Group Project

March 13, 2016

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Introduction to Our Technique

The approach we will be taking will have to do with going into detail with Ensembles. Instead of having an ensemble of only one type of learner, we decided to have an ensemble with three different types of learners. These learners will be decision trees, K-Nearest-Neighbours, and support vector machines.

We will each choose our own type of learner and perform our own experiments to try and maximize the accuracy of our learner's predictions. After we have chosen our optimal learners, we will combine them into an ensemble of size 24 (8 learners of each type) and predict the test data using this new ensemble.

We will use existing packages provided by sklearn library. We will use the methods provided by the library and explore additional techniques to supplement these methods to vary the complexities of the models that result.

In [1]: ########Libraries Used Throughout The Code:#########

```
import numpy as np
import matplotlib.pyplot as plt
import matplotlib.patches as mpatches
import mltools as ml
import mltools.dtree as dtree
import mltools.logistic2 as lcs2
import sklearn
from sklearn import svm
from sklearn import preprocessing
%matplotlib inline
```

We will used the provided Kaggle data in our class Kaggle Competition.

Since the Kaggle data has 91 dimensions, one of the type of learners we chose was a support vector machine, as they perform well in high dimensionality. First, we tested different types of kernels to find which

one performed the best. It is obvious that linear kernel was not going to estimate the data correctly, so a linear kernal was not tested.

```
In [3]: # First we scale the data for the SVM
        XiTe = preprocessing.scale(Xte)
        XiTr = preprocessing.scale(Xtr)
In [4]: # Test accuracy of each type of SVM
        kernels = ['rbf', 'sigmoid', 'poly']
        for k in kernels:
            clf = svm.SVR(kernel=k)
            clf.fit(XiTr[:10000,],Ytr[:10000])
            YhatTrain = clf.predict(Xtr)
            YhatTest = clf.predict(Xte)
            MSEtrain = np.mean((Ytr - YhatTrain)**2)
            MSEtest = np.mean((Yte - YhatTest)**2)
            print("SVM with {} as kernal".format(k))
            print("\tMSE of training data: " + str(MSEtrain))
            print("\tMSE of test data: " + str(MSEtest))
SVM with rbf as kernal
       MSE of training data: 0.694313336835
        MSE of test data: 0.720472853771
SVM with sigmoid as kernal
        MSE of training data: 0.786630556699
        MSE of test data: 0.821199078992
SVM with poly as kernal
        MSE of training data: 7.17265890208e+27
        MSE of test data: 7.2573234728e+27
```

Based on the data, the RBF kernel had the best performance. Now, since SVM can take a long time with large amounts of data, we will see how long an SVM with a RBF kernel takes with different subsets of data and it's performance on said data.

```
In [28]: import time
         clf = svm.SVR(kernel='rbf')
         Xi = preprocessing.scale(X)
         for i in [5000,10000,20000,40000,60000]:
             print("Data for {} data points".format(i))
             t0 = time.time()
             clf.fit(Xi[:i,],Y[:i])
             print("\tTraining: {:.2f} seconds".format(time.time()-t0))
             t0 = time.time()
             Yhat = clf.predict(X[:i,])
             print("\tPredicting: {:.2f} seconds".format(time.time()-t0))
             MSE = np.mean((Y[:i] - Yhat)**2
             print("\tMSE of data: {:.2f}".format(MSE))
Data for 5000 data points
        Training: 3.24 seconds
        Predicting: 1.79 seconds
```

MSE of data: 0.71 Data for 10000 data points

Training: 12.60 seconds Predicting: 6.96 seconds

MSE of data: 0.70 Data for 20000 data points

Training: 180.33 seconds Predicting: 28.04 seconds

MSE of data: 0.70

Data for 40000 data points
Training: 800 83 see

Training: 890.83 seconds Predicting: 112.21 seconds

MSE of data: 0.69 Data for 60000 data points

Training: 3303.78 seconds Predicting: 245.58 seconds

MSE of data: 0.70

Based on these results, it is best to train on a subset of the data of size 20000 since the accuracy is not improved much after 20000 data points but is taking much longer. This number will be used when training the final data in the ensemble.

Since the SVM learner seems to be underfitting, I have decided to try and increase the value of the penalty (parameter C) and increasing the value of gamma, the kernel coefficient, to try and increase the complexity to see if it would do better. The following are the result of the test:

Data for C=0.1

MSE for training data: 0.61558137485 MSE for test data: 1.33784400503

Data for C=0.5

MSE for training data: 0.537344845127 MSE for test data: 1.19044021805

Data for C=1

MSE for training data: 0.497589581634 MSE for test data: 1.06304257652

Data for C=20

MSE for training data: 0.393138627212 MSE for test data: 0.822942582304

Data for C=50

MSE for training data: 0.403543750388 MSE for test data: 0.829147117563

Data for gamma=0.1

MSE for training data: 0.665729419872 MSE for test data: 1.23645277441

Data for gamma=0.5

MSE for training data: 0.728801622922

MSE for test data: 1.2831635786 Data for gamma=1

MSE for training data: 0.729558885724

MSE for test data: 1.28298434064

Data for gamma=10

MSE for training data: 0.729791044544 MSE for test data: 1.28297833932

Data for gamma=20

MSE for training data: 0.729791076776 MSE for test data: 1.28297833925

Changing these parameters did not do much to make the learner better. Since it seems to be underfitting, we tried to make an AdaBoost ensemble of SVMs to try and remedy the fact that it is underfitting.

This resulted in an MSE of 0.807 for the test data. Since we were not able to improve on the SVM model, we decided to drop it entirely on our final ensemble.

Here we will be learning on decision trees. We try to tackle the problem of overfitting, which occurs with overcomplex trees. These trees can become unstable because small variations in the data might result in a completely different tree being generated. Luckily, we can avoid this problem by either setting the maximum depth of the tree or setting the minimum number of samples required at a leaf node. We proceed to learning a decision tree regressor on the data, and specifying a maximum depth of 20. We predict on the training data and the testing data and obtain the mean squared averages for both data sets. The mean squared averages are displayed as follows. Next, we specify a variety of maximum depths for the decision trees, ranging anywhere from 1 - 19. We want to find out how adjusting the maximum depths changes the complexities of the trees, and when they begin to overfit. Which maximum depth best handles the problem of overfitting? For each maximum depth, we learn a decision tree regressor and calculate the mean squared average of the training data we predicted on, and also calculate the mean squared average of the testing data we predicted on.

```
In [10]:
         #########Code for the Decision Tree:#########
         #Imported Library for Decision Trees
         from sklearn import tree
         #Decision Tree Learner INFO...
         # Decision Tree Variables
         depth = 20 # the maxth depth of the decision tree
         nodes = 8 # the minimum number of data to split node
         #learn a decision tree regressor on data, specify max depth of 20
         learner = tree.DecisionTreeRegressor(max_depth=20)
         learner.fit(Xtr, Ytr)
         YhatTrain = learner.predict(Xtr)
         YhatTest = learner.predict(Xte)
         MSETrain = np.mean((Ytr - YhatTrain)**2)
         MSETest = np.mean((Yte - YhatTest)**2)
         print('{}{}'.format("MSE for training data: ", MSETrain))
         print('{}{}'.format("MSE for testing data: ", MSETest))
         #try different ranges of maximum depths
         for depth in range(20):
             learner = tree.DecisionTreeRegressor(max depth = depth+1)
             learner.fit(Xtr, Ytr)
             Yhat_train = learner.predict(Xtr)
             Yhat_test = learner.predict(Xte)
             mseTrain = np.mean((Ytr - Yhat_train)**2)
             mseTest = np.mean((Yte - Yhat_test)**2)
             print("Depth {:02d} --> mse train: {}, mse validation: {}".format(depth+1
         , mseTrain, mseTest))
```

```
MSE for training data: 0.0357911570162
MSE for testing data: 0.712112171593
Depth 01 --> mse train: 0.557937953351, mse validation: 0.57414551383
Depth 02 --> mse train: 0.505146872922, mse validation: 0.519822936078
Depth 03 --> mse train: 0.472694791396, mse validation: 0.483718589446
Depth 04 --> mse train: 0.451999080531, mse validation: 0.465591656337
Depth 05 --> mse train: 0.434261655119, mse validation: 0.455494004267
Depth 06 --> mse train: 0.418033072227, mse validation: 0.446586295059
Depth 07 --> mse train: 0.397270867629, mse validation: 0.434863566073
Depth 08 --> mse train: 0.377611804684, mse validation: 0.438300262859
Depth 09 --> mse train: 0.353281895038, mse validation: 0.443817028614
Depth 10 --> mse train: 0.32512500598, mse validation: 0.467904685154
Depth 11 --> mse train: 0.2920177674, mse validation: 0.487800548445
Depth 12 --> mse train: 0.25566965481, mse validation: 0.525525238568
Depth 13 --> mse train: 0.216916242863, mse validation: 0.555434048757
Depth 14 --> mse train: 0.178732828402, mse validation: 0.586412749513
Depth 15 --> mse train: 0.143080936773, mse validation: 0.61639887708
Depth 16 --> mse train: 0.11355919281, mse validation: 0.638016511422
Depth 17 --> mse train: 0.0871032692009, mse validation: 0.671957807537
Depth 18 --> mse train: 0.0657788229549, mse validation: 0.685808352885
Depth 19 --> mse train: 0.0492075353514, mse validation: 0.709960727505
Depth 20 --> mse train: 0.0357897739192, mse validation: 0.712080417074
```

We stick to specifying a maximum depth of 20, and proceed to learning on decision trees with that fixed depth, and we adjust the next parameter vital to creating the decision trees we want. The next parameter we adjust is the minimum number of samples required at a leaf node, otherwise known as the min_samples_leaf parameter. We learn on a range from 2^3, up to 2^12 minimum number of samples required at a leaf node. We predict on the training data and on the testing data, and we calculate their mean squared averages. The mean squared averages of the training and testing data are displayed as follows.

```
In [5]: for nodes in range(3, 13):
             learner = tree.DecisionTreeRegressor(max depth = 20, min samples leaf = 2
        **nodes)
             learner.fit(Xtr, Ytr)
            Yhat_train = learner.predict(Xtr)
             Yhat_test = learner.predict(Xte)
             mseTrain = np.mean((Ytr - Yhat_train)**2)
             mseTest = np.mean((Yte - Yhat test)**2)
             print("2^{} data at leaf node --> mse train: {}, mse validation: {}".form
        at(nodes, mseTrain, mseTest))
        2<sup>3</sup> data at leaf node --> mse train: 0.159069620689, mse validation: 0.592962
        93253
        2<sup>4</sup> data at leaf node --> mse train: 0.238927974161, mse validation: 0.515825
        198467
        2<sup>5</sup> data at leaf node --> mse train: 0.303901057014, mse validation: 0.457419
        046287
        2<sup>6</sup> data at leaf node --> mse train: 0.348795257147, mse validation: 0.428665
        71325
        2<sup>7</sup> data at leaf node --> mse train: 0.376680650869, mse validation: 0.424888
        119888
        2^8 data at leaf node --> mse train: 0.398765237031, mse validation: 0.429117
        852099
        2<sup>9</sup> data at leaf node --> mse train: 0.414565160586, mse validation: 0.438279
        933896
        2^10 data at leaf node --> mse train: 0.431358990435, mse validation: 0.45011
        4946475
        2^11 data at leaf node --> mse train: 0.452946540527, mse validation: 0.46886
        867678
        2^12 data at leaf node --> mse train: 0.47046962598, mse validation: 0.483359
        063336
```

We decide to choose 2'8 minimum number of samples required at each leaf node, based on the

results of the mean squared averages from the training and testing data.

Now we proceed to storing our individual learners in an ensemble. We set the size of the ensemble to 20 because we will be storing 20 different learners within the ensemble. We will store the KNN and the decision tree learners in the ensemble. Unfortunately, we will not be able to store the Support Vector learners in this ensemble. We create 10 different decision tree learners. Using a maximum depth of 20 and 8 minimum samples required at each leaf node as our favorite parameters, we add each of the learners to the ensemble of different learners. Finally, we proceed to storing our individual learners in an ensemble. We set the size of the ensemble to 20 because we will be storing 20 different learners within the ensemble. We will store the KNN and the decision tree learners in the ensemble. Unfortunately, we will not be able to store the Support Vector learners in this ensemble.

from sklearn.ensemble import BaggingRegressor ensemble = [None]*10 #for now store 10 learners in ensemble before combining all learners # Create learners and add to dtree ensemble #dtLearners = BaggingRegressor(tree.DecisionTreeRegressor(max_depth = 20, min samples leaf = 2**8) M = Xtr.shape[0]Me = Xte.shape[0]YtrHat = np.zeros((M, 10))YteHat = np.zeros((Me, 10))for 1 in range(10): #add 10 decision tree learners to the ensemble Xi, Yi = ml.bootstrapData(Xtr, Ytr, M) ensemble[1] = BaggingRegressor(tree.DecisionTreeRegressor(max_depth=20, mi n samples leaf = 2**8)) ensemble[1].fit(Xi, Yi) Ythat = ensemble[1].predict(Xtr) Yehat = ensemble[1].predict(Xte)

```
from sklearn.neighbors import KNeighborsRegressor #Imported Library for KNN
       Regressor
       from sklearn.ensemble import BaggingRegressor
       #print Xtr.shape, Ytr.shape
       #Create a list to store the top KnnRegressor Performers
       topKnnPerformers = [None]*10
       #Predict and print the MSE for training and test data on knnRegressors from 1
       to 20 Nearest Neighbors
       for K in range(1, 21):
           knnRegressor = KNeighborsRegressor(n_neighbors = K)
           yhatTr = knnRegressor.fit(Xtr, Ytr).predict(Xtr)
           yhatVa = knnRegressor.fit(Xte, Yte).predict(Xte)
           print "MSE for K = {:>2} Neighbors Error(Training) = {:>7.4f}
                                                                             Err
       or(Validation) = {:>7.4f}".format(
                   K, np.mean((Ytr - yhatTr)**2), np.mean((Yte - yhatVa)**2))
           #Since the first 10 regressors
           if (K <= 10):
               topKnnPerformers[K-1] = knnRegressor
```

MSE for K = 1 = 0.0000	Neighbors	<pre>Error(Training) =</pre>	0.0000	Error(Validation)
	Neighbors	<pre>Error(Training) =</pre>	0.1863	Error(Validation)
	Neighbors	<pre>Error(Training) =</pre>	0.2555	Error(Validation)
	Neighbors	<pre>Error(Training) =</pre>	0.2932	Error(Validation)
	Neighbors	<pre>Error(Training) =</pre>	0.3175	Error(Validation)
MSE for $K = 6$ = 0.3676	Neighbors	<pre>Error(Training) =</pre>	0.3352	Error(Validation)
	Neighbors	<pre>Error(Training) =</pre>	0.3486	Error(Validation)
MSE for $K = 8$ = 0.3965	Neighbors	<pre>Error(Training) =</pre>	0.3578	Error(Validation)
MSE for $K = 9$ = 0.4033	Neighbors	<pre>Error(Training) =</pre>	0.3655	Error(Validation)
MSE for $K = 10$ = 0.4087	Neighbors	<pre>Error(Training) =</pre>	0.3719	Error(Validation)
MSE for $K = 11$ = 0.4145	Neighbors	<pre>Error(Training) =</pre>	0.3771	Error(Validation)
MSE for $K = 12$ = 0.4190	Neighbors	<pre>Error(Training) =</pre>	0.3812	Error(Validation)
MSE for $K = 13$ = 0.4221	Neighbors	<pre>Error(Training) =</pre>	0.3855	Error(Validation)
MSE for $K = 14$ = 0.4259	Neighbors	<pre>Error(Training) =</pre>	0.3895	Error(Validation)
MSE for $K = 15$ = 0.4289	Neighbors	<pre>Error(Training) =</pre>	0.3928	Error(Validation)
MSE for $K = 16$ = 0.4311	Neighbors	<pre>Error(Training) =</pre>	0.3964	Error(Validation)
MSE for $K = 17$	Neighbors	<pre>Error(Training) =</pre>	0.3993	Error(Validation)

= 0.4340Error(Training) = 0.4020Error(Validation) MSE for K = 18 Neighbors = 0.4370Error(Training) = 0.4041Error(Validation) MSE for K = 19 Neighbors = 0.4393MSE for K = 20 Neighbors Error(Training) = 0.4062Error(Validation) = 0.4417

In [7]: print(topKnnPerformers) #checking to see if the first 10 knn neighbors were s tored in the list

```
[KNeighborsRegressor(algorithm='auto', leaf size=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=1, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=2, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=3, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=4, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=5, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=6, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf_size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=7, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=8, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=9, p=2,
          weights='uniform'), KNeighborsRegressor(algorithm='auto', leaf size
=30, metric='minkowski',
          metric_params=None, n_jobs=1, n_neighbors=10, p=2,
          weights='uniform')]
```

Using the KNeighborsRegressor from the sklearn library a knnregressor learner was constructed

with neighbors ranging from 1 to 20. It was expected, as the commentary mentioned on the project writeup that the KNN learners would take some time to train and predict. The process was ended up being fast (30 minutes - 1 hour) and the resulting MSE was very low giving us good results for the FULL data set. As the results are seen above, the error on the training data was < ~.41 and < ~.44 for the test data. Because the model starts underfitting for larger values of K we decided to move forward with neighbors less than K = 10, meaning taking the top 10 performers to store into the ensemble.

```
In [ ]: topEnsemblePerformers = [None]*10
        for i, regressor in enumerate(topKnnPerformers):
            knnEnsemble = BaggingRegressor(base_estimator = topKnnPerformers[i], n_es
        timators = 10)
            yhatTr = knnEnsemble.fit(Xtr, Ytr).predict(Xtr)
            yhatVa = knnEnsemble.fit(Xte, Yte).predict(Xte)
            print "MSE for Ensemble with a K = {:>2} Neighbors Regressor
                                                                             Error(Tr
        aining) = {:>7.4f}
                               Error(Validation) = {:>7.4f}".format(
                    regressor.n_neighbors, np.mean((Ytr - yhatTr)**2), np.mean((Yte -
        yhatVa)**2))
```

MSE for Ensemble with a K = 1 Neighbors Regressor Error(Training) = 0.0955 Error(Validation) = 0.1055

MSE for Ensemble with a K = 2 Neighbors Regressor Error(Training) = 0.1729 Error(Validation) = 0.1936

MSE for Ensemble with a K = 3 Neighbors Regressor Error(Training) = 0.2359 Error(Validation) = 0.2596

Playing around with the BaggingRegressor library, there was no way to store the 10

topKnnPerformers into one Ensemble so we tested storing 10 of each topKnnPerformers and train/predicted to see the results. We found that since it was taking too long to output the MSE and this approach just did not make sense, so we did not move forward with it. However, it is worth noting the drop in MSE for the 3 outputs that did end up printing as shown above. Afterwards, reading through library documentation we moved forward with another approach below.

```
In [18]: knnEnsemble = BaggingRegressor(n_estimators = len(topKnnPerformers))
         knnEnsemble.estimators = topKnnPerformers
         yhatTr = knnEnsemble.fit(Xtr, Ytr).predict(Xtr)
         yhatVa = knnEnsemble.fit(Xte, Yte).predict(Xte)
         print "MSE for Ensemble with top 10 KnnRegressors
                                                               Error(Training) = {:>7.
                 Error(Validation) = {:>7.4f}".format(
         4f}
                 np.mean((Ytr - yhatTr)**2), np.mean((Yte - yhatVa)**2))
         MSE for Ensemble with top 10 KnnRegressors
                                                        Error(Training) = 0.0719
         Error(Validation) = 0.0773
```

MSE for Ensemble with top 10 KnnRegressors Error(Training) = 0.0719 Error(Validation) = 0.0773

Since BaggingRegressors library had an attribute to store subset of estimators, we decided to take our list of topPerformers trained previously and store this list to that attribute. This allowed us to train and predicted on the entire list of learners rather than having to indivudally store them in a BaggingRegressor with 10 copies of each knnRegressor. After training and predicting on this entire list of already well performing learners, the result was an even lower MSE!

In [9]: #Combining the Ensemble **#To output to Kaggle** #Based off of our results above we will use the optimum values given out by t he knnEnsemble above as well as #store the optimum Decision Tree leaf value of 2^8 into its own Ensemble repr edict and average it with knnEnsemble prediction avgPredTeList = [] optimumKnnEnsemble = BaggingRegressor(n_estimators = len(topKnnPerformers)) optimumDtEnemble = BaggingRegressor(base estimator = tree.DecisionTreeRegressor(max depth = 20, min samples 1 eaf = 2**8), n_estimators = 10) KnnTr = optimumKnnEnsemble.fit(Xtr, Ytr) DtTr = optimumDtEnemble.fit(Xtr, Ytr)

The code above takes the Knn values based on our previous test runs earlier of KNearestNieghbors of up to 10 as well as the optimum Decsion Tree Ensemble based on our previous findings of min samples leafs @ 2\% and retrained aftering storing each to their own Ensemble.

```
In [15]: yhatKnnTr = KnnTr.predict(Xe1)
         yhatDtTr = DtTr.predict(Xe1)
         avgPredTeList = (yhatKnnTr + yhatDtTr) / 2
         print(len(avgPredTeList))
         fh = open('predictions.csv','w') # open file for upload
         fh.write('ID, Prediction\n') # output header line
         for i, yi in enumerate(avgPredTeList):
             fh.write('{}, {}\n'.format(i+1, yi)) # output each prediction
         fh.close() # close the file
```

40000

Because these were our top learners we decided to take the predictions of both the Decision Tree Ensemble and the K nearest Ensemble and average these values to upload to Kaggle. Because of the large amount of time it takes to build these learners and the Ensemble version of these learners, we were only able to upload our predictions once and not reupload with improved prediction quality.

Conclusion

In conclusion, since our document ipython notebook size was exceeding far beyond 6 pages and we were not sure if our writeup was suppose to be in a seperate document, we decided to do our write up as we were writing and testing our code (as shown above). At the beginning we had decided to go more into detail with Ensembles by using learners that we did not go into detail with in class. This enabled us to use 3 different types of learners to equally balance the workload. Rodrigo, started with the intent of using Neural Networks until he realized that they only work with

classification models. He decide to switch to Support Vector Machines, also a learner we did not apply in our homeworks. Because the Support Vector Machine took too long to train on the entire dataset, a smaller subset was used. The result of predicting on these smaller subset were very high MSEs for both the training and test data. Because the SVM was constantly underfitting and unable to improve we came to the decision of removing it from the final ensemble. The decision tree, the learner tested by Mayra, had mixed performances. Aftering training and prediction on the decision tree using the library it looked as the depth increased the MSE decreased. Iterating thru different values of the leaf node parameter, Mayra found that the decision tree at leaf node 2⁸ had the optimum learner so we decided to store this learner into the final ensemble. The third learner, explored by Muzamil, was going into detail with K Nearest Nieghbors, but as apposed to using classifiers as in the previous homework, he used the library's knnRegressor model. He decided to train the KNN mode of up to 20 neighbors and the performance, as seen above, turned out to be very good. It was noticed, as expected that the more neighbors you have the more the model was starting to underfit as seen in the increase in the MSEs. He decided to take knnRegressors from 2-10 neighbors to store into the Ensemble. At first an attempt was made to try to use these 10 different learners in the baggingRegressor library but the only way to do this was store multiple of the same learners into baggingRegressor meaning he would have had to create 10 different ensembles to train on. After looking more into the library an attribute in the baggingRegressor was discovered where a list of sublearners can be set. He decided to take this approach and train the ensemble on the list of 10 KnnRegressors of up to 10 neighbors. The training and predicted process ended up being quick and the MSE was even less! We decided to take the predictions along with the optimum Decision Tree learner to upload to Kaggle. Overall, our final goal of creating an emsemble containing 7 or so of each our learners did not go as planned so we decided to choose the 2 learners and use 10 of each of those. There were many times that we could not see the results of our trainers due to very long run times. If something were to be changed in order to get the results we intended, we could probably use partitions of the dataset whenever possible.