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Machine Learning Seminar Project Data Analysis for Diabetes Prediction Using Health Indicators

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CONTENT

1 Introduction

1.1 Dataset description

This dataset contains detailed health indicators collected from a large population and is designed to support the analysis and prediction of diabetes risk. It includes a variety of columns describing lifestyle factors and demographic data, such as high blood pressure (HighBP), high cholesterol (HighChol), body mass index (BMI), smoking and alcohol use, physical activity, diet, mental and physical health status, as well as gender, age, education level, and income.

The main variable of interest is Diabetes_binary, which indicates whether an individual has been diagnosed with diabetes (1) or not (0). This dataset can be used to develop classification models aimed at predicting diabetes risk based on known behavioral and health-related factors.

index	feature	description
Diabetes_binary	0 = no diabetes, 1 = diabetes	Diabetes status
HighBP	0 = no high BP, 1 = high BP	High blood pressure
HighChol	0 = no high cholesterol, 1 = high cholesterol	High cholesterol
CholCheck	0 = no cholesterol check in 5 years, 1 = yes	Cholesterol check
BMI	Continuous	Body Mass Index
Smoker	0 = no, 1 = yes	Smoked at least 100 cigarettes in life
Stroke	0 = no, 1 = yes	Ever had a stroke
HeartDiseaseorAttack	0 = no, 1 = yes	Heart disease or heart attack history
PhysActivity	0 = no, 1 = yes	Physical activity in past 30 days (not job-related)
Fruits	0 = no, 1 = yes	Consumes fruit 1+ times per day
Veggies	0 = no, 1 = yes	Consumes vegetables 1+ times per day
HvyAlcoholConsump	0 = no, 1 = yes	Heavy alcohol consumption
AnyHealthcare	0 = no, 1 = yes	Healthcare coverage
NoDocbcCost	0 = no, 1 = yes	Couldn't see a doctor due to cost
GenHlth	1 = excellent, 5 = poor	General health status
MentHlth	1-30 days	Poor mental health days in the past 30 days
PhysHlth	1-30 days	Physical illness/injury days in past 30 days
DiffWalk	0 = no, 1 = yes	Difficulty walking or climbing stairs
Sex	0 = female, 1 = male	Gender
Age	1 = 18-24, $9 = 60-64$, $13 = 80$ or older	Age category
Education	1-6	Education level
Income	1-8	Income scale

Table 1: List of all attributes in the dataset

1.2 Motivation of the study

The motivation of this study is based on three main questions related to understanding the risk factors for diabetes and how these factors can be used to predict the likelihood of developing the disease:

- 1. How can machine learning models be used to predict diabetes risk more accurately? This broader question relates to using advanced techniques to help create a reliable model for predicting diabetes. Machine learning can uncover relationships and patterns that may not be immediately apparent using traditional methods.
- 2. Can a subset of factors be used to accurately predict whether an individual has diabetes? An important question is whether only a few factors, such as BMI and Age, can be used to make an accurate prediction of diabetes, simplifying the prediction process without losing too much accuracy.

Through these questions, the study aims to improve our understanding of the factors influencing diabetes risk and to develop a simple yet accurate method for predicting who may be at risk.

1.3 Selection of algorithms for the dataset

For this dataset, where the target variable is Diabetes_binary (diabetes status), three well-known machine learning algorithms have been chosen: MLP (Multilayer Perceptron), Autoencoder, and I TRETI. These algorithms are suitable for this type of problem for various reasons:

• MLP (Multilayer Perceptron):

MLP is a type of neural network that uses multiple processing layers and is excellent for handling classification and regression problems. This algorithm is powerful for capturing complex relationships between different features. For our dataset, it is well-suited to model the connections between health indicators and diabetes risk, as it can effectively analyze factors such as BMI, Age, Physical_Health, Smoking, and other related variables.

Autoencoder:

An Autoencoder is a type of neural network used primarily for unsupervised learning and dimensionality reduction. It can help identify hidden patterns in the data and is useful for reducing the number of features while maintaining the underlying structure. For our dataset, it can assist in detecting underlying factors contributing to diabetes risk by learning an efficient representation of the data. Autoencoders are also useful for feature extraction, especially in cases where the data is high-dimensional.

• Naive Bayes:

Naïve Bayes is a probabilistic supervised learning algorithm based on Bayes Theorem, assuming that all features are independent of one another given the target class (in this case, diabetes status). Although this assumption of independence rarely holds true in real-world data, Naïve Bayes often performs remarkably well, especially in binary classification problems with multiple features. For our dataset, this algorithm is particularly well-suited due to its simplicity, fast training time, and effectiveness even with high-dimensional data.

2 Preprocessing

At this part of the project, we begin with the data preprocessing phase. This step is necessary to prepare the dataset for machine learning, ensuring the data is ready for analysis and modeling.

2.1 Data cleaning

2.1.1 Removing Duplicates

For this step, we first analyze if there are any duplicate rows in the dataset using the following Python code:

```
import pandas as pd

# Load the dataset

df = pd.read_csv('datasets/diabetes_binary_5050split_health_indicators_BRFSS2015.csv')

# Check for duplicate rows

duplicates = df[df.duplicated()]

# Display the result

if duplicates.empty:
    print("No duplicates found!")

else:
    print(f"Found {duplicates.shape[0]} duplicate rows.")

print(duplicates)
```

After running the code, we found that there are 1635 duplicate rows. However, since these duplicates could potentially represent individuals who share the same health indicators (such as age, BMI, or smoking status), and not necessarily represent data entry errors, we decided not to remove them. This is because, in medical datasets, it is common for multiple individuals to have identical data points across various features. Therefore, removing these rows could lead to loss of valid data.

```
Found 1635 duplicate rows.
```

As shown, there are 1635 duplicate rows, but they are not removed to ensure we preserve all relevant data for analysis.

2.1.2 Handling missing values

In this step, we aim to check whether the dataset contains any missing values. Missing data can impact the accuracy of machine learning models, so identifying and addressing it is an important part of the data preprocessing process. Below is the Python code used for this task:

Finally, we used the z-score method, which was more suitable for our dataset. This method found 1,927 outlier records, which is only 2.7% of the total dataset. Since this number was small, we thought it was reasonable to remove these records without further analysis, as we still had enough data left to work with.

```
import pandas as pd

2

# Load the dataset

df = pd.read_csv('datasets/diabetes_binary_5050split_health_indicators_BRFSS2015.csv')

# Check for missing values

missing_values = df.isnull().sum()

# Display missing values per column

print(missing_values)
```

The output after executing the script was:

```
Diabetes_binary
2 HighBP
3 HighChol
                             0
                             0
4 CholCheck
                             0
5 BMI
6 Smoker
7 Stroke
                             0
8 HeartDiseaseorAttack
                            Ω
                             0
9 PhysActivity
10 Fruits
11 Veggies
                             0
12 HvyAlcoholConsump
                             0
13 AnyHealthcare
                             0
14 NoDocbcCost
                             0
15 GenHlth
16 MentHlth
                             0
17 PhysHlth
                             Ω
                             0
18 DiffWalk
19 Sex
                             0
                             0
20 Age
21 Education
                             0
22 Income
```

As we can see, all columns in the dataset have a count of 0 missing values. This means the dataset is complete in terms of data presence, and no additional cleaning or imputation for missing values is necessary.

2.1.3 Handling outliers

3 Diabetes_binary

4 HighBP

5 HighChol

The first step we used to analyze the outlier or noise values was checking the min, max, average and mode of each attribute. From these data we can conclude that there are no noises since all the min-max fields are within the range declared on the metadata.

Next, we applied the interquartile range (IQR) method to identify outliers. However, this approach was not very effective, as it flagged 40,205 records as containing at least one outlier attribute—more than half of the dataset.

Because this didn't seem like a logical result, we decided to ignore this method and try something else.

The next test was the z-score method and this one was far more reasonable giving 1927 outlier lines. Since that would be less than 3% of the dataset, it is acceptable to remove the outlier lines completely. That left our dataset with 68,765 remainin records, a considerable number of rows.

```
1 import pandas as pd
2 import numpy as np
3 from scipy.stats import zscore
5 df = pd.read_csv('datasets/diabetes_binary_5050split_health_indicators_BRFSS2015.csv')
7 ###### Making a summary of the fields ######
8 min_vals = df.min()
9 max_vals = df.max()
10 avg_vals = df.mean()
mode_vals = df.mode().iloc[0]
summary_df = pd.DataFrame({
     'Min': min_vals,
      'Max': max_vals,
14
      'Mean': avg_vals,
15
      'Mode': mode_vals
16
17 })
18 print("###### Summary of the dataframe ######")
19 print(summary_df)
print("\n###### Checking for outliers with Interquartile Range (IQR) method ######")
Q1 = df.quantile(0.25)
Q3 = df.quantile(0.75)
_{25} IQR = Q3 - Q1
outliers = (df < (Q1 - 1.5 * IQR)) | (df > (Q3 + 1.5 * IQR))
outlier_counts = outliers.sum(axis=1)
28 print("Rows with >=1 outliers:", (outlier_counts >= 1).sum())
30
31 print("\n###### Checking for outliers with Z-Score Method (Assumes Normal Distribution) ######")
z_scores = df.select_dtypes(include='number').apply(zscore)
outliers = (z_scores.abs() > 5)
print("Outlier rows (Z-score):", outliers.any(axis=1).sum())
36
37 ###### Removing the outliers ######
rows_to_remove = outliers.any(axis=1)
39 cleaned_df = df[~rows_to_remove]
40 cleaned_df.to_csv('datasets/dataset_without_outliers.csv', index=False)
41 print("Cleaned dataset saved as 'dataset_without_outliers.csv'")
1 ###### Summary of the dataframe ######
                         Min
                              Max
                                         Mean Mode
```

0.0

1.0

0.500000

0.563458

0.0

0.0

1.0

0.0 1.0 0.525703 1.0

1.0

```
6 CholCheck
                         0.0 1.0 0.975259 1.0
7 BMI
                        12.0 98.0 29.856985 27.0
8 Smoker
                         0.0 1.0 0.475273 0.0
                               1.0
                          0.0
                                      0.062171 0.0
Stroke
                                1.0
1.0
10 HeartDiseaseorAttack 0.0
                                       0.147810
                                                   0 0
10 HeartDiseascol
11 PhysActivity 0.0
0.0
                                       0.703036
                                1.0
                                      0.611795
                                                   1.0
                         0.0 1.0 0.788774
                                                  1.0
13 Veggies

      14
      HvyAlcoholConsump
      0.0
      1.0
      0.042721
      0.0

      15
      AnyHealthcare
      0.0
      1.0
      0.954960
      1.0

16 NoDocbcCost
                         0.0 1.0 0.093914 0.0
17 GenHlth
                         1.0 5.0 2.837082 3.0
                                       3.752037
18 MentHlth
                          0.0 30.0
                                                   0.0
19 PhysHlth
                          0.0 30.0
                                       5.810417
                                                   0.0
20 DiffWalk
                          0.0
                                 1.0
                                       0.252730
                               1.0
                         0.0
                                      0.456997
21 Sex
                                                   0.0
                         1.0 13.0 8.584055 10.0
22 Age
23 Education
                         1.0 6.0 4.920953 6.0
24 Income
                         1.0 8.0 5.698311 8.0
26 ####### Checking for outliers with Interquartile Range (IQR) method #######
27 Rows with >=1 outliers: 40205
29 ###### Checking for outliers with Z-Score Method (Assumes Normal Distribution) #######
30 Outlier rows (Z-score): 1927
31 Cleaned dataset saved as 'dataset_without_outliers.csv'
```

2.2 Data integration

In this project, all the required data is already combined into a single CSV file. Therefore, no additional integration from multiple sources is necessary. The dataset is self-contained and ready for the next steps of data preprocessing.

2.3 Data transformation

In this step, we apply feature scaling to the dataset in order to standardize the features. Scaling is necessary because some machine learning algorithms, such as the ones we have chosen for this analysis, are sensitive to the scale of the data. These algorithms work better when all features have a similar range.

2.3.1 Data scaling

In this step, we apply feature scaling to the dataset in order to standardize the features. Scaling ensures that all features have a similar range. The StandardScaler from the sklearn.preprocessing library is used to perform scaling.

The following Python code demonstrates the process of scaling the features of the dataset:

```
from sklearn.preprocessing import StandardScaler
import pandas as pd

# Load the dataset
df = pd.read_csv('datasets/dataset_without_outliers.csv')

# Separate features and target
X = df.drop(columns=['Diabetes_binary']) # Features
y = df['Diabetes_binary'] # Target

# Initialize and apply the scaler
scaler = StandardScaler()
X_scaled = scaler.fit_transform(X)
```

```
# Convert scaled features back to DataFrame
%_scaled_df = pd.DataFrame(X_scaled, columns=X.columns)

# Add the target column back
%_scaled_df['Diabetes_binary'] = y.values

# Save to a new CSV file
% X_scaled_df.to_csv('datasets/diabetes_scaled.csv', index=False)

print("Scaled dataset saved as 'datasets/diabetes_scaled.csv'.")
```

This code scales all the features of the dataset using the StandardScaler, which standardizes the features by removing the mean and scaling to unit variance. The scaled features are then saved in a new CSV file called diabetes_scaled.csv.

3 Prepare for analysis

3.1 Correlation between features

Understanding how features interact within the dataset is a key step before building predictive models. One effective way to explore these interactions is through correlation analysis. This technique reveals the strength and direction of linear relationships between variables, helping us detect redundant features, highlight those most relevant to the target variable, and optimize the feature set for better model efficiency and accuracy. By refining the input space early on, we set a strong foundation for the performance of our machine learning models.

One important aspect of correlation analysis is examining how strongly each feature relates to the target variable, in this case, Diabetes_binary. Identifying variables that show a higher correlation with the target can guide us in selecting the most predictive features for our model. On the other hand, features that exhibit very weak relationships with the target may contribute little to the predictive power and can be excluded to simplify the model. Additionally, correlation analysis helps reveal whether certain features are highly related to each other. When two features show a strong correlation such as above 0.5 it may indicate redundancy. Retaining both could introduce multicollinearity, which not only complicates model interpretation but can also impair the algorithm's performance. In such cases, it's often beneficial to remove one of the correlated features to streamline the dataset and enhance model robustness.

To calculate the correlation between features, we used the corr() function from the pandas library. We visualized the results using a Heatmap. A heatmap is a graphical representation that uses color gradients to indicate the strength of relationships in a two-dimensional matrix. This is especially useful for identifying patterns and connections between features

The implementation can be seen in the code below, and the visual results are presented in Figure 1:

```
import pandas as pd
import seaborn as sns
import matplotlib.pyplot as plt

df = pd.read_csv('datasets/diabetes_binary_5050split_health_indicators_BRFSS2015.csv')

# Compute correlation matrix
corr_matrix = df.corr()

# Plot the heatmap
plt.figure(figsize=(12, 10))
sns.heatmap(corr_matrix, annot=True, fmt=".2f", cmap="coolwarm", square=True)
plt.title("Correlation Matrix Heatmap")
plt.tight_layout()
plt.savefig("report/images/diabetes_correlation_matrix.png")
plt.close()
```

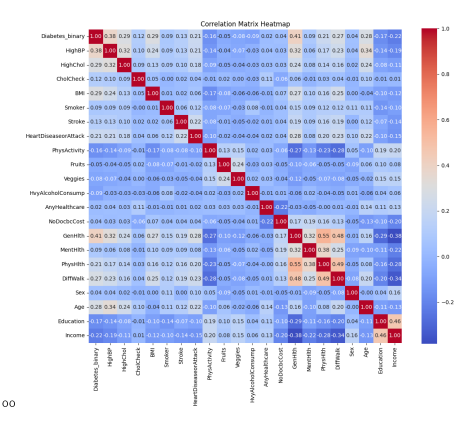


Figure 1: Correlation matrix between features.

From the heatmap visualization, we observe that some features are more strongly correlated with each other. We then filtered the feature pairs that have an absolute correlation higher than 0.45. The implementation of this filtering process in Python is shown below:

The results are displayed in Table 2:

Feature Selection Strategy

To optimize the feature set for model performance, we implemented a two-step strategy:

Feature 1	Feature 2	Correlation
GenHlth	PhysHlth	0.552757
PhysHlth	DiffWalk	0.487976
GenHlth	DiffWalk	0.476639
Education	Income	0.460565

Table 2: Feature pairs with correlation c > |0.45|

1. Removal of weakly correlated features

To address weak correlation with the target variable, we will remove features exhibiting an absolute Pearson correlation coefficient of less than 0.1 with the 'Diabetes_binary' column. This step aims to eliminate features with a minimal linear relationship to the outcome we are trying to predict, potentially reducing noise and simplifying the model.

2. Mitigation of multicollinearity

Subsequently, to mitigate the issue of multicollinearity among the remaining features, we will identify pairs of features with a high absolute Pearson correlation coefficient (greater than 0.45) as shown in table 2. For each such pair, one of the features will be removed. The decision on which feature to remove will be based on its correlation with the target variable; the feature with the weaker absolute correlation to 'Diabetes_binary' will be prioritized for removal. This strategy aims to reduce redundancy in the feature set, which can lead to instability and interpretability issues in some models, without significantly sacrificing predictive power related to the target.

The implementation process in Python is shown below:

```
1 import pandas as pd
2 import numpy as np
4 # Load the dataset
5 df = pd.read_csv('datasets/diabetes_binary_5050split_health_indicators_BRFSS2015.csv')
6 target = 'Diabetes_binary'
8 # Step 1: Remove features with very low correlation to the target
abs_corr_target = df.corr()[target].abs()
10 low_corr_features = abs_corr_target[(abs_corr_target < 0.1) & (abs_corr_target.index != target)].
      index.tolist()
11 df_dropped_low = df.drop(columns=low_corr_features)
# Step 2: Identify and handle multicollinearity (high inter-feature correlation)
corr_remaining = df_dropped_low.corr()
upper_triangle = np.triu(np.ones(corr_remaining.shape), k=1).astype(bool)
_{17} # Extract high-correlation feature pairs (|correlation| > 0.45)
18 high_corr_pairs = (
     corr_remaining.where(upper_triangle)
20
     .stack()
     .reset_index()
22 )
23 high_corr_pairs.columns = ['Feature 1', 'Feature 2', 'Correlation']
24 filtered_high_corr = high_corr_pairs[high_corr_pairs['Correlation'].abs() > 0.45]
# Drop the less important feature (based on correlation with target) from each high-correlation
     pair
27 drop_multicollinearity = set()
abs_corr_remaining_target = df_dropped_low.corr()[target].abs()
```

```
30 for _, row in filtered_high_corr.iterrows():
31
      f1, f2 = row['Feature 1'], row['Feature 2']
      if f1 != target and f2 != target:
          corr_f1_target = abs_corr_remaining_target.get(f1, 0)
          corr_f2_target = abs_corr_remaining_target.get(f2, 0)
34
          if corr_f1_target < corr_f2_target:</pre>
36
              drop_multicollinearity.add(f1)
37
          else:
              drop_multicollinearity.add(f2)
38
40 # --- Step 3: Final dataset after feature selection ---
41 df_final = df_dropped_low.drop(columns=list(drop_multicollinearity))
42 df_final.to_csv('datasets/datasetF.csv', index=False)
44 # Output summary
print("Low correlation features:", low_corr_features)
print("\nHigh correlation pairs (> 0.45):")
47 print(filtered_high_corr.sort_values(by='Correlation', ascending=False))
48 print("\nFeatures dropped for multicollinearity:", list(drop_multicollinearity))
49 print("\nFinal features:", df_final.columns.tolist())
```

4 Algorithm implementation

Next, we will implement the three chosen algorithms: MLP (Multilayer Perceptron), Autoencoder, and [Third Algorithm]. These algorithms have been selected due to their suitability for classification problems and their ability to handle structured and complex datasets. First, we will provide a brief introduction to each algorithm, highlighting their key characteristics. After that, we will proceed with the implementation and performance analysis for our dataset.

4.1 Multilayer Perceptron (MLP)

The **Multilayer Perceptron (MLP)** is a core model in the field of machine learning, used for both classification and regression tasks. MLP consists of three primary layers: the **input layer**, one or more **hidden layers**, and the **output layer**. Each layer contains multiple neurons, and neurons from one layer are fully connected to neurons in the next layer. This dense connectivity allows MLP to capture complex relationships in data, particularly non-linear ones, which are common in many real-world applications [1].

In the context of our dataset, which includes health-related features such as BMI, Age, Physical_Health, and Sleep_Time, MLP is well-suited to identify hidden patterns and make predictions about whether an individual has diabetes or not. The network learns from the data through a process called *backpropagation*, where errors from the output are propagated back through the network to adjust the weights of the connections between neurons. This process is combined with *gradient descent*, an optimization technique that minimizes the prediction error by adjusting the weights during each iteration of training. Over time, the model learns the optimal weights, improving its ability to make accurate predictions [2].

The basic structure of an MLP is illustrated in the figure below. The input features, such as BMI and Age, are fed into the input layer. From there, they pass through one or more hidden layers, where neurons transform the data by applying learned weights and activation functions. The transformed data then flows to the output layer, which produces the final prediction, such as whether an individual has diabetes.

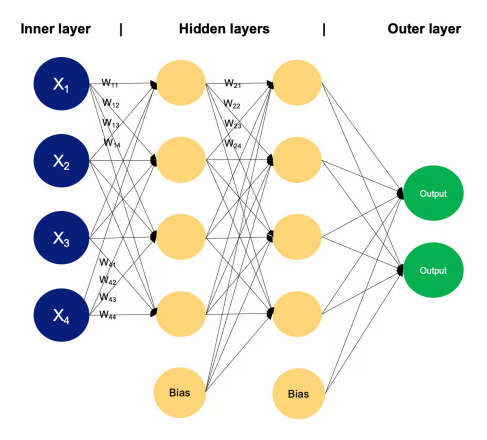


Figure 2: Basic structure of a Multilayer Perceptron (MLP) [3].

In the image, you can observe the flow of information from the input layer through the hidden layers to the output layer. Each layer plays a critical role in transforming the input data to make sense of the patterns and relationships. The neurons in the hidden layers learn features in the data, and as the network adjusts its weights through training, it becomes more accurate in predicting the target outcome.

MLPs are particularly effective in tasks where the relationships between input features and the target variable are non-linear, as is the case in health-related predictions like diabetes classification [1]. This model's ability to learn from complex and large datasets allows it to handle a wide variety of problems, from image recognition to medical diagnoses.

What makes MLPs particularly well-suited for our dataset is their capacity to model intricate relationships between multiple features simultaneously. By learning these relationships, MLPs can make accurate predictions, even when the data contains complex, interdependent factors. This makes MLP an ideal choice for predicting the likelihood of diabetes based on a variety of health indicators [3].

4.1.1 Implementation and Evaluation

The MLP model was implemented using the scikit-learn library, leveraging two hidden layers with sizes 100 and 50 respectively. The activation function used was ReLU, and stochastic gradient descent was chosen as the optimizer with adaptive learning rate and a maximum of 500 iterations.

Before finalizing the model, we performed hyperparameter tuning using GridSearchCV to identify the best combination of parameters. This ensures that the model generalizes well to unseen data. The best parameters found were:

```
{'mlp_activation': 'relu', 'mlp_alpha': 0.001, 'mlp_hidden_layer_sizes': (100, 50),
'mlp_learning_rate': 'adaptive', 'mlp_max_iter': 500, 'mlp_solver': 'sgd'}
```

```
import pandas as pd
2 import numpy as np
3 import matplotlib.pyplot as plt
5 from sklearn.model_selection import train_test_split, cross_validate
6 from sklearn.neural_network import MLPClassifier
7 from sklearn.metrics import (
     accuracy_score, precision_score, recall_score,
      f1_score, confusion_matrix, ConfusionMatrixDisplay,
9
      classification_report
10
11 )
13 # 1. Load data
df = pd.read_csv("datasets/diabetes_scaled.csv")
16 # 2. Features & Labels
17 X = df.drop("Diabetes_binary", axis=1)
y = df["Diabetes_binary"]
20 # 3. Train/Test
21 X_train = pd.read_csv("datasets/X_train.csv")
22 X_test = pd.read_csv("datasets/X_test.csv")
y_train = pd.read_csv("datasets/y_train.csv")
y_test = pd.read_csv("datasets/y_test.csv")
26 # 4. Define MLP model
27 mlp = MLPClassifier(
     hidden_layer_sizes=(100, 50),
28
     activation='relu',
     solver='sgd',
30
     alpha=0.001,
31
      learning_rate='adaptive',
      max_iter=500,
34
      random_state=42
35 )
36
37 # 5. Train model
mlp.fit(X_train, y_train)
40 # 6. Predict on test set
41 y_pred = mlp.predict(X_test)
# Plot loss curve for full feature set
44 plt.figure(figsize=(8, 4))
45 plt.plot(mlp.loss_curve_)
46 plt.title("Loss Curve (Full Feature Set)")
47 plt.xlabel("Iterations")
48 plt.ylabel("Loss")
49 plt.grid(True)
50 plt.show()
52 # 7. Evaluate metrics
print("Performance on Full Feature Set:")
print(f"Accuracy: {accuracy_score(y_test, y_pred):.4f}")
55 print(f"Precision: {precision_score(y_test, y_pred):.4f}")
56 print(f"Recall: {recall_score(y_test, y_pred):.4f}")
57 print(f"F1 Score: {f1_score(y_test, y_pred):.4f}")
59 # 8. Confusion matrix visualization
60 cm = confusion_matrix(y_test, y_pred)
61 disp = ConfusionMatrixDisplay(confusion_matrix=cm, display_labels=mlp.classes_)
62 disp.plot(cmap=plt.cm.Blues)
```

```
plt.title("Confusion Matrix (Full Feature Set)")
64 plt.show()
65
66 # 9. Cross-validation without plot
67 cv_results = cross_validate(
      mlp , X_train , y_train ,
69
       cv=5,
       scoring=["accuracy", "precision", "recall", "f1"],
       return_train_score=True
72 )
74 print("\nCross-Validation Mean Scores (5-Fold):")
75 for metric in ["accuracy", "precision", "recall", "f1"]:
       test_score = np.mean(cv_results[f'test_{metric}'])
76
       train_score = np.mean(cv_results[f'train_{metric}'])
78
       print(f"{metric.capitalize()}: Train = {train_score:.4f}, Test = {test_score:.4f}")
print("\nDetailed Classification Report (Full Feature Set):")
81 print(classification_report(y_test, y_pred, target_names=["No Diabetes", "Diabetes"]))
** # === Performance on Selected Features + Interaction Terms ===
# Select features of interest + target
86 df_interact = df[["GenHlth", "PhysHlth", "DiffWalk", "Education", "Income", "Diabetes_binary"]].
       copy()
87
88 # Add interaction features
89 df_interact["GenHlth_x_PhysHlth"] = df_interact["GenHlth"] * df_interact["PhysHlth"]
90 df_interact["PhysHlth_x_DiffWalk"] = df_interact["PhysHlth"] * df_interact["DiffWalk"]
91 df_interact["GenHlth_x_DiffWalk"] = df_interact["GenHlth"] * df_interact["DiffWalk"]
92 df_interact["Education_x_Income"] = df_interact["Education"] * df_interact["Income"]
94 # Prepare features and labels
95 X_interact = df_interact.drop("Diabetes_binary", axis=1)
96 y_interact = df_interact["Diabetes_binary"]
98 # Train/Test split
99 X_train_i, X_test_i, y_train_i, y_test_i = train_test_split(
      X_interact, y_interact, test_size=0.2, stratify=y_interact, random_state=42
101 )
103 # Define MLP model
104 mlp_interact = MLPClassifier(
      hidden_layer_sizes=(100, 50),
      activation = 'relu',
      solver='sgd',
      alpha=0.001,
108
      learning_rate='adaptive',
109
      max_iter=500,
110
      random_state=42
112
114 # Train model
mlp_interact.fit(X_train_i, y_train_i)
116
117 # Predict on test set
y_pred_i = mlp_interact.predict(X_test_i)
119
# Plot loss curve for selected features + interaction terms
plt.figure(figsize=(8, 4))
plt.plot(mlp_interact.loss_curve_)
plt.title("Loss Curve (Interactions Model)")
```

```
plt.xlabel("Iterations")
plt.ylabel("Loss")
plt.grid(True)
127 plt. show()
129 # Evaluate metrics
print("\nPerformance on Selected Features + Interactions:")
print(f"Accuracy: {accuracy_score(y_test_i, y_pred_i):.4f}")
print(f"Precision: {precision_score(y_test_i, y_pred_i):.4f}")
print(f"Recall: {recall_score(y_test_i, y_pred_i):.4f}")
print(f"F1 Score: {f1_score(y_test_i, y_pred_i):.4f}")
# Confusion matrix visualization
cm_i = confusion_matrix(y_test_i, y_pred_i)
138 disp_i = ConfusionMatrixDisplay(confusion_matrix=cm_i, display_labels=mlp_interact.classes_)
disp_i.plot(cmap=plt.cm.Greens)
plt.title("Confusion Matrix (Interactions Model)")
141 plt.show()
143 print("\nDetailed Classification Report (Interactions Model):")
144 print(classification_report(y_test_i, y_pred_i, target_names=["No Diabetes", "Diabetes"]))
```

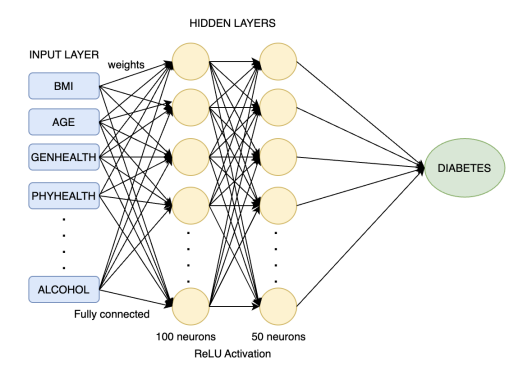


Figure 3: Our structure of a Multilayer Perceptron (MLP).

Figure 3 illustrates the architecture of the Multilayer Perceptron used in our implementation. It includes an input layer that receives features such as BMI, Age, and GenHealth, followed by two hidden layers with 100 and 50 neurons respectively. These layers apply ReLU activation and are fully connected. The final layer outputs a binary prediction indicating whether the individual is diabetic or not.

4.1.2 Results on Full Feature Set

Performance on Full Feature Set:

Accuracy: 0.7526 Precision: 0.7362 Recall: 0.8011 F1 Score: 0.7673

This baseline performance shows that the MLP model handles the full feature set well, achieving over 75% accuracy and strong recall (80.11%). Precision and F1-score are also relatively balanced, which means the model is both identifying and distinguishing cases effectively.

Next, we look at cross-validation results to assess the model's ability to generalize across different data splits:

Cross-Validation Mean Scores (5-Fold):
Accuracy: Train = 0.7558, Test = 0.7484
Precision: Train = 0.7359, Test = 0.7295
Recall: Train = 0.8113, Test = 0.8040
F1: Train = 0.7718, Test = 0.7649

These results confirm that the model generalizes well, as train and test scores remain consistent across folds. Importantly, test recall remains high (80.40%), reinforcing the model's reliability in correctly identifying diabetic cases in unseen data. This cross-validation step validates the single-run results and supports the model's robustness.

Now we present the full classification report to explore class-wise performance in more detail:

Detailed Classification Report (Full Feature Set):

	precision	recall	f1-score	support
No Diabetes	0.77	0.70	0.74	6753
Diabetes	0.74	0.80	0.77	7000
accuracy			0.75	13753
macro avg	0.75	0.75	0.75	13753
weighted avg	0.75	0.75	0.75	13753

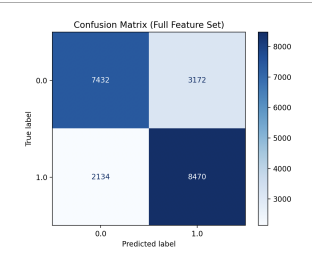


Figure 4: Confusion Matrix for MLP on Full Feature Set

From this report, it's evident that the model performs slightly better at detecting positive (diabetic) cases than negative ones, with a higher recall for the "Diabetes" class (80%). This is desirable in our context: it is far more critical to minimize false negatives (undiagnosed diabetic patients) than false positives. Therefore, recall is the metric we prioritize most.

4.1.3 Results on Selected Features with Interaction Terms

Performance on Selected Features + Interactions:

Accuracy: 0.6917 Precision: 0.6633 Recall: 0.8007 F1 Score: 0.7256

While accuracy and precision have dropped in this simplified model, recall has remained nearly identical (80.07%) to the full-feature model. This suggests that the reduced feature set is still effective in identifying diabetic patients, even if it leads to more false positives (lower precision).

We now show the detailed classification report for this model:

Detailed Classification Report (Interactions Model):

	precision	recall	f1-score	support
No Diabetes	0.74	0.58	0.65	6753
Diabetes	0.66	0.80	0.73	7000
accuracy			0.69	13753
macro avg	0.70	0.69	0.69	13753
weighted avg	0.70	0.69	0.69	13753

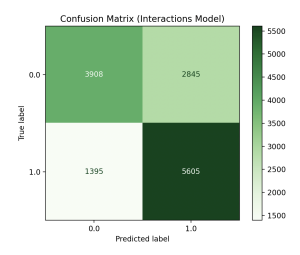


Figure 5: Confusion Matrix for MLP on Selected Features + Interaction Terms

Here again, recall for diabetic cases remains at 80%, but performance for non-diabetic predictions drops noticeably (recall of 58%). In practice, this means more people without diabetes are misclassified as diabetic. While this may raise false alarms, it is an acceptable trade-off in medical diagnostics where missing true cases is more dangerous than flagging potential ones.

In both models, recall is prioritized above all, aligning with our goal of reducing undiagnosed diabetic cases. The full-feature model is preferred for balanced performance, but the interaction-based model still holds value in more constrained environments.

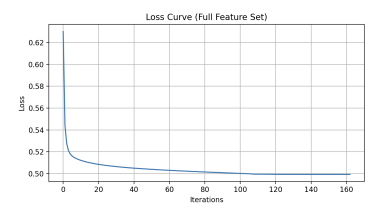


Figure 6: Training loss curve for the MLP model over 500 iterations.

The loss curve for the diabetes dataset shows a rapid initial decrease followed by convergence, indicating stable training and successful optimization.

4.2 Autoencoder

4.3 Naive Bayes

The Naive Bayes algorithm offers a straightforward yet often surprisingly effective approach to supervised learning, particularly for classification tasks like predicting diabetes status. At its core, it leverages Bayes' Theorem, a fundamental concept in probability that allows us to update the probability of an event based on new evidence. What makes Naive Bayes "naive" is its key simplifying assumption: it treats every feature in the dataset as if it were completely independent of all other features, given the class label (in our case, whether an individual has diabetes or not). The algorithm works by learning the conditional probability of each feature value for each class. For instance, it calculates the probability of having a certain BMI range given that the individual has diabetes, and the probability of having the same BMI range given that they do not have diabetes. It does this for all features in our dataset. When presented with a new individual's health profile, Naive Bayes uses these learned probabilities, along with the overall prevalence of diabetes in the training data (the prior probability), to calculate the probability of that individual belonging to each class (diabetes or no diabetes). The class with the highest calculated probability is then assigned as the prediction. The mathematical foundation lies in Bayes' Theorem:

$$P(\mathsf{Class}|\mathsf{Features}) = \frac{P(\mathsf{Class}) \times P(\mathsf{Features}|\mathsf{Class})}{P(\mathsf{Features})}$$

The "naive" independence assumption simplifies the term P(Features|Class) into the product of the individual feature probabilities:

$$P(x_1, x_2, ..., x_n | Class) = P(x_1 | Class) \times P(x_2 | Class) \times ... \times P(x_n | Class)$$

For our study, Naive Bayes presents a compelling option due to its speed, ease of implementation, and ability to handle datasets with numerous features. Even though the health indicators in our dataset are likely not entirely independent, the algorithm's empirical success in various domains suggests it can still provide valuable insights and potentially strong predictive performance for identifying diabetes risk. Its simplicity also makes it a good baseline model against which to compare the performance of more complex algorithms like MLP and Autoencoder.

5 Result interpretation

References

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