Mario: A Machine Learning Classification Approach

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Abstract

The purpose of this project was to construct a classifer capable of successfully navigating a level of Super Mario Brothers. To this end, a means of representing the data describing a level is presented. The ability of various classifying algorithms, including support vector machines, k-nearest neighbors, naïve Bayes, reduced error pruning trees and random forests, to properly classify new data is then evaluated.

1 Introduction

This project represents a departure from the common data sets of the machine learning world. The authors have eschewed corpi and proteins in lieu of something a bit closer to their hearts. To this end, we've repurposed the classification powers of the machine learning world for use with a classic platforming game from our youth, Super Mario Brothers(Mario). Despite the levity of the topic, the underlying algorithms are the same as the ones of the many problems in the field. Indeed, if we consider the Mario world a rough approximation of our own then it's easy to see the relevance of studying a simpler version of what some might consider the holy grail of artificial intelligence: human level interaction with arbitrary environments.

With such a goal in mind, the work here explores treating world interaction as a classification problem. Specifically, we want to classify every possible state in the world to some optimal action. Taking such an action would lead to a new state and a new action, and by continuing in this fashion an agent explores the world in the manner it was trained.

The present work describes and evaluates the application of popular machine learning algorithms to our lofty task. We start with a discussion of the of previous work in the mario AI area. In general, no standing body of data exists for the Mario domain. To overcome this obstacle we developed a system for generating a dataset corresponding to optimal actions in response to the current

environment. A description of this process and various attempted optimizations appears in Section 3. With data in hand, the authors attempt to create a Mario agent by applying popular machine learning algorithms to said data. Each classifier is described in turn in Section 4. Section 5 evaluates the previously described classifiers using common performance metrics as well as actual performance on unknown Mario levels. We end with a discussion of future work and a conclusion.

1.1 Division of Work

1.1.1 Erek

Leveraging some prior experience with the Mario Benchmark software, Erek created a means of mining data from the Mario being played by an expert agent. He also handled changes to the data miner that produced other datasets such as the 5x5 grid. To aid with feature selection, he implemented two DistanceMetrics (JaccardIndex and SymmetricalUncertainty). Erek was additionally responsible for the svm and random forest classifiers. Finally, he implemented the changes allowing classifier agents to be tested on the benchmark system.

1.1.2 Nick

Working with the raw 4697 dimension data obtained by Erek's datamining software, Nick handled running feature selection. Additionally, he worked on a generic one-versus-one classifier implementation and was responsible for the knn, naïve Bayes and REP tree classifiers. Finally, he compiled the classifier performance data that was gathered over the course of the project.

2 Previous Work

The present work is the first to attempt to solve Mario as a classification problem. That said, in recent years there has been a surge of attempts to develop Mario playing agents. The interest started with a MarioAI contest in 2009 which solicited all computer agent solutions to the Mario problem. Interestingly, the report from said contest was dominated by handcoded planning, such as A* [2]. In fact, due to the ability to reverse engineer the simulation engine, such agents reduced the problem to one of planning and easily solved it. A scattering of learning algorithms including neural networks and genetic algorithms were submitted but performed poorly. Later, learning algorithms were given their own category in which they only had to learn a single repeated environment. With such a constraint, learning algorithms were able to reach near optimal performance [1]. In contrast, our work rejects the notion that Mario is only solvable with complete prior knowledge of the the world. However, we also respect the vastness of the problem space and the failures of past learning problems in getting lost in its many plateaus. By creating an agent which trains

on data generated from an expert player we greatly shink the problem space while maintaining some amount of generality.

3 Data Generation

It is a nice metephor to imagine a untrained novice classifier viewing instance after instance of data derived from studying an expert agent. In order to make such a thought reality, one not only needs a source of expert data to mine but also needs to massage it into a convenient format for current machine learning problems. The following subsections will describe the initial state of the data, the conversion to a more general form, and finally performance optimizations.

3.1 MarioAI Benchmark

The MarioAI Benchmark [4] was developed in order to facilitate the developing and testing of AI agents. It placed the original software in a wrapper which the creators felt would allow easier creation of all kinds of agents. For instance, while the various locations of objects in the world are represented as continuous variables, the benchmark wraps the low level data in a representation which discretizes the data into a convenient grid like format. Each cell of the grid contains a integer relating to its contents. Furthermore, the agent has access to public information outside the grid, including whether it is alive or dead, on the ground, and it's status(size, powers). Given this information, the agent is expected to produce an action which coressponds to the buttons that might be pressed by a human playing the original game. In order to gather data, we use a script which records the state and action pairs generated as a trained agent [1] plays expertly. By modifying the parameters of generated environments, millions of unique data points can be generated in minutes.

3.2 Data Conversion

Obviously, the representation has an analog to common formats used in machine learning. If we consider each grid space as feature then we can use it directly as a feature vector into a learning algorithm. Furthermore, we can simply append the extra public information to the end to produce a complete vector. Unfortunately, while this would be a valid input to most algorithms, the results would likely underperform our expectations.

The first problem is that the values chosen to represent the contents of each feature are seemingly random. The scattering of the values are such that the features appear continuous upon first glance. Unfortunately, upon further inspection it becomes obvious that the specific values a feature may take have no deeper meaning and would lead to poor results in the many algorithms requiring placing the vectors in some measured space. To solve this problem, we identified

all possible values and grouped them into 13 cateogories. For consistent weighting of every discrete value, we expanded each feature into 13 binary attributes. At this point, the data is ready to be exported for classifier consumption. Using the default settings of the benchmark (19x19 grid) and including a few of the most important state flags we end up with a feature vector in 4697 dimensions.

3.3 Optimizations

Given the data described so far, it's possible to apply common algorithms and get valid results. Unfortunately, for many software packages such a large data set in this many dimensions is simply intractable. For example, knn scales terribly with large numbers of dimensions. In order to take advantage of such classifiers, we applied various feature reduction strategies.

The primary method we used was a greedy forward selection algorithm which considered each attribute compared with the class distribution. The software we used allowed specifying the distance metric for comparing the attribute vector with the class vector, but with our data changing the metric had little effect on which features were selected. Examples of metrics include Euclidean distance, the Pearson coefficient, and symmetrical uncertainty. Unfortunately, this construction doesn't reduce redundancy amongst the features which leads to the top of the feature rankings being dominated by useful but correlated attributes. As an extreme example, the top two features included a state flag "isMarioOnGround" and a grid attribute "is grid space below Mario solid ground." Regardless of shortcomings, creating datasets with 1,5, and 10 percent features allowed tractable classification.

The second feature selection optimization is simply one of reasoning. We realized that the most important features are the ones closest to the Mario agent. With this in mind, we produced data sets that reduced the grid size from 19x19 to 5x5. This not only allowed algorithms to complete training faster, but it also allowed them to focus on the most important features.

The final optimization dealt with the problem of noise. The most advanced Mario agents will sometimes make erratic actions in the interest of exploring the worlds and finding hidden secrets. The subset of possible environments used for the present work, however, required very little exploration. Upon such realization, the parameters of observed expert were changed such that almost all erratic behavior was removed. This allowed nosie sensitive algorithms like SVM to work better.

4 Classifiers

4.1 Software

4.1.1 Libraries

We considered several libraries for this project and ultimately settled on Java ML [3]. Our main requirements were that it support the various classifiers we wished to use, have an implementation of feature selection, and be capable of interfacing with the MarioAI Benchmark. We looked at using weka [5], Java ML [3], scikit-learn [6], and libsvm [7].

Weka is a relatively full featured library written in java, and as such was one of the most attractive libraries available. It is well used and contains all of the types of classifiers we hoped to use. The major drawback surrounding the use of weka was the data format it required. Programatically it was rather complicated to setup data on the fly, and translating the datafiles we had already created into the proper format was a taxing proposition.

Scikit, written in python, was considered due to the ease of data manipulation in python and the fact that both of us were comfortable with the language. The major drawback of this is that we would have to find a way to export all of our classifiers and import them into the java based MarioAI Benchmark. The libsym integration with scikit was very appealing and was one of the major reasons we considered it.

Libsvm was a great choice because its widespread use and its proficient handling of various multiclass support vector machines. What made us shy away from direct use of libsvm was that all three other libraries had wrappers for libsvm that would allow us to reap its benefits from within a single library that contained all of our classifiers.

We ultimately chose Java ML due to its simple data interface, native java compatibility, and the fact that it wrapped both Weka and libsvm functionality. This enabled us to use libsvm for our support vector classifiers, wrapped Weka classifiers for RE Ptrees, naïve Bayes, knn and random forests, and forward and backwards feature selection.

4.2 knn

4.2.1 Reasons for selection and expectations

We chose to include k nearest neighbors as one of our algorithms because we thought that the large search space would help it identify similarities between attributes in our dataset. Hopes were high for the reduced feature sets as they should only be looking at the attributes that have an impact on the action to be taken for a given board configuration. We expected knn to train quickly and

respond slowly, making it less than ideal for actual play. However, the simplicity of the algorithm would give us a good baseline for what is possible.

4.2.2 Parameter selection

knn had very few parameters to select: the number, k, of neighbors to look at and the distance measure that would be used to find the nearest neighbors. First, due to the scarcity of data for some of our classes, we chose to experiment with k from one to five. We then selected a few distance measures to test: Euclidian, symmetric uncertainty and Jaccard indexing. We then ran a grid search using five-fold cross validation to determine the best set. The metrics we measured to compare the success of classifiers with one another were accuracy, precision, f1 and recall.

4.3 SVM

4.3.1 Reason for selection and expectation

SVMs are considered amongst the best methods for dealing with extremely high dimensional data. While not as tailored toward multi-class algorithms such as ours, the most popular SVM implementation featured multi-class support.

The expectation was that SVMs would be able to handle the data best unadultered, but without an easy way to visualize the data it's hard to predict possible difficulties.

4.3.2 Parameter Selection

Grid Search is expensive for multiclass SVMs but its simplicity allowed for a parallel implementation which excused the naive algorithm. A standard search for C and gamma with an RBF kernel was done with 6 values for each parameter. A second search for C values for linear kernel was also done over 6 parameters. Performance was tested with 2-fold cross validation. This is a valid choice due to the large data size.

4.4 Naïve Bayes

4.4.1 Reason for selection and expectation

Naïve Bayes was an attractive classifier to use since it allowed for priors, allowed us to view a probability distribution on optimal actions, and yielded classifications relatively quickly. We expected it to perform decently with the full ensemble of data, but really shine with a reduced dataset that hopefully proved to have a better signal to noise ratio.

4.4.2 Parameter selection

The wrapped implementation of naïve Bayes provided by Java ML required very few parameters. We were given the option of using a Laplace correction as a prior and the use of logarithmic results. As our dataset was sparse, we were appreciative of the ability to make the classifier aware of this in order to improve performance. Due to the very limited number of parameters, our grid search for optimal parameters focused on whether or not to include a Laplace correction and the size of dataset on which the classifier was trained.

4.5 REP Trees

4.5.1 Reason for selection and expectation

REP (Reduced Error Pruning) trees, while not specifically covered in class, were used as a representative of the various tree classification methods. This algorithm builds a decision/regression tree and internally splits the dataset into folds in order to perform reduced error pruning. The key motivation for including REP trees in our ensemble of classifiers was to have a tree that actively prunes itself as they are purported to be especially good at classifying categorical data. We expected the tree to take longer to train than knn or naïve Bayes, but to respond to requests quickly.

4.5.2 Parameter selection

The REP tree algorithm we utilized required a few parameters – the minimum number of records per leaf, the number of folds for pruning, and the maximum depth. Due to the size and imbalance of our data, we decided to retain most of the default settings of three folds, no maximum depth and two instances per leaf. Our experimentation with REP trees focused on the amount of data provided for training and its impact on the accuracy of the classifier.

4.6 Random Forest

4.6.1 Reason for selection and expectation

Random Forest was our choice for an ensemble classifier. It's incredibly simple to implement which means it was found in almost every library. We expected above average performance due to how well trees preform on categorization without applying sometimes arbitrary metrics. Moreover, we were encouraged by reports of how well this algorithm handled large datasets.

4.6.2 Parameter selection

The standard parameters for this function are number of trees and number of features to sample. There is no guiding principle for comprehensive parameter search for these parameters however so simple heuristics were used. The number of trees was chosen to be the square root of the data, and the number of dimensions was chosen to be log_2 of the dimensionality.

5 Results

5.1 Metrics

In evaluating our methods we decided to use a combination of accuracy, precision, recall and f1 characteristics to evaluate our cross validation results for knn, SVM and naïve Bayes. As the tree algorithms do not measure these statistics, we used root mean squared error as our metric.

Similarly, a metric was needed to compare results from runs through the Mario Benchmark. The metrics picked for that were a combination of a weighted fitness score that the benchmarking program calculated and a scalar progress count that represented how far through the level the classifier controlled Mario was able to get. The following analyses are based on these metrics.

5.2 knn

5.2.1 Cross Validation

Running five-fold cross validation on knn revealed interesting results. Cross validation revealed that 3 was the optimal choice for k. The first thing we noticed was with respect to the distance measures – Euclidean distance was worse than the Jaccard index for out data. In cross validation, the best results achieved with Euclidean distance were precision of 0.645, accuracy of .851, recall of .0327 and F1 measure of .0472. Due to many classes not being chosen, precision could not be averaged across the datasets for a few configurations. The results are below.

Table 1: Cross validation results for knn

Data Size	Feature Reduction	Precision	Accuracy	Recall	F1 Measure
20K	99%	0.915	0.989	0.0597	0.0625
20K	95%	0.920	0.990	0.0599	0.0625
20K	90%	0.917	0.989	0.0598	0.0625
2.5K	5x5	0.837	0.988	0.101	0.0957
2.5K	5x5	0.882	0.991	0.101	0.0956

5.2.2 Mario Benchmark

As expected, the knn algorithm was incredibly slow to respond to move requests. Additionally, as eluded to by the invalid precision results on some of the cross validation, the majority of responses consisted of holding down the forward and jump buttons continually in a manner that prevented Mario from doing more than one jump per level. As such, the knn classifiers never finished a level. The best performance of the classifier was on the simplest level on the lowest

difficulty with a weighted fitness of 1488. It was able to make its way through 26.9% of the level.

5.3 SVM

5.3.1 Cross Validation

Cross validation for our support vector machines was done with two-fold validation, as the size of our datasets made it very costly time-wise to perform more than two-fold validation. A grid search was done for parameters using both linear and RBF kernels. The best linear svm was produced by C=.01 with a score of 0.9842, and the best RBF svm was produced by $\gamma=.5$ and C=10 with a score of 0.59. The scores given here are f1 measures. In future trainings, cross validation correlated exactly with success in the Mario benchmark.

5.3.2 Mario Benchmark

With data reduction of 99% (the 1% dataset), the support vector machine was incapable of learning to pass the simplest level which requires running forward and jumping. It consistently collided with a ledge and got stuck until time ran out. With data reductions of 95% or less, the SVM was capable of learning to run forward and jump repeatedly, allowing it to pass the simplest levels. Moreover, if features chosen carefully, only two were required to match this behavior. Both the linear and RBF SVMs finished completed levels that contained no gaps, enemies or blocks, and was capable of finishing the easiest level that included gaps. However if the dataset size was reduced to 5000, then it reverted to it's previous subpar performance.

5.4 Naïve Bayes

5.4.1 Cross Validation

Cross validation of the various naïve Bayes classifiers alerted us that it would not be a very good player. For an example of the performance as evaluated by five-fold cross validation, the results for 20,000 datapoints with various levels of feature reduction are shown below.

Table 2: Cross validation results for Naïve Bayes

Data Size	Feature Reduction	Precision	Accuracy	Recall	F1 Measure
20K	99%	0.133	0.967	0.0751	0.0973
20K	95%	0.077	0.931	0.0565	0.0935
20K	90%	0.082	0.943	0.0617	0.0864

5.4.2 Mario Benchmark

None of the naıı̈ve Bayes classifiers were able to complete a level on the benchmark. The highest fitness value found was shared between four different parameter settings. Classifiers trained with the 5x5 dataset with both 5000 and 10,000 datapoints were capable of scoring a 1913 on the easiest level. Likewise, Classifiers trained on the 5% and 1% datasets with 20,000 datapoints achieved the same score.

This alludes to an interesting threshold difference between the reduced datasets and the 5x5 grid data. Reduced datasets required four times as many datapoints to achieve the same level of success as the 5x5 grid with a Bayesian classifier. Intuitively this does not make sense, as the length of the feature vectors in the 1% and 5x5 datasets were about the same.

5.5 REP Trees

5.5.1 Cross Validation

As previously mentioned, cross validation of REP trees was performed by measuring the root mean squared error. Using the various levels of feature reduction and changing the number of folds used by the algorithm did not change this metric. This makes sense because it should choose the optimal features to look at and use as few of them as possible. The root mean squared error of our REP trees was 0.0353.

5.5.2 Mario Benchmark

REP trees were one of the few classifiers to successfully complete the clear levels with ledges consistently while showing the lowest threshold for datapoints required to reach optimality.

Table 3: REP tree classifiers capable of scoring the highest score

Data Size	Feature Reduction	Score
500	5x5	6568
500	95%	6568
500	90%	6568

Also of note is that with 99% feature reduction, the REP trees were unable to complete the level. This suggests that a key feature for learning to rest between jumps is discarded with the move from 95% to 99%.

5.6 Random Forests

5.6.1 Cross Validation

There was no cross validation of Random Forests but every training is able to produce an out-of-bag error rating which is unbiased. Unfortunately, no it was nearly impossible to correlate accuracy with any parameter. Due to this variance, a few classifiers were trained with accuracies as high as .65 but such classifiers could not be produced with consistency. Furthermore, the implementation we used was unfortunately not able to cope with large data sizes and thus fewer tweaks were able to be tried.

5.6.2 Mario Benchmark

While it's not able to meet the standard of some of the other algorithms, it performs at levels well above chance. Given the mediocre accuracy ratings reported at build time this is not surprising. It tends to start levels intelligently, but quickly degrades to simply walking forward thus getting stuck on ledges.

5.7 Overall

The overall performance of the classifiers is best communicated through graphical representations of the weighted fitness score of the best classifier of each category graphed against each other. It is very interesting to note the similarities in performance between the REP tree algorithm and the SVM, whose scores are nearly identical for every configuration. This leads us to believe that both algorithms found the ideal means of separating our data.

As can be seen in Figure 1, REP tree and SVM are indistinguishable and highly successful with simple levels devoid of gaps, blocks or enemies. Random forests, knn and naïve Bayes all struggle, with the best and most inconsistent of them being the random forest.

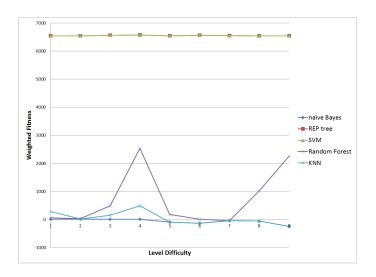


Figure 1: Graph of performance on basic levels by classifier

For levels including gaps (Figure 2), REP tree and SVM succeed at the easiest level and remain the most successfull as difficulty increases. Random forest is once again very erratic in its performance while knn and naïve Bayes fluctuate but remain the worst performers.

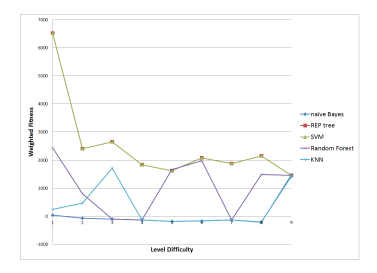


Figure 2: Graph of performance on levels with gaps by classifier

When enemies are added to the levels, none of the classifiers can successfully navigate a level. As can be seen in Figure 3, the classifiers converge to a point for the highest difficulty and the general general trend is a decrease in score as

difficulty increases. This suggests that the presence of enemies is an equalizer between the usual highly successful REP tree and SVM and the rest of the classifiers.

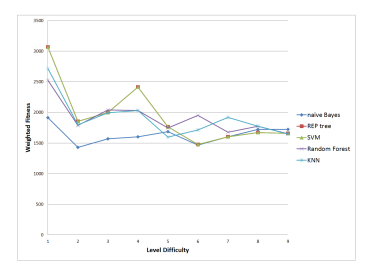


Figure 3: Graph of performance on levels with enemies by classifier

When both enemies and blocks are present, all classifiers have an eratic performance. The REP tree and SVM pair are still the best performing overall. Random forest shows a bit of promise for the middle difficulties, even doing better than REP trees on level six. Overall, performance with enemies and blocks is unreliable and a bit random as can be seen in Figure 4.

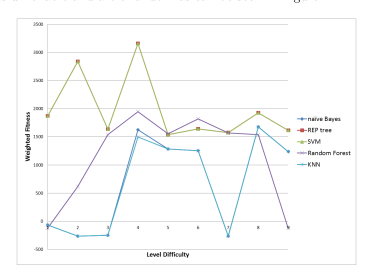


Figure 4: Graph of performance on levels with enemies and blocks by classifier

The final situation the classifiers were tested on was levels with enemies, blocks and gaps present. As can be seen in Figure 5, the classifiers finally become more stratified than in the previous four examples. SVM and REP tree are still identical and the best overall with a sharp decline for difficulties greater than 1, but random forests now does consistently better than naïve Bayes or knn. As this level is the most like levels that would appear in the real Mario games, this is the most interesting one to see, and really signifies the difficulty of a learning agent successfully navigating a level.

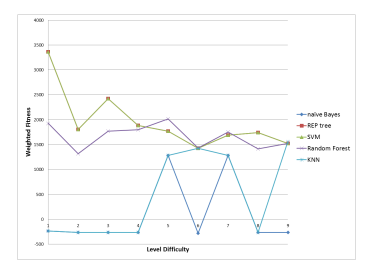


Figure 5: Graph of performance on levels with enemies, blocks and gaps by classifier

6 Future Work

Having examined the source code of the Java ML library that we used, the implementation of some algorithms was not entirely correct. Unfortunately we realized this with too little time to switch to a new library and re-run all of our tests, so the first thing we would do if we were to continue this avenue of exploration would be to validate these results using a second library, such as Weka.

Further improvements could be attempted by implementing adjustments for the sampling imbalance of our data, such as the SMOTE algorithm.

Priors could be better enforced by weighting the value of data gathered from nearest Mario higher than that gathered from across the screen. Various examples of this would be starting with the weight of Mario's square being 100 and decreasing with weight as a function inversely proportional to the distance from Mario. We attempted to do this in a small fashion with the 5x5 dataset, though it would be interesting to see additional approaches.

7 Conclusion

Through this exercise, we have determined that at this time, the classifiers we tested are not suitable for playing a level of Mario that contains gaps, blocks or enemies. At their best, they are capable of immitating a basic forward jumping agent that continually jumps as it runs across the level. At their worst, classifiers run into walls for minutes on end or walk back and forth through the same two squares. Additional work must be done for classifiers to stand a chance against Mario agents programmed to use reinforcement learning, genetic algorithms or modified A*.

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