

Machine Madness - A Machine Learning approach to predicting March Madness Brackets

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Abstract

March Madness is the NCAA Men's Division I Basketball Championship tournament that happens every March. The tournament is organized by the National Collegiate Athletic Association (NCAA). It has 64 tournament matches. The aim of this project is to come up with a good machine learning based model has the best classification accuracy in predicting the march madness bracket.

1 Introduction

Since 1939, the best colleges and universities across the United States have participated in a yearly tournament called the NCAA Men's Basketball Championship. This basketball tournament has become one of the most popular and famous sporting tournaments in the United States. Millions of people around the country participate in contests in which the participants make their best guesses on who will win each game throughout the tournament. These types of contests have become so popular that more than 11 million brackets were filled out on ESPN.com in 2014. One billion dollars was even being rewarded to anyone who achieved a "perfect bracket" (guessed every game correctly). Every game is unpredictable, and the teams that are supposedly the "better team" sometimes end up losing. This is called an upset in basketball lingo, and happens regularly throughout the tournament. Because of these upsets, it can be difficult to correctly guess the winner of each game. The tournament can be so unpredictable that the time period over which the tournament runs has been termed March Madness. Since there are 64 games played in the NCAA tournament, it is nearly impossible to predict a perfect bracket. High Point Enterprise, a morning paper from North Carolina, stated that "you have a much greater chance of winning the lottery, shooting a hole-in-one in golf or being struck by lightning". They estimated that the chances of predicting a perfect bracket are 1 in 9.2 quintillion. It became clear that developing a model that provided perfect win/loss classification was unrealistic,

so instead we focused on improving the prediction accuracy of individual games. The problem at hand is not classification of individual teams, but rather predicting the outcome of a match between any two teams.

1.1 Objectives

Our goal was to identify key factors in predicting NCAA tournament wins and to find a model that would perform well in the Kaggle competition. The Kaggle competition had the following stages:

- Predict outcomes of the 2015 NCAA Tournament
- Evaluate performance against benchmarks
- Evaluate log loss with actual outcomes

For each stage we submitted a list \hat{y} of probabilities (values between 0 and 1) that each team in the tournament would defeat every other team in the tournament, regardless of whether this match-up actually occurs. For this year, this was $m = 2278$ predictions. We were judged based on the log-loss $L(y - \hat{y})$, or the predictive binomial deviance, of the games that actually occurred.

$$L(y|\hat{y}) = -1/n * \sum_{i=1}^n [y_i \cdot \log(\hat{y}_i + (1 - y_i) \cdot \log(1 - \hat{y}_i))]$$

where n is the actual number of games played in the tournament (67), y_i is the actual binary outcome of each game, and \hat{y}_i is the corresponding submitted probability. If the opponents in game i are teams A and B, then $\hat{y}_i(A, B)$ is the predicted probability that team A beats team B, and $\hat{y}_i(B, A) = 1 - \hat{y}_i(A, B)$

The goal is to find a set of predictions \hat{y} that minimizes $L(y - \hat{y})$ for the unknown outcome vector y . This scoring method heavily penalizes being simultaneously confident and wrong. If we say Team A will beat Team B with probability 1 and Team B wins, then our log-loss is infinite (although Kaggle bounded the log-loss function to avoid infinite scores). At the same time, it is important to balance between being too conservative and too confident. If we are too

conservative (e.g. $\hat{y} \approx 0.5$), then we will never accrue points, but if we are too confident, we make ourselves vulnerable to huge losses.[1]

Table 1: Dataset provided by Kaggle

| Field | Description |
|----------------|---|
| season | indicates the year in which the tournament was played |
| dayzero | tells you the date corresponding to daynum=0 during that season. |
| region W/X/Y/Z | the four regions in the final tournament are always named W X Y and Z. |
| wscore | this identifies the number of points scored by the winning team. |
| lteam | this identifies the id number of the team that lost the game. |
| lscore | this identifies the number of points scored by the losing team. |
| numot | this indicates the number of overtime periods in the game - an integer 0 or higher. |
| wloc | this identifies the location of the winning team. If the winning team was the home team - this value will be H. |
| wfgm | field goals made |
| wfga | field goals attempted |
| wfgm3 | three pointers made |
| wfga3 | three pointers attempted |
| wftm | free throws made |
| wfta | free throws attempted |
| wor | offensive rebounds |
| wdr | defensive rebounds |
| wast | assists |
| wto | turnovers |
| wstl | steals |
| wblk | blocks |
| wpf | personal fouls |

2 Challenges

2.1 Upsets

Every game is unpredictable, and the teams that are supposedly the "better team" sometimes end up losing. This is called an upset in basketball lingo, and happens regularly throughout the tournament. Because of these upsets, it can be difficult to correctly guess the winner of each game. The tournament can be so unpredictable.[2]

2.2 Chance of predicting a perfect bracket

Since there are 64 games played in the NCAA tournament, the odds of forecasting a perfect bracket are astronomical. Each game win/lose has a probability of 1/2. The complexity of the bracket prediction is $1/2 \cdot 1/2 \cdot 1/2$ (63 times) which is 1 in *approx* 9.2 quintillion chance. This is lower than the chance of winning a lottery.

2.3 Unpredictable events

There are multiple events that happen during a match that cannot be predicted. The best example being injury of a key player. A new player being added to a team is one other such example. Playing at home ground vs playing elsewhere sometimes also affects the outcome of the match.

2.4 Team Dynamics

The dynamics of a team change yearly. New players could be added, older key players could be removed. This makes it difficult to compare the performance of a new team to a old team as the logistics and features of an older team do not represent the newer team accurately.

3 Data

The main source of data is Kaggle. The data is made available as .csv files. Kaggle provides regular season wins, losses, point differences, dates, and game locations (home/away) as well as tournament wins, losses, point differences, seeds, and dates of games. This data comes from the season and tournament matches for the years 2005-2015. [1]. The data provided by Kaggle mainly consists of match statistics (seasons and tournaments) and team information. We extract average team statistics from this data. As a result we broadly end up with two types of data: [1]

3.1 Match Statistics

The match statistics include each team's performance for that individual match instance. This type of data is available for all the matches in all the seasons and tournaments. *Table 1* provides the format of this data.

| teamId | season | wscore | wfgm | wfga | wfgm3 |
|--------|--------|-------------------|--------------------|--------------------|--------------------|
| 1102 | 2003 | 68.75 | 22.58333333333333 | 40.0 | 10.0 |
| 1103 | 2003 | 87.76923076923076 | 30.0 | 55.384615384615394 | 5.461538461538462 |
| 1104 | 2003 | 74.70588235294117 | 25.823529411764714 | 58.352941176470594 | 7.0588235294117645 |
| 1105 | 2003 | 79.42857142857143 | 25.57142857142857 | 61.85714285714285 | 9.142857142857142 |
| 1106 | 2003 | 68.30769230769229 | 24.76923076923077 | 53.846153846153854 | 5.846153846153847 |

Table 2: Snapshot of the extracted team level data

| team1 | team2 | wscore | wfgm | lscore | lfgm |
|-------|-------|--------------|--------------|--------------|--------------|
| 1411 | 1421 | -2.205128205 | -1.61965812 | 1.5625 | 1.354166667 |
| 1112 | 1436 | 14.41052632 | 4.482105263 | 16.8 | 5.366666667 |
| 1113 | 1272 | 5.896135266 | 2.403381643 | -5.151515152 | -0.651515152 |
| 1141 | 1166 | 2.658170915 | -1.250374813 | -4.75 | -3.5 |
| 1143 | 1301 | -1.793650794 | 1.904761905 | 6.083333333 | 4.416666667 |
| 1140 | 1163 | -11.14699793 | -6.538302277 | -0.125 | -3.416666667 |

Table 3: Snapshot of the training data used in the models

3.2 Team Statistics

Using the match statistics provided by Kaggle, we extract team level statistics for an entire season for all the teams. This represents a cumulative performance of a team for the previous four seasons starting from the year before the target season that we want to predict for. Some of the most important team statistics are:

- Two point field goals made
Three point attempts
- Assists per game
- Average scoring margin
- Blocks for season
- Defensive rating
- Field goal percentage
- Total free throw attempts

3.3 Scale of the Data

Kaggle provides data from the NCAA seasons and tournaments between 2005 and 2015. The statistics below give an idea about the scale of the dataset.

- Number of seasons: 10 seasons
Number of tournaments: 10 tournaments
- Number of matches in each season: 5000
- Number of matches in each tournament: 64
- Total number of matches in the training set 50,640
- Total number of matches in testing set: 64
- Total number of columns in the training set/testing set number of data fields or features ≈ 40

4 Implementation

Here we describe the implementation details that we used in our approach to predict the outcomes of matchups.

4.1 Features

As described in the section 3.1 and 3.2 the data provided by Kaggle has a lot statistics compiled for every basketball game, so it was difficult to decide which ones to use. Hence we used all the features initially to construct a training set.

4.1.1 Feature Representation

Using the match level statistics provided by Kaggle, we extracted the team level data for every team in every season. Table 2 shows a small snapshot of this data, but it only contains 4 of the 33 features that we actually use in the dataset. Each of the values for the teams represent the average statistic of that team for that specific season. For example, 'wscore' represents the average points a team scored in a game in which it was the winning team. A statistic starting with a 'w' represents that statistic when the team was a winning team. Similarly, the 'l' represents a statistic when the team was a losing team.

Using the above team level statistic data and the matchup data, we create the training set for the machine learning models. For every match up, we look at the teams involved and their corresponding statistics in the team level statistics that we created previously. For each team, we look at all the matches they won, and compute the average of the 'w' statistics. Similarly, we look at the matches that the team lost and compute the average of the 'l' statistics for that team.

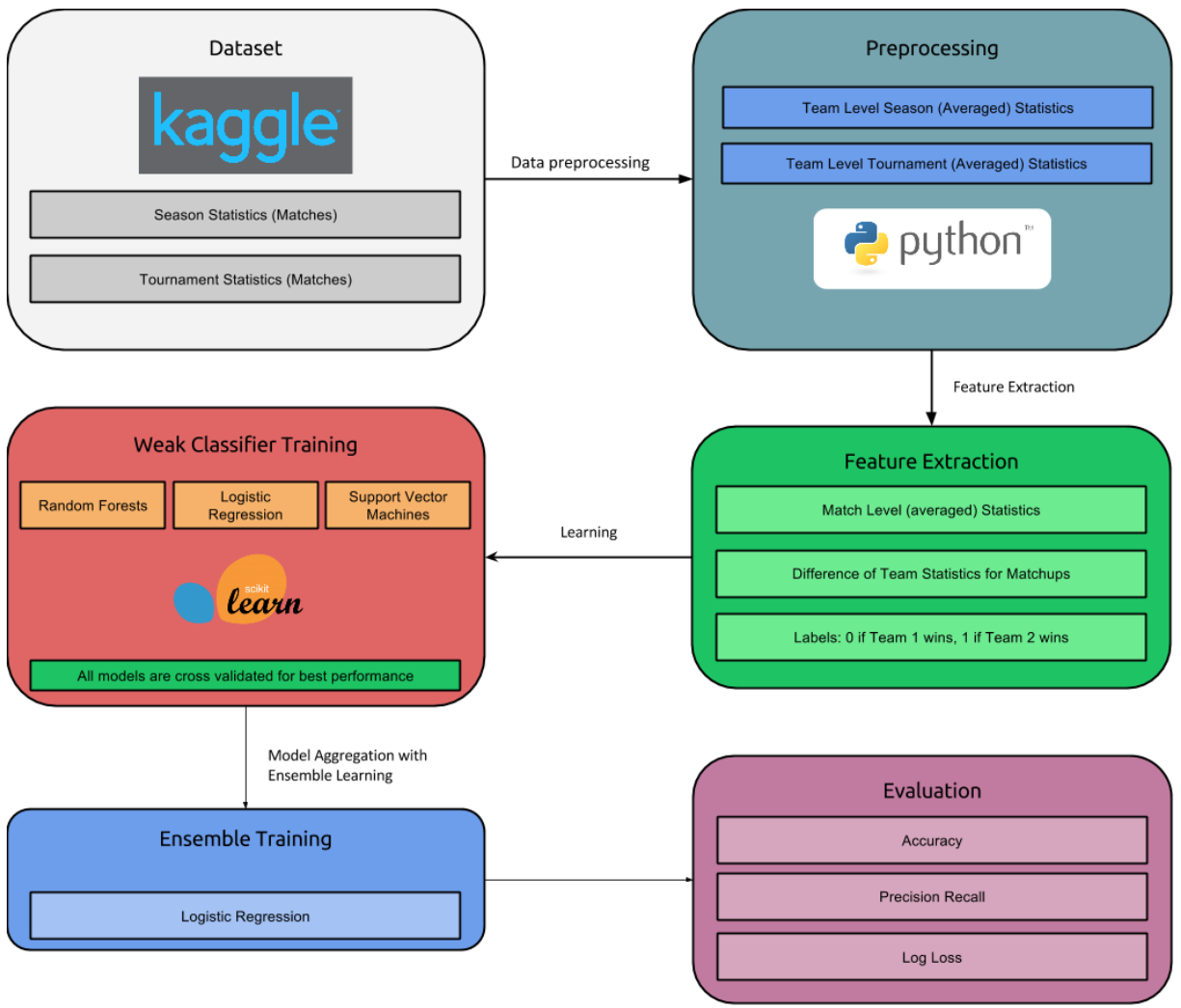


Figure 1: Architecture

We then combine the two vectors for a matchup for a specific team which results in the team vector representing the average statistics of the teams involved in the matchup. This results in two team vectors for every matchup. We can use two approaches to represent these vectors as data points in our training dataset, as follows:

- **Vector of differences:** Each match is represented by the vector of difference between the individual team vectors. This essentially means that the training vector/data point is represented as:

$$X = \text{mod}(\text{TeamAStats} - \text{TeamBStats}).$$

The result or the label vector Y is represented by 0/1. The result is 1 if Team A beats Team B and 0 if Team B beats Team A.

- **Concatenated Vector of both the team statistics:** In this representation the statistics of a match is represented by keeping both Team A and Team B statistics in the vector next to each other. Hence the training vector

$$X = [\text{TeamAStats}, \text{TeamBStats}].$$

The result or the label set representation is same as the previous case.

In our implementation, we use the first 'Vector of differences' approach to represent each data point for the training model. When we consider a match between two teams, we sort them by their ids and then take their differences. As mentioned above, if the first team wins, we label the outcome as 0, and 1 if the second team wins. This gives us a distribution of target labels to predict.

4.1.2 Additional Features

We considered applying a few additional ideas to apply different feature extraction algorithms to the existing team and match statistics and consider the output from the algorithm as a feature itself.

Team Rank A method similar to PageRank algorithm called the GEM method was applied to the NCAA dataset [3]. The margin of victory ($v1 - v2$) is used as the weight the "link" between two football teams, where $v1$ and $v2$ are the teams' scores against

each other. Then the process same as PageRank is followed to make it row stochastic. This will help us obtain the final ranking. Below is an example of the team Rank calculation.

$$H = \begin{matrix} & \begin{matrix} \text{NO} & \text{PHL} & \text{ATL} & \text{CAR} \end{matrix} \\ \begin{matrix} \text{NO} \\ \text{PHL} \\ \text{ATL} \\ \text{CAR} \end{matrix} & \begin{pmatrix} 0 & 0 & 0 & 13 \\ 26 & 0 & 0 & 0 \\ 11 & 27 & 0 & 9 \\ 10 & 28 & 8 & 0 \end{pmatrix} \end{matrix}.$$

Figure 2: Team Rank Matrix

This team rank matrix is represented as a graph.

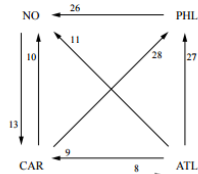


Figure 3: Team Rank graph

Next, we continue to make it row stochastic and follow the PageRank algorithm to get the final ranking. We obtain (0.330 0.252 0.087 0.332) for the PageRank vector, which produces the ranking 1. CAR, 2. NO, 3. PHL, 4. ATL.

4.1.3 Statistical Features

Statistical Features on Player Statistics: One more extra feature that was added to the training set at the list of top players that belong to every team. Only the top players that are playing a particular match are considered in the training set.

5 Architecture

The project can be divided into six important parts; Data collection, Data Preprocessing, Feature extraction, Weak Classifier Learning, Ensemble Training, Prediction/Evaluation. Figure 1 is a pictorial representation of the architecture of the project. The first three parts of the architecture are explained in Section 4. The remaining are explained in the following sections.

6 Machine learning Models

For reasons stated above in section 3, we wanted to use many different machine learning techniques and leverage them against one another so that we would gain the benefits from each and hopefully mitigate the weaknesses of each. We used Support Vector Machines, Random Forests, and Logistic Regression for the base level models.

6.1 Support Vector Machines

Each game in the training set X has a series of characteristics based on the difference in team statistics

and a label y_i (win or loss). These can be plotted in a p -dimensional Euclidean space. The support vector machine then finds the hyperplanes $w\Delta X - b = 1$ and $w\Delta X = 0$ with the greatest distance between them that partitions the space such that wins and losses are separated (where w is the normal vector to the plane and $\|w\|/b$ is the offset of the hyperplane from the origin along w .)[4]. Let the output of an SVM be $f(X)$. Then

$$\hat{y}_{i1}(A, B) = \frac{Pr(y_i = 1|f(X))}{1/1 + \exp(Q * f(X) + V)},$$

where Q and V are parameters chosen through maximum likelihood estimation from a training set made up of coordinates $(f(X)_i, y_i)$. [4]

6.2 Decision Trees and Random Forests

Decision trees split a data set using a set of binary rules that make up the branches of the tree. Sets further down in the tree contain increasingly similar class labels. The objectives are to produce the purest sets possible and to be able to classify any object based on comparison of features to the rules in the tree. A random forest is a bootstrap of the decision tree. Like in the standard bootstrap method, the available training data can be resampled to build many different decision trees. Additionally, the random forest method takes a random subset of available predictors to build its new decision tree. By resampling both the cases and the input variables, many decision trees are grown. Final classification decisions are made based on a majority rule of the many decision trees. Given a decision tree with height k , let $Pr(r_d)$ be the probability associated with node d on branch path r . Then the probability of success for a new case is:

$$y_i(A, B) = \prod_d Pr(r_d)$$

For the random forest of m trees where $Pr(r_{(d,j)})$ is the probability associated with node d on branch path r on decision tree j , the probability of success for a new case is:

$$y_i(A, B) = \sum_j \prod_d Pr(r_{(d,j)})$$

6.3 Logistic Regression

Logistic regression is a statistical method for analyzing a dataset in which there are one or more independent variables that determine an outcome. The outcome is measured with a dichotomous variable (in which there are only two possible outcomes). In logistic regression, the dependent variable is binary or dichotomous, i.e. it only contains data coded as 1 (TRUE, success) or 0 (FALSE, failure). In our case the dependent variable is the outcome of the match and independent variables are the features themselves. Logistic Regression uses a logit function which

measures the log odds of Team A winning a match against Team B:

$$\text{logit}(p) = \ln(p/(1-p))$$

where P is the probobality of A winning.

6.4 Ensemble Model

Ensemble methods are commonly used in classification problems. These methods allow predictions based on different models to be combined, often using a type of voting schema, to determine one final prediction. The benefit to using an ensemble method is that we can leverage the information gleaned from different methods, combining the benefits from each method while hopefully negating or minimizing the flaws in each. Ensemble methods have been used to combine predictions from different machine learning techniques in order to increase stability and to more accurately classify test cases. The combination of methods can also reduce the variance of the predictions as they are less sensitive to irregularities in a training set.[4] In order to ensure that our predictions were in (0,1), we fit a logistic regression model where each explanatory variable is a prediction from one machine learning technique, and the response is the binary outcome. Our model takes the form:

$$y_i(A,B) = 1/e^{-(\beta_0+\beta_1y_1(A,B)+..\beta_3y_3(A,B)+\varepsilon)}$$

where y_1 is from Support Vector machines, y_2 is from Random Forests and y_3 from Logistic Regression.

7 Evaluation

To evaluate the performance of our models, we use a number of performance statistics. Primarily, we use:

| Model | Dataset | Accuracy |
|------------------------|----------|----------|
| Random Forest | Training | 0.97 |
| Logistic Regression | Training | 0.76 |
| Support Vector Machine | Training | 1.0 |
| Random Forest | Testing | 0.61 |
| Logistic Regression | Testing | 0.59 |
| Ensemble | Testing | 0.64 |

Table 4: Snapshot of the extracted team level data

- Accuracy
- Precision
- Recall
- F1 Measure
- Log Loss

7.1 Performance

From the tables that follow, we see that our classifiers perform very well when we test the learned model on

the training data. On the testing data, the classifiers do not perform as good as they did on the training data, but they still perform fairly well, definitely better than a random coin flip.

| | precision | recall | f1-score | support |
|-------------|-----------|--------|----------|---------|
| 0 | 0.99 | 0.96 | 0.98 | 109 |
| 1 | 0.96 | 0.99 | 0.97 | 92 |
| avg / total | 0.98 | 0.98 | 0.98 | 201 |

Table 5: Random Forest on Training Data

| | precision | recall | f1-score | support |
|-------------|-----------|--------|----------|---------|
| 0 | 0.76 | 0.79 | 0.78 | 103 |
| 1 | 0.77 | 0.74 | 0.76 | 98 |
| avg / total | 0.77 | 0.77 | 0.77 | 201 |

Table 6: Logistic Regression on Training Data

| | precision | recall | f1-score | support |
|-------------|-----------|--------|----------|---------|
| 0 | 1.00 | 1.00 | 1.00 | 106 |
| 1 | 1.00 | 1.00 | 1.00 | 95 |
| avg / total | 1.00 | 1.00 | 1.00 | 201 |

Table 7: Support Vector Machines on Training Data

| | precision | recall | f1-score | support |
|-------------|-----------|--------|----------|---------|
| 0 | 0.79 | 0.62 | 0.70 | 48 |
| 1 | 0.38 | 0.58 | 0.46 | 19 |
| avg / total | 0.67 | 0.61 | 0.63 | 67 |

Table 8: Random Forest on Testing Data

| | precision | recall | f1-score | support |
|-------------|-----------|--------|----------|---------|
| 0 | 0.55 | 0.68 | 0.61 | 31 |
| 1 | 0.66 | 0.53 | 0.58 | 36 |
| avg / total | 0.61 | 0.60 | 0.60 | 67 |

Table 9: Logistic Regression on Testing Data

| | precision | recall | f1-score | support |
|-----------|-----------|--------|----------|---------|
| 0 | 0.74 | 0.67 | 0.70 | 42 |
| 1 | 0.52 | 0.60 | 0.56 | 25 |
| avg/total | 0.65 | 0.64 | 0.65 | 67 |

Table 10: Ensemble Learning on Testing Data

We see from the above tables that the random forest classifier performs the best among the individual classifiers on the testing data and the ensemble learner does make a few gains over the random forest classifier. Thereby it does improve the prediction ability by combining the underlying base models.

Although, we observe that since the models perform extremely well on the training data and not as well on the testing data. This is primarily because the models overfit to the training data.

7.2 Log Loss

In addition to the performance metrics, we also evaluated our performance on a log loss function, provided by Kaggle to evaluate how confident our model makes predictions. The loss function is:

$$L(y|\hat{y}) = -1/n * \sum_{i=1}^n [y_i.log(\hat{y}_i + (1 - y_i).log(1 - \hat{y}_i))]$$

where, y represents the actual 1 or 0 outcomes and \hat{y} represents model prediction probability values.

| Benchmark | Score |
|--------------------|------------|
| Winners | 0.529 |
| Seed Benchmarks | 0.6 |
| All 0.5 Benchmarks | 0.693 |
| All 0 Benchmarks | 19.19 |
| OUR PERFORMANCE | 0.62444662 |

Table 11: Log Loss Benchmarks

We see from the table below that, although we are way off the best performing score, we still did better than 2 of the benchmarks. It seems like just following the seed predictions leads to better performance than our classifier. These issues and areas of improvement are discussed in the following section.

8 Conclusion

Even with large amounts of statistical information, there are some parts of a basketball game you simply cannot predict. Injuries cannot be forecast, and the mental state of players playing the game cannot be observed. However, these things can drastically affect the outcome of a basketball match. Perfect brackets will therefore continue to be impossible to calculate,

no matter how much data is provided to the learning models.

However, we believe that our classification accuracy can improve to above 75% if the proper features are used in training. Vital statistical information we were unable to gather include the following:

- Venue of the match including the distance
- Importance of a match-up
- Average number of players who play consistently

All of this data is expected to play a role in how teams perform. However, this type of data was out of our scope to accumulate. Organizations with more access to this type of data will be able to study whether these features have a profound effect on the success of a basketball team. This research would also benefit greatly with more data. It is recommended that much more data be gathered to help avoid overfit. However, since basketball continually changes throughout the years, using data from more than ten years back is also useless. We would also recommend experimenting with using several feature reduction techniques together as opposed to using the reduction techniques one by one. This is something we did not explore as a part of the project.

The overfitting problem can be solved by dividing the training set into 2 parts - training and evaluation. In this way the model can be built using training set, and fine tuned using evaluation set. It can also be reduced by using some kind of regularisation in the model. It could L1 or L2 regularisation depending on the machine learning model and parameters.