

# Diabetes Prediction

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## Summary

In this project we attempt to build a model to predict diabetes disease. We compared a decision tree model and naive bayes model and found the decision tree is stronger in this context. We used f2-score as our scoring function because detecting diabetes is the priority: a false negative could be much worse than a false positive.

In the test dataset: the decision tree model correctly detected 8283 of 10604 positive cases (recall rate is about 78%). This result does come at a fairly significant cost in terms of false positives (precision rate is about 30%) with 19650 false positives. Depending on the actual cost of false positive this may need significant improvement to be a viable screening model.

## Introduction

In Canada and the USA approximately 10% of people are living with diabetes. In Canada in 2023 approximately 3.7 million people were living with diabetes and in the USA in 2021 approximately 38.4 million people were living with diabetes. (“Snapshot of Diabetes in Canada, 2023” (2023)) In the USA it is the 8th leading cause of death. (Rios et al. (2017)) Globally an estimated 44% of people living with diabetes are undiagnosed. (Stafford et al. (2025))

In this project we try to predict diabetes disease based on common health factors. A reliable model could help to prescreen people and recommend following up with a physician for people who are at risk. Given the large number of people living with undiagnosed diabetes this could potentially have a significant positive impact on world health.

The analysis uses the American CDC Behavioural Risk Factor Surveillance System (BRFSS) 2015 Diabetes Health Indicators dataset (UCI ID 891), containing 253,680 survey responses with 21 health-related features and a binary diabetes outcome (0 = no diabetes/pre-diabetes, 1 = diabetes). (Dane and Teboul (2021))

## Methods

### Packages

This analysis was performed in Python 3.11.6 (*Python 3.11.6 Documentation* 2021-2025). Additionally, here is a list of the Python packages used within the analysis with brief explanation:

Table 1: Table of Python packages used

Package	Version	Use case	Reference
<code>numpy</code>	<a href="#">1.26.4</a>	General analysis use	<i>NumPy Documentation</i> (2008-2022)
<code>pandas</code>	<a href="#">2.1.2</a>	Data management/processing	team (2020), McKinney (2010)
<code>pandera</code>	<a href="#">0.27.0</a>	Data validation	Bantilan (2020)
<code>altair</code>	<a href="#">5.1.2</a>	Generating plots	VanderPlas et al. (2018), Satyanarayanan et al. (2017)
<code>scikit-learn</code>	<a href="#">1.3.2</a>	Model creation and evaluation	Pedregosa et al. (2011)
<code>ucimlrepo</code>	<a href="#">0.0.7</a>	Data extraction	Kelly, Longjohn, and Nottingham (2021)
<code>deepchecks</code>	<a href="#">0.18.1</a>	Data validation	Chorev et al. (2022)
<code>click</code>	<a href="#">8.3.1</a>	Script tool	Pallets (2020)

Package	Version	Use case	Reference
quarto	1.8.26	Report creation	Allaire et al. (2025)
tabulate	0.9.0	Table formatting	Astanin (2025)

## Modeling Approach

The data were split 70/30 into training and test sets with stratification on the target. Two classifiers were trained and tuned using 5-fold cross-validated grid search with **f2-score** as the scoring metric. We chose to use f2-score because it is most important to not miss true positives.

### 1. Decision Tree (class\_weight='balanced')

Hyperparameters: max\_depth: {6,8,10,12,14}, min\_samples\_leaf: {175, 200, 225, 250}

**Best parameters:** max\_depth=10, min\_samples\_leaf=225

**Best CV f2-score = 0.587**

### 2. Bernoulli Naive Bayes (with StandardScaler preprocessing)

Hyperparameters: alpha: {1e-3, 1e-2, 1e-1, 1e0, 1e1, 1e2, 1e3, 1e4}

**Best parameters:** alpha=0.001

**Best CV f2-score = 0.46**

## Results

Table 2: Scores for models.

Unnamed:		Test Accuracy	Test f2-score	Test recall	Test precision
0	Model	0.706	0.587	0.784	0.293
0	Decision Tree				
1	Naive Bayes	0.814	0.46	0.489	0.373

## EDA

### Data Summary

Table 3: Description of the training data.

Unnamed: 0	count	mean	std	min	25%	50%	75%	max
HighBP	177576.0	0.4293	0.495	0.0	0.0	0.0	1.0	1.0

Table 3: Description of the training data.

HighChol	177576.0	0.4228	0.494	0.0	0.0	0.0	1.0	1.0
CholCheck	177576.0	0.9627	0.1896	0.0	1.0	1.0	1.0	1.0
BMI	177576.0	28.385	6.5916	12.0	24.0	27.0	31.0	98.0
Smoker	177576.0	0.4433	0.4968	0.0	0.0	0.0	1.0	1.0
Stroke	177576.0	0.0407	0.1976	0.0	0.0	0.0	0.0	1.0
HeartDiseaseorAttack	177576.0	0.094	0.2919	0.0	0.0	0.0	0.0	1.0
PhysActivity	177576.0	0.7563	0.4293	0.0	1.0	1.0	1.0	1.0
Fruits	177576.0	0.6351	0.4814	0.0	0.0	1.0	1.0	1.0
Veggies	177576.0	0.8122	0.3905	0.0	1.0	1.0	1.0	1.0
HvyAlcoholConsump	177576.0	0.0563	0.2305	0.0	0.0	0.0	0.0	1.0
AnyHealthcare	177576.0	0.9513	0.2153	0.0	1.0	1.0	1.0	1.0
NoDocbcCost	177576.0	0.0845	0.2782	0.0	0.0	0.0	0.0	1.0
GenHlth	177576.0	2.5128	1.0685	1.0	2.0	2.0	3.0	5.0
MentHlth	177576.0	3.1833	7.4056	0.0	0.0	0.0	2.0	30.0
PhysHlth	177576.0	4.2542	8.7248	0.0	0.0	0.0	3.0	30.0
DiffWalk	177576.0	0.1682	0.374	0.0	0.0	0.0	0.0	1.0
Sex	177576.0	0.4408	0.4965	0.0	0.0	0.0	1.0	1.0
Age	177576.0	8.0321	3.0558	1.0	6.0	8.0	10.0	13.0
Education	177576.0	5.049	0.9863	1.0	4.0	5.0	6.0	6.0
Income	177576.0	6.0514	2.0729	1.0	5.0	7.0	8.0	8.0
diabetes	177576.0	0.1393	0.3463	0.0	0.0	0.0	0.0	1.0

Table 3 displays a numerical distribution of the dataset, where as Table 4 lists the first 5 rows transposed of the training dataset as example observations used. Both tables have been transposed from their original form for more syntactical presentation. All features of the dataset consist entirely of integer values, and by all count values being identically 177576 (the number of observations in the training dataset), there exist no null values, making the preprocessing step of the analysis relatively simple.

Table 4: First few rows of the training data as example observations.

	0	1	2	3	4
HighBP	0	0	1	0	1
HighChol	1	0	1	0	0
CholCheck	1	1	1	1	1
BMI	23	25	28	25	30
Smoker	0	0	0	0	1
Stroke	0	0	0	0	0

Table 4: First few rows of the training data as example observations.

	0	1	2	3	4
HeartDiseaseorAttack	0	0	0	0	0
PhysActivity	1	1	1	0	1
Fruits	1	1	0	1	1
Veggies	1	1	1	1	1
HvyAlcoholConsump	0	0	0	0	0
AnyHealthcare	1	1	1	1	1
NoDocbcCost	0	0	0	0	1
GenHlth	1	3	2	2	4
MentHlth	0	0	15	0	30
PhysHlth	0	30	2	0	15
DiffWalk	0	0	0	0	0
Sex	0	0	1	0	0
Age	10	12	6	8	8
Education	6	6	5	5	4
Income	8	7	6	7	4
diabetes	0	0	0	0	0

## Visualizations

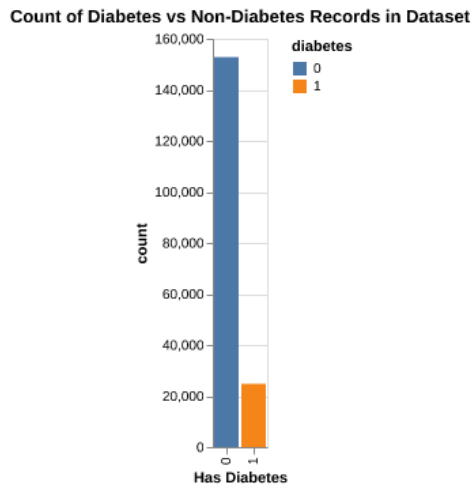


Figure 1: Frequency bar graph of the labels.

For this supervised learning problem, all observations are labelled as either 0, indicating a negative label for no diabetes, or 1, indicating a positive label for either diabetes or pre-diabetes.

From Figure 1, the class distribution is imbalanced with roughly 14% of observations labelled positive for diabetes. This motivates using F2 score as our metric; in this data, a false negative is more problematic than a false positive, thus recall rate is prioritized over precision rate. The exact equation for the F2 score used within model evaluation is as follows:

$$F_2 = \frac{5 * \text{precision} * \text{recall}}{(4 * \text{precision} + \text{recall})}$$

Continuing the EDA, Figure 2 displays the distribution of all binary features, where the labels  $f = a$ ,  $d = b$  ( $a, b \in \{0, 1\}$ ) indicate the frequency that an observation had value  $a$  in the given feature, and value  $b$  in the diabetes label.

### Bar Plots of Binary Features

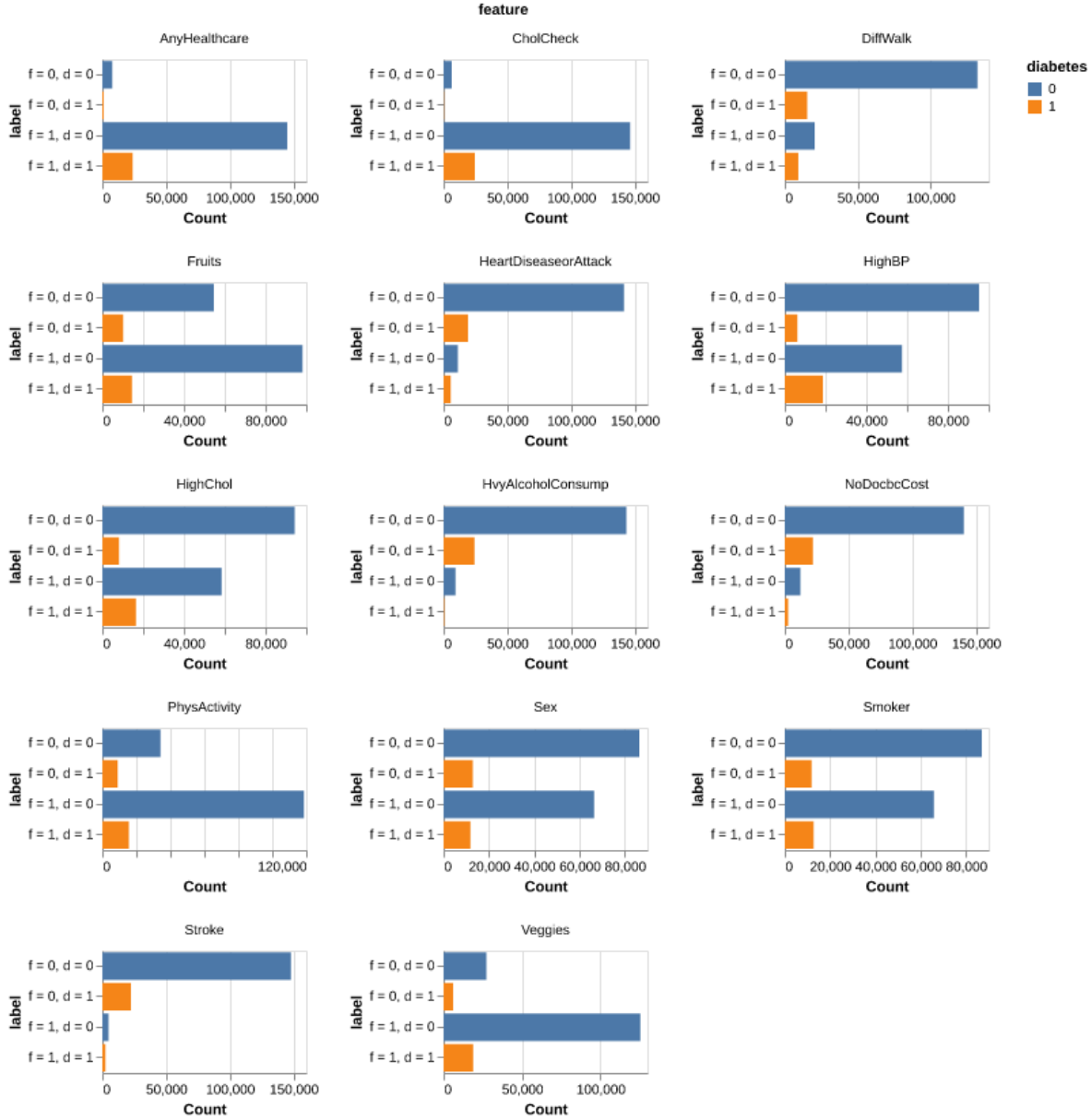


Figure 2: Frequency bar graphs of the binary features.

There is a significant variance in the distributions of the binary features; **Smoker** and **Sex** have similar number of observations where  $f = 0$  compared to  $f = 1$ , whereas a feature like **Stroke** is heavily weighted to  $f = 0$ , indicating very few observations indicate they have had a stroke before. Due to the high number of observations in the dataset, even where there are imbalances, we can expect there to be enough observations with either a positive (1) or negative (0) value

for any given binary feature that should prevent low sample size bias. `HighBP`, `HighChol`, and `Smoker` all have more observations with  $f = 0$  than  $f = 1$ , but in all of those features, the frequency of  $f = 1$ ,  $d = 1$  labels exceed that of  $f = 0$ ,  $d = 1$  labels, which indicate that we can expect that the model will indicate an association with these risks factors and a risk of diabetes.

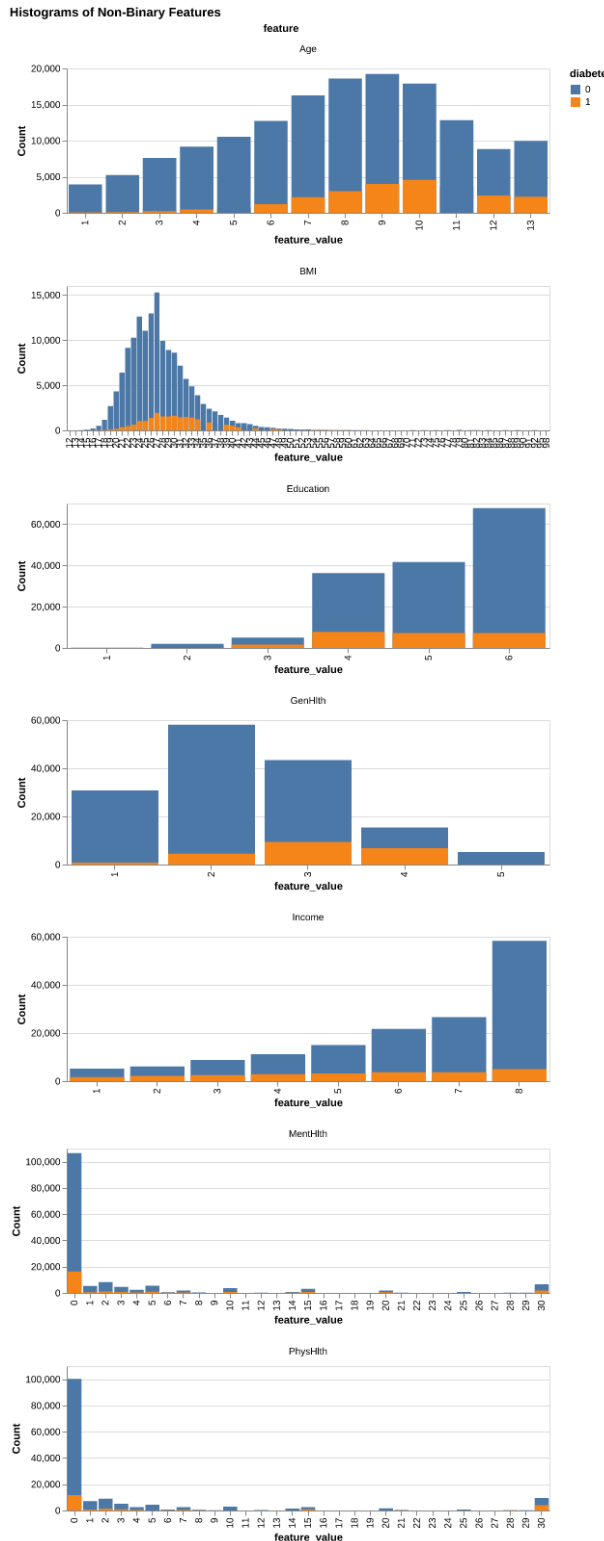


Figure 3: Histograms of the non-binary numeric features.

Boxplots for Non-Binary Features (Sample size n = 1000)

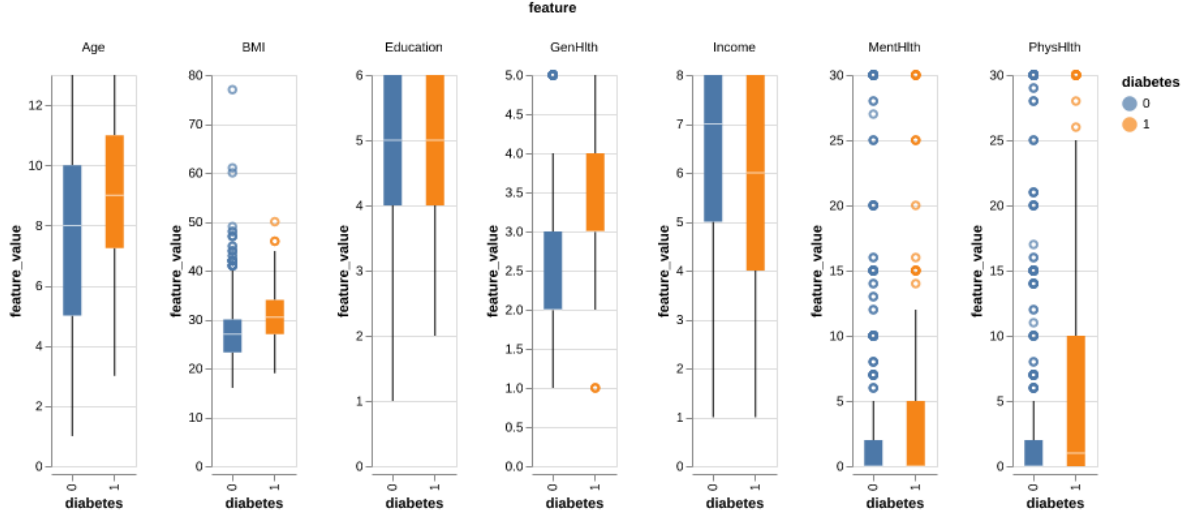


Figure 4: Boxplots of the non-binary numeric features.

Figure 4 takes a random sample of 1000 observations so that the boxplot is not overly populated and that the density of the points visibly correlates with Figure 3 more closely. From analyzing both charts, we can make the following brief observations about each non-binary feature:

- **Age** has been encoded to a 13-point scale that is somewhat normally distributed with a slight negative skew.
- **BMI** is a normally distributed feature with several extreme upper outliers that go up to 98. A BMI of 98 for someone that is 5 feet tall would result in a weight of roughly 500 pounds Keys et al. (2014), which is an outlier, but still reasonably possible that this is considered an accurate observation.
- **Education** is on a 6-point scale, while **GenHlth** is on a 5-point scale. Both of these features have the option to be preprocessed categorically, but since these are ordinal in nature, using a standard scaler on these features is a reasonable option this analysis goes with.
- **MentHlth** and **PhysHlth** have the majority of observations with a value of 0

Most importantly, with all of the varying scales these features are on, it indicates that a `StandardScaler()` can be used on all numerical features to normalize the values of each feature into identical normal distributions.

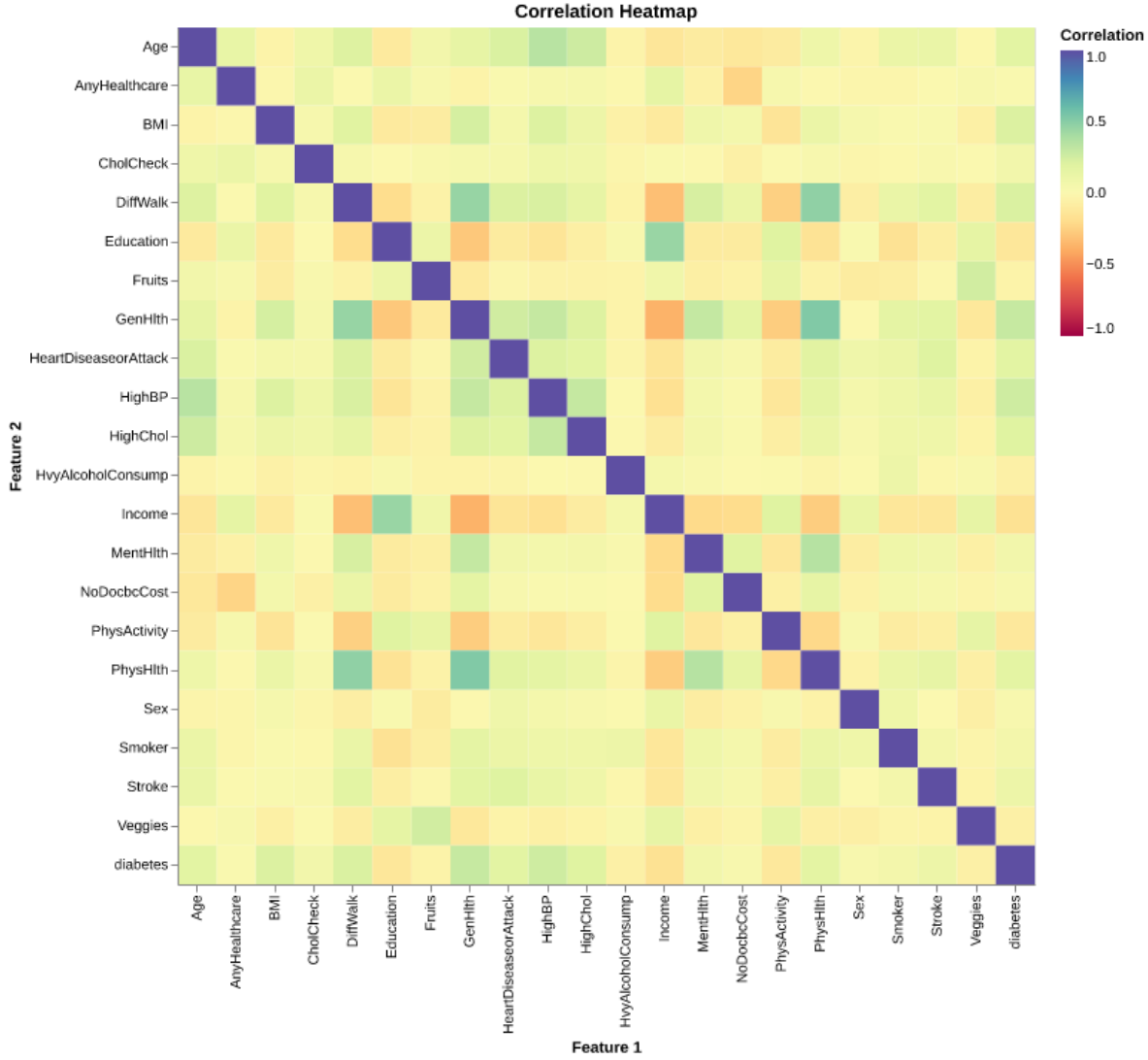


Figure 5: Feature-feature correlation plot of all features.

Lastly with Figure 5, the main observation is that the strongest non-trivial feature-feature correlation does not exceed an absolute magnituded of 0.7. There are several significantly positively correlated feature pairs (PhysHlth-GenHlth, PhysHlth-DiffWalk, Income-Education, GenHlth-DiffWalk) and several significantly negative correlated feature pairs (DiffWalk-Income, GenHlth-Income). Noticably, Income, DiffWalk, and GenHlth appear in both the most positively and most negatively correlated feature pairs and have higher overall correlations with other features compared to other features such as Sex, Smoker, Stroke, and Veggies from the overall darker shades of green and orange in the correlation heatmap. With the diabetes feature, it is most positively correlated with GenHlth and most negatively correlated

with `Income`.

## Modeling

### Model Fitting

The data were split 70/30 into training and test sets with stratification on the target. Two classifiers were trained and tuned using 5-fold cross-validated grid search with **f2-score** as the scoring metric. We chose to use f2-score because it is most important to not miss true positives.

#### 1. Decision Tree (class\_weight='balanced')

Hyperparameters: max\_depth: {6,8,10,12,14}, min\_samples\_leaf: {175, 200, 225, 250}

**Best parameters:** max\_depth=10, min\_samples\_leaf=225

**Best CV f2-score = 0.587**

#### 2. Bernoulli Naive Bayes (with StandardScaler preprocessing)

Hyperparameters: alpha: {1e-3, 1e-2, 1e-1, 1e0, 1e1, 1e2, 1e3, 1e4}

**Best parameters:** alpha=0.001

**Best CV f2-score = 0.46**

## Results

### Performance Table

The following table summarizes the performance metrics for our two models evaluated on the test set:

Table 5: Comparison of model performance metrics on the test set

Unnamed:					
	Model	Test Accuracy	Test f2-score	Test recall	Test precision
0	Decision Tree	0.706	0.587	0.784	0.293
1	Naive Bayes	0.814	0.46	0.489	0.373

As shown in Table 5, the Decision Tree model demonstrates superior performance across all evaluation metrics compared to the Naive Bayes model. The Decision Tree achieves higher accuracy, f2-score, recall, and precision scores, making it the optimal choice for our diabetes prediction task.

## Performance Visualization

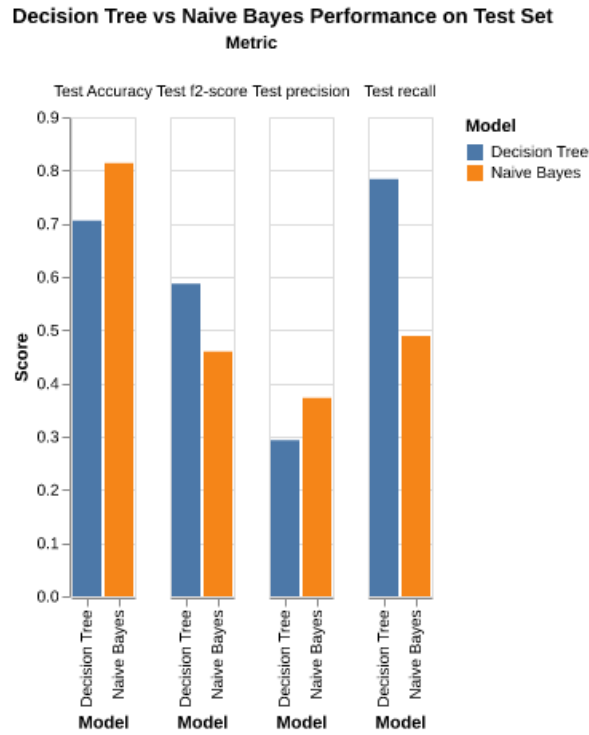


Figure 6: Comparison of model performance metrics on the test set

The performance comparison in Figure 6 demonstrates that the Decision Tree model outperforms the Naive Bayes model across all evaluation metrics.

## Confusion Matrix

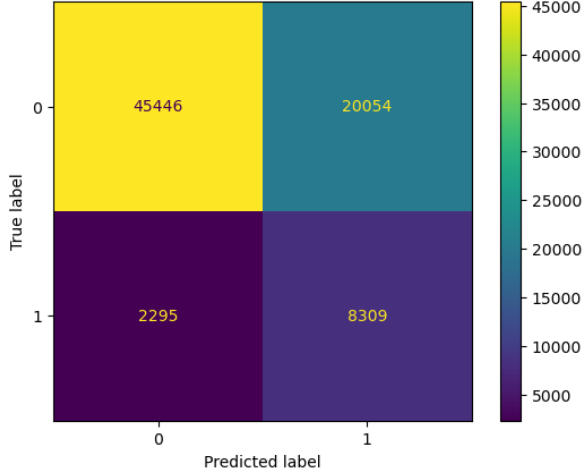


Figure 7: Confusion matrix for the best performing model (Decision Tree)

The confusion matrix in Figure 7 shows the detailed classification performance of our best model. The model correctly identified a significant portion of the test cases, demonstrating strong predictive capability.

## Discussion

The current performance of the Decision Tree model is likely already good enough to offer some benefit in the real world given the large number of people with undiagnosed diabetes. However, the recall score of 78.4% could likely be improved, and the precision score of 29.3% definitely leaves something to be desired.

We were surprised by the high rate of false positives shown in Figure 7, which might be an indication of how many non-diabetic people are at risk. Further improvements to predicting diabetes could likely be found by trying a wider variety of model types and using a wider hyperparameter search, and possibly through more feature engineering. A future study could be done to find a smaller set of the most easy to obtain features, as such a model would be more usable by the average person.

Some work is needed to determine this smaller number of easy to obtain features that doesn't significantly reduce model performance. Another question is whether a regression model could be made that predicts a person's risk as a percent chance of developing diabetes, though longitudinal data might be required for this type of prediction.

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