EPL Match Prediction Report

I. Definition

Project Overview

For this capstone project, I will use an English Premier League Football dataset from http://football-data.co.uk/englandm.php. This is a binary classification problem similar to the CharityML in Supervised Learning section of the MLND. The legal sports-betting market in the U.S. was worth an estimated USD270 million in 2017 -- with another USD2.5 billion to USD3 billion in black market betting, according to research firm Eilers & Krejcik Gaming, LLC.

Even though it is not legalized in many other countries like India, but for my own interest, I want to build a predictive model capable of predicting if the home team will win a football match. Usually betting is conducted with human instincts but now we can use some machine learning algorithm to predict the result of the future matches also.

There are reports related to sports prediction using machine learning. Some of them I have listed below:

<u>Using Machine Learning to Predict the Outcome of English County twenty over</u>
 <u>Cricket Matches</u>

"It is possible to predict the winner of English county twenty-twenty cricket games in almost two-thirds of instances."

https://gz.com/233830/world-cup-germany-argentina-predictions-microsoft/

For the 2014 World Cup, Bing correctly predicted the outcomes for all of the 15 games in the knockout round.

Predicting Football Results With Statistical Modelling

"Something that becomes clear from the results is that Twitter contains enough information to be useful for predicting outcomes in the Premier League"

I am a regular viewer of football around the globe, my favorite club is Manchester United and I follow English Premier League religiously and now I am excited I can use my knowledge of machine learning to have some fun with data.

Problem Statement

The problem is to use the existing dataset of EPL obtained through http://football-data.co.uk/englandm.php and use it to train some supervised learning algorithms to predict the matches of EPL. I want to predict whether a home team is gonna win the match by training some supervised learning algorithms. This is a pure binary classification problem and II will use a Logistic Regression model as a benchmark model. Then I will use Support Vector Machine machine algorithm and finally I will implement Ensemble Learning to beat my benchmark model score. This dataset has enough points to train a machine learning algorithm & predict results on the data.

Metrics

I will be using accuracy_score & f1_score from sklearn.metrics to evaluate both the benchmark and my final model. The goal of this project is to predict the winner of 'Home Team' accurately. So accuracy as a metric to evaluate a model's performance is appropriate. However, predicting a team is not going to win is not that much important, hence, a model's ability to precisely predict the winner of a 'home team' is more important than the model's ability to recall those teams.

For this reason, we can use the f1-score as a metric which considers both precision & recall.

Let's discuss these two metrics in detail:

		Predicted class	
90000000000000000000000000000000000000		Class = Yes	Class = No
Actual Class	Class = Yes	True Positive	False Negative
	Class = No	False Positive	True Negative

True positive and true negatives are the observations that are correctly predicted and therefore shown in green. We want to minimize false positives and false negatives so they are shown in red color. These terms are a bit confusing. So let's take each term one by one and understand it fully.

True Positives (TP) - These are the correctly predicted positive values which mean that the value of the actual class is yes and the value of the predicted class is also yes.

E.g. if the actual class value indicates that if a home team is going to win and predicted class tells you the same thing.

True Negatives (TN) - These are the correctly predicted negative values which mean that the value of the actual class is no and value of the predicted class is also no. E.g. if actual class says the home team didn't win and predicted class tells you the same thing.

False positives and false negatives, these values occur when your actual class contradicts with the predicted class.

False Positives (FP) – When actual class is no and predicted class is yes. E.g. if actual class says the home team didn't win but predicted class tells you that this passenger will survive.

False Negatives (FN) – When actual class is yes but predicted class in no. E.g. if the actual class value indicates that the home team wins and predicted class tells you that away team is going to win.

Once you understand these four parameters then we can calculate Accuracy, Precision, Recall and F1 score.

Accuracy - Accuracy is the most intuitive performance measure and it is simply a ratio of correctly predicted observation to the total observations. One may think that, if we have high accuracy then our model is best. Yes, accuracy is a great measure but only when you have symmetric datasets where values of false positive and false negatives are almost the same. Therefore, you have to look at other parameters to evaluate the performance of your model.

Accuracy = TP+TN/TP+FP+FN+TN

Precision - Precision is the ratio of correctly predicted positive observations to the total predicted positive observations. The question that this metric answer is of all passengers that labeled as survived, how many actually survived? High precision relates to the low false positive rate.

Precision = TP/TP+FP

Recall (Sensitivity) - Recall is the ratio of correctly predicted positive observations to all observations in actual class - yes.

Recall = TP/TP+FN

F1 score - F1 Score is the weighted average of Precision and Recall. Therefore, this score takes both false positives and false negatives into account. Intuitively it is not as easy to understand as accuracy, but F1 is usually more useful than accuracy, especially if you have an uneven class distribution. Accuracy works best if false positives and false negatives have a similar cost. If the cost of false positives and false negatives are very different, it's better to look at both Precision and Recall. I

F1 Score = 2*(Recall * Precision) / (Recall + Precision)

II. Analysis

Exploratory Data Analysis of EPL sessions

In the <u>Datasets</u> I have downloaded all the sessions data separately from the http://football-data.co.uk/englandm.php . Then I consolidated all the datasets into a final dataset called My Capstone Dataset.csv file.

Dataset Exploration:

	FTR	HTP	ATP	НМ1	HM2	нмз	AM1	AM2	АМЗ	HTGD	ATGD	DiffFormPts	DiffLP
30	Н	1.25	1.00	D	D	W	D	W	L	0.50	0.25	0.25	-16.0
31	NH	0.75	0.25	L	L	W	D	L	L	-0.50	-0.75	0.50	-2.0
32	Н	1.00	1.00	L	D	W	D	W	L	0.00	0.25	0.00	-3.0
33	NH	0.75	0.50	L	L	W	D	L	D	-0.25	-0.25	0.25	3.0
34	NH	1.00	1.50	D	L	W	W	W	L	0.00	0.75	-0.50	3.0

The original dataset has 6080 data points and 42 input variables, I reduced the dimension to 12 features which are relevant to predict the target variable which is **FTR** (**Full Time Result.**)

Feature set Explanation:

Input Feature Set Explanation

- HTP Home team points
- ATP Away team points
- HTGD Home team goal difference
- ATGD away team goal difference
- DiffFormPts Difference in points
- DiffLP Difference in last years prediction
- HM Home Match
- AM Away Match

Target Variable

FTR: Full-Time Result (H=Home Win, D=Draw, A=Away Win)

Let's explore this data a little bit

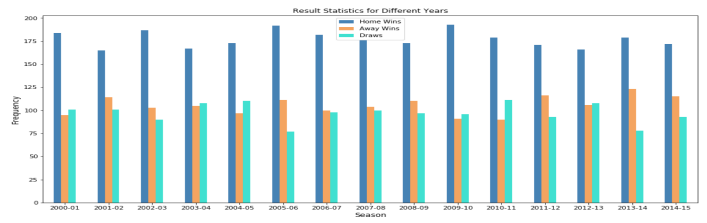
```
#what is the win rate for the home team?
# Total number of matches.
n_matches = data.shape[0]
# Calculate number of features. -1 because we are saving one as the target variable (win/lose/draw)
n_{features} = data.shape[1] - 1
# Calculate matches won by home team.
n_homewins = len(data[data.FTR == 'H'])
# Calculate win rate for home team.
win rate = (float(n homewins) / (n matches)) * 100
# Print the results
print "Total number of matches: {}".format(n_matches)
print "Number of features: {}".format(n_features)
print "Number of matches won by home team: {}".format(n_homewins)
print "Win rate of home team: {:.2f}%".format(win_rate)
Total number of matches: 5600
Number of features: 12
Number of matches won by home team: 2603
Win rate of home team: 46.48%
```

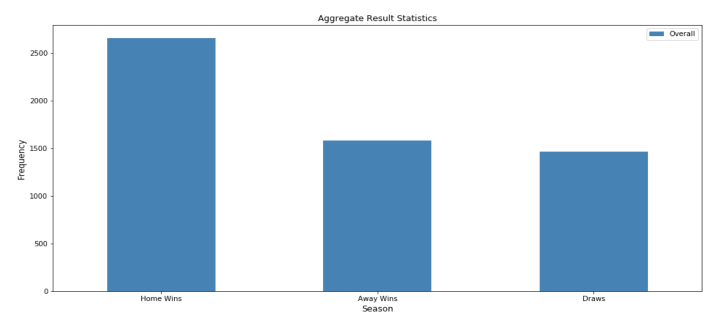
Now I will use all separate sessions CSV files to find answers to some interesting questions listed below:

How many matches have been won/drawn by home teams?

What is the win percentage for home teams & away teams?

While searching for above two answers I will create some visualization for clear understanding.





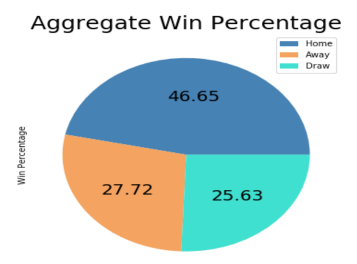
Displaying the overall results

	Home wins	Away Wins	Draws		
Overall	2659	1580	1461		

So our first question was How many matches have been won/drawn by home teams?

From the above visualization, we can clearly see more matches have won by the home teams only. It is too common in football matches as home team can feel advantage while

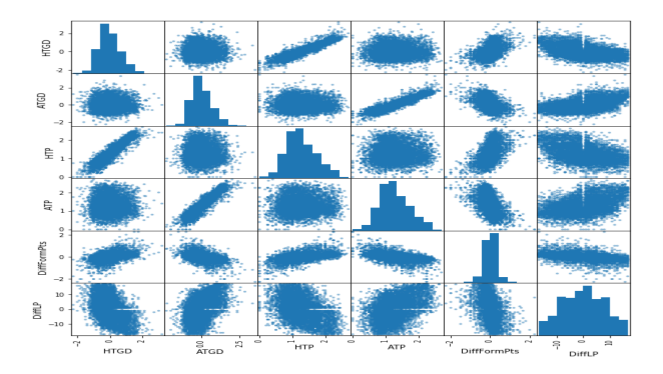
the home crowd is supporting behind them and pitch conditions and other playing factors are also in favor of them. Now, let's visualize the win percentage for Home & Away teams.



So from above visualization also we can see 46% times Home team wins, 28% time Away team wins and 25% time matches are drawn. So the majority of the times home team usually wins.

Visualizing distribution of data

The scatter matrix is plotting each of the columns specified against each other column. You would have observed that the diagonal graph is defined as a histogram, which means that in the section of the plot matrix where the variable is against itself, a histogram is plotted. Scatter plots show how much one variable is affected by another. The relationship between two variables is called their correlation negative vs positive correlation.



Algorithms & Techniques

Some questions I think are right can be asked which are listed below:

- What model should we use?
- What are the features (the aspects of a game) that matter the most to predicting a team wins?
- Does being the home team give a team the advantage?

I will use three *supervised learning* algorithm listed below to train and predict on my dataset. I will deploy a train_test_split to shuffle & split data into a training & testing set and will compare the metrics of *three* learning algorithms and will choose the best one amongst them. Finally, I will perform a hyperparameter tuning to optimize my final classifier. Then I will try some prediction on my test data set.

Support Vector Machine(SVM):

What is a Support Vector Machine?

"Support Vector Machine" (SVM) is a supervised machine learning algorithm which can be used for both classification or regression challenges. However, it is mostly used in classification problems. In this algorithm, we plot each data item as a point in n-dimensional space (where n is a number of features you have) with the value of each feature being the value of a particular coordinate. Then, we perform classification by finding the hyperplane that differentiates the two classes very well. Support Vectors are simply the co-ordinates of individual observation. Support Vector Machine is a frontier which best segregates the two classes (hyper-plane/ line).

How does SVM works?

Support vector machines focus only on the points that are the most difficult to tell apart, whereas other classifiers pay attention to all of the points.

The intuition behind the support vector machine approach is that if a classifier is good at the most challenging comparisons (the points in B and A that are closest to each other in Figure 2), then the classifier will be even better at the easy comparisons (comparing points in B and A that are far away from each other.

In SVM, it is easy to have a linear hyperplane between two classes. But, another burning question which arises is, should we need to add this feature manually to have a hyperplane. No, SVM has a technique called the kernel trick. These are functions which take low dimensional input space and transform it to a higher dimensional space i.e. it converts not separable problem to separable problem, these functions are called kernels. It is most useful in non-linear separation problem. Simply put, it does some extremely complex data transformations, then find out the process to separate the data based on the labels or outputs you've defined

Tuning the parameters of SVM:

Tuning parameters value for machine learning algorithms effectively improves the model performance. Let's look at the list of parameters available with SVM

```
sklearn.svm.SVC(C=1.0, kernel='rbf', degree=3, gamma=0.0, coef0=0.0, shrinking=True, probability=Fals e,tol=0.001, cache_size=200, class_weight=None, verbose=False, max_iter=-1, random_state=None)
```

I am going to discuss one parameter having a higher impact on model performance, "kernel".

This is a pure classification problem and SVM is well suited for classification, so we can use SVM to predict whether a home team is going to win or not.

we have various options available with the kernel, like, "linear", "rbf", "poly" and others (default value is "rbf"). Here "rbf" and "poly" are useful for non-linear hyperplane. I will use 'rbf' as my data points are non-linear here and features are only 12.

Reference:

https://stats.stackexchange.com/questions/23391/how-does-a-support-vector-machine-svm-work

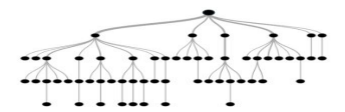
Random Forest:

What is Random Forest?

Random Forest is a versatile machine learning method capable of performing both regression and classification tasks. It also undertakes dimensional reduction methods, treats missing values, outlier values and other essential steps of data exploration, and does a fairly good job. It is a type of ensemble learning method, where a group of weak models combines to form a powerful model.

What is a Decision Tree?

A decision tree is a type of supervised learning algorithm (having a predefined target variable) that is mostly used in classification problems. It works for both categorical and continuous input and output variables. In this technique, we split the population or sample into two or more homogeneous sets (or sub-populations) based on most significant splitter/differentiator in input variables.



How Does Random Forest Work?

Random forest is like a bootstrapping algorithm with Decision tree (CART) model. Say, we have 1000 observation in the complete population with 10 variables. Random forest tries to build multiple CART model with a different sample and different initial variables. For instance, it will take a random sample of 100 observation and 5 randomly chosen initial variables to build a CART model. It will repeat the process (say) 10 times and then make a final prediction on each observation. Final prediction is a function of each prediction. This final prediction can simply be the mean of each prediction.

In our case only 12 input features are available and Random Forest can perform very well by using the decision tree(CART) model.

Reference:

https://www.analyticsvidhya.com/blog/2014/06/introduction-random-forest-simplified/

XGBoost:

What is boosting?

Boosting is an ensemble technique in which the predictors are not made independently, but sequentially. This technique employs the logic in which the subsequent predictors learn from the mistakes of the previous predictors. Therefore, the observations have an unequal probability of appearing in subsequent models and ones with the highest error appear most. (So the observations are not chosen based on the bootstrap process, but based on the error). The predictors can be chosen from a range of models like decision trees, regressors, classifiers etc. Because new predictors are learning from mistakes committed by previous predictors, it takes less time/iterations to reach close to actual predictions. But we have to choose the stopping criteria carefully or it could lead to overfitting on training data. Gradient Boosting is an example of a boosting algorithm.

What is gradient boosting?

Gradient boosting is a machine learning technique for regression and classification problems, which produces a prediction model in the form of an ensemble of weak prediction models, typically decision trees. (Wikipedia definition).

The objective of any supervised learning algorithm is to define a loss function and minimize it. Let's see how maths work out for Gradient Boosting algorithm. Say we have the mean squared error (MSE) as loss defined as:

$$Loss = MSE = \sum (y_i - y_i^p)^2$$

where, y_i = ith target value, y_i^p = ith prediction, $L(y_i, y_i^p)$ is Loss function

We want our predictions, such that our loss function (MSE) is minimum. By using gradient descent and updating our predictions based on a learning rate, we can find the values where MSE is minimum.

$$y_i^p = y_i^p + \alpha * \delta \sum_i (y_i - y_i^p)^2 / \delta y_i^p$$
 which becomes, $y_i^p = y_i^p - \alpha * 2 * \sum_i (y_i - y_i^p)$

where, α is learning rate and $\sum (y_i - y_i^p)$ is sum of residuals

So, we are basically updating the predictions such that the sum of our residuals is close to 0 (or minimum) and predicted values are sufficiently close to actual values.

What is XGBoost?

XGBoost is an algorithm that has recently been dominating applied machine learning and Kaggle competitions for structured or tabular data. XGBoost is an implementation of gradient boosted decision trees designed for speed and performance. XGBoost stands for eXtreme Gradient Boosting.

As Tianqi Chen said:

The name xgboost, though, actually refers to the engineering goal to push the limit of computations resources for boosted tree algorithms. Which is the reason why many people use xgboost

Generally, XGBoost is fast. Really fast when compared to other implementations of gradient boosting.

XGBoost is consists of classification & regression tree and works well for both problems. In our case also it is a classification problem & I am sure xgboost will have good performance

Reference:

https://machinelearningmastery.com/gentle-introduction-xgboost-applied-machine-learning/

Benchmark

I will use an untuned Logistic Regression classifier from 'sklearn' to benchmark my final result. When the outcome (dependent variable) has only a limited number of possible values (in our cases hometeam win or loss), Logistic Regression is used when the response variable is categorical in nature.

III. Methodology

Data Preprocessing:

- We want to do is say given all of those other features and try to predict the full-time result (FTR)
- We will separate the data into feature set and the target variable

Then we are gonna standardize it, which means it's all gonna be on the same scale. It means we want all of our data to be in integer format and we want it all to be on the same scale, so it's not like we have like one feature is in the hundreds of thousands and then the other feature is in between 1-10. It will improve the prediction capability of our model.

Once we have standardized our data, then we're going t add these features which are the last three wins for both sides (HM1, HM2, HM2 | AM1, AM2, AM3).

Now if we can look back at the data some of the data was categorical.

	FTR	HTP	ATP	НМ1	HM2	нмз	AM1	AM2	АМЗ	HTGD	ATGD	DiffFormPts	DiffLP
30	Н	1.25	1.00	D	D	W	D	W	L	0.50	0.25	0.25	-16.0
31	NH	0.75	0.25	L	L	W	D	L	L	-0.50	-0.75	0.50	-2.0
32	Н	1.00	1.00	L	D	W	D	W	L	0.00	0.25	0.00	-3.0
33	NH	0.75	0.50	L	L	W	D	L	D	-0.25	-0.25	0.25	3.0
34	NH	1.00	1.50	D	L	W	W	W	L	0.00	0.75	-0.50	3.0

We want all of our data to be a number. We want it to be some continuous variables, there should be no discrete numbers. We are gonna pre-process those features by creating a new data frame and find those feature columns that are categorical by saying if it's data type is equal to an object instead of an integer so that we remove all categorical features. We will only have one categorical variable that is our label FTR. Below is our dataset with all continuous variables:

reacure values.	Feature	values:
-----------------	---------	---------

	HTP	ATP	HM1_D	HM1_L	HM1_W	HM2_D	HM2_L	HM2_W	HM3_D	HM3_L	 AM2_D	AM2_L	AM2_W	AM3_D	AM3_L	AM3_W	HTG
30	-0.043829	-0.611968	1	0	0	1	0	0	0	0	 0	0	1	0	1	0	0.75371
31	-1.120644	-2.238746	0	1	0	0	1	0	0	0	 0	1	0	0	1	0	-0.73708
32	-0.582236	-0.611968	0	1	0	1	0	0	0	0	 0	0	1	0	1	0	0.00831
33	-1.120644	-1.696487	0	1	0	0	1	0	0	0	 0	1	0	1	0	0	-0.36438
34	-0.582236	0.472551	1	0	0	0	1	0	0	0	 0	0	1	0	1	0	0.00831

5 rows × 24 columns

Now all input features are continuous variables, now we can train and we can split our model into a training & testing set.

Implementation:

For the next step, we are gonna build this model. I have defined some helper functions in which are gonna help us train the model.

Here I defined three functions which are listed below & explained:

train classifier()

This function will take the classifier name & X_train & y_train variable as argument and will train the model. It will return the time taken to train the classifier.

predict_labels()

It will take classifier name, features, and target as arguments. This function will predict using a fit classifier based on f1-score. It will return the time taken to predict and also the f1-score & the accuracy score.

train_predict()

This is a cumulative function of the above two functions, here I will train and predict the model. It will take classifier, X_train, y_train, X_test, y_test as arguments.

Initial Model Evaluation:

Here I will define three of my chosen classifier along with my benchmark model (Logistic Regression) and will call train_predict() function which will help me reporting the training & prediction time on both train & test set along with the f1-score & accuracy score.

Next, I have attached the output of my four classifier

7571 0.680360360360 9692 4238 0.986306306306 0.694954954954954954 4054 F1 score and accuracy score for training set: 0.6205 , 0.6654. Made predictions in Made 4238 0.986306306306 0.69495495495495495495495495495495495495495	Training a LogisticRegres sion using a training set size of 5550 Trained model in 0.0260 seconds Made predictions in 0.0040 seconds. 0.621561035255	Training a SVC using a training set size of 5550 Trained model in 1.8139 seconds Made predictions in 1.0184 seconds. 0.620453572956	Training a RandomForestCl assifier using a training set size of 5550 Trained model in 0.0799 seconds Made predictions in 0.0120 seconds.	Training a XGBClassifier using a training set size of 5550 Trained model in 0.4477 seconds Made predictions in 0.0220 seconds.
seconds. F1 score and seconds. seconds. F1 score and accuracy score F1 score and F1 score and	Made predictions in 0.0040 seconds. 0.621561035255 7571 0.665405405405 4054 F1 score and accuracy score for training set: 0.6216, 0.6654. Made predictions in 0.0000 seconds.	predictions in 1.0184 seconds. 0.620453572956 7822 0.680360360360 3604 F1 score and accuracy score for training set: 0.6205 , 0.6804. Made predictions in 0.0110 seconds. F1 score and	Made predictions in 0.0120 seconds. 0.985378991919 9692 0.986306306306 3063 F1 score and accuracy score for training set: 0.9854, 0.9863. Made predictions in 0.0020 seconds.	Made predictions in 0.0220 seconds. 0.652147113211 4238 0.694954954954 955 F1 score and accuracy score for training set: 0.6521, 0.6950. Made predictions in 0.0020 seconds.

	0.6818 ,		for test set:
0.6957 ,		0.6923 ,	0.7451 ,
0.7200.		0.6800.	0.7400.

When I coded the program for this section initially, surprisingly I got highest f1-score & accuracy score in my benchmark model which is Logistic Regression and XGBoost was performing worst. Then I went through my dataset and found that if I am including initial 3 weeks result then the data is linearly separable and Logistic regression was performing well, Then I removed those initial three weeks data as EPL matches are too unpredictable in the sessions starting phase and will not help us to predict the future matches.

Refinement

- From the above result, my benchmark model *Logistic Regression's* f1 score & accuracy score on the test set is 0.6957, 0.7200 respectively.
- Even though Random Forest performed very well on training data but failed on test data
- It is clearly visible from the above result that XgBoost is the best model for this problem.

Tuning the parameters of XGBoost

GBDT Hyper Parameter Tuning

Hyper Parameter	Tuning Approach	Range	Note
# of Trees	Fixed value	100-1000	Depending on datasize
Learning Rate	Fixed => Fine Tune	[2 - 10] / # of Trees	Depending on # trees
Row Sampling	Grid Search	[.5, .75, 1.0]	
Column Sampling	Grid Search	[.4, .6, .8, 1.0]	
Min Leaf Weight	Fixed => Fine Tune	3/(% of rare events)	Rule of thumb
Max Tree Depth	Grid Search	[4, 6, 8, 10]	
Min Split Gain	Fixed	0	Keep it 0

Best GBDT implementation today: https://github.com/tqchen/xqboost by **Tianqi Chen** (U of Washington)



Now that we know the XGBoost is the best model, let's optimize this model. In this case, I have optimized it by tuning the hyperparameter. From scikit-learn, Grid search module was imported which is basically a brute forcing method, we can brute force all the possible combinations of all the hyperparameters. First I will create an initial set of hyperparameters here and will initialize the xgboost classifier and then we will make f1-score function given the initial parameters we defined earlier. It will find the ideal set of parameters and will report the increased f1-score & accuracy score.

IV. Results

My final tuned XGBoost classifier is producing below result:

On the test set, the accuracy score is **80**% and we can easily validate my final trained classifier is performing better than any other classifier I introduced earlier. Only RandomForest classifier was performing well on training set but on the test set my final XGBoost model is more optimized and which is more important for the prediction job.

Final Model Evaluation

Let's find out the difference between optimized & unoptimized metrics of my final XGBoost classifier on the test set

Metric	Unoptimized Model	Optimized Model
Accuracy Score	0.74	0.80
F-Score	0.74	0.78

My **benchmark model** *Logistic Regression* f_score & Accuracy Score was 0.6957, 0.7200 respectively on the test set.

Robustness of the model:

To test the robustness of my final classifier I introduced a never seen dataset to my final classifier. It is located at Datasets folder in the name of test.csv. It is the data of 2015-16 sessions of EPL. I performed the same pre-processing on this data and fed it to my optimized xgboost classifier. Below is the result snippet:

```
# Lets predict the result on the training data
f1, acc = predict_labels(clf, X_train, y_train)
print ("F1 score and accuracy score for training set: {:.4f} , {:.4f}.".format(f1 , acc))
# Lets predict the result on the test data
f1, acc = predict_labels(clf, X_test, y_test)
print ("F1 score and accuracy score for test set: {:.4f} , {:.4f}.".format(f1 , acc))

Made predictions in 0.0330 seconds.
F1 score and accuracy score for training set: 0.8339 , 0.8576.
Made predictions in 0.0030 seconds.
F1 score and accuracy score for test set: 0.8095 , 0.8400.
```

To my surprise it, the classifier accuracy & f1-score reported beyond 80%. So my final classifier performance is not getting affected by any perturbations.

Justification

By using my final model we can predict the results of an EPL game which is statistically more significant than pure guessing the result of a match with human instinct.

Conclusion

Free Form Visualization

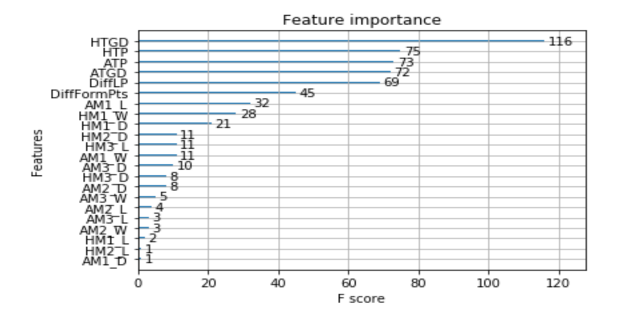
While exploring the data I guess below three features are more important to predict the future match result:

- HTP Home team points
- HTGD Home team goal difference
- DiffFormPts Difference in points

Let me check with the feature importance of XGBClassifier.

```
clf = xgb.XGBClassifier(seed=2)
clf.fit(X_train, y_train)
# TODO: Extract the feature importances using .feature_importances_
importances = clf.feature_importances_

from xgboost import plot_importance
# plot feature importance
plot_importance(clf)
plt.show()
```



Seems like HTGD(Home Team Goal Difference) is the most determining factor in our future match prediction. HTP(Home Team Points) and ATP(Away Team Points) also have a higher importance to predict FTR. It is true in every football match if one team has a higher goal difference then it means the team is scoring more goals in regular interval. By using the above visualization we can clearly see this feature is a determining factor to predict the result of a match

Summary & Reflection

I have completed the entire football match prediction project and let's summarize the entire process

- In this project first I performed exploratory data analysis(EDA) with some simple visualization where it was clearly displayed that in EPL home team wins.
- Then I dropped all irrelevant features from the data and took only 12 input features to do the prediction job.
- A scatter matric was visualized to show the correlation between the input features.
- Then we pre-processed the data by converting all categorical features into numerical features and scaled all the data by using the sklearn scaling method in the preprocessing module.
- After pre-processing, the dataset was shuffled & split into a training & testing set.
- Three helper functions were defined to report the prediction time, f1-score & accuracy score on training & testing data.
- All classifier was compared and the best classifier was chosen on the basis of performance on the test set. After that best classifier which was XGBoost was

- tuned by using Grid search method from sklearn and the improved score was reported.
- I have tested the robustness of the model by introducing a never seen model to the final classifier & reported the score. Surprisingly, it performed better.
- Finally, I have used feature importance of xgboost classifier and found the most important feature HTGD (Home Team Goal Difference), which is the single most important feature in the prediction of winning of a home team.

Improvement

- Possible improvement can be a web app taking the input of the two teams & predicting the winner based on the model trained above.
- We can implement more data features like twitter sentiment analysis on pre-match analysis by training a deep neural network. For this purpose, we can use millions of data points affecting a football match result. In FIFA 2018 world cup bing tried a similar thing by predicting some matches result accurately.
- Possibly a neural network with a much bigger feature rich dataset can predict more accurately than my final model.