### **Dhruv Ragunathan**

#### **Contact Information**

• Linkedin: <a href="https://www.linkedin.com/in/dhruv-ragunathan-908993b1/">https://www.linkedin.com/in/dhruv-ragunathan-908993b1/</a>)

• Github: https://github.com/dragunat2016 (https://github.com/dragunat2016)

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# **Overview**



The Behavioral Risk Factor Surveillance System (BRFSS) is the nation's premier system of health-related telephone surveys that collects state data about U.S. residents regarding their health-related risk behaviors, chronic health conditions, and use of preventive services.

Established in 1984 with 15 states, BRFSS now collects data in all 50 states as well as the District of Columbia and three U.S. territories. BRFSS completes more than 400,000 adult interviews each year, making it the largest continuously conducted health survey system in the world.

Researchers have seen the opportunity to apply machine learning algorithms to make predictions on the data, since it was a feature rich dataset with hundreds-of-thousands of records.

# **Business Objectives**



We have been tasked by the CDC to create models from previous BRFSS data that predicts diabetes. The CDC wants to help the people it surveys and alert them if they are at risk for diabetes given their survey results. Long-term the CDC would like to publish an application to Americans allowing them to fill out a form with questions on their vitals like BMI and habits such as exercise. Upon completing the form, the CDC would send back a diabetic risk to the person.

The motivation behind this is that diabetes is one of the most prevalent and costly diseases in the USA. Currently, 38 million people have diabetes of which 9 million are undiagnosed. When considering the precursor, prediabetes, that number jumps to 98 million people.

Diabetic patients are more likely to visit the emergency department and require expensive treatments and medications for their life. Reducing diabetes across the country would greatly improve the quality of life of millions of Americans.

Accuracy and precision are our primary metrics of evaluation. Accuracy defines the number of correct predictions made by the model over the total number of predictions. Precision defines the number of True positive identified over the true positive plus the false positive rate.

Optimizing on these two metrics should reduce the amount of false positives we encounter. We want to avoid false positives because they could result in unnecessary outreach and wasting resources. We will still record and review other metrics such as F1 score, ROC-AUC, and recall to review in-case these metrics are even for some models.

We will also be incorporating the "run time" of the model in our evaluation. Run time is the amount of time it takes to train and test the model.

A final model evaluation will be made by some heuristic combination of the accuracy, precision, and time it takes model too run. Any gains in accuracy and precision need to justify the time it takes to train and use the model.

### **Data Overview**

#### Source

The 2015 data is available on this link from the CDC's website. The table with all the responses and the key donoting the data terms are also available. The link to the survey questions is <a href="https://www.cdc.gov/brfss/questionnaires/pdf-ques/2015-brfss-questionnaire-12-29-14.pdf">https://www.cdc.gov/brfss/questionnaires/pdf-ques/2015-brfss-questionnaire-12-29-14.pdf</a>)

The page on the CDC's website containing the data is <a href="here">here</a> (https://www.cdc.gov/brfss/annual data/annual data.htm).

The data on the CDC's page is in an ASCII format and hard too decode with time constraints. We found a CSV version of that data on Kaggle. The download link for the CSV is specifically <a href="https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system">https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system</a>).

Full Link: <a href="https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system">https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system</a>)

### **Limitations**

This is survey data where the user responses were segmented into several categories.

So the following limitations apply:

- Survey respondants may not be comfortable revealing sensitive information over the phone even if the response is anonymous.
- Many respondants who answer "no" for diabetes may actually have diabetes, but were not diagnosed. Note: That there was a significant imbalance of diabetes/pre-diabetes versus those who stated that they do not have the condition.
- Many variables that are continuous in nature were treated as ordinal in the study such as income and age. These variables were treated as ordinal as part of the models.

# **Data Preparation**

The steps for data preparation and cleaning were done in this <u>notebook</u> (<u>notebooks/Data Cleaning.ipynb</u>) for the sake of simplifying the main notebook.

This is the short version of the data cleaning process. For more detail please click the link above.

### **High - Level Process**

- · Selected for columns related to diabetes
- Dropped columns with significant data missing
- · Reviewed the data in the features.
  - Values within features that corresponded to information like 'N/A', 'Refused', 'Didn't Know' were dropped.
  - Values were transformed to be more ordinal
- · Combined Diabetes and Prediabetes data

Addressed class imbalance by making the diabetes/non-diabetes records 50-50

```
In [1]:
            import numpy as np
         2
            import pandas as pd
         3 import seaborn as sns
         4 import matplotlib.pyplot as plt
            import warnings
            warnings.filterwarnings("ignore")
            import pickle
In [2]:
            from sklearn.model_selection import train_test_split, GridSearchCV,
         2 from sklearn.preprocessing import StandardScaler, OneHotEncoder, Fun
         3 from sklearn.impute import SimpleImputer
         4 from sklearn.compose import ColumnTransformer
         5 from sklearn.linear_model import LogisticRegression
         6 from sklearn.svm import SVC
         7 from sklearn.ensemble import RandomForestClassifier, GradientBoostin
         8 from sklearn.svm import LinearSVC
         9 from sklearn.tree import DecisionTreeClassifier
        10 from sklearn.naive bayes import GaussianNB
        11 from sklearn.neighbors import KNeighborsClassifier
        12 from sklearn.metrics import confusion matrix, ConfusionMatrixDisplay
        13 from sklearn.compose import ColumnTransformer
        14 from sklearn.pipeline import Pipeline
        15 from sklearn import metrics
        16 from xgboost import XGBClassifier
        17 from datetime import datetime as dt
        18 random state=42
```

In [3]:	1	<pre>diab_df = pd.read_csv('diabetes_binary_5050_DR_BRFSS2015.csv')</pre>	
	2		
	3	diab_df.head()	

#### Out[3]:

	Diabetes_binary	HighBP	Asthma	HighChol	CholCheck	ВМІ	Smoker	Stroke	HeartDiseaseor
0	0.0	0.0	0.0	0.0	1.0	20.0	0.0	0.0	
1	0.0	0.0	1.0	1.0	1.0	32.0	1.0	0.0	
2	0.0	1.0	0.0	0.0	1.0	50.0	1.0	0.0	
3	0.0	1.0	0.0	1.0	1.0	27.0	0.0	0.0	
4	0.0	1.0	0.0	1.0	1.0	14.0	1.0	0.0	

5 rows × 26 columns

```
In [4]: 1 diab_df.info()
```

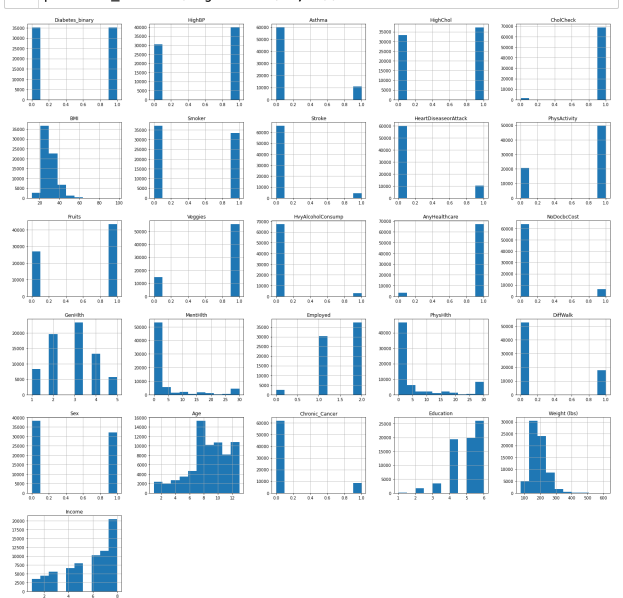
<class 'pandas.core.frame.DataFrame'>
RangeIndex: 70252 entries, 0 to 70251
Data columns (total 26 columns):

#	Column	-	ıll Count	Dtype
0	Diabetes_binary	70252	non-null	float64
1	HighBP	70252		float64
2	Asthma		non-null	float64
3	HighChol		non-null	float64
4	CholCheck		non-null	float64
5	BMI		non-null	float64
6	Smoker		non-null	float64
7	Stroke		non-null	float64
8	HeartDiseaseorAttack	70252		float64
9	PhysActivity	70252	non-null	float64
10	Fruits	70252		float64
11	Veggies	70252	non-null	float64
12	HvyAlcoholConsump	70252	non-null	float64
13	AnyHealthcare	70252	non-null	float64
14	NoDocbcCost	70252	non-null	float64
15	GenHlth	70252	non-null	float64
16	MentHlth	70252	non-null	float64
17	Employed	70252	non-null	float64
18	PhysHlth	70252	non-null	float64
19	DiffWalk	70252	non-null	float64
20	Sex	70252	non-null	float64
21	Age	70252	non-null	float64
22	Chronic_Cancer	70252	non-null	float64
23	Education	70252	non-null	float64
24	Weight (lbs)	70252	non-null	float64
25	Income	70252	non-null	float64
dtype	es: float64(26)			
	42 0 45			

dtypes: float64(26) memory usage: 13.9 MB

# **Exploratory Data Analysis**

#### 1 p = diab\_df.hist(figsize = (26,26))



We can see a few interesting trends from the various histograms. First the diabetes versus nondiabetes is balanced as designed in the data cleaning process.

Second, High Blood pressure is also near balanced.

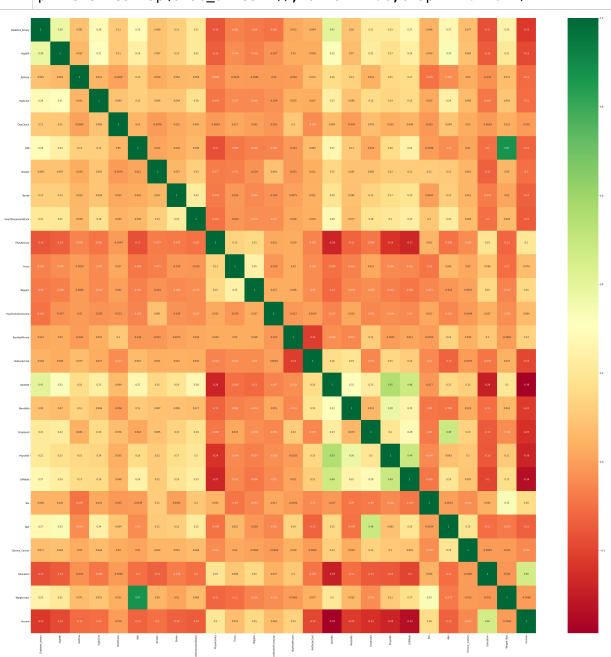
Weight is centered around near 200 points, which tracks on average.

There are more females than males in this study.

Higher incomes are mostly represented in the study. This could imply that the study is biased towards collecting data for those of a higher income. This would make sense since higher income individuals are more likely too have landlines.

Similarly, variables that show co-morbities such as stroke, heart disease, and chronic cancer victims are not represented well in the data.

In [6]: 1 plt.figure(figsize=(50,50))
2 p = sns.heatmap(diab\_df.corr(), annot=True,cmap ='RdYlGn')



The vast majority of variables are not correlated with one another. This makes this data set could for modeling and less likely for overfitting/multicolinearity.

However, there is one exception. That being BMI and Weight. Since BMI is calculated from Weight this is not suprising.

To reduce the possibility of overfitting, we will drop the weight column. We chose to drop weight instead of BMI because BMI is more correlated with diabetes than weight is (0.29 vs 0.25). Therefore, dropping BMI as a feature would reduce the accuracy of the model more than weight would.

This data-driven decisions tracks with intuition. BMI is a better metric of determining how

# **Modeling**

Sections include

- · Scaled Data for Model
- Ran Baseline Model
- Ran Additional Models
- Tuned best performing model from 'Additional Models' section
- Created a neural network since literature implied it was the best performing model for this use-case

### **Scaling Data**

Using Standard Scaler to scale the data

```
In [11]:
           1 | X_scaled = sc_X.fit_transform(X)
             X scaled = pd.DataFrame(X scaled,columns=X.columns)
           2
           3 X_scaled
```

#### Out[11]:

		HighBP	Asthma	HighChol	CholCheck	ВМІ	Smoker	Stroke	HeartDiseaseor
	0	-1.14055	-0.425492	-1.057809	0.156285	-1.379308	-0.948568	-0.257453	-0.4
	1	-1.14055	2.350221	0.945350	0.156285	0.301592	1.054221	-0.257453	-0.4
	2	0.87677	-0.425492	-1.057809	0.156285	2.822943	1.054221	-0.257453	-0.4
	3	0.87677	-0.425492	0.945350	0.156285	-0.398783	-0.948568	-0.257453	2.0
	4	0.87677	-0.425492	0.945350	0.156285	-2.219758	1.054221	-0.257453	-0.4
7	0247	-1.14055	-0.425492	0.945350	0.156285	1.001967	-0.948568	-0.257453	-0.4
7	0248	-1.14055	-0.425492	0.945350	0.156285	-0.118633	1.054221	-0.257453	2.0
7	0249	0.87677	2.350221	0.945350	0.156285	-0.678933	-0.948568	-0.257453	2.0
7	0250	0.87677	-0.425492	0.945350	0.156285	-1.659458	-0.948568	-0.257453	-0.4
7	0251	0.87677	-0.425492	0.945350	0.156285	-0.678933	-0.948568	-0.257453	2.0

70252 rows × 24 columns

```
In [12]:
             X_train, X_test, y_train, y_test = train_test_split(X_scaled,y, test)
```

```
In [76]:
```

```
# Pickle data to run models in other notebooks
1
2
3
  with open('Variables/X_train.pickle', 'wb') as xtr:
      pickle.dump(X_train,xtr)
4
5
6
```

```
In [77]:
             #Store other variables
           1
           2
           3
           4
           5
             with open('Variables/X_test.pickle', 'wb') as xtst:
           6
                  pickle.dump(X_test,xtst)
           7
           8
             with open('Variables/y_train.pickle', 'wb') as ytr:
           9
                  pickle.dump(y_train,ytr)
          10
          11
          12
             with open('Variables/y_test.pickle', 'wb') as ytst:
                  pickle.dump(y_test,ytst)
          13
          14
          15
          16
```

#### **Baseline Model**

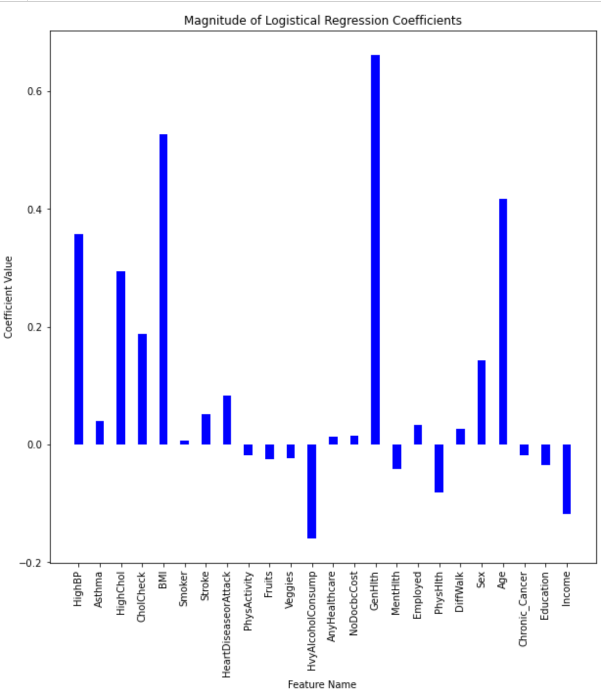
- Start with baseline Logistic Regression
- Train data
- Make predictions from test data set
- Review metrics such as accuracy, recall, precision, ROC-AUC, and F1
- · Review features

```
In [15]:
             # Baseline Model is a logistic regression
             lr model = LogisticRegression()
           3
             lr model.fit(X train,y train)
Out[15]: LogisticRegression()
In [16]:
          1
             lr_preds = lr_model.predict(X_train)
           3 | Ir train acc = round(metrics.accuracy score(y train, Ir preds), 3)
In [17]:
           1 print('Training Accuracy score is ',lr_train_acc)
         Training Accuracy score is 0.745
In [18]:
           1
             # Predictions from testing data set
           2
           3
             y_pred = lr_model.predict(X_test)
In [19]:
          1
           2
             lr_acc = metrics.accuracy_score(y_test, y_pred)
             lr rec = recall score(y test, y pred)
             lr_prec = precision_score(y_test, y_pred)
             lr_roc_auc = roc_auc_score(y_test, y_pred)
             lr_F1 = f1_score(y_test,y_pred)
             print('Accuracy: ',lr_acc)
             print('Recall: ', lr rec)
             print('Precision', lr_prec)
         10
             print('ROC - AUC', lr_roc_auc)
         11
             print('F1 Score', lr_F1)
         Accuracy: 0.7494128531777098
         Recall: 0.7691867943404316
         Precision 0.7384742041712404
         ROC - AUC 0.7494927450579391
```

Let's take a look at the features this model prioritized.

F1 Score 0.7535176758837943

```
In [20]:
           1
             fig = plt.figure(figsize = (10, 10))
           2
           3
              feature_name = X_train.columns
             coef_val = lr_model.coef_[0]
           4
           5
           6
             # creating the bar plot
           7
              plt.bar(feature_name, coef_val, color ='blue',
           8
                      width = 0.4)
           9
              plt.xlabel("Feature Name")
          10
              plt.ylabel("Coefficient Value")
          11
             plt.title("Magnitude of Logistical Regression Coefficients")
          12
          13
              plt.xticks(rotation=90)
             plt.show()
          14
```

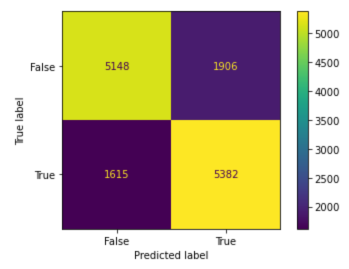


We can see that the feature given the most importance was GenHlth.

Other top features were High Blood Pressure, BMI, and Age.

Interestingly, heavy alcohol consumption did not positively affect diabetes correlation. Even though intuitively, one would think that more alcohol means more calories/sugar, which means higher likelyhood for diabetes.

```
In [21]:
              cm = confusion_matrix(y_test, y_pred)
           1
           2
             lr_TN, lr_FP, lr_FN, lr_TP = confusion_matrix(y_test, y_pred).ravel(
           3
             print('True Positive(TP)
                                        = ', lr_TP)
           4
           5
             print('False Positive(FP) = '
                                            , lr FP)
                                        = '
                                            , lr_TN)
              print('True Negative(TN)
              print('False Negative(FN) = ', lr_FN)
         True Positive(TP)
                                5382
                             =
         False Positive(FP) =
                                1906
         True Negative(TN)
                                5148
         False Negative(FN) =
                                1615
```



The confusion matrix above identifies similar amounts of true positives and true negatives. In addition, it also identified a similar number of false positives and false negatives.

- True Positive(TP) = 5320
- False Positive(FP) = 1982
- True Negative(TN) = 5033
- False Negative(FN) = 1716

These numbers are not too bad for a baseline model. The training and testing accuracy were similar, 74% indicating that the model is not overfitting the data. Let's see if we can use other models to improve these metrics from a baseline of 74%.

#### **Additional Models**

- Ran additional models such as Random Forest, XGB, Deicison Tree Classier, GaussianNB, and KNeighbors
- · Reviewed metrics and selected one model for tuning
- Tunned the XGB model under hyper parameter tuning.
- Created confusion matrices, calculated metrics, and compared performance

```
In [24]:
          1
             # loop over each classifier to evaluate poerformance
          2
          3
             train_acc, test_acc, rec, prec, F1, Roc_Auc,trained_model,run_time =
          4
          5
             for model name in model arr.keys():
          6
          7
                 model = model_arr[model_name]
          8
          9
                 start = dt.now()
         10
         11
                 # Fit the classifier
         12
                 trained_model[model_name] = model.fit(X_train, y_train)
                                                                             #Store
         13
         14
                 #Find training accuracy
         15
                 y_train_pred = model.predict(X_train)
         16
         17
         18
                 # Make predictions
         19
                 y_pred = model.predict(X_test)
         20
         21
                  running secs = (dt.now() - start).seconds
         22
         23
                 # Calculate metrics
         24
                 train acc[model name] = accuracy score(y train,y train pred)
         25
                 test_acc[model_name] = accuracy_score(y_test, y_pred)
         26
                  rec[model_name] = recall_score(y_test, y_pred)
         27
                 prec[model_name] = precision_score(y_test, y_pred)
                 F1[model_name] = f1_score(y_test,y_pred)
         28
                 Roc_Auc[model_name] = roc_auc_score(y_test,y_pred)
         29
         30
                  run time[model name] = running secs
```

```
In [25]: 1 measures = pd.DataFrame(index=model_arr.keys(), columns=['Training A
2 measures['Training Accuracy'] = train_acc.values()
3 measures['Testing Accuracy'] = test_acc.values()
4 measures['Recall'] = rec.values()
5 measures['Precision'] = prec.values()
6 measures['F1 Score'] = F1.values()
7 measures['Roc-AUC Score'] = Roc_Auc.values()
8 measures['Runtime (s)'] = run_time.values()
9 measures
```

#### Out[25]:

	Training Accuracy	Testing Accuracy	Recall	Precision	F1 Score	Roc-AUC Score	Runtime (s)
Logistical Regression	0.745076	0.749413	0.769187	0.738474	0.753518	0.749493	0
Random Forest	0.997313	0.742652	0.780620	0.724115	0.751307	0.742805	7
Decision Tree Classifer	0.997331	0.655327	0.653566	0.654033	0.653799	0.655320	0
XGB Classifier	0.791214	0.748345	0.788767	0.728389	0.757376	0.748509	5
svc	0.769719	0.753541	0.806917	0.727765	0.765300	0.753756	350
GaussianNB	0.714614	0.717743	0.703016	0.722638	0.712692	0.717683	0
KNeighbors	0.796107	0.709487	0.734886	0.697788	0.715857	0.709589	275

The dispararity between the training and testing accuracy above for Random Forest and Decision Tree Classifier indicates that those models are highly overfit. Especially, the Decision Tree Classifier which had the lowest testing accuracy but a near 100% training accuracy.

The testing accuracies of the rest of the models were similar. SVC and XGB have the highest accuracies and have very close metrics to one another.

The only differences is that XGB has a marginally higher precision and SVC has a higher recall by 1% and a ROC-AUC score and accuracy score. Based on these metrics alone, it would make sense to chose SVC over XGB.

However, XGB runs significantly faster than SVC. In fact, XGB ran ~80 times faster than SVC. Note: Times may vary depending on machine. Since its significantly easier to use.

Ultimately, all of these models fall short of logistical's regressions accuracy to runtime ratio. Of all the models that ran in 0 seconds, logistical regression had the highest accuracy/precision.

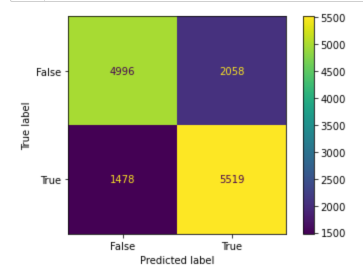
That said, XGB has the potential to improve on these numbers through hyper-parameter tuning. We will be using this model for further analysis to try on improving on these results.

```
In [26]: 1 xgb = trained_model['XGB Classifier']
```

```
In [27]:
             # Create a confusion matrix to visualize results
          1
          2
          3
             y_pred_xgb = xgb.predict(X_test)
          4
          5
             cm = confusion_matrix(y_test, y_pred_xgb)
             xgb_TN, xgb_FP, xgb_FN, xgb_TP = confusion_matrix(y_test, y_pred_xgb
          6
          7
          8
             print('True Positive(TP) = ', xqb TP)
             print('False Positive(FP) = '
                                           , xgb_FP)
          9
             print('True Negative(TN) = '
                                           , xgb_TN)
          10
             print('False Negative(FN) = ', xqb FN)
          11
         True Positive(TP)
                            =
                                5519
         False Positive(FP) =
                                2058
         True Negative(TN)
                                4996
         False Negative(FN) =
                                1478
```

#### In [28]:

```
28]: 1 # Plot Results
2
3 xgb_cm_matrix = confusion_matrix(y_test,y_pred_xgb)
4
5 xgb_cm_display = ConfusionMatrixDisplay(confusion_matrix = xgb_cm_ma
6
7 xgb_cm_display.plot()
8 plt.show()
```



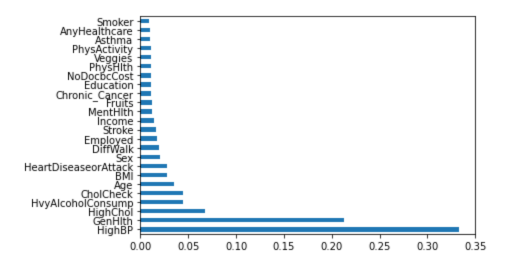
The confusion matrix above shows a high number of true positives/true negatives compared to the false positives/negatives. Let's see how many more correct prediction it made compared to the baseline model.

```
In [29]:
              print('True Positive(TP)
           1
                                            , xqb TP)
           2
             print('False Positive(FP) =
                                            , xgb_FP)
                                            , xgb_TN)
           3
              print('True Negative(TN)
                                         =
              print('False Negative(FN) = '
                                            , xgb_FN)
         True Positive(TP)
                             =
                                5519
         False Positive(FP) =
                                2058
         True Negative(TN)
                                4996
         False Negative(FN) =
                                1478
```

The xgboost model made 15 more correct predictions than the baseline model.

Let's take a look at what features XGB deemed important.

#### Out[31]: <AxesSubplot:>



Interestingly, the model put the highest weight on blood pressure by a significant margin. Almost 4 times higher than the next parameter of general health. This find tracks well with medical knowledge that high blood pressure and diabetes often are caused by unhealthy diet/health maintenance.

### **Hyper Parameter Tuning**

Now that we have picked a model to further investigate, let's see if we can improve our accuracy through hyper parameter tuning. There are different methods for hyper parameter tuning such as grid searching and random search, but here we will use bayesian optimization. From research, we determined that this method is generally more successful than the others.

Due too time and resources constraints we will use this method instead of trying several and comparing the results.

Source: Hyperparameter Optimization for Machine Learning Models Based on Bayesian Optimization, Wu et. al. <u>link</u>

(https://www.sciencedirect.com/science/article/pii/S1674862X19300047)

```
In [32]:
              from bayes_opt import BayesianOptimization
In [33]:
           1
              from sklearn.model selection import cross val score
In [34]:
           1
              from hyperopt import fmin, tpe, hp
In [35]:
           1
             #Function that takes in parameters for xgboost and returns the highe
           2
           3
             def xgboost_hyper_param(learning_rate,
           4
                                       n_estimators,
           5
                                       max depth,
           6
                                       subsample,
           7
                                       gamma):
           8
           9
                  max depth = int(max depth)
          10
                  n_estimators = int(n_estimators)
          11
          12
                  clf = XGBClassifier(
          13
                      max depth=max depth,
          14
                      learning_rate=learning_rate,
          15
                      n estimators=n estimators,
          16
                      gamma=gamma)
          17
                  return np.mean(cross_val_score(clf, X_train, y_train, cv=3, scor
          18
          19
In [36]:
             # Parameters for xgboost model. Start with arbirtrary parameter value
           1
           2
           3
             pbounds = {
                  'learning_rate': (0.01, 1.0),
           4
           5
                  'n_estimators': (100, 1000),
           6
                  'max_depth': (3,10),
           7
                  'subsample': (1.0, 1.0),
           8
                  'gamma': (0, 5)}
           9
```

```
In [37]:
            #Instantiate the Optimizer
          1
          2
          3
            optimizer = BayesianOptimization(
          4
                f=xgboost_hyper_param,
          5
                pbounds=pbounds
          6
             )
In [38]:
          1
             optimizer.maximize(
                 init_points=2,
          2
          3
                n_iter=3,
          4
             )
                                            | learni... | max depth | n esti...
             iter
                       target
                                    gamma
          subsample |
          1
                      0.8072
                                | 1.012
                                            0.4912
                                                       | 6.282
                                                                   896.6
          1.0
          2
                      0.8087
                                0.03199
                                            0.2878
                                                       | 5.184
                                                                   | 566.6
          1.0
          3
                      0.8174
                                0.4783
                                            0.353
                                                       | 5.156
                                                                   567.6
         1.0
                                2.406
          4
                      0.8272
                                            0.3556
                                                       4.402
                                                                   1 124.7
          1.0
                                                                   | 715.8
          5
                      0.7818
                                | 1.233
                                            0.8737
                                                       | 6.646
         1.0
                      _____
In [39]:
            optimizer.max
Out[39]: {'target': 0.8272277614471464,
          'params': {'gamma': 2.4060556409414273,
           'learning_rate': 0.3556184841702696,
           'max_depth': 4.402240692590198,
           'n estimators': 124.67046120487794,
           'subsample': 1.0}}
In [40]:
            #parameters are in the 'params' keys
          1
          2
          3
            xgb_best_params = optimizer.max['params']
          4
            xgb_best_params
Out[40]: {'gamma': 2.4060556409414273,
          'learning_rate': 0.3556184841702696,
          'max_depth': 4.402240692590198,
          'n_estimators': 124.67046120487794,
          'subsample': 1.0}
```

```
In [41]:
             # Create new classier with the best params
          1
          2
           3
             gamma = xgb_best_params['gamma']
             learning rate = xqb best params['learning rate']
             max depth = int(round(xgb best params['max depth'])) # Needs to be a
             n estimators = int(round(xgb best params['n estimators'])) # Needs t
          7
          8
           9
             xqb tuned = XGBClassifier(gamma = gamma,learning rate=learning rate,
In [42]:
             # Fit tuned model on training data
          1
           2
           3
             start = dt.now()
           4
             xgb_tuned.fit(X_train,y_train)
Out[42]: XGBClassifier(base score=0.5, booster='gbtree', colsample bylevel=1,
                       colsample_bynode=1, colsample_bytree=1, gamma=2.406055640
         9414273,
                       gpu_id=-1, importance_type='gain', interaction_constraint
         s='',
                       learning rate=0.3556184841702696, max delta step=0, max d
         epth=4,
                       min_child_weight=1, missing=nan, monotone_constraints
         ='()',
                       n_estimators=125, n_jobs=0, num_parallel_tree=1, random_s
         tate=0,
                       reg_alpha=0, reg_lambda=1, scale_pos_weight=1, subsample=
         1.0,
                       tree_method='exact', validate_parameters=1, verbosity=Non
         e)
In [43]:
             # Make predictions. Do the same for training data to determine if th
           1
           2
          3
             y_pred_tuned_train = xgb_tuned.predict(X_train)
             y pred tuned = xgb tuned.predict(X test)
           5
```

running secs xgb = (dt.now() - start).seconds

```
In [44]:
             xgb_tnd_trn_acc = accuracy_score(y_train,y_pred_tuned_train)
          1
          2
             xgb_tnd_tst_acc = accuracy_score(y_test, y_pred_tuned)
          3
             xgb_tnd_rec = recall_score(y_test, y_pred_tuned)
             xgb_tnd_rec_prec = precision_score(y_test, y_pred_tuned)
             xgb tnd rec roc auc = roc auc score(y test, y pred tuned)
             xgb tnd rec F1 = f1 score(y test,y pred tuned)
          7
          8
             print('Training Accuracy: ',xgb_tnd_trn_acc)
             print('Testing Accuracy: ',xgb_tnd_tst_acc)
             print('Recall: ',xgb_tnd_rec)
         10
             print('Precision', xgb_tnd_rec_prec)
         11
             print('ROC - AUC',xgb_tnd_rec_roc_auc)
         12
             print('F1 Score',xgb_tnd_rec_F1)
         13
```

Training Accuracy: 0.757566591341791
Testing Accuracy: 0.7544658743149953
Recall: 0.7981992282406746
Precision 0.7326511871966418
ROC - AUC 0.7546425684724779
F1 Score 0.7640218878248974

It appears the parameters did not change the results significantly. For better visualization let's use a confusion matrix.

```
In [45]:
             cm xgb tnd = confusion matrix(y test, y pred tuned)
             TN xgb tnd, FP xgb tnd, FN xgb tnd, TP xgb tnd = confusion matrix(y
          2
          3
             print('True Positive(TP) = ', TP xgb tnd)
             print('False Positive(FP) = ', FP_xgb_tnd)
             print('True Negative(TN) = ', TN_xgb_tnd)
             print('False Negative(FN) = ', FN_xgb_tnd)
         True Positive(TP) =
                               5585
         False Positive(FP) =
                               2038
         True Negative(TN) =
                               5016
         False Negative(FN) = 1412
In [46]:
          1 # Difference between first tuning iteration and baseline model
            lr_TP + lr_TN - TP_xgb_tnd - TN_xgb_tnd
          3
```

This tuning actually reduced the number of correct predictions the model makes.

Out[46]: -71

```
In [47]:
             # Add these values to our model dictionary
             # Since pandas does not allow you to add rows without removing the i
             # we need to recreate the table again
          5
             model_name = 'XGB Tuned 1'
             model_arr['XGB Tuned 1'] = xgb_tuned
          7
          8
             train acc[model name] = xqb tnd trn acc
             test acc[model name] = xqb tnd tst acc
             rec[model_name] = xgb_tnd_rec
         10
             prec[model name] = xqb tnd rec prec
         11
         12
             F1[model name] = xqb tnd rec F1
             Roc_Auc[model_name] = xgb_tnd_rec_roc_auc
         13
             run_time[model_name] = running_secs_xgb
In [48]:
             measures = pd.DataFrame(index=model_arr.keys(), columns=['Training A
             measures['Training Accuracy'] = train_acc.values()
          2
             measures['Testing Accuracy'] = test acc.values()
             measures['Recall'] = rec.values()
```

measures['Precision'] = prec.values()
measures['F1 Score'] = F1.values()

measures['Roc-AUC Score'] = Roc\_Auc.values()
measures['Runtime (s)'] = run\_time.values()

#### Out[48]:

measures

	Training Accuracy	Testing Accuracy	Recall	Precision	F1 Score	Roc-AUC Score	Runtime (s)
Logistical Regression	0.745076	0.749413	0.769187	0.738474	0.753518	0.749493	0
Random Forest	0.997313	0.742652	0.780620	0.724115	0.751307	0.742805	7
Decision Tree Classifer	0.997331	0.655327	0.653566	0.654033	0.653799	0.655320	0
XGB Classifier	0.791214	0.748345	0.788767	0.728389	0.757376	0.748509	5
SVC	0.769719	0.753541	0.806917	0.727765	0.765300	0.753756	350
GaussianNB	0.714614	0.717743	0.703016	0.722638	0.712692	0.717683	0
KNeighbors	0.796107	0.709487	0.734886	0.697788	0.715857	0.709589	275
XGB Tuned 1	0.757567	0.754466	0.798199	0.732651	0.764022	0.754643	5

Let's see if we can further improve them by increasing the bounds and also by increasing the number of iterations the optimizer runs over.

```
In [49]:
              pbounds2 = {
           1
           2
                  'learning_rate': (0.01, 0.6),
           3
                  'n_estimators': (100, 300),
           4
                  'max_depth': (3,7),
                  'subsample': (1.0, 1.0), # Won't allow values over 1.0
           5
                  'qamma': (5, 20)}
           6
           7
           8
           9
              optimizer2 = BayesianOptimization(
                  f=xgboost_hyper_param,
          10
                  pbounds=pbounds
          11
          12
              )
```

#### In [50]:

```
# Increased init points and n iter by 1 from previous tuning
1
2
3
  optimizer2.maximize(
4
       init_points=4,
5
       n_iter=5,
6
```

iter   subsample	target   	gamma	learni	max_depth	n   n_esti
1	0.7889	1.224	0.8298	6.204	706.0
1.0		1 4 624		l	1 705 2
2   1.0	0.8289	4.624	0.1121	5.794	705.2
3	0.813	1.293	0.7179	5.142	988.3
1.0					
4	0.7824	0.9487	0.6586	9.097	517.2
1.0   5	   0.8232	3.325	0.3872	7.412	790.9
1.0		1 01020	1 01007	, ,,,==	1
6	0.8287	4.448	0.1623	3.624	635.5
1.0	   0.8267	3.234	0.5833	3.322	531.9
1.0	0.0207	3.234	0.3033	3:322	551.9
8	0.8162	0.3406	0.8856	3.381	645.7
1.0	706	. 1 110		. 0 503	1 702 1
9   1.0	0.796	1.448	0.4718	9.502	782.1
110	 ========	=========	========	=========	========

\_\_\_\_\_

```
In [51]:
           1
             xgb_best_params2 = optimizer2.max['params']
           2 xgb_best_params2
```

```
Out[51]: {'gamma': 4.624140496883825,
          'learning_rate': 0.11214395671681449,
          'max_depth': 5.794307663328713,
          'n_estimators': 705.2226048065555,
          'subsample': 1.0}
```

```
In [52]:
             # Create new classier with the best params
          1
          2
             gamma = xgb best params2['gamma']
             learning_rate = xgb_best_params2['learning_rate']
          3
             max depth = int(round(xgb best params2['max depth'])) # Needs to be
             n estimators = int(round(xgb_best_params2['n_estimators'])) # Needs
          6
          7
             xqb tuned 2 = XGBClassifier(gamma = gamma,learning rate=learning rat
In [53]:
          1
             start = dt.now()
          2
             xgb_tuned_2.fit(X_train,y_train)
          3
Out[53]: XGBClassifier(base_score=0.5, booster='gbtree', colsample_bylevel=1,
                       colsample_bynode=1, colsample_bytree=1, gamma=4.624140496
         883825.
                       gpu id=-1, importance type='gain', interaction constraint
         s='',
                       learning rate=0.11214395671681449, max delta step=0, max
         depth=6,
                       min_child_weight=1, missing=nan, monotone_constraints
         ='()',
                       n estimators=705, n jobs=0, num parallel tree=1, random s
         tate=0,
                       reg alpha=0, reg lambda=1, scale pos weight=1, subsample=
         1.0,
                       tree_method='exact', validate_parameters=1, verbosity=Non
         e)
In [54]:
             y_pred_tuned_2_train = xgb_tuned_2.predict(X_train)
          2
             y pred tuned 2 = xgb tuned 2.predict(X test)
          3
             running_secs_xgb_2 = (dt.now() - start).seconds
In [55]:
             xgb_tnd_2_trn_acc = accuracy_score(y_train,y_pred_tuned_2_train)
          1
          2
             xgb_tnd_2_tst_acc = accuracy_score(y_test, y_pred_tuned_2)
          3
             xgb_tnd_2_rec = recall_score(y_test, y_pred_tuned_2)
             xqb tnd 2 rec prec = precision score(y test, y pred tuned 2)
             xgb_tnd_2_rec_roc_auc = roc_auc_score(y_test, y_pred_tuned_2)
             xgb_tnd_2_rec_F1 = f1_score(y_test,y_pred_tuned_2)
          7
             print('Training Accuracy: ',xgb_tnd_2_trn_acc)
             print('Testing Accuracy: ',xgb_tnd_2_tst_acc)
             print('Recall: ',xgb_tnd_2_rec)
         10
         11
             print('Precision', xgb_tnd_2_rec_prec)
         12
             print('ROC - AUC',xgb_tnd_2_rec_roc_auc)
             print('F1 Score',xgb tnd 2 rec F1)
         Training Accuracy: 0.7644347965338695
         Testing Accuracy: 0.7538965198206533
         Recall: 0.7974846362726883
         Precision 0.7321873769846476
```

ROC - AUC 0.7540726271808579 F1 Score 0.7634423313722808

```
In [56]:
             # Add these values to our model dictionary
          2
             # Since pandas does not allow you to add rows without removing the i
             # we need to recreate the table again
          5
             model_name = 'XGB Tuned 2'
             model arr['XGB Tuned 2'] = xgb tuned 2
          6
          7
          8
             train acc[model name] = xgb tnd 2 trn acc
             test acc[model name] = xqb tnd 2 tst acc
             rec[model_name] = xgb_tnd_2_rec
         10
             prec[model_name] = xgb_tnd_2_rec_prec
         11
         12
             F1[model name] = xgb tnd 2 rec F1
             Roc_Auc[model_name] = xgb_tnd_2_rec_roc_auc
         13
         14
             run_time[model_name] = running_secs_xgb_2
```

## In [57]:

```
measures = pd.DataFrame(index=model_arr.keys(), columns=['Training A measures['Training Accuracy'] = train_acc.values()
measures['Testing Accuracy'] = test_acc.values()
measures['Recall'] = rec.values()
measures['Precision'] = prec.values()
measures['F1 Score'] = F1.values()
measures['Roc-AUC Score'] = Roc_Auc.values()
measures['Runtime (s)'] = run_time.values()
measures
```

#### Out[57]:

	Training Accuracy	Testing Accuracy	Recall	Precision	F1 Score	Roc-AUC Score	Runtime (s)
Logistical Regression	0.745076	0.749413	0.769187	0.738474	0.753518	0.749493	0
Random Forest	0.997313	0.742652	0.780620	0.724115	0.751307	0.742805	7
Decision Tree Classifer	0.997331	0.655327	0.653566	0.654033	0.653799	0.655320	0
XGB Classifier	0.791214	0.748345	0.788767	0.728389	0.757376	0.748509	5
svc	0.769719	0.753541	0.806917	0.727765	0.765300	0.753756	350
GaussianNB	0.714614	0.717743	0.703016	0.722638	0.712692	0.717683	0
KNeighbors	0.796107	0.709487	0.734886	0.697788	0.715857	0.709589	275
XGB Tuned 1	0.757567	0.754466	0.798199	0.732651	0.764022	0.754643	5
XGB Tuned 2	0.764435	0.753897	0.797485	0.732187	0.763442	0.754073	38

These numbers seem slightly better than the initial Xgboost model as well as the baseline. Given the magnitude of data we are working over, over 10,000 records, the gains are marginal at best.

However, interestingly it took less time for the second tuned model to evaluate. The second tuning showed good improvements.

With more computation resources, it would be interesting to see how much higher we can increase the accuracy of the model.

```
In [58]:
              cm_xgb_tnd = confusion_matrix(y_test, y_pred_tuned_2)
           1
           2
              TN_xgb_tnd2, FP_xgb_tnd2, FN_xgb_tnd2, TP_xgb_tnd2 = confusion_matri
           3
              print('True Positive(TP) = ', TP_xgb_tnd2)
           5
              print('False Positive(FP) = '
                                              , FP_xgb_tnd2)
              print('True Negative(TN) = ', TN_xgb_tnd2)
              print('False Negative(FN) = ', FN_xgb_tnd2)
          True Positive(TP)
                                 5580
          False Positive(FP) =
                                 2041
          True Negative(TN)
                                 5013
          False Negative(FN) =
                                 1417
In [59]:
              cm_xgb_tnd = confusion_matrix(y_test,y_pred_tuned_2)
           1
           2
           3
              cm_xgb_tnd = ConfusionMatrixDisplay(confusion_matrix = cm_xgb_tnd, d
           4
           5
              cm_xgb_tnd.plot()
              plt.show()
                                                 5500
                                                 5000
                      5013
                                    2041
            False
                                                 4500
                                                 4000
          True label
                                                - 3500
                                                 3000
                                                - 2500
                      1417
                                    5580
             True
                                                 2000
                                                 1500
```

The tuned model has performed slightly better than the baseline and initial XGBoost model. Since the percentages are small, let's see how many correct predictions this translates too.

True

False

Predicted label

```
In [60]:
             # Find the difference in correct predictions made between the tuned
           1
          2
             # Correct Predictions are defined as the number of TP + TN
           3
             lr_corr_pred = lr_TP + lr_TN # Correct number of predictions made by
           4
           5
             xgb_corr_pred = xgb_TP + xgb_TN # Correct number of predictions mad
          6
          7
             xgb_tnd_corr_pred = TP_xgb_tnd + TN_xgb_tnd
          8
          9
             diff_preds_1 = xgb_corr_pred - lr_corr_pred
         10
             diff_preds_2 = xgb_tnd_corr_pred - xgb_corr_pred
         11
             diff preds 3 = xgb tnd corr pred - lr corr pred
         12
         13
         14
             print("The initial XGBoost model made", diff_preds_1, "more correct pr
             print("The tuned XGBoost model made",diff_preds_2,"more correct pred
         15
             print("The tuned XGBoost model made",diff_preds_3,"more correct pred
         17
         18
```

The initial XGBoost model made -15 more correct predictions than the baseline model.

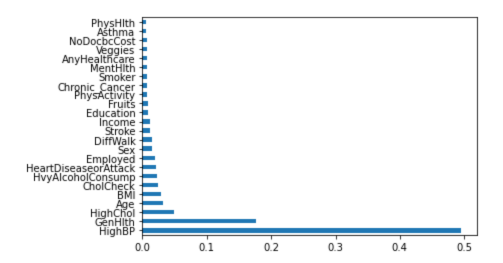
The tuned XGBoost model made 86 more correct predictions than the initial XGBoost model.

The tuned XGBoost model made 71 more correct predictions than the basel ine model.

Through our iterative modeling process we are increasing the accuracy of our model. However, these increases are marginal at best over a dataset that has tens of thousands of values.

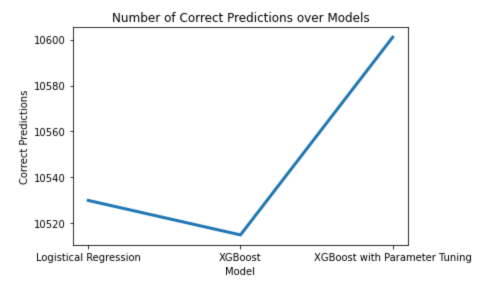
It's unclear if the time and effort spent on tuning the model is worth the gain in accuracy.





There does not seem to be a huge difference in the features. Though the coefficient for HighBP increased and the rest decreased.

```
In [62]:
             x_axis = ["Logistical Regression", "XGBoost", "XGBoost with Paramete
           1
           2
             y_axis = [lr_corr_pred, xgb_corr_pred, xgb_tnd_corr_pred]
           3
             plt.plot(x_axis,y_axis,linewidth = 3)
           4
              plt.xlabel('Model')
           5
              plt.ylabel('Correct Predictions')
           7
              plt.title('Number of Correct Predictions over Models')
           8
           9
             # Show the plot
          10
              plt.show()
```



## **Using Neural Network Models on the Data**

In addition to our own iterative modeling, we wanted to research the techniques experts were finding to be the most accurate in predicting diabetes.

We found several articles that found neural networks to provide the best model including one that used a dataset from a previous BRFSS dataset in a previous year.

The following sources evaluated the implementing different machine learning models on diabetes data. They concluded that neural networks were the best model when evaluating based on accuracy.

- Building Risk Prediction Models for Type 2 Diabetes Using Machine Learning Techniques,
   Xie et. al. link (https://www.cdc.gov/pcd/issues/2019/19 0109.htm)
  - This article used the 2014 data from the survey to create these models.
- Cardiovascular complications in a diabetes prediction model using machine learning: a systematic review, Kee et. al. <u>link (https://link.springer.com/article/10.1186/s12933-023-01741-7)</u>

We created our own neural network based on the data. Due to the size and amount of text generated by neural networks, we ran them on a different notebook. We saved the best model and loaded it here to create the confusion matrix, graphs, etc.

The analysis and notebook containing the optimization of the neural network is <a href="here">here</a> (notebooks/Neural Network Modeling.ipynb)

The neural networks architecture is:

#### Neural

- 3 dense layers
  - 40 neurons in the first layer
  - 20 neurons in the second
  - 10 neurons in the third
- relu activation
- Use sigmoid curve
- · Early Stopping

```
In [63]:
              import keras
           1
             from keras import models
           2
In [64]:
             nn_model = keras.models.load_model('Neural_Network')
In [65]:
           1
             # Predict on training data
           2
             # The data of y_preds_nn is float not binary 0/1 so we cannot compar
           3
             y_pred_nn = nn_model.predict(X_test)
           4
           5
             y_pred_nn
Out[65]: array([[0.6546868],
                 [0.5212415]
                 [0.71433425],
                 [0.02479693],
                 [0.44503072],
                 [0.0013229 ]], dtype=float32)
In [66]:
             # We will round y_preds_nn to 0 or 1 depending on if it's above or b
           1
           2
           3
             y_pred_nn_rnd = np.around(y_pred_nn,0)
           4
             y_pred_nn_rnd
Out[66]: array([[1.],
                 [1.],
                 [1.],
                 . . . ,
                 [0.],
                 [0.],
                 [0.]], dtype=float32)
```

```
In [67]:
             # Calculate metrics below
          1
          2
            nn trn acc = 0.7578 # Pulled from neural network notebook
          3
             nn_tst_acc = accuracy_score(y_test, y_pred_nn_rnd)
             nn rec = recall_score(y_test, y_pred_nn_rnd)
             nn rec prec = precision score(y test, y pred nn rnd)
          7
             nn_rec_roc_auc = roc_auc_score(y_test, y_pred_nn_rnd)
          8
             nn rec F1 = f1 score(y test,y pred nn rnd)
             print('Training Accuracy: ',nn_trn_acc)
         10
             print('Testing Accuracy: ',nn tst acc)
         11
         12
             print('Recall: ',nn_rec)
         13
             print('Precision', nn_rec_prec)
            print('ROC - AUC',nn rec roc auc)
             print('F1 Score',nn_rec_F1)
```

Training Accuracy: 0.7578 Testing Accuracy: 0.7598747420112447 Recall: 0.7884807774760612 Precision 0.7444339495344757 ROC - AUC 0.7599903178562614 F1 Score 0.7658245419211548

```
In [68]:
             # Add these values to our model dictionary
          2
             # Since pandas does not allow you to add rows without removing the i
             # we need to recreate the table again
          5
             model_name = 'Neural Network'
             model arr[model name] = nn model
          7
             train_acc[model_name] = nn_trn_acc
          8
             test acc[model name] = nn tst acc
             rec[model_name] = nn_rec
         10
             prec[model name] = nn rec prec
         12 F1[model name] = nn rec F1
             Roc Auc[model name] = nn rec roc auc
         13
         14
             run_time[model_name] = 48 # Pulled from neural network notebook. Thi
```

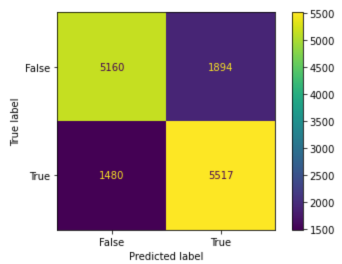
```
In [69]: 1  measures = pd.DataFrame(index=model_arr.keys(), columns=['Training A
2  measures['Training Accuracy'] = train_acc.values()
3  measures['Testing Accuracy'] = test_acc.values()
4  measures['Recall'] = rec.values()
5  measures['Precision'] = prec.values()
6  measures['F1 Score'] = F1.values()
7  measures['Roc-AUC Score'] = Roc_Auc.values()
8  measures['Runtime (s)'] = run_time.values()
9  measures
```

#### Out[69]:

	Training Accuracy	Testing Accuracy	Recall	Precision	F1 Score	Roc-AUC Score	Runtime (s)
Logistical Regression	0.745076	0.749413	0.769187	0.738474	0.753518	0.749493	0
Random Forest	0.997313	0.742652	0.780620	0.724115	0.751307	0.742805	7
Decision Tree Classifer	0.997331	0.655327	0.653566	0.654033	0.653799	0.655320	0
XGB Classifier	0.791214	0.748345	0.788767	0.728389	0.757376	0.748509	5
svc	0.769719	0.753541	0.806917	0.727765	0.765300	0.753756	350
GaussianNB	0.714614	0.717743	0.703016	0.722638	0.712692	0.717683	0
KNeighbors	0.796107	0.709487	0.734886	0.697788	0.715857	0.709589	275
XGB Tuned 1	0.757567	0.754466	0.798199	0.732651	0.764022	0.754643	5
XGB Tuned 2	0.764435	0.753897	0.797485	0.732187	0.763442	0.754073	38
Neural Network	0.757800	0.759875	0.788481	0.744434	0.765825	0.759990	48

```
True Positive(TP) = 5517
False Positive(FP) = 1894
True Negative(TN) = 5160
False Negative(FN) = 1480
```

```
In [71]:
             cm_nn = confusion_matrix(y_test,y_pred_nn_rnd)
           1
           2
             cm_nn = ConfusionMatrixDisplay(confusion_matrix = cm_nn, display_lab
           3
           4
           5
             cm nn.plot()
             plt.show()
```



The neural network identified slightly more true positives and true negatives.

```
In [72]:
             # Find the difference in correct predictions between all models
          2
             # Correct Predictions are defined as the number of TP + TN
             lr corr pred = lr TP + lr TN # Correct number of predictions made by
             xqb corr pred = xqb TP + xqb TN # Correct number of predictions mad
             xgb_tnd_corr_pred = TP_xgb_tnd + TN_xgb_tnd # Correct number of pred
             nn corr pred = TP nn + TN nn # Correct number of predictions made by
          7
          9
             diff preds 1 = xgb corr pred - lr corr pred
         10
             diff_preds_2 = xgb_tnd_corr_pred - xgb_corr_pred
             diff_preds_3 = xgb_tnd_corr_pred - lr_corr_pred
         11
         12
             diff preds 4 = nn corr pred - xqb tnd corr pred
         13
         14
         15
             print("The initial XGBoost model made", diff_preds_1, "more correct pr
             print("The tuned XGBoost model made",diff_preds_2,"more correct pred
         16
             print("The tuned XGBoost model made", diff preds 3, "more correct pred
         17
             print("The neural network made", diff_preds_4, "more correct predictio")
```

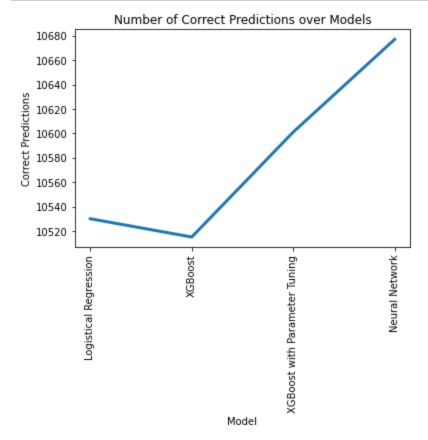
The initial XGBoost model made -15 more correct predictions than the ba seline model.

The tuned XGBoost model made 86 more correct predictions than the initi al XGBoost model.

The tuned XGBoost model made 71 more correct predictions than the basel ine model.

The neural network made 76 more correct predictions than the tuned XGBo ost model.

```
In [73]:
             x_axis = ["Logistical Regression", "XGBoost", "XGBoost with Paramete
           1
           2
             y_axis = [lr_corr_pred, xgb_corr_pred, xgb_tnd_corr_pred,nn_corr_pre
           3
             plt.plot(x_axis,y_axis,linewidth = 3)
             plt.xlabel('Model')
           5
             plt.ylabel('Correct Predictions')
             plt.title('Number of Correct Predictions over Models')
           7
             plt.xticks(rotation=90)
           8
          10
             # Show the plot
             plt.show()
          11
```



Interestingly, like the paper the neural network is the most accurate model. We did not seem the same level of difference in accuracy that was observed in the paper (82% vs 79%), however, we may not have the computing resources to add more layers and create a denser neural network.

```
In [74]: 1 # Calculate percentage increase in accuracy between the most accurat
2 print("The neural network is", round((nn_corr_pred - lr_corr_pred)/di
```

The neural network is 0.21 % more accurate than the base model

Though iterative modeling, we've improved the efficacy of our model by 125 predictions. This represents an increase of around ~0.2%.

### **Final Model Evaluation**

For the final model, we are recommending our baseline model the logistic regression model for use by the CDC. Even though we iteratively improved the accuracy and precision metrics across the XGB, tuned models, and neural network, the increase in these metrics is not worth the time and resources it takes to train and tune these models.

We only used a small sample of the data available in the study in our training/testing (31K vs 440K records). Deploying this model across data that goes into the hundreds of thousands or millions of records if you consider the previous years data, may not be economical if you consider time, computational resources, and FTEs it takes.

Logistical Regression probably gives the CDC what they need to reasonably determine the likelyhood of diabetes with a limited budget.

In [75]:

1 measures

#### Out[75]:

	Training Accuracy	Testing Accuracy	Recall	Precision	F1 Score	Roc-AUC Score	Runtime (s)
Logistical Regression	0.745076	0.749413	0.769187	0.738474	0.753518	0.749493	0
Random Forest	0.997313	0.742652	0.780620	0.724115	0.751307	0.742805	7
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XGB Classifier	0.791214	0.748345	0.788767	0.728389	0.757376	0.748509	5
svc	0.769719	0.753541	0.806917	0.727765	0.765300	0.753756	350
GaussianNB	0.714614	0.717743	0.703016	0.722638	0.712692	0.717683	0
KNeighbors	0.796107	0.709487	0.734886	0.697788	0.715857	0.709589	275
XGB Tuned 1	0.757567	0.754466	0.798199	0.732651	0.764022	0.754643	5
XGB Tuned 2	0.764435	0.753897	0.797485	0.732187	0.763442	0.754073	38
Neural Network	0.757800	0.759875	0.788481	0.744434	0.765825	0.759990	48

### Recommendations

- When optimizing for resource costs, logistical regression is the best choice.
- The top features across the models for predicting diabetes were high blood pressure, BMI, general activity.
- High Blood Pressure was the best predictor on whether a patient has diabetes/prediabetes. The CDC needs to consider campaigns and educational outreach informing people on the importance of regularly checking their blood pressure.

# **Future Projects**

- Go through the data sets the BRFSS has produced through the years. Measure the rate of diabetes and other factors to predict the trends across the country of these sicknesses.
- Use the model to create an application on the CDC's website that allows a person to enter their data and get a diabetic risk score.

# **Reproduction Steps**

- Download CSV from this link: <a href="https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system">https://www.kaggle.com/datasets/cdc/behavioral-risk-factor-surveillance-system</a>)
- 2. Save CSV to file and run steps from the data cleaning notebook
- 3. Run the main notebook to the neural networks page.
- 4. Run the neural networks notebook
- 5. Continue running the main notebook to the end.