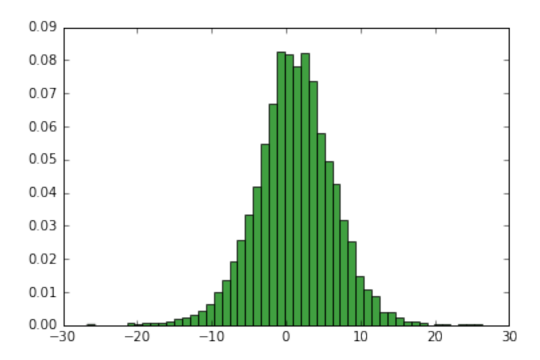
## Regression

## March 24, 2016

```
In [229]: ##importing the required libariries##
          import pandas as pd
          import numpy as np
          import matplotlib.pyplot as plt
         from sklearn import svm
         from sklearn import preprocessing
         from sklearn.cross_validation import train_test_split
         from sklearn.decomposition import PCA
         from sklearn.ensemble.forest import RandomForestRegressor
         from sklearn.cross_validation import cross_val_score
          import operator
          from sklearn.metrics import mean_squared_error
          %pylab inline
Populating the interactive namespace from numpy and matplotlib
In [222]: ##importing the data into a pandas data frame##
          train_DF=pd.read_csv('codetest_train.txt', delimiter='\t')
In [223]: ##getting the statistics of the target values to get a better underestanding
          #of the proper learning method##
         n, bins, patches = plt.hist(train_DF['target'], 50, normed=1, facecolor='green', alpha=0.75)
         train_DF['target'].describe()
Out [223]: count
                  5000.000000
         mean
                     1.143878
                     5.259896
         std
         min
                   -26.705570
         25%
                    -2.034383
         50%
                     1.166835
         75%
                     4.439549
                     26.347818
         max
         Name: target, dtype: float64
```



```
In [224]: ##splitting the data into train/test set to have a set for performance evaluation##
          X_train, X_test, y_train, y_test = train_test_split(train_DF.iloc[:,1:255], \
                                                              train_DF['target'], test_size=0.2)
In [225]: ##data_refine function will: 1-find the non-numeric columns, 2- fill the NaN
          #in non-numeric columns with the most used element 3- transform the non-numeric
          #columns to numeric values using LabelEncoder method from SKLearn, 4- fill the
          #NaN values in the numeric columns using the mean value of them##
          def data_refine(train_DF):
              #1-find the non-numeric columns#
              real_or_str=train_DF.applymap(np.isreal).all(0)
              non_num_feature=real_or_str[~real_or_str].keys()
              le = preprocessing.LabelEncoder()
              for col in non_num_feature:
                  #2- fill the NaN in non-numeric columns with the most used element#
                  train_DF[col].fillna(train_DF[col].describe().top, inplace=True)
                  #3- transform the non-numeric columns to numeric values using
                  #LabelEncoder method from SKLearn#
                  train_DF[col] = le.fit_transform(train_DF[col])
              #4- fill the NaN values in the numeric columns using the mean value of them
              train_DF.fillna(train_DF.mean(), inplace=True)
              return train_DF
          X_train_refined=data_refine(X_train)
          X_test_refined=data_refine(X_test)
In [226]: ##Using a Support Vector Regressor as our learning technique##
          clf = svm.SVR(C=1, epsilon=0.2)
          clf.fit(X_train_refined, y_train)
          print ("MSE on the training set is:\n"), mean_squared_error(y_train, \
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clf.predict(X_train_refined))
          print ("MSE on the test set is:\n"), mean_squared_error(y_test, \
                                                  clf.predict(X_test_refined))
MSE on the training set is:
12.0869268614
MSE on the test set is:
17.2529650053
In [227]: ##Using another learning technique (Random Forrest Regressor) on our data##
          rfr = RandomForestRegressor(n_estimators=10, random_state=0)
          rfr.fit(X_train_refined, y_train)
          print ("MSE on the training set is:\n"), mean_squared_error(y_train, \
                                                  rfr.predict(X_train_refined))
          print ("MSE on the test set is:\n"), mean_squared_error(y_test, \
                                              rfr.predict(X_test_refined))
MSE on the training set is:
2.49144749533
MSE on the test set is:
15.9314309093
```

As we can see, both of these models tend to overfit the training set. Linear Regressors/K Nearest Neighbor/Neural Networks yield the same performance and all tend to overfit as well. We will try two different method to overcome overfitting.

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1- We will try to use PCA to reduce the number of features
2- We will pick Random Forrest Regressor as our model and using the k-fold cross
validation technique fine tune the parameters of the SVR.
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```
With 50 features, 0.281870 of variance is preserved and the MSE on
With 100 features, 0.509171 of variance is preserved and the MSE on
With 150 features, 0.702820 of variance is preserved and the MSE on
With 200 features, 0.866501 of variance is preserved and the MSE on
With 200 features, 0.866501 of variance is preserved and the MSE on
The test set is 36.695022
the test set is 36.236900
the test set is 36.795260
```

We can see that almost all of the columns are containing significant amount of variance/data and even using 200 features we are still losing around 15% of the data. Also the performance of our model doesn't improve by using less features and it still overfit. So, next we try to fine tune the parameters of our RFR model using a 5-fold cross-validation

As we can see the performance improvement using a larger number of trees was marginal