoding (OHE), would need to be performed on our categorical columns. For additional details on our OHE process, please see the sub-header of this case study titled, One Hot Encoding. The next step performed in pre-processing was running the command diabetes_data.describe() to view the summary statistics of our data frame. Output from this command showed that several columns

contained outliers. This was something we remained cognizant of throughout our analysis. Finally, to view the distributions of our categorical columns with missing values, we created count plots. Visualizing these columns was important, as it helped us determine what data imputation method was most

appropriate for our data. Output from our count plots showed that all seven of our columns with missing data contained non-normal distributions (Exhibit 1.0). Since all seven columns were of the categorical data type, we noted that imputing these columns with either the mean or median value would be appropriate.

Exhibit 1.0: Count Plot of Payer_Code

Percentage of Missing Values

39.55%

Column Name

2.23% race 96.85% weight payer_code medical_specialty 49.08% 0.02% diag_1 diag_2 0.35% diag_3 1.39%

Exhibit 1.1: Percent Missing Values

DATA IMPUTATION:

Upon reviewing our full data set and calculating the percentage of missing values that existed in our categorical columns, we proceeded to impute our missing values.

this was complete, we re-calculated the sum of missing values in our columns to validate that no data loss had occurred in our conversion process. When we considered imputing the columns: race, payer_code, medical_specialty, diag_1, diag_2, and diag_3, we tried two different approaches. One

approach was imputing these columns with the mode of each column, and the second approach was leaving the columns as is with missing values. We fit our Logistic

All the missing values in our data set were denoted by: ? . Since computers cannot impute data with special characters, we converted the question marks to "NaN". Once

Regression model on both approaches and found that our performance results were negligible. Consequently, we decided to leave the columns with missing values, as we felt this represented our data the best. The first column we chose to impute was our weight column. Given that 96% of the data in our weight column were missing, we chose to drop the column from our

data set. Next, we imputed the columns: diabetesMed, change, and readmitted with values of 0 and 1. This was done to simplify OHE as these columns had a maximum of

three classes. Please note that although the column: readmitted contains three classes, we chose to convert it to a binary variable as we are only concerned with whether a patient has been readmitted or not. Exhibit 1.2 details our conversion process of these columns: Column Name Original Classes Data Dictionary for Converted Classes

diabetesMed

Column Name

payer_code

0=No

		res	i=res		
	Change	Ch No	0=No 1=Yes		
	readmitted	NO <30 >30	0=No 1=Yes		
Exhibit 1.2: Imputation process for the columns: "diabetesMed", "change", and" readmitted"					

When viewing the shape of our data set, we recognized that if we one hot encoded all 37 of our categorical variables, that our data set would be extremely wide. As a

RE-CODING CATEGORICAL COLUMNS:

result, we decided to reduce the classes in each categorical variable by specifying a threshold for infrequent observations. Exhibit 1.3 details the thresholds that were chosen for each variable, as well as explanations as to why thresholds were chosen.

Explanation

~90% of our data falling into the top 7 classes

Selected Threshold

0.02

medical_specialty	0.03	~85% of our data falling into the top 5 classes		
max_glu_serum	0.02	~96% of our data falling into the top 2 classes		
A1Cresult	0.08	~91% of our data falling into the top 2 classes		
metformin	0.1	~98% of our data falling into the top 2 classes		
repalglinide	0.01	~99% of our data falling into the top 2 classes		
nateglinide	0.9	~99% of our data falling into the top class		
chloropropamide	0.9	~99% of our data falling into the top class		
glimepiride	0.9	~95% of our data falling into the top class		
glipizide	0.1	~97 of our data falling into the top 2 classes		
glyburide	0.09	~98 of our data falling into the top 2 classes		
pioglitazone	0.06	~98 of our data falling into the top 2 classes		
rosiglitazone	0.05	~98% of our data falling into the top 2 classes		
acarbose	0.9	~99% of our data falling into the top class		
miglitol	0.9	~99% of our data falling into the top class		
tolazamide	0.0004	~99% of our data falls into the top class		
glyburide_metformin	0.9	~99% of our data falling into the class "No"		
diag_1	0.0075	many of the columns contained values <= 0.000010		
diag_2	0.0075	many of the columns contained values <= 0.000010		
diag_3	0.0075	many of the columns contained values <= 0.000010		
Exhibit 1.3: Detailed Recoding Threshold				

Once we imputed our missing values and re-coded our categorical columns we separated our diabetes data set into two variables, one containing all our numeric columns and the other containing all our categorical columns. Next, using Pandas' get dummies function, we one hot encoded our categorical columns and joined our one hot

encoded data to our numeric columns to arrive at our final full data set.

modeling assumptions.

assumption was met.

ONE HOT ENCODING:

Please note that for modeling, we scaled our non-hot encoded data to ensure that our full data set was on the same scale. For additional modeling details, please see our sub-header below titled "Modeling."

EXPLORATORY DATA ANALYSIS: For our exploratory data analysis (EDA), we began by viewing histograms and pair plots of our data (Exhibits 1.4 & 1.5). Two takeaways from these visualizations included:

Exhibit 1.4: Encounter ID Density Plot

Exhibit 1.5: Admission Type ID Density Plot

After reviewing our histograms and pair plots, we assessed multicollinearity in our data by creating a correlation plot. Seeing that no columns had a correlation coefficient

2. Our variables were not on the same scale.

1. Many of our numeric columns exhibited non-normal distributions.

of 1.0, we chose not to remove any columns from our data set. At this point we were satisfied with our data and began evaluating if our data met our Logistic Regression

The three key assumptions of Logistic Regression models include: 1. Independent variables have a linear relationship to the log loss of the response. 2. Absence of multicollinearity.

Exhibit 1.7: Log Odds for Num Meds Feature

Exhibit 1.8: Snapshot of several pair plots generated from our data

Seeing that our log odds plots showed that our independent variables had a linear relationship to the log loss our response, we deemed that our first assumption was met.

3. Lack of outliers. To assess our first assumption, we created two log odds linear plots of our response variable versus the independent variables (Exhibit 1.6): "time_in_hospital" and "num_medications"

ASSUMPTIONS OF LOGISTIC REGRESSION MODELS:

Exhibit 1.6: Log Odds for Time in Hospital

plots were built on un-scaled data, we proceeded in our analysis assuming that this assumption was met.

To address our multicollinearity assumption, we created a correlation plot. Given that no columns had a correlation coefficient of 1.0, we proceeded assuming that this

Exhibit 1.9: Addressing our Logistic Regression outlier assumption with pair plots

The final assumption we addressed was lack of outliers. (Exhibit 1.8). As illustrated in our pair plots below, we did see that our data contained outliers. Given that the pair

3. MODEL BUILDING & RESULTS The use of Sklearn's LogisticRegression was used to model the data for this case study.

Models use: 10-fold Cross validation (Kfold), random_seed = 0, and max_iter = 50000, and scoring metric of Negative Mean Absolute Error

After preprocessing, EDA, and scaling the data, modeling was able to begin. To determine the best hyperparameters that we should use, we needed to iterate through

several of sklearn's modules. We began with utilizing GridSearchCV, but due to the shape of our data and inability to scale our CPU, GPU, and Memory for the

experimental and model_selection packages we were able to utilize HalvingRandomSearchCV to obtain good, but potentially not the best hyperparameters

needs of this project, GridSearchCV was unable to complete and we needed to try other methods of tuning hyperparameters. With the use of Skelearn's

for this model. HalvingRandomSearchCV combines the idea of HalvingSearchCV and RandomizedSearchCV. HalvingSearchCV works by modeliong all potential candidates with less data and selects half of the best performing models to add additional resources and data until a "best" model is output. RandomizedSearchCV randomly picks candidate modles from the grid to model.

We passed the below parameters into HalvingRandomSearchCV and the best model outputs were the following: **Halving Random Search CV Parameters:**

np.logspace(-3,3,7),

np.arange(0.0, 1.0, 0.1),

'solver': ['saga'], ['elasticnet'], 'penalty': [1e-9,1e-8,1e-7,1e-6,1e-5, 1e-4, 1e-3, 1e-2, 1e-1] "tol": **Best Model Output:**

"l1_ratio":

"n_jobs":

"penalty":

"solver":

"l1_ratio":

"C":

"C":

Model HalvingRandomSearchCV:

"tol": 1e-09 **ElasticNetCV with GridSearchCV Tuned Parameters:**

1

2

3

1.0

0.2

-1

'saga'

'elasticnet'

25.6763 0.00533915

0.00650811

After performing HalvingRandomSearchCV to tune the model parameters, Sklearn's cross_validate was used to validate the model and determine final performance. The results of all 10 folds are below with a MAE score of 0.431234. We modeled several different metrics, but determined that MAE was the best metric for determine the best model. Model results for the 10 fold Cross Validation are below in Exhibit 2.0. CV Run fit_time score_time estimator test_score train_score

LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.432642 -0.431231 4 5 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.427729 -0.431613 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.425806 -0.432023 6 0.00782394 7 -0.431848 0.00559735 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.426985 8 25.2208 0.008039 -0.430757 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.437991 9 0.006809 -0.431226 25.2794 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.4318 MEAN 24.7177 0.00664113 -0.431234 -0.431372 Exhibit 2.0: 10 Fold Cross Validation Model Results **Feature Importance:** After completing the Logistic Regression modeling, we wanted to determine which features were the most important in predicting readmission. These values are showin in Exhibit 2.1.

25.5363 0.00609469 LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.431463

LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.431561

LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.432544

LogisticRegression(I1_ratio=0.2, n_jobs=-1, penalty='elasticnet',random_state=0, solver='saga', tol=1e-09) -0.433821

-0.431449

-0.431384

-0.431187

-0.431002

Exhibit 2.1: Top 10 Features in Logistic Regression Model

The top 10 features in the model were:

In conclusion, after significant updates to thresholds and hyperparameters, we have determined that logistic regression does not properly model this data due to inablilty for the coefficient's to converge. Potential models that would be better for this data set would include decision trees or any sort of gradient boosting.

Attached in file CS2.ipynb