# Identifying Fraud from Enron Emails and Financial Data

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## Project Overview

Enron Corporation was an American energy, commodities, and services company, which was formed in 1985 by, Kenneth Lay. This corporation filed for bankruptcy on December 2001. Enron case is famous for being the largest bankruptcy reorganization the biggest audit failure in American history at that time. In this project, we are asked to use the Enron email data and identify employees who may have committed fraud and were involved in the Enron scam, using supervised machine learning.

Machine learning algorithms are very useful to accomplish these kinds of goals, because their output is a machine system that builds models without demanding human involvement which would be really time consuming. The goal is to make a model that uses some financial and email data to identify Persons of Interest among the executives. The Person of Interest (POI) is defined as “*individuals who were indicted, reached a settlement or plea deal with the government, or testified in exchange for prosecution immunity*”. The mentioned information became public after the federal investigation of Enron scandal. Enron Email Dataset was prepared by the CALO Project in MIT and is available here: (<https://www.cs.cmu.edu/~./enron/>).

This data has financial and email features as well as POI label which shoes whether an executive was POI as well. The features are:

* **Financial features:** salary, deferral\_payments, total\_payments, loan\_advances, bonus, restricted\_stock\_deferred, deferred\_income, total\_stock\_value, expenses, exercised\_stock\_options, other', long\_term\_incentive, restricted\_stock, director\_fees
* **Email features:** to\_messages, email\_address, from\_poi\_to\_this\_person, from\_messages, from\_this\_person\_to\_poi, shared\_receipt\_with\_poi

## Data Cleaning and Outliers

The first step is to know the data and find and handle possible problems. First, I took a look at the name of the executives. Over all there are 146 executives in this data which 18 of them are POIs. The last two names **Total** and **The Travel Agency In The Park**are different from the rest. Total is the sum of all financial features therefore does not represent any executive just like The Travel Agency In The Park. Therefore, I excluded them from the data.

Each executive is presented by 21 features. I went through them and using my intuition and some online research about the Enron scandal, I excluded some of them. At the end, I chose a mixture of financial and email data to be used in the feature selection part. This table shows all the available features and the one that I used. In the table below, the red ones are the excluded ones:

| **Feature's Name** | **Included?** | **Feature's Name** | **Included?** | **Feature's Name** | **Included?** |
| --- | --- | --- | --- | --- | --- |
| bonus | Yes | exercised\_stock\_options | Yes | deferral\_payments | Yes |
| from\_this\_person\_to\_poi | Yes | from\_poi\_to\_this\_person | Yes | salary | Yes |
| email\_address | No | deferred\_income | No | director\_fees | No |
| restricted\_stock\_deferred | Yes | total\_payments | Yes | from\_messages | No |
| loan\_advances | No | expenses | No | other | No |
| shared\_receipt\_with\_poi | No | to\_messages | No | long\_term\_incentive | No |
| total\_stock\_value | No | restricted\_stock | No |  | |

I also looked into each features value to see whether there are missing values or anything out of ordinary. Of course there are features with very different value range, especially in financial features. However, I'm not going to exclude those extremely high values because many of them like extremely high bonuses actually can be an indicator of the fraud. Another step that I took was scaling the data. As mentioned before, there are two types of variables: financial and email. Email data has smaller numbers compared to financial data. Therefore, I've decided to scale them all.

| **Features** | **Values** | **Features** | **Values** |
| --- | --- | --- | --- |
| poi | 144 non-null | bonus | 81 non-null |
| salary | 94 non-null | salary\_to\_avg | 94 non-null |
| deferral\_payments | 38 non-null | total\_payments | 123 non-null |
| restricted\_stock\_deferred | 17 non-null | exercised\_stock\_options | 101 non-null |
| from\_poi\_to\_this\_person | 86 non-null | from\_this\_person\_to\_poi | 86 non-null |

In addition to the mentioned features, I made a new feature by dividing the salary of each executive to the average of all salaries. This ratio shows how ones salary is way above the average.

## Classifier Selection

Before the tuning step, I ran Gaussian Naive Bayes, K-Neighbors, and Decision Tree using their default parameters. While Gaussian Naive Bayes results were really underwhelming, the other two algorithms showed very good results.

| **Classifier** | **Accuracy** | **Recall** | **Precision** |
| --- | --- | --- | --- |
| Gaussian Naive Bayes | 0.28 | 0.16 | 1.0 |
| K-Neighbors | 0.82 | 0.34 | 0.25 |
| Decision Tree | 0.82 | 0.4 | 0.5 |

As you can see, with the default parameters, decision tree achieved the best accuracy, recall and precision. However, K-Neighbors has the same accuracy and the other two metrics are not as bad as the Gaussian Naive Bayes. Based on these results, I decided to move forward with both K-Neighbors and Decision Tree and improve both of them with scaling and feature selection.

## Tuning

Tuning the parameters of a machine learning algorithm means to manually change and optimize the values of them in order to enable the algorithm to perform the best. These parameters have a default value that will be applied if no other values have been assigned manually. For example, K-Neighbors has a parameter named weights, with the default value of 'uniform'. This parameter defines the weighting system for the points. The default value of 'uniform' means all points in each neighborhood has the same weight. However, this parameter can take another pre-defined value: 'distance'. This value, ’distance’, assigns the inverse of each point's distance as their weight. Instead of doing tuning one by one, I used the GridSearch and pipeline.

Pipeline allows a sequence of procedures chained together. GridSearch uses a grid of parameters and tries all possible combinations and returns the combination of parameters that have the maximum score. Using these two, I tried to find the best combination of methods and features to get the best results. For example, for K Neighbors Classifier I used the following pipeline and GridSearch:

pipe\_KNC = Pipeline([

('reduce\_dim', SelectKBest()),

('scaler', MinMaxScaler()),

('classify', KNeighborsClassifier())

grid\_KNC = GridSearchCV(pipe\_KNC, cv=sss, param\_grid=param\_grid\_KNC, scoring='recall')

Since there are some negative values present in the data and the scale of numbers among features are different, I took advantage of Feature scaling method: MinMaxScaler. This function uses the following formula to scale down all values into a number between 0 and 1.

Here, I chose K-Neighbors and Decision Tree for tuning. Here are the parameters that I chose for the GridSearch to tune:

**K-Neighbors**

| Parameter | Description | Values |
| --- | --- | --- |
| leaf\_size | Number of cases or observations in that leaf. | 30, 50 |
| n\_neighbors | Number of neighbors to use by default for k\_neighbors queries | 5, 10 |
| p | Power parameter for the Minkowski metric | 1, 2 |
| weights | Weighting system for the points | Distance, Uniform |
| algorithm | Algorithm used to compute the nearest neighbors | Auto, Ball\_Tree, Kd\_Tree, Brute |

**Decision Tree**

| Parameter | Description | Values |
| --- | --- | --- |
| criterion | The function to measure the quality of a split | Gini, Balanced |
| class\_weight | Weights associated with classes | None, Balanced |
| min\_samples\_split | The minimum number of samples required to split an internal node | 2, 3, 4, 5 |
| max\_leaf\_nodes | Grow a tree with max\_leaf\_nodes in best-first fashion | None, 5 |
| min\_samples\_leaf | The minimum number of samples required to be at a leaf node: | 4, 2 |

## Validation

The goal of a machine learning created model, is to predict the probability of occurring an event. Measuring the quality of the model is to see how well the model does this prediction, which is known as model validation. In order to do so, we need to split the dataset into training and test. The training data is used to build the model while the test data is used to measure the performance of the model. If we use the whole dataset to build and test the model, we may over fit it. One sign of over fitting is when the model performs well on the training and very poorly in validation (on test data). Keeping the training and test data separate and logically proportioned, is the best way to avoid over fitting. However, if your dataset is imbalanced (number of positive cases is much less than the number of negative cases), splitting your dataset into training and test is not the best idea because there is a chance that none of your positive cases falls in the test set. To overcome this problem, I used StratifiedShuffleSplit that iteratively provides train and split indices for 100 times.

Accuracy, recall and precision are the validation metric that I focused on for choosing my algorithm. these three metrics are using values from confusion matrix to evaluate a n algorithm. Accuracy shows the how many of the predicted labels were predicted correctly. In other words, accuracy is:

accuracy = (true positives + true negatives) / total predictions

However, depending on the scenario that we want to predict, the false positive or false negative predictions might be far more important for us. For example, when it comes to medical tests, specially in the first steps, the preference is to predict one might has a type of cancer and rule it out by further tests (false positive) rather than missing someone who has cancer and finding out about it when it is too late (false negative). For these cases, we can take advantage of two other metrics: precision and recall.

precision = true positives / (true positives + false positives)

recall = true positives / (true positives + false negatives)

The after tuning results shows although K neighbors has slightly higher accuracy after tuning, Decision Tree has much better Precision, Recall, F1, and F2. Therefore, I chose it as the final tuned algorithm.

| Classifier | Accuracy | Precision | Recall | F1 | F2 |
| --- | --- | --- | --- | --- | --- |
| KNeighborsClassifier | 0.87 | 0.54 | 0.18 | 0.27 | 0.21 |
| DecisionTreeClassifier | 0.82 | 0.35 | 0.43 | 0.38 | 0.41 |

The results of Decision Tree modes show the accuracy of the model is 82%. it mean out of 100 predictions, 82 of them will be predicted correctly. The precision is 35% which means 35% of the predictions that were identified as POIs, were actually POIs. Finally, the 43% value for Recall the model catches 43% of the POIs.

The SelectKBest chose 6 best features to be included in the model. As it is shown in the following table bonus, from\_poi\_to\_this\_person, and exercised\_stock\_options have significantly higher importance score compared to the rest:

* bonus, 0.41040635
* salary, 0.10045741
* salary\_to\_avg, 0.05440033
* total\_payments, 0.069393
* exercised\_stock\_options, 0.17840767
* from\_poi\_to\_this\_person, 0.18693525

## Discussion

In order to measure the effects of the added feature (salary\_to\_avg), I ran two sets of experiments: one with the added feature and one without it. Here is the summary of the results:

|  |  |  |  |
| --- | --- | --- | --- |
| **Type of Experiment** | **Accuracy** | **Precision** | **Recall** |
| Without the New Feature | 0.81207 | 0.33320 | 0.40900 |
| With the New Feature | 0.81820 | 0.35096 | 0.42800 |

The new feature slightly improved the validation results. So I would say choosing to make and keep “salary\_to\_avg” was a good call.