Personalized cancer diagnosis

1. Business Problem

1.1. Description

Source: https://www.kaggle.com/c/msk-redefining-cancer-treatment/

Data: Memorial Sloan Kettering Cancer Center (MSKCC)

Download training_variants.zip and training_text.zip from Kaggle.

Context:

Source: https://www.kaggle.com/c/msk-redefining-cancer-treatment/discussion/35336#198462

Problem statement:

Classify the given genetic variations/mutations based on evidence from text-based clinical literature.

1.2. Source/Useful Links

Some articles and reference blogs about the problem statement

- 1. https://www.forbes.com/sites/matthewherper/2017/06/03/a-new-cancer-drug-helped-almost-everyone-who-took-it-almost-heres-what-it-teaches-us/#2a44ee2f6b25)
- 2. https://www.youtube.com/watch?v=UwbuW7oK8rk)
- 3. https://www.youtube.com/watch?v=qxXRKVompI8 (https://www.youtube.com/watch?v=qxXRKVompI8)

1.3. Real-world/Business objectives and constraints.

- No low-latency requirement.
- Interpretability is important.
- Errors can be very costly.
- Probability of a data-point belonging to each class is needed.

2. Machine Learning Problem Formulation

2.1. Data

2.1.1. Data Overview

- Source: https://www.kaggle.com/c/msk-redefining-cancer-treatment/data (https://www.kaggle.com/c/msk-redefining-cancer-treatment/data)
- We have two data files: one conatins the information about the genetic mutations and the other contains the clinical evidence (text) that human experts/pathologists use to classify the genetic mutations.
- Both these data files are have a common column called ID
- Data file's information:
 - training_variants (ID , Gene, Variations, Class)
 - training_text (ID, Text)

2.1.2. Example Data Point

training_variants

ID,Gene,Variation,Class 0,FAM58A,Truncating Mutations,1 1,CBL,W802*,2 2,CBL,Q249E,2

...

training_text

ID, Text

0||Cyclin-dependent kinases (CDKs) regulate a variety of fundamental cellular processes. CDK10 stands out as one of the last orphan CDKs for which no activating cyclin has been identified and no kinase activity revealed. Previous work has shown that CDK10 silencing increases ETS2 (v-ets erythroblastosis virus E26 oncogene homolog 2)-driven activation of the MAPK pathway, which confers tamoxifen resistance to breast cancer cells. The precise mechanisms by which CDK10 modulates ETS2 activity, and more generally the functions of CDK10, remain elusive. Here we demonstrate that CDK10 is a cyclin-dependent kinase by identifying cyclin M as an activating cyclin. Cyclin M, an orphan cyclin, is the product of FAM58A, whose mutations cause STAR syndrome, a human developmental anomaly whose features include toe syndactyly, telecanthus, and anogenital and renal malformations. We show that STAR syndrome-associated cyclin M mutants are unable to interact with CDK10. Cyclin M silencing phenocopies CDK10 silencing in increasing c-Raf and in conferring tamoxifen resistance to breast cancer cells. CDK10/cyclin M phosphorylates ETS2 in vitro, and in cells it positively controls ETS2 degradation by the proteasome. ETS2 protein levels are increased in cells derived from a STAR patient, and this increase is attributable to decreased cyclin M levels. Altogether, our results reveal an additional regulatory mechanism for ETS2, which plays key roles in cancer and development. They also shed light on the molecular mechanisms underlying STAR syndrome. Cyclindependent kinases (CDKs) play a pivotal role in the control of a number of fundamental cellular processes (1). The human genome contains 21 genes encoding proteins that can be considered as members of the CDK family owing to their sequence similarity with bona fide CDKs, those known to be activated by cyclins (2). Although discovered almost 20 y ago (3, 4), CDK10 remains one of the two CDKs without an identified cyclin partner. This knowledge gap has largely impeded the exploration of its biological functions. CDK10 can act as a positive cell cycle regulator in some cells (5, 6) or as a tumor suppressor in others (7, 8). CDK10 interacts with the ETS2 (v-ets erythroblastosis virus E26 oncogene homolog 2) transcription factor and inhibits its transcriptional activity through an unknown mechanism (9). CDK10 knockdown derepresses ETS2, which increases the expression of the c-Raf protein kinase, activates the MAPK pathway, and induces resistance of MCF7 cells to tamoxifen (6). ...

2.2. Mapping the real-world problem to an ML problem

2.2.1. Type of Machine Learning Problem

There are nine different classes a genetic mutation can be classified into => Multi class classification problem

2.2.2. Performance Metric

Source: https://www.kaggle.com/c/msk-redefining-cancer-treatment#evaluation (https://www.kaggle.com/c/msk-redefining-cancer-treatment#evaluation)

Metric(s):

- Multi class log-loss
- Confusion matrix

2.2.3. Machine Learing Objectives and Constraints

Objective: Predict the probability of each data-point belonging to each of the nine classes.

Constraints:

- Interpretability
- · Class probabilities are needed.
- Penalize the errors in class probabilites => Metric is Log-loss.
- No Latency constraints.

2.3. Train, CV and Test Datasets

Split the dataset randomly into three parts train, cross validation and test with 64%,16%, 20% of data respectively

3. Exploratory Data Analysis

```
In [0]: import pandas as pd
        import matplotlib.pyplot as plt
        import re
        import time
        import warnings
        import numpy as np
        from nltk.corpus import stopwords
        from sklearn.decomposition import TruncatedSVD
        from sklearn.preprocessing import normalize
        from sklearn.feature_extraction.text import CountVectorizer
        from sklearn.manifold import TSNE
        import seaborn as sns
        from sklearn.neighbors import KNeighborsClassifier
        from sklearn.metrics import confusion_matrix
        from sklearn.metrics.classification import accuracy_score, log_loss
        from sklearn.feature extraction.text import TfidfVectorizer
        from sklearn.linear_model import SGDClassifier
        from imblearn.over_sampling import SMOTE
        from collections import Counter
        from scipy.sparse import hstack
        from sklearn.multiclass import OneVsRestClassifier
        from sklearn.svm import SVC
        from sklearn.model_selection import StratifiedKFold
        from collections import Counter, defaultdict
        from sklearn.calibration import CalibratedClassifierCV
        from sklearn.naive_bayes import MultinomialNB
        from sklearn.naive_bayes import GaussianNB
        from sklearn.model_selection import train_test_split
        from sklearn.model_selection import GridSearchCV
        import math
        from sklearn.metrics import normalized_mutual_info_score
        from sklearn.ensemble import RandomForestClassifier
        warnings.filterwarnings("ignore")
        from mlxtend.classifier import StackingClassifier
        from sklearn import model_selection
        from sklearn.linear_model import LogisticRegression
```

3.1. Reading Data

2 2

3 3

CBL

CBL

CBL

Q249E

N454D

L399V

3.1.1. Reading Gene and Variation Data

training/training_variants is a comma separated file containing the description of the genetic mutations used for training. Fields are

- ID: the id of the row used to link the mutation to the clinical evidence
- Gene: the gene where this genetic mutation is located
- Variation : the aminoacid change for this mutations
- Class: 1-9 the class this genetic mutation has been classified on

3.1.2. Reading Text Data

```
In [9]: # note the seprator in this file
         data_text =pd.read_csv("training_text",sep="\|\|",engine="python",names=["ID","TEXT"],skiprows=1)
         print('Number of data points : ', data_text.shape[0])
         print('Number of features : ', data_text.shape[1])
         print('Features : ', data_text.columns.values)
         data_text.head()
         Number of data points : 3321
         Number of features : 2
         Features : ['ID' 'TEXT']
Out[9]:
            ID
                                                   TEXT
                 Cyclin-dependent kinases (CDKs) regulate a var...
          0 0
          1 1
                   Abstract Background Non-small cell lung canc...
          2 2
                   Abstract Background Non-small cell lung canc...
          3 Recent evidence has demonstrated that acquired...
            4 Oncogenic mutations in the monomeric Casitas B...
```

3.1.3. Preprocessing of text

```
In [10]: | # loading stop words from nltk library
         import nltk
         nltk.download("stopwords")
         stop_words = set(stopwords.words('english'))
         def nlp_preprocessing(total_text, index, column):
              if type(total_text) is not int:
                 string = ""
                  # replace every special char with space
                 total_text = re.sub('[^a-zA-Z0-9\n]', ' ', total_text)
                  # replace multiple spaces with single space
                 total_text = re.sub('\s+',' ', total_text)
                  # converting all the chars into lower-case.
                  total_text = total_text.lower()
                  for word in total text.split():
                  # if the word is a not a stop word then retain that word from the data
                      if not word in stop_words:
                          string += word + " "
                  data_text[column][index] = string
```

[nltk_data] Downloading package stopwords to /root/nltk_data...
[nltk_data] Unzipping corpora/stopwords.zip.

Time took for preprocessing the text : 253.356001 seconds

```
In [11]: # Text processing stage.
    start_time = time.clock()
    for index, row in data_text.iterrows():
        if type(row['TEXT']) is str:
            nlp_preprocessing(row['TEXT'], index, 'TEXT')
        else:
            print("there is no text description for id:",index)
        print('Time took for preprocessing the text :',time.clock() - start_time, "seconds")

        there is no text description for id: 1109
        there is no text description for id: 1277
        there is no text description for id: 1407
        there is no text description for id: 1639
        there is no text description for id: 2755
```

```
In [12]: | # Merging both gene_variations and text data based on ID
           result = pd.merge(data_variants, data_text,on='ID', how='left')
           result.head()
Out[12]:
              ID
                                    Variation Class
                                                                                         TEXT
                     Gene
            0
               0
                  FAM58A
                           Truncating Mutations
                                                       cyclin dependent kinases cdks regulate variety...
               1
                      CBL
                                      W802*
                                                  2
                                                      abstract background non small cell lung cancer...
               2
            2
                      CBL
                                      Q249E
                                                      abstract background non small cell lung cancer...
                                                    recent evidence demonstrated acquired uniparen...
            3
               3
                      CBL
                                      N454D
                                       L399V
                                                    oncogenic mutations monomeric casitas b lineag...
              4
                      CBL
           result[result.isnull().any(axis=1)]
Out[13]:
                    ID
                         Gene
                                         Variation Class TEXT
                                          S1088F
            1109 1109
                        FANCA
                                                          NaN
                                                       1
            1277 1277 ARID5B Truncating Mutations
                                                          NaN
            1407 1407
                        FGFR3
                                           K508M
                                                          NaN
                                      Amplification
            1639 1639
                          FLT1
                                                          NaN
                                           G596C
            2755 2755
                         BRAF
                                                          NaN
 In [0]: | result.loc[result['TEXT'].isnull(), 'TEXT'] = result['Gene'] +' '+result['Variation']
In [15]: | result[result['ID']==1109]
Out[15]:
                    ID
                         Gene
                               Variation Class
                                                        TEXT
            1109 1109 FANCA
                                             1 FANCA S1088F
                                S1088F
```

3.1.4. Test, Train and Cross Validation Split

3.1.4.1. Splitting data into train, test and cross validation (64:20:16)

```
In [16]: | result.Gene = result.Gene.str.replace('\s+', '_')
         result.Variation = result.Variation.str.replace('\s+', '_')
         y_true = result[['Class']]
         x_true = result.drop(['Class'], axis=1)
         print("Feature columns in dataset: ")
         print(x_true.head())
         print()
         print("Target columns in dataset: ")
         print(y_true.head())
         Feature columns in dataset:
            ID ...
                                                                   TEXT
                ... cyclin dependent kinases cdks regulate variety...
                     abstract background non small cell lung cancer...
         1
                     abstract background non small cell lung cancer...
                     recent evidence demonstrated acquired uniparen...
                     oncogenic mutations monomeric casitas b lineag...
         [5 rows x 4 columns]
         Target columns in dataset:
            Class
                2
         1
         2
                2
         3
                3
 In [0]: | # Split the data into test and train by maintaining same distribution of output varaible 'y_true' [stratify=y_true]
         x_train, x_test, y_train, y_test = train_test_split(x_true, y_true, stratify=y_true, test_size=0.2)
         # Split the train data into train and cross validation by maintaining same distribution of output varaible 'y_train'
          [stratify=y_train]
         x_train, x_cv, y_train, y_cv = train_test_split(x_train, y_train, stratify=y_train, test_size=0.2)
```

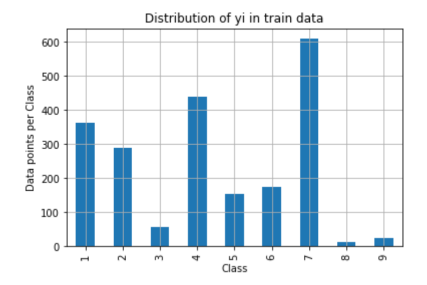
```
In [18]: print('Number of data points in train data:', x_train.shape[0])
    print('Number of data points in test data:', x_test.shape[0])
    print('Number of data points in cross validation data:', x_cv.shape[0])

Number of data points in train data: 2124
    Number of data points in test data: 665
```

3.1.4.2. Distribution of y_i's in Train, Test and Cross Validation datasets

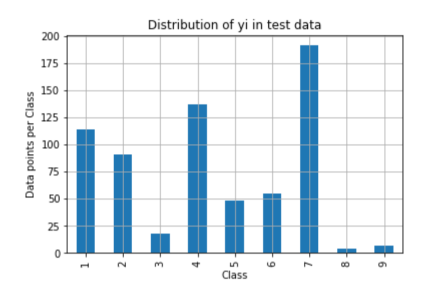
Number of data points in cross validation data: 532

```
In [21]: | def plot_distribution(class_distribution,title,xlabel,ylabel):
             class_distribution.plot(kind='bar')
             plt.xlabel(xlabel)
             plt.ylabel(ylabel)
             plt.title(title)
             plt.grid()
             plt.show()
         # it returns a dict, keys as class labels and values as the number of data points in that class
         train_class_distribution = y_train['Class'].value_counts().sort_index()
         test_class_distribution = y_test['Class'].value_counts().sort_index()
         cv_class_distribution = y_cv['Class'].value_counts().sort_index()
         plot_distribution(train_class_distribution,
                           'Distribution of yi in train data',
                           'Class',
                           'Data points per Class')
         # ref: argsort https://docs.scipy.org/doc/numpy/reference/generated/numpy.argsort.html
         # -(train class_distribution.values): the minus sign will give us in decreasing order
         sorted_yi = np.argsort(-train_class_distribution.values)
         for i in sorted_yi:
             print('Number of data points in class', i+1, ':',train_class_distribution.values[i],
                    '(', np.round((train_class_distribution.values[i]/x_train.shape[0]*100), 3), '%)')
         print('-'*80)
         plot_distribution(test_class_distribution,
                           'Distribution of yi in test data',
                           'Class',
                           'Data points per Class')
         # ref: argsort https://docs.scipy.org/doc/numpy/reference/generated/numpy.argsort.html
         # -(test_class_distribution.values): the minus sign will give us in decreasing order
         sorted_yi = np.argsort(-test_class_distribution.values)
         for i in sorted_yi:
             print('Number of data points in class', i+1, ':',test_class_distribution.values[i],
                    '(', np.round((test_class_distribution.values[i]/x_test.shape[0]*100), 3), '%)')
         print('-'*80)
         plot_distribution(cv_class_distribution,
                           'Distribution of yi in cross validation data',
                           'Class',
                           'Data points per Class')
         # ref: argsort https://docs.scipy.org/doc/numpy/reference/generated/numpy.argsort.html
         # -(cv_class_distribution.values): the minus sign will give us in decreasing order
         sorted_yi = np.argsort(-cv_class_distribution.values)
         for i in sorted_yi:
             print('Number of data points in class', i+1, ':',cv_class_distribution.values[i],
                    '(', np.round((cv_class_distribution.values[i]/x_cv.shape[0]*100), 3), '%)')
```

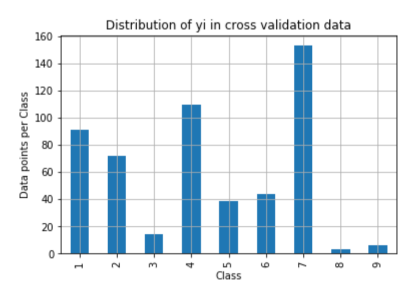


```
Number of data points in class 7 : 609 ( 28.672 %)
Number of data points in class 4 : 439 ( 20.669 %)
Number of data points in class 1 : 363 ( 17.09 %)
Number of data points in class 2 : 289 ( 13.606 %)
Number of data points in class 6 : 176 ( 8.286 %)
Number of data points in class 5 : 155 ( 7.298 %)
Number of data points in class 3 : 57 ( 2.684 %)
Number of data points in class 9 : 24 ( 1.13 %)
Number of data points in class 8 : 12 ( 0.565 %)
```

-----/



Number of data points in class 7 : 191 (28.722 %) Number of data points in class 4 : 137 (20.602 %) Number of data points in class 1 : 114 (17.143 %) Number of data points in class 2 : 91 (13.684 %) Number of data points in class 6 : 55 (8.271 %) Number of data points in class 5 : 48 (7.218 %) Number of data points in class 3 : 18 (2.707 %) Number of data points in class 9 : 7 (1.053 %) Number of data points in class 8 : 4 (0.602 %)



```
Number of data points in class 7 : 153 ( 28.759~\%) Number of data points in class 4 : 110~(~20.677~\%) Number of data points in class 1 : 91~(~17.105~\%) Number of data points in class 2 : 72~(~13.534~\%) Number of data points in class 6 : 44~(~8.271~\%) Number of data points in class 5 : 39~(~7.331~\%) Number of data points in class 3 : 14~(~2.632~\%) Number of data points in class 9 : 6~(~1.128~\%) Number of data points in class 8 : 3~(~0.564~\%)
```

In a 'Random' Model, we generate the '9' class probabilites randomly such that they sum to 1.

```
In [0]: def plot_matrix(matrix,labels):
            plt.figure(figsize=(20,7))
            sns.heatmap(matrix, annot=True, cmap="Y1GnBu", fmt=".3f", xticklabels=labels, yticklabels=labels)
            plt.xlabel('Predicted Class')
            plt.ylabel('Original Class')
            plt.show()
        # This function plots the confusion matrices given y_i, y_i_hat.
        def plot_confusion_matrix(test_y, predict_y):
            cm = confusion_matrix(test_y, predict_y)
            \# C = 9,9 matrix, each cell (i,j) represents number of points of class i are predicted class j
            recall_table =(((cm.T)/(cm.sum(axis=1))).T)
            # How did we calculateed recall_table :
            # divide each element of the confusion matrix with the sum of elements in that column
            \# C = [[1, 2],
                  [3, 4]]
            # C.T = [[1, 3],
                     [2, 4]]
            # C.sum(axis = 1) axis=0 corresonds to columns and axis=1 corresponds to rows in two diamensional array
            \# C.sum(axix = 1) = [[3, 7]]
            \# ((C.T)/(C.sum(axis=1))) = [[1/3, 3/7]
                                        [2/3, 4/7]]
            \# ((C.T)/(C.sum(axis=1))).T = [[1/3, 2/3]
                                        [3/7, 4/7]]
            # sum of row elements = 1
            precision_table =(cm/cm.sum(axis=0))
            # How did we calculateed precision_table :
            # divide each element of the confusion matrix with the sum of elements in that row
            \# C = [[1, 2],
                  [3, 4]]
            # C.sum(axis = 0) axis=0 corresonds to columns and axis=1 corresponds to rows in two diamensional array
            # C.sum(axix = 0) = [[4, 6]]
            \# (C/C.sum(axis=0)) = [[1/4, 2/6],
                                   [3/4, 4/6]]
            labels = [1,2,3,4,5,6,7,8,9]
            print()
            print("-"*20, "Confusion matrix", "-"*20)
            plot_matrix(cm,labels)
            print("-"*20, "Precision matrix (Column Sum=1)", "-"*20)
            plot_matrix(precision_table,labels)
            print("-"*20, "Recall matrix (Row sum=1)", "-"*20)
            plot_matrix(recall_table,labels)
```

```
In [23]: # We need to generate 9 numbers and the sum of numbers should be 1
         # one solution is to genarate 9 numbers and divide each of the numbers by their sum
         # ref: https://stackoverflow.com/a/18662466/4084039
         test_data_len = x_test.shape[0]
         cv_data_len = x_cv.shape[0]
         # we create a output array that has exactly same size as the CV data
         cv_predicted_y = np.zeros((cv_data_len,9))
         for i in range(cv_data_len):
             rand_probs = np.random.rand(1,9)
             cv_predicted_y[i] = ((rand_probs/sum(sum(rand_probs)))[0])
         print("Log loss on Cross Validation Data using Random Model",log_loss(y_cv,cv_predicted_y, eps=1e-15))
         # Test-Set error.
         # We create a output array that has exactly same as the test data
         test_predicted_y = np.zeros((test_data_len,9))
         for i in range(test_data_len):
             rand_probs = np.random.rand(1,9)
             test_predicted_y[i] = ((rand_probs/sum(sum(rand_probs)))[0])
         print("Log loss on Test Data using Random Model",log_loss(y_test,test_predicted_y, eps=1e-15))
         predicted_y =np.argmax(test_predicted_y, axis=1)
         plot_confusion_matrix(y_test, predicted_y+1)
```

----- Confusion matrix -----

11.000 11.000 11.000 12.000 11.000 6.000 9.000 8.000 11.000 11.000 10.000 7.000 - 20 1.000 2.000 2.000 1.000 1.000 2.000 3.000 4.000 2.000 12.000 12.000 Original Class 9.000 4.000 7.000 4.000 6.000 10.000 3.000 2.000 3.000 7.000 5.000 6.000 3.000 6.000 8.000 9.000 3.000 8.000 - 10 24.000 27.000 23.000 26.000 1.000 0.000 0.000 1.000 1.000 0.000 1.000 0.000 0.000 0.000 0.000 0.000 1.000 1.000 1.000 1.000 1.000 2.000 Predicted Class ----- Precision matrix (Columm Sum=1) ------0.157 0.149 0.143 0.151 - 0.30 0.129 0.090 0.138 0.114 0.149 0.119 0.097 0.015 0.031 0.029 0.013 0.012 0.027 0.036 0.055 0.028 - 0.24 Original Class -0.18 0.134 0.062 0.100 0.053 0.071 0.135 0.036 0.027 0.042 0.077 0.086 0.040 0.071 0.123 0.111 0.104 0.041 0.095 - 0.12 0.308 0.343 - 0.06 0.015 0.000 0.000 0.013 0.012 0.012 0.000 0.000 0.000 0.000 0.000 0.013 0.012 0.014 0.012 0.014 0.028 - 0.00 Predicted Class ----- Recall matrix (Row sum=1) ------0.123 0.096 0.096 0.132 0.096 0.105 0.096 0.114 - 0.25 0.088 0.121 0.066 0.099 0.121 0.110 0.077 0.111 0.111 0.056 0.056 0.111 0.111 0.056 - 0.20 0.088 0.102 0.088 0.124 0.117 0.095 0.109 0.131 - 0.15 0.083 0.083 0.125 0.062 0.042 0.062 0.091 0.109 0.055 0.109 -0.10 0.089 0.105 0.126 0.105 0.120 0.136 0.079 0.099 - 0.05 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.286 - 0.00 4 6 7 8 9 Predicted Class

3.3 Univariate Analysis

```
In [0]: | # code for response coding with Laplace smoothing.
        # alpha : used for laplace smoothing
        # feature: ['gene', 'variation']
        # df: ['x_train', 'x_test', 'x_cv']
        # algorithm
        # -----
        # Consider all unique values and the number of occurances of given feature in train data dataframe
        # build a vector (1*9), the first element = (number\ of\ times\ it\ occurred\ in\ class1 + 10*alpha / number\ of\ time\ it\ occurred
        rred in total data+90*alpha)
        # gv_dict is like a look up table, for every gene it store a (1*9) representation of it
        # for a value of feature in df:
        # if it is in train data:
        # we add the vector that was stored in 'gv_dict' look up table to 'gv_fea'
        # if it is not there is train:
        # we add [1/9, 1/9, 1/9, 1/9, 1/9, 1/9, 1/9, 1/9] to 'qv_fea'
        # return 'gv_fea'
        # get_gv_fea_dict: Get Gene varaition Feature Dict
        def get_gv_fea_dict(alpha, feature, df):
            # value_count: it contains a dict like
            # print(train_df['Gene'].value_counts())
            # output:
                  {BRCA1
                               174
            #
                    TP53
                               106
                   EGFR
            #
                               86
                   BRCA2
                                75
            #
                    PTEN
                   KIT
            #
                                61
                   BRAF
            #
                                60
                   ERBB2
                                 47
                   PDGFRA
                     ...}
            # print(train_df['Variation'].value_counts())
            # output:
            # {
            # Truncating_Mutations
                                                     63
                                                     43
            # Deletion
            # Amplification
                                                     43
            # Fusions
                                                     22
            # Overexpression
                                                     3
            # E17K
                                                      3
            # Q61L
                                                      3
            # S222D
                                                      2
            # P130S
            # ...
            # }
            value_count = x_train[feature].value_counts()
            # gv_dict : Gene Variation Dict, which contains the probability array for each gene/variation
            gv_dict = dict()
            # denominator will contain the number of time that particular feature occured in whole data
            for i, denominator in value_count.items():
                # vec will contain (p(yi==1/Gi)) probability of gene/variation belongs to perticular class
                # vec is 9 diamensional vector
               vec = []
                for k in range(1,10):
                   # print(train_df.loc[(train_df['Class']==1) & (train_df['Gene']=='BRCA1')])
                            ID Gene Variation Class
                   # 2470 2470 BRCA1
                                                   S1715C 1
                   # 2486 2486 BRCA1
                                                    S1841R
                                                               1
                   # 2614 2614 BRCA1
                                                      M1R
                                                   L1657P
                   # 2432 2432 BRCA1
                                                               1
                   # 2567 2567 BRCA1
                                                   T1685A
                                                                1
                                                   E1660G
                    # 2583 2583 BRCA1
                                                                 1
                   # 2634 2634 BRCA1
                                                     W1718L
                    # cls_cnt.shape[0] will return the number of rows
                    cls_cnt = x_train.loc[(y_train['Class']==k) & (x_train[feature]==i)]
                    # cls_cnt.shape[0](numerator) will contain the number of time that particular feature occured in whole dat
                    vec.append((cls_cnt.shape[0] + alpha*10)/ (denominator + 90*alpha))
                # we are adding the gene/variation to the dict as key and vec as value
                gv dict[i]=vec
            return gv dict
        # Get Gene variation feature
        def get_gv_feature(alpha, feature, df):
            # print(gv_dict)
                 {'BRCA1': [0.20075757575757575, 0.037878787878788, 0.0681818181818177, 0.13636363636363635, 0.25, 0.1931
        81818181818, 0.03787878787878788, 0.0378787878788, 0.037878787878788],
                  'TP53': [0.32142857142857145, 0.061224489795918366, 0.061224489795918366, 0.27040816326530615, 0.0612244897
        95918366, 0.066326530612244902, 0.051020408163265307, 0.051020408163265307, 0.056122448979591837],
                  'EGFR': [0.0568181818181816, 0.21590909090909091, 0.0625, 0.0681818181818177, 0.0681818181818177, 0.0
        625, 0.34659090909090912, 0.0625, 0.056818181818181816],
```

```
'BRCA2': [0.13333333333333333, 0.060606060606060608, 0.0606060606060608, 0.078787878787878782, 0.13939393
93939394, 0.34545454545454546, 0.060606060606060608, 0.06060606060608, 0.060606060606060608],
          'PTEN': [0.069182389937106917, 0.062893081761006289, 0.069182389937106917, 0.46540880503144655, 0.075471698
113207544, 0.062893081761006289, 0.069182389937106917, 0.062893081761006289, 0.062893081761006289],
          'KIT': [0.066225165562913912, 0.25165562913907286, 0.072847682119205295, 0.072847682119205295, 0.0662251655
62913912, 0.066225165562913912, 0.27152317880794702, 0.066225165562913912, 0.066225165562913912],
          'BRAF': [0.066666666666666666, 0.17999999999999, 0.073333333333334, 0.073333333333334, 0.093333333
333333338, 0.0800000000000000000, 0.2999999999999, 0.066666666666666, 0.0666666666666666,
   #
   gv_dict = get_gv_fea_dict(alpha, feature, df)
   # value_count is similar in get_gv_fea_dict
   value_count = x_train[feature].value_counts()
   # gv_fea: Gene_variation feature, it will contain the feature for each feature value in the data
   gv_fea = []
    # for every feature values in the given data frame we will check if it is there in the train data then we will add
the feature to gv_fea
   # if not we will add [1/9,1/9,1/9,1/9,1/9,1/9,1/9,1/9] to gv_fea
   for index, row in df.iterrows():
        if row[feature] in dict(value_count).keys():
            gv_fea.append(gv_dict[row[feature]])
        else:
            gv_fea.append([1/9,1/9,1/9,1/9,1/9,1/9,1/9,1/9])
             gv_fea.append([-1,-1,-1,-1,-1,-1,-1,-1])
   return gv_fea
```

when we caculate the probability of a feature belongs to any particular class, we apply laplace smoothing

• (numerator + 10*alpha) / (denominator + 90*alpha)

3.2.1 Univariate Analysis on Gene Feature

Q1. Gene, What type of feature it is?

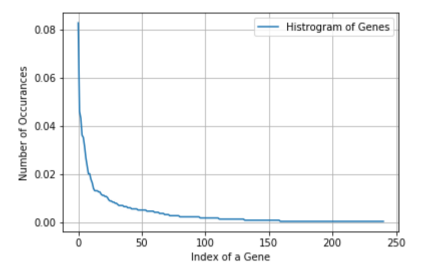
Ans. Gene is a categorical variable

Q2. How many categories are there and How they are distributed?

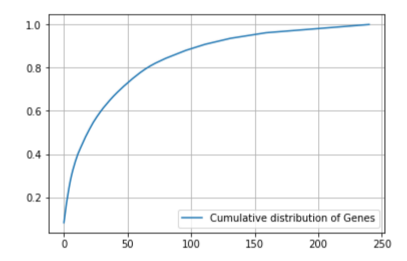
```
In [25]: | unique_genes = x_train['Gene'].value_counts()
         print('Number of Unique Genes :', unique_genes.shape[0])
         # the top 10 genes that occured most
         print(unique_genes.head(10))
         Number of Unique Genes: 241
         BRCA1
                   176
         TP53
                    98
         EGFR
                    92
         BRCA2
                    77
         PTEN
                    75
         KIT
                    67
         BRAF
                    57
         ERBB2
                    50
                    43
         ALK
         PDGFRA
                    43
         Name: Gene, dtype: int64
In [26]: print("Ans: There are", unique_genes.shape[0], "different categories of genes in the train data, and they are distibut
         ed as follows",)
```

Ans: There are 241 different categories of genes in the train data, and they are distibuted as follows

```
In [27]: s = sum(unique_genes.values);
h = unique_genes.values/s;
plt.plot(h, label="Histrogram of Genes")
plt.xlabel('Index of a Gene')
plt.ylabel('Number of Occurances')
plt.legend()
plt.grid()
plt.show()
```



```
In [28]: c = np.cumsum(h)
    plt.plot(c,label='Cumulative distribution of Genes')
    plt.grid()
    plt.legend()
    plt.show()
```



Q3. How to featurize this Gene feature ?

Ans.there are two ways we can featurize this variable check out this video: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/handling-categorical-and-numerical-features/

- 1. One hot Encoding
- 2. Response coding

We will choose the appropriate featurization based on the ML model we use. For this problem of multi-class classification with categorical features, one-hot encoding is better for Logistic regression while response coding is better for Random Forests.

```
In [0]: #response-coding of the Gene feature
    # alpha is used for laplace smoothing
    alpha = 1
    # train gene feature
    train_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_train))
    # test gene feature
    test_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_test))
# cross validation gene feature
    cv_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_cv))
```

```
In [30]: print("train_gene_feature_responseCoding is converted feature using respone coding method. The shape of gene feature:"
   , train_gene_feature_responseCoding.shape)
```

train_gene_feature_responseCoding is converted feature using respone coding method. The shape of gene feature: (2124, 9)

```
In [0]: # one-hot encoding of Gene feature.
gene_vectorizer = TfidfVectorizer()
    train_gene_feature_onehotCoding = gene_vectorizer.fit_transform(x_train['Gene'])
    test_gene_feature_onehotCoding = gene_vectorizer.transform(x_test['Gene'])
    cv_gene_feature_onehotCoding = gene_vectorizer.transform(x_cv['Gene'])
```

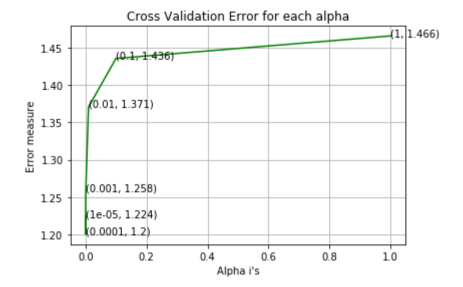
Q4. How good is this gene feature in predicting y_i?

240)

There are many ways to estimate how good a feature is, in predicting y_i. One of the good methods is to build a proper ML model using just this feature. In this case, we will build a logistic regression model using only Gene feature (one hot encoded) to predict y_i.

```
In [34]: alpha = [10 ** x for x in range(-5, 1)] # hyperparam for SGD classifier.
         # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear model.SGDClassifi
         er.html
         # default parameters
         # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
         # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
         # class weight=None, warm start=False, average=False, n iter=None)
         # some of methods
         # fit(X, y[, coef_init, intercept_init, ...])
                                                       Fit linear model with Stochastic Gradient Descent.
         # predict(X)
                       Predict class labels for samples in X.
         #-----
         # video Link:
         cv_log_error_array=[]
         for i in alpha:
             clf = SGDClassifier(alpha=i, penalty='12', loss='log', random_state=42)
             clf.fit(train_gene_feature_onehotCoding, y_train)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_gene_feature_onehotCoding, y_train)
             predict_y = sig_clf.predict_proba(cv_gene_feature_onehotCoding)
             cv_log_error_array.append(log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
             print('For values of alpha = ', i, "The log loss is:",log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],np.round(txt,3)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = SGDClassifier(alpha=alpha[best alpha], penalty='12', loss='log', random state=42)
         clf.fit(train_gene_feature_onehotCoding, y_train)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_gene_feature_onehotCoding, y_train)
         predict_y = sig_clf.predict_proba(train_gene_feature_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_gene_feature_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_gene_feature_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

```
For values of alpha = 1e-05 The log loss is: 1.2236781162199728
For values of alpha = 0.0001 The log loss is: 1.199893678134948
For values of alpha = 0.001 The log loss is: 1.2576124856590776
For values of alpha = 0.01 The log loss is: 1.3706776000961103
For values of alpha = 0.1 The log loss is: 1.435762543658923
For values of alpha = 1 The log loss is: 1.4659788505597753
```



```
For values of best alpha = 0.0001 The train log loss is: 0.994115089745061

For values of best alpha = 0.0001 The cross validation log loss is: 1.199893678134948

For values of best alpha = 0.0001 The test log loss is: 1.1739628975567062
```

Q5. Is the Gene feature stable across all the data sets (Test, Train, Cross validation)?

Ans. Yes, it is. Otherwise, the CV and Test errors would be significantly more than train error.

3.2.2 Univariate Analysis on Variation Feature

Q7. Variation, What type of feature is it?

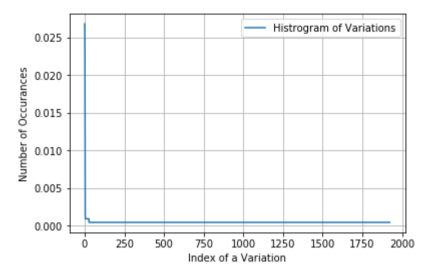
Ans. Variation is a categorical variable

Q8. How many categories are there?

```
In [36]: unique_variations = x_train['Variation'].value_counts()
         print('Number of Unique Variations :', unique_variations.shape[0])
         # the top 10 variations that occured most
         print(unique_variations.head(10))
         Number of Unique Variations: 1923
         Truncating_Mutations
                                       57
                                       56
         Deletion
         Amplification
                                       47
         Fusions
                                       20
         G12V
                                        3
                                        2
         G67R
                                        2
         P34R
         E542K
                                        2
         Promoter_Hypermethylation
                                        2
         R170W
         Name: Variation, dtype: int64
In [37]: print("Ans: There are", unique_variations.shape[0] ,
                "different categories of variations in the train data, and they are distibuted as follows",)
```

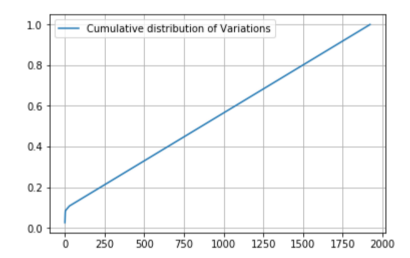
Ans: There are 1923 different categories of variations in the train data, and they are distibuted as follows

```
In [38]: s = sum(unique_variations.values);
h = unique_variations.values/s;
plt.plot(h, label="Histrogram of Variations")
plt.xlabel('Index of a Variation')
plt.ylabel('Number of Occurances')
plt.legend()
plt.grid()
plt.show()
```



```
In [39]: c = np.cumsum(h)
    print(c)
    plt.plot(c,label='Cumulative distribution of Variations')
    plt.grid()
    plt.legend()
    plt.show()
```

[0.02683616 0.05320151 0.07532957 ... 0.99905838 0.99952919 1.]



Q9. How to featurize this Variation feature?

Ans.There are two ways we can featurize this variable check out this video: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/handling-categorical-and-numerical-features/

- 1. One hot Encoding
- 2. Response coding

We will be using both these methods to featurize the Variation Feature

```
In [0]: # alpha is used for laplace smoothing
alpha = 1

# train gene feature
train_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_train))

# test gene feature
test_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_test))

# cross validation gene feature
cv_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_cv))
```

train_variation_feature_responseCoding is a converted feature using the response coding method. The shape of Variation feature: (2124, 9)

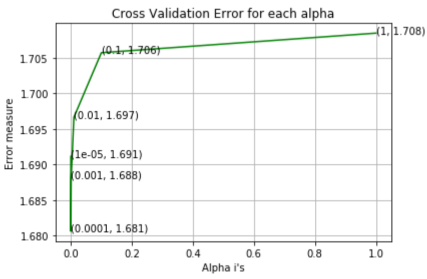
```
In [0]: # one-hot encoding of variation feature.
    variation_vectorizer = TfidfVectorizer()
    train_variation_feature_onehotCoding = variation_vectorizer.fit_transform(x_train['Variation'])
    test_variation_feature_onehotCoding = variation_vectorizer.transform(x_test['Variation'])
    cv_variation_feature_onehotCoding = variation_vectorizer.transform(x_cv['Variation'])
```

train_variation_feature_onehotEncoded is converted feature using the onne-hot encoding method. The shape of Variation feature: (2124, 1949)

Q10. How good is this Variation feature in predicting y_i?

Let's build a model just like the earlier!

```
In [44]: | alpha = [10 ** x for x in range(-5, 1)]
         # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear model.SGDClassifi
         er.html
         # default parameters
         # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
         # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
         # class_weight=None, warm_start=False, average=False, n_iter=None)
         # some of methods
         # fit(X, y[, coef_init, intercept_init, ...])
                                                       Fit linear model with Stochastic Gradient Descent.
         # predict(X)
                       Predict class labels for samples in X.
         #-----
         # video link:
         cv log error array=[]
         for i in alpha:
             clf = SGDClassifier(alpha=i, penalty='12', loss='log', random_state=42)
             clf.fit(train_variation_feature_onehotCoding, y_train)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_variation_feature_onehotCoding, y_train)
             predict_y = sig_clf.predict_proba(cv_variation_feature_onehotCoding)
             cv_log_error_array.append(log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
             print('For values of alpha = ', i, "The log loss is:",log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],np.round(txt,3)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train_variation_feature_onehotCoding, y_train)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_variation_feature_onehotCoding, y_train)
         predict_y = sig_clf.predict_proba(train_variation_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The train log loss is:",log_loss(y_train, predict_y, labels=c
         lf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_variation_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The cross validation log loss is:",log_loss(y_cv, predict_y,
         labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_variation_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The test log loss is:",log_loss(y_test, predict_y, labels=clf
         .classes_, eps=1e-15))
         For values of alpha = 1e-05 The log loss is: 1.691139709117014
         For values of alpha = 0.0001 The log loss is: 1.6806145419757827
         For values of alpha = 0.001 The log loss is: 1.6880558987263015
         For values of alpha = 0.01 The log loss is: 1.6966520646198406
         For values of alpha = 0.1 The log loss is: 1.7057342175637962
         For values of alpha = 1 The log loss is: 1.7084878704532445
                        Cross Validation Error for each alpha
                                                          (1, 1.708)
```



```
For values of best alpha = 0.0001 The train log loss is: 0.7456543819919453

For values of best alpha = 0.0001 The cross validation log loss is: 1.6806145419757827

For values of best alpha = 0.0001 The test log loss is: 1.717870407542839
```

Q11. Is the Variation feature stable across all the data sets (Test, Train, Cross validation)?

Ans. Not sure! But lets be very sure using the below analysis.

3.2.3 Univariate Analysis on Text Feature

```
1. How many unique words are present in train data?
```

- 2. How are word frequencies distributed?
- 3. How to featurize text field?
- 4. Is the text feature useful in predicitng y_i?
- 5. Is the text feature stable across train, test and CV datasets?

Name: TEXT, dtype: object

lung adenocarcinoma comprised distinct mutatio...

```
In [0]: def top_tfidf_feats(row, features, top_n=25):
              ''' Get top n tfidf values in row and return them with their corresponding feature names.'''
             topn_ids = np.argsort(row)[::-1][:top_n]
             top_feats = [(features[i], row[i]) for i in topn_ids]
             df = pd.DataFrame(top_feats)
             df.columns = ['feature', 'tfidf']
             return df
         def top_mean_feats(Xtr, features, min_tfidf=0.1, grp_ids=None, top_n=25):
              ''' Return the top n features that on average are most important amongst documents in rows
                  indentified by indices in grp_ids. '''
             if grp_ids:
                 D = Xtr[grp_ids].toarray()
             else:
                 D = Xtr.toarray()
             D[D < min_tfidf] = 0</pre>
             tfidf_means = np.mean(D, axis=0)
             return top_tfidf_feats(tfidf_means, features, top_n)
 In [0]: # building a CountVectorizer with all the words that occured minimum 3 times in train data
         text_vectorizer = TfidfVectorizer(min_df=3)
         train_text_feature_onehotCoding = text_vectorizer.fit_transform(x_train['TEXT'])
         # getting top 1000 feature names (words)
         train_text_features = top_mean_feats(train_text_feature_onehotCoding,
                                                          text_vectorizer.get_feature_names(),
                                                          top_n=1000)['feature'].tolist()
In [51]: | # train_text_feature_onehotCoding.sum(axis=0).A1 will sum every row and returns (1*number of features) vector
         train_text_fea_counts = train_text_feature_onehotCoding.sum(axis=0).A1
         train_text_fea_counts
Out[51]: array([9.31658646, 9.06079609, 0.04509122, ..., 0.03620948, 0.02890577,
                0.07312701])
In [52]: | # zip(list(text_features),text_fea_counts) will zip a word with its number of times it occured
         text_fea_dict = dict(zip(list(train_text_features),train_text_fea_counts))
         print("Total number of unique words in train data :", len(train_text_features))
         Total number of unique words in train data : 1000
 In [0]: | dict_list = []
         # dict_list =[] contains 9 dictoinaries each corresponds to a class
         for i in range(1,10):
             cls_text = x_train[y_train['Class']==i]
             # build a word dict based on the words in that class
             dict_list.append(extract_dictionary_paddle(cls_text))
             # append it to dict_list
         # dict_list[i] is build on i'th class text data
         # total_dict is buid on whole training text data
         total_dict = extract_dictionary_paddle(x_train)
         confuse_array = []
         for i in train_text_features:
             ratios = []
             \max val = -1
             for j in range(0,9):
                  ratios.append((dict_list[j][i]+10 )/(total_dict[i]+90))
             confuse_array.append(ratios)
         confuse_array = np.array(confuse_array)
 In [0]: | #response coding of text features
         train_text_feature_responseCoding = get_text_responsecoding(x_train)
         test_text_feature_responseCoding = get_text_responsecoding(x_test)
         cv_text_feature_responseCoding = get_text_responsecoding(x_cv)
 In [0]: # https://stackoverflow.com/a/16202486
         # we convert each row values such that they sum to 1
         train_text_feature_responseCoding = (train_text_feature_responseCoding.T/train_text_feature_responseCoding.sum(axis=1)
         )).T
```

test_text_feature_responseCoding = (test_text_feature_responseCoding.T/test_text_feature_responseCoding.sum(axis=1)).T

cv_text_feature_responseCoding = (cv_text_feature_responseCoding.T/cv_text_feature_responseCoding.sum(axis=1)).T

```
In [0]: # don't forget to normalize every feature
    train_text_feature_onehotCoding = normalize(train_text_feature_onehotCoding, axis=0)

# we use the same vectorizer that was trained on train data
    test_text_feature_onehotCoding = text_vectorizer.transform(x_test['TEXT'])
    # don't forget to normalize every feature
    test_text_feature_onehotCoding = normalize(test_text_feature_onehotCoding, axis=0)

# we use the same vectorizer that was trained on train data
    cv_text_feature_onehotCoding = text_vectorizer.transform(x_cv['TEXT'])
    # don't forget to normalize every feature
    cv_text_feature_onehotCoding = normalize(cv_text_feature_onehotCoding, axis=0)
```

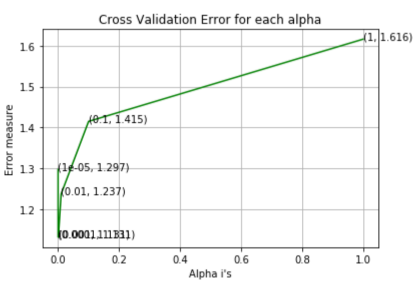
```
In [0]: #https://stackoverflow.com/a/2258273/4084039
sorted_text_fea_dict = dict(sorted(text_fea_dict.items(), key=lambda x: x[1] , reverse=True))
sorted_text_occur = np.array(list(sorted_text_fea_dict.values()))
```

In [58]: # Number of words for a given frequency.
print(Counter(sorted_text_occur))

```
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```

```
In [59]: # Train a Logistic regression+Calibration model using text features whicha re on-hot encoded
         alpha = [10 ** x for x in range(-5, 1)]
         # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear_model.SGDClassifi
         er.html
         # -----
         # default parameters
         # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
         # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
         # class_weight=None, warm_start=False, average=False, n_iter=None)
         # some of methods
         # fit(X, y[, coef_init, intercept_init, ...])
                                                      Fit linear model with Stochastic Gradient Descent.
         \# predict(X) Predict class labels for samples in X.
         # video link:
         #-----
         cv_log_error_array=[]
         for i in alpha:
             clf = SGDClassifier(alpha=i, penalty='12', loss='log', random_state=42)
             clf.fit(train_text_feature_onehotCoding, y_train)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_text_feature_onehotCoding, y_train)
             predict_y = sig_clf.predict_proba(cv_text_feature_onehotCoding)
             cv_log_error_array.append(log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
             print('For values of alpha = ', i, "The log loss is:",log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],np.round(txt,3)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train_text_feature_onehotCoding, y_train)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_text_feature_onehotCoding, y_train)
         predict_y = sig_clf.predict_proba(train_text_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The train log loss is:",log_loss(y_train, predict_y, labels=c
         lf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_text_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The cross validation log loss is:",log_loss(y_cv, predict_y,
         labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_text_feature_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The test log loss is:",log_loss(y_test, predict_y, labels=clf
         .classes_, eps=1e-15))
         For values of alpha = 1e-05 The log loss is: 1.2970145428445017
         For values of alpha = 0.0001 The log loss is: 1.1311437101849808
         For values of alpha = 0.001 The log loss is: 1.1307292395835367
         For values of alpha = 0.01 The log loss is: 1.236725105270069
         For values of alpha = 0.1 The log loss is: 1.4148159910785088
         For values of alpha = 1 The log loss is: 1.6163539150773092
```



```
For values of best alpha = 0.001 The train log loss is: 0.6143981572645738

For values of best alpha = 0.001 The cross validation log loss is: 1.1307292395835367

For values of best alpha = 0.001 The test log loss is: 1.088223763464001
```

Q. Is the Text feature stable across all the data sets (Test, Train, Cross validation)?

Ans. Yes, it seems like!

```
In [0]: def get_intersec_text(df):
             df_text_vec = TfidfVectorizer(min_df=3)
             df_text_fea = df_text_vec.fit_transform(df['TEXT'])
             df_text_features = top_mean_feats(df_text_fea,
                                                df_text_vec.get_feature_names(),
                                                top_n=1000)['feature'].tolist()
             df_text_fea_counts = df_text_fea.sum(axis=0).A1
             df_text_fea_dict = dict(zip(list(df_text_features),df_text_fea_counts))
             len1 = len(set(df_text_features))
             len2 = len(set(train_text_features) & set(df_text_features))
             return len1,len2
In [61]: len1,len2 = get_intersec_text(x_test)
         print(np.round((len2/len1)*100, 3), "% of word of test data appeared in train data")
         len1,len2 = get_intersec_text(x_cv)
         print(np.round((len2/len1)*100, 3), "% of word of Cross Validation appeared in train data")
         61.8 % of word of test data appeared in train data
         59.4 % of word of Cross Validation appeared in train data
```

4. Machine Learning Models

```
In [0]: #Data preparation for ML models.

#Misc. functionns for ML models

def predict_and_plot_confusion_matrix(train_x, train_y,test_x, test_y, clf):
        clf.fit(train_x, train_y)
        sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
        sig_clf.fit(train_x, train_y)
        pred_y = sig_clf.predict(test_x)

# for calculating log_loss we will provide the array of probabilities belongs to each class
        print("Log loss:",log_loss(test_y, sig_clf.predict_proba(test_x)))
# calculating the number of data points that are misclassified
        print("Number of mis-classified points:", np.count_nonzero((pred_y- test_y))/test_y.shape[0])
        plot_confusion_matrix(test_y, pred_y)
```

```
In [0]: def report_log_loss(train_x, train_y, test_x, test_y, clf):
    clf.fit(train_x, train_y)
    sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
    sig_clf.fit(train_x, train_y)
    sig_clf_probs = sig_clf.predict_proba(test_x)
    return log_loss(test_y, sig_clf_probs, eps=1e-15)
```

```
In [0]: | # this function will be used just for naive bayes
        # for the given indices, we will print the name of the features
        # and we will check whether the feature present in the test point text or not
        def get_impfeature_names(indices, text, gene, var, no_features):
            gene_count_vec = TfidfVectorizer()
            var_count_vec = TfidfVectorizer()
            text_count_vec = TfidfVectorizer(min_df=3)
            gene_vec = gene_count_vec.fit(x_train['Gene'])
            var_vec = var_count_vec.fit(x_train['Variation'])
            text_vec = text_count_vec.fit(x_train['TEXT'])
            fea1_len = len(gene_vec.get_feature_names())
            fea2_len = len(var_count_vec.get_feature_names())
            word_present = 0
            for i,v in enumerate(indices):
                 if (v < fea1_len):</pre>
                     word = gene_vec.get_feature_names()[v]
                     yes_no = True if word == gene else False
                     if yes_no:
                         word_present += 1
                         print(i, "Gene feature [{}] present in test data point [{}]".format(word,yes_no))
                 elif (v < fea1_len+fea2_len):</pre>
                     word = var_vec.get_feature_names()[v-(fea1_len)]
                     yes_no = True if word == var else False
                     if yes_no:
                         word present += 1
                         print(i, "variation feature [{}] present in test data point [{}]".format(word,yes_no))
                 else:
                     word = text_vec.get_feature_names()[v-(fea1_len+fea2_len)]
                     yes_no = True if word in text.split() else False
                     if yes_no:
                         word_present += 1
                         print(i, "Text feature [{}] present in test data point [{}]".format(word,yes_no))
            print("Out of the top ",no_features," features ", word_present, "are present in query point")
```

Stacking the three types of features

```
In [0]: | # merging gene, variance and text features
        # building train, test and cross validation data sets
        \# a = [[1, 2],
              [3, 4]]
        #b = [[4, 5],
             [6, 7]]
        # hstack(a, b) = [[1, 2, 4, 5],
                        [ 3, 4, 6, 7]]
        train_gene_var_onehotCoding = hstack((train_gene_feature_onehotCoding,train_variation_feature_onehotCoding))
        test_gene_var_onehotCoding = hstack((test_gene_feature_onehotCoding,test_variation_feature_onehotCoding))
        cv_gene_var_onehotCoding = hstack((cv_gene_feature_onehotCoding,cv_variation_feature_onehotCoding))
        train_x_onehotCoding = hstack((train_gene_var_onehotCoding, train_text_feature_onehotCoding)).tocsr()
        train_y = np.array(list(y_train['Class']))
        test_x_onehotCoding = hstack((test_gene_var_onehotCoding, test_text_feature_onehotCoding)).tocsr()
        test_y = np.array(list(y_test['Class']))
         cv_x_onehotCoding = hstack((cv_gene_var_onehotCoding, cv_text_feature_onehotCoding)).tocsr()
        cv_y = np.array(list(y_cv['Class']))
        train_gene_var_responseCoding = np.hstack((train_gene_feature_responseCoding,train_variation_feature_responseCoding))
        test_gene_var_responseCoding = np.hstack((test_gene_feature_responseCoding,test_variation_feature_responseCoding))
        cv_gene_var_responseCoding = np.hstack((cv_gene_feature_responseCoding,cv_variation_feature_responseCoding))
        train x responseCoding = np.hstack((train gene var responseCoding, train text feature responseCoding))
        test_x_responseCoding = np.hstack((test_gene_var_responseCoding, test_text_feature_responseCoding))
        cv_x_responseCoding = np.hstack((cv_gene_var_responseCoding, cv_text_feature_responseCoding))
```

```
In [66]: print("One hot encoding features :")
    print("(number of data points * number of features) in train data = ", train_x_onehotCoding.shape)
    print("(number of data points * number of features) in test data = ", test_x_onehotCoding.shape)
    print("(number of data points * number of features) in cross validation data =", cv_x_onehotCoding.shape)

One hot encoding features :
    (number of data points * number of features) in train data = (2124, 55087)
    (number of data points * number of features) in test data = (665, 55087)
    (number of data points * number of features) in cross validation data = (532, 55087)

In [67]: print(" Response encoding features :")
    print("(number of data points * number of features) in train data = ", train_x_responseCoding.shape)
    print("(number of data points * number of features) in cross validation data = ", cv_x_responseCoding.shape)

    Response encoding features :
    (number of data points * number of features) in train data = (2124, 27)
    (number of data points * number of features) in test data = (665, 27)
    (number of data points * number of features) in test data = (665, 27)
    (number of data points * number of features) in cross validation data = (532, 27)
```

4.1. Base Line Model

4.1.1. Naive Bayes

4.1.1.1. Hyper parameter tuning

```
In [68]: | # find more about Multinomial Naive base function here http://scikit-learn.org/stable/modules/generated/sklearn.naive_
         bayes.MultinomialNB.html
         # -----
         # default paramters
         # sklearn.naive_bayes.MultinomialNB(alpha=1.0, fit_prior=True, class_prior=None)
         # some of methods of MultinomialNB()
         # fit(X, y[, sample_weight])
                                      Fit Naive Bayes classifier according to X, y
         # predict(X) Perform classification on an array of test vectors X.
         # predict_log_proba(X) Return log-probability estimates for the test vector X.
         # -----
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/naive-bayes-algorithm-1/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # -----
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         # -----
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/naive-bayes-algorithm-1/
         alpha = [0.00001, 0.0001, 0.001, 0.1, 1, 10, 100,1000]
         cv_log_error_array = []
         for i in alpha:
             print("for alpha =", i)
             clf = MultinomialNB(alpha=i)
             clf.fit(train_x_onehotCoding, train_y)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_x_onehotCoding, train_y)
             sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
             cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
             # to avoid rounding error while multiplying probabilites we use log-probability estimates
             print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         fig, ax = plt.subplots()
         ax.plot(np.log10(alpha), cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],str(txt)), (np.log10(alpha[i]),cv_log_error_array[i]))
         plt.grid()
         plt.xticks(np.log10(alpha))
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = MultinomialNB(alpha=alpha[best_alpha])
         clf.fit(train_x_onehotCoding, train_y)
         sig clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_onehotCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The train log loss is:",log_loss(y_train, predict_y, labels=c
         lf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
         print('For values of best alpha =
                                           ', alpha[best_alpha], "The cross validation log loss is:",log_loss(y_cv, predict_y,
         labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_x_onehotCoding)
         print('For values of best alpha = ', alpha[best_alpha], "The test log loss is:",log_loss(y_test, predict_y, labels=clf
         .classes_, eps=1e-15))
```

for alpha = 1e-05

Log Loss: 1.2582343535110632

for alpha = 0.0001

Log Loss : 1.2531643211398023

for alpha = 0.001

Log Loss: 1.240590195050788

for alpha = 0.1

Log Loss : 1.244706999926573 for alpha = 1

Log Loss: 1.2805027591793159 for alpha = 10

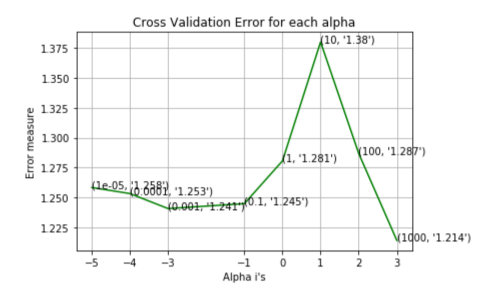
Log Loss : 1.3800673906727279

for alpha = 100

Log Loss: 1.2867822733133558

for alpha = 1000

Log Loss : 1.2139322601905145



For values of best alpha = 1000 The train log loss is: 0.9423970058439604 For values of best alpha = 1000 The cross validation log loss is: 1.2139322601905145

For values of best alpha = 1000 The test log loss is: 1.192912559131826

4.1.1.2. Testing the model with best hyper paramters

```
In [69]: | # find more about Multinomial Naive base function here http://scikit-learn.org/stable/modules/generated/sklearn.naive_
         bayes.MultinomialNB.html
         # -----
         # default paramters
         # sklearn.naive_bayes.MultinomialNB(alpha=1.0, fit_prior=True, class_prior=None)
         # some of methods of MultinomialNB()
         # fit(X, y[, sample_weight])
                                      Fit Naive Bayes classifier according to X, y
         \# predict(X) Perform classification on an array of test vectors X.
         # predict_log_proba(X) Return log-probability estimates for the test vector X.
         # -----
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/naive-bayes-algorithm-1/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # ------
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         clf = MultinomialNB(alpha=alpha[best_alpha])
         clf.fit(train_x_onehotCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig clf.fit(train_x_onehotCoding, train_y)
         sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
         # to avoid rounding error while multiplying probabilites we use log-probability estimates
         print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         print("Number of missclassified point :", np.count_nonzero((sig_clf.predict(cv_x_onehotCoding)- cv_y))/cv_y.shape[0])
         plot_confusion_matrix(cv_y, sig_clf.predict(cv_x_onehotCoding.toarray()))
```

Number of missclassified point : 0.43796992481203006

----- Confusion matrix -----43.000 0.000 0.000 20.000 2.000 1.000 25.000 0.000 0.000 - 125 1.000 13.000 0.000 1.000 0.000 0.000 57.000 0.000 0.000 0.000 0.000 4.000 2.000 0.000 0.000 8.000 0.000 0.000 - 100 65.000 15.000 0.000 0.000 3.000 0.000 27.000 0.000 0.000 Original Class - 75 5.000 2.000 0.000 7.000 7.000 6.000 12.000 0.000 0.000 5.000 0.000 0.000 5.000 2.000 24.000 8.000 0.000 0.000 - 50 1.000 6.000 0.000 1.000 0.000 0.000 145.000 0.000 0.000 - 25 0.000 1.000 0.000 0.000 0.000 0.000 2.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 4.000 0.000 2.000 -0 Predicted Class ----- Precision matrix (Columm Sum=1) ------1.0 0.000 0.194 0.125 0.032 0.087 0.000 0.010 0.000 0.014 0.000 0.198 0.000 - 0.8 0.000 0.000 0.039 0.125 0.000 0.028 0.000 0.214 0.000 0.188 0.000 0.094 0.000 - 0.6 Original Class 0.071 0.091 0.068 0.438 0.194 0.042 0.000 - 0.4 0.071 0.000 0.049 0.125 0.028 0.000 0.014 0.273 0.010 0.000 0.000 0.000 - 0.2 0.000 0.045 0.000 0.000 0.000 0.007 0.000 0.000 0.000 0.000 0.000 0.000 0.014 1.000 - 0.0 Predicted Class ----- Recall matrix (Row sum=1) ------0.000 0.000 0.220 0.022 0.011 0.275 0.000 0.000 - 0.8 0.014 0.181 0.000 0.014 0.000 0.000 0.000 0.000 0.286 0.000 0.000 0.000 0.143 0.000 0.000 0.000 - 0.6 0.136 0.000 0.000 0.027 0.000 0.245 0.000 0.000 0.179 0.128 0.051 0.000 0.179 0.154 0.308 0.000 0.000 0.4 0.114 0.000 0.000 0.114 0.045 0.182 0.000 0.000 0.007 0.039 0.000 0.007 0.000 0.000 0.948 0.000 0.000 - 0.2 0.333 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.333 0.000 0.000 0.000

Predicted Class

'n

- 0.0

í

2

Predicted Class : 4 Predicted Class Probabilities: [[3.167e-01 9.800e-03 2.000e-03 5.684e-01 1.110e-02 1.110e-02 7.990e-02 9.000e-04 1.000e-04]] Actual Class: 4 _____ 10 Text feature [proteins] present in test data point [True] 11 Text feature [activity] present in test data point [True] 12 Text feature [protein] present in test data point [True] 14 Text feature [experiments] present in test data point [True] 15 Text feature [acid] present in test data point [True] 16 Text feature [mammalian] present in test data point [True] 17 Text feature [whether] present in test data point [True] 18 Text feature [shown] present in test data point [True] 19 Text feature [function] present in test data point [True] 20 Text feature [loss] present in test data point [True] 21 Text feature [indicated] present in test data point [True] 22 Text feature [results] present in test data point [True] 23 Text feature [important] present in test data point [True] 24 Text feature [whereas] present in test data point [True] 25 Text feature [amino] present in test data point [True] 26 Text feature [determined] present in test data point [True] 27 Text feature [described] present in test data point [True] 30 Text feature [two] present in test data point [True] 32 Text feature [ability] present in test data point [True] 33 Text feature [expressed] present in test data point [True] 34 Text feature [functions] present in test data point [True] 35 Text feature [also] present in test data point [True] 36 Text feature [missense] present in test data point [True] 37 Text feature [type] present in test data point [True] 39 Text feature [partially] present in test data point [True] 40 Text feature [retained] present in test data point [True] 41 Text feature [reduced] present in test data point [True] 44 Text feature [standard] present in test data point [True] 45 Text feature [thus] present in test data point [True] 46 Text feature [levels] present in test data point [True] 49 Text feature [lower] present in test data point [True] 50 Text feature [containing] present in test data point [True] 52 Text feature [vitro] present in test data point [True] 53 Text feature [determine] present in test data point [True] 54 Text feature [mutations] present in test data point [True] 55 Text feature [see] present in test data point [True] 56 Text feature [bind] present in test data point [True] 57 Text feature [30] present in test data point [True] 58 Text feature [related] present in test data point [True] 60 Text feature [wild] present in test data point [True] 61 Text feature [purified] present in test data point [True] 62 Text feature [indicate] present in test data point [True] 63 Text feature [may] present in test data point [True] 64 Text feature [analyzed] present in test data point [True] 65 Text feature [although] present in test data point [True] 68 Text feature [either] present in test data point [True] 69 Text feature [system] present in test data point [True] 70 Text feature [germline] present in test data point [True] 71 Text feature [suppressor] present in test data point [True] 72 Text feature [suggest] present in test data point [True] 73 Text feature [discussion] present in test data point [True] 74 Text feature [using] present in test data point [True] 75 Text feature [three] present in test data point [True] 76 Text feature [percentage] present in test data point [True] 77 Text feature [expression] present in test data point [True] 78 Text feature [result] present in test data point [True] 79 Text feature [made] present in test data point [True] 80 Text feature [mm] present in test data point [True] 82 Text feature [performed] present in test data point [True] 85 Text feature [previously] present in test data point [True] 86 Text feature [contribute] present in test data point [True] 87 Text feature [transfected] present in test data point [True] 89 Text feature [tris] present in test data point [True] 90 Text feature [sds] present in test data point [True] 91 Text feature [effects] present in test data point [True] 93 Text feature [marked] present in test data point [True] 94 Text feature [buffer] present in test data point [True] 96 Text feature [functional] present in test data point [True] 97 Text feature [yielded] present in test data point [True] 101 Text feature [indicates] present in test data point [True] 102 Text feature [possible] present in test data point [True] 103 Text feature [introduction] present in test data point [True] 107 Text feature [cells] present in test data point [True] 109 Text feature [transfection] present in test data point [True] 110 Text feature [associated] present in test data point [True] 111 Text feature [analysis] present in test data point [True] 113 Text feature [terminal] present in test data point [True] 114 Text feature [addition] present in test data point [True] 115 Text feature [high] present in test data point [True] 116 Text feature [one] present in test data point [True] 117 Text feature [yeast] present in test data point [True] 118 Text feature [site] present in test data point [True]

119 Text feature [critical] present in test data point [True]

```
120 Text feature [several] present in test data point [True]
121 Text feature [however] present in test data point [True]
122 Text feature [low] present in test data point [True]
123 Text feature [reported] present in test data point [True]
125 Text feature [tested] present in test data point [True]
126 Text feature [referred] present in test data point [True]
128 Text feature [within] present in test data point [True]
129 Text feature [therefore] present in test data point [True]
130 Text feature [vector] present in test data point [True]
131 Text feature [possibility] present in test data point [True]
134 Text feature [substitutions] present in test data point [True]
135 Text feature [suggesting] present in test data point [True]
136 Text feature [resulting] present in test data point [True]
137 Text feature [lack] present in test data point [True]
138 Text feature [found] present in test data point [True]
139 Text feature [generated] present in test data point [True]
141 Text feature [correspond] present in test data point [True]
142 Text feature [similar] present in test data point [True]
144 Text feature [fact] present in test data point [True]
145 Text feature [required] present in test data point [True]
147 Text feature [role] present in test data point [True]
150 Text feature [suggested] present in test data point [True]
151 Text feature [nonsense] present in test data point [True]
153 Text feature [assay] present in test data point [True]
154 Text feature [control] present in test data point [True]
155 Text feature [general] present in test data point [True]
156 Text feature [full] present in test data point [True]
157 Text feature [could] present in test data point [True]
158 Text feature [note] present in test data point [True]
159 Text feature [previous] present in test data point [True]
161 Text feature [derived] present in test data point [True]
162 Text feature [affected] present in test data point [True]
163 Text feature [bars] present in test data point [True]
165 Text feature [heterologous] present in test data point [True]
167 Text feature [inactivated] present in test data point [True]
168 Text feature [binding] present in test data point [True]
170 Text feature [indicating] present in test data point [True]
172 Text feature [phenotype] present in test data point [True]
173 Text feature [show] present in test data point [True]
174 Text feature [acids] present in test data point [True]
175 Text feature [diminution] present in test data point [True]
180 Text feature [essential] present in test data point [True]
181 Text feature [human] present in test data point [True]
182 Text feature [involved] present in test data point [True]
186 Text feature [amount] present in test data point [True]
187 Text feature [rather] present in test data point [True]
191 Text feature [loops] present in test data point [True]
193 Text feature [regions] present in test data point [True]
194 Text feature [together] present in test data point [True]
196 Text feature [co] present in test data point [True]
197 Text feature [exist] present in test data point [True]
198 Text feature [four] present in test data point [True]
199 Text feature [lacking] present in test data point [True]
200 Text feature [10] present in test data point [True]
202 Text feature [relative] present in test data point [True]
203 Text feature [covering] present in test data point [True]
204 Text feature [presented] present in test data point [True]
205 Text feature [properties] present in test data point [True]
206 Text feature [mutant] present in test data point [True]
207 Text feature [directly] present in test data point [True]
209 Text feature [min] present in test data point [True]
210 Text feature [according] present in test data point [True]
212 Text feature [dependent] present in test data point [True]
213 Text feature [included] present in test data point [True]
214 Text feature [washed] present in test data point [True]
216 Text feature [localization] present in test data point [True]
217 Text feature [importance] present in test data point [True]
219 Text feature [cycle] present in test data point [True]
220 Text feature [cellular] present in test data point [True]
223 Text feature [half] present in test data point [True]
225 Text feature [15] present in test data point [True]
227 Text feature [might] present in test data point [True]
228 Text feature [50] present in test data point [True]
229 Text feature [recent] present in test data point [True]
230 Text feature [representative] present in test data point [True]
231 Text feature [distinct] present in test data point [True]
234 Text feature [including] present in test data point [True]
235 Text feature [lysates] present in test data point [True]
236 Text feature [exception] present in test data point [True]
237 Text feature [predicted] present in test data point [True]
239 Text feature [40] present in test data point [True]
240 Text feature [anti] present in test data point [True]
241 Text feature [fig] present in test data point [True]
243 Text feature [ml] present in test data point [True]
245 Text feature [view] present in test data point [True]
246 Text feature [data] present in test data point [True]
247 Text feature [specific] present in test data point [True]
249 Text feature [probably] present in test data point [True]
```

```
251 Text feature [affect] present in test data point [True]
253 Text feature [respectively] present in test data point [True]
256 Text feature [key] present in test data point [True]
257 Text feature [mammals] present in test data point [True]
258 Text feature [mutation] present in test data point [True]
259 Text feature [multiple] present in test data point [True]
260 Text feature [furthermore] present in test data point [True]
261 Text feature [test] present in test data point [True]
263 Text feature [compared] present in test data point [True]
266 Text feature [extensive] present in test data point [True]
267 Text feature [slide] present in test data point [True]
271 Text feature [variety] present in test data point [True]
274 Text feature [changes] present in test data point [True]
276 Text feature [revealed] present in test data point [True]
281 Text feature [green] present in test data point [True]
282 Text feature [different] present in test data point [True]
283 Text feature [mutagenesis] present in test data point [True]
284 Text feature [sigma] present in test data point [True]
286 Text feature [inactivate] present in test data point [True]
288 Text feature [used] present in test data point [True]
290 Text feature [comprehensive] present in test data point [True]
292 Text feature [mutants] present in test data point [True]
301 Text feature [figure] present in test data point [True]
302 Text feature [42] present in test data point [True]
303 Text feature [cell] present in test data point [True]
305 Text feature [frequently] present in test data point [True]
306 Text feature [remarkably] present in test data point [True]
307 Text feature [well] present in test data point [True]
311 Text feature [findings] present in test data point [True]
314 Text feature [effect] present in test data point [True]
320 Text feature [necessary] present in test data point [True]
321 Text feature [additional] present in test data point [True]
324 Text feature [provide] present in test data point [True]
325 Text feature [include] present in test data point [True]
328 Text feature [dna] present in test data point [True]
416 Text feature [majority] present in test data point [True]
420 Text feature [mediated] present in test data point [True]
422 Text feature [caused] present in test data point [True]
424 Text feature [level] present in test data point [True]
426 Text feature [tumor] present in test data point [True]
427 Text feature [reading] present in test data point [True]
428 Text feature [italics] present in test data point [True]
431 Text feature [substantially] present in test data point [True]
436 Text feature [due] present in test data point [True]
441 Text feature [25] present in test data point [True]
446 Text feature [gene] present in test data point [True]
448 Text feature [28] present in test data point [True]
450 Text feature [present] present in test data point [True]
451 Text feature [escherichia] present in test data point [True]
452 Text feature [centrifugation] present in test data point [True]
455 Text feature [would] present in test data point [True]
456 Text feature [part] present in test data point [True]
458 Text feature [table] present in test data point [True]
462 Text feature [100] present in test data point [True]
463 Text feature [remains] present in test data point [True]
464 Text feature [cerevisiae] present in test data point [True]
465 Text feature [red] present in test data point [True]
466 Text feature [plasmids] present in test data point [True]
468 Text feature [consistent] present in test data point [True]
469 Text feature [even] present in test data point [True]
470 Text feature [reference] present in test data point [True]
471 Text feature [obtained] present in test data point [True]
473 Text feature [presence] present in test data point [True]
475 Text feature [make] present in test data point [True]
478 Text feature [contrast] present in test data point [True]
480 Text feature [since] present in test data point [True]
481 Text feature [proposed] present in test data point [True]
482 Text feature [many] present in test data point [True]
483 Text feature [observation] present in test data point [True]
484 Text feature [families] present in test data point [True]
486 Text feature [display] present in test data point [True]
487 Text feature [quantified] present in test data point [True]
489 Text feature [arrest] present in test data point [True]
491 Text feature [based] present in test data point [True]
492 Text feature [lysis] present in test data point [True]
494 Text feature [product] present in test data point [True]
495 Text feature [finally] present in test data point [True]
500 Text feature [association] present in test data point [True]
501 Text feature [edta] present in test data point [True]
502 Text feature [detected] present in test data point [True]
505 Text feature [27] present in test data point [True]
506 Text feature [largedownload] present in test data point [True]
507 Text feature [domain] present in test data point [True]
508 Text feature [least] present in test data point [True]
510 Text feature [studies] present in test data point [True]
512 Text feature [negative] present in test data point [True]
513 Text feature [equal] present in test data point [True]
514 Text feature [unclear] present in test data point [True]
```

```
515 Text feature [distribution] present in test data point [True]
516 Text feature [added] present in test data point [True]
517 Text feature [hypothesis] present in test data point [True]
518 Text feature [observed] present in test data point [True]
519 Text feature [methods] present in test data point [True]
520 Text feature [directed] present in test data point [True]
521 Text feature [another] present in test data point [True]
527 Text feature [study] present in test data point [True]
528 Text feature [partial] present in test data point [True]
529 Text feature [appears] present in test data point [True]
530 Text feature [normal] present in test data point [True]
533 Text feature [regulation] present in test data point [True]
537 Text feature [region] present in test data point [True]
540 Text feature [equally] present in test data point [True]
541 Text feature [limited] present in test data point [True]
542 Text feature [without] present in test data point [True]
543 Text feature [showed] present in test data point [True]
544 Text feature [five] present in test data point [True]
545 Text feature [relatively] present in test data point [True]
547 Text feature [2000] present in test data point [True]
548 Text feature [subcellular] present in test data point [True]
549 Text feature [separated] present in test data point [True]
550 Text feature [color] present in test data point [True]
552 Text feature [centrifuged] present in test data point [True]
553 Text feature [spectrum] present in test data point [True]
554 Text feature [decreased] present in test data point [True]
558 Text feature [assessment] present in test data point [True]
560 Text feature [eight] present in test data point [True]
561 Text feature [significant] present in test data point [True]
564 Text feature [plasmid] present in test data point [True]
566 Text feature [antibodies] present in test data point [True]
567 Text feature [precise] present in test data point [True]
568 Text feature [represent] present in test data point [True]
571 Text feature [growth] present in test data point [True]
572 Text feature [allows] present in test data point [True]
573 Text feature [26] present in test data point [True]
574 Text feature [syndromes] present in test data point [True]
576 Text feature [followed] present in test data point [True]
578 Text feature [http] present in test data point [True]
580 Text feature [taken] present in test data point [True]
581 Text feature [et] present in test data point [True]
583 Text feature [nevertheless] present in test data point [True]
585 Text feature [single] present in test data point [True]
587 Text feature [released] present in test data point [True]
590 Text feature [1a] present in test data point [True]
591 Text feature [highly] present in test data point [True]
592 Text feature [almost] present in test data point [True]
596 Text feature [assessed] present in test data point [True]
599 Text feature [divergent] present in test data point [True]
601 Text feature [stability] present in test data point [True]
604 Text feature [increased] present in test data point [True]
610 Text feature [confirmed] present in test data point [True]
611 Text feature [considered] present in test data point [True]
612 Text feature [saccharomyces] present in test data point [True]
613 Text feature [31] present in test data point [True]
614 Text feature [al] present in test data point [True]
615 Text feature [autosomal] present in test data point [True]
616 Text feature [induced] present in test data point [True]
617 Text feature [total] present in test data point [True]
618 Text feature [reflect] present in test data point [True]
622 Text feature [verified] present in test data point [True]
624 Text feature [endogenous] present in test data point [True]
625 Text feature [frameshift] present in test data point [True]
627 Text feature [blue] present in test data point [True]
628 Text feature [materials] present in test data point [True]
632 Text feature [nuclear] present in test data point [True]
633 Text feature [expected] present in test data point [True]
637 Text feature [displayed] present in test data point [True]
638 Text feature [briefly] present in test data point [True]
641 Text feature [larger] present in test data point [True]
642 Text feature [certain] present in test data point [True]
646 Text feature [cloned] present in test data point [True]
647 Text feature [us] present in test data point [True]
649 Text feature [identified] present in test data point [True]
651 Text feature [37] present in test data point [True]
652 Text feature [approximately] present in test data point [True]
653 Text feature [sufficient] present in test data point [True]
654 Text feature [characterized] present in test data point [True]
656 Text feature [complete] present in test data point [True]
657 Text feature [length] present in test data point [True]
658 Text feature [investigated] present in test data point [True]
659 Text feature [independent] present in test data point [True]
660 Text feature [sequence] present in test data point [True]
661 Text feature [higher] present in test data point [True]
662 Text feature [stained] present in test data point [True]
664 Text feature [lost] present in test data point [True]
667 Text feature [figs] present in test data point [True]
669 Text feature [cancer] present in test data point [True]
```

```
671 Text feature [refers] present in test data point [True]
673 Text feature [deletion] present in test data point [True]
675 Text feature [interesting] present in test data point [True]
677 Text feature [29] present in test data point [True]
681 Text feature [indeed] present in test data point [True]
684 Text feature [supporting] present in test data point [True]
686 Text feature [genetic] present in test data point [True]
687 Text feature [active] present in test data point [True]
688 Text feature [complex] present in test data point [True]
691 Text feature [cannot] present in test data point [True]
695 Text feature [significantly] present in test data point [True]
697 Text feature [less] present in test data point [True]
701 Text feature [conclusions] present in test data point [True]
702 Text feature [resultant] present in test data point [True]
704 Text feature [dual] present in test data point [True]
705 Text feature [moreover] present in test data point [True]
706 Text feature [conditions] present in test data point [True]
708 Text feature [notion] present in test data point [True]
710 Text feature [among] present in test data point [True]
715 Text feature [38] present in test data point [True]
716 Text feature [supported] present in test data point [True]
717 Text feature [predisposition] present in test data point [True]
718 Text feature [identical] present in test data point [True]
719 Text feature [likely] present in test data point [True]
720 Text feature [1b] present in test data point [True]
722 Text feature [yet] present in test data point [True]
723 Text feature [unable] present in test data point [True]
725 Text feature [disruption] present in test data point [True]
726 Text feature [must] present in test data point [True]
729 Text feature [seven] present in test data point [True]
731 Text feature [seems] present in test data point [True]
732 Text feature [dominant] present in test data point [True]
734 Text feature [33] present in test data point [True]
737 Text feature [affecting] present in test data point [True]
741 Text feature [ca] present in test data point [True]
742 Text feature [play] present in test data point [True]
743 Text feature [sites] present in test data point [True]
744 Text feature [give] present in test data point [True]
745 Text feature [incubation] present in test data point [True]
748 Text feature [develop] present in test data point [True]
749 Text feature [48] present in test data point [True]
750 Text feature [central] present in test data point [True]
754 Text feature [transformants] present in test data point [True]
755 Text feature [suggests] present in test data point [True]
756 Text feature [bold] present in test data point [True]
757 Text feature [first] present in test data point [True]
758 Text feature [coli] present in test data point [True]
759 Text feature [3b] present in test data point [True]
760 Text feature [20] present in test data point [True]
765 Text feature [common] present in test data point [True]
767 Text feature [located] present in test data point [True]
770 Text feature [manipulation] present in test data point [True]
771 Text feature [explained] present in test data point [True]
773 Text feature [recently] present in test data point [True]
781 Text feature [3a] present in test data point [True]
782 Text feature [lead] present in test data point [True]
786 Text feature [shows] present in test data point [True]
789 Text feature [motif] present in test data point [True]
790 Text feature [large] present in test data point [True]
791 Text feature [modified] present in test data point [True]
792 Text feature [functionally] present in test data point [True]
793 Text feature [following] present in test data point [True]
798 Text feature [grown] present in test data point [True]
799 Text feature [binds] present in test data point [True]
800 Text feature [systems] present in test data point [True]
802 Text feature [via] present in test data point [True]
803 Text feature [pellets] present in test data point [True]
804 Text feature [null] present in test data point [True]
806 Text feature [300] present in test data point [True]
807 Text feature [substitution] present in test data point [True]
810 Text feature [often] present in test data point [True]
811 Text feature [apparent] present in test data point [True]
813 Text feature [disease] present in test data point [True]
816 Text feature [virtue] present in test data point [True]
819 Text feature [yeasts] present in test data point [True]
825 Text feature [target] present in test data point [True]
827 Text feature [support] present in test data point [True]
828 Text feature [little] present in test data point [True]
831 Text feature [nine] present in test data point [True]
832 Text feature [failed] present in test data point [True]
835 Text feature [consequence] present in test data point [True]
839 Text feature [12] present in test data point [True]
846 Text feature [carried] present in test data point [True]
847 Text feature [construction] present in test data point [True]
850 Text feature [despite] present in test data point [True]
853 Text feature [eliminated] present in test data point [True]
858 Text feature [provided] present in test data point [True]
859 Text feature [occur] present in test data point [True]
```

```
868 Text feature [consequences] present in test data point [True]
870 Text feature [24] present in test data point [True]
872 Text feature [groups] present in test data point [True]
874 Text feature [exerts] present in test data point [True]
877 Text feature [selected] present in test data point [True]
880 Text feature [90] present in test data point [True]
884 Text feature [cdna] present in test data point [True]
886 Text feature [overnight] present in test data point [True]
887 Text feature [close] present in test data point [True]
889 Text feature [understanding] present in test data point [True]
891 Text feature [occurs] present in test data point [True]
894 Text feature [corresponding] present in test data point [True]
895 Text feature [distributed] present in test data point [True]
899 Text feature [tumorigenesis] present in test data point [True]
903 Text feature [express] present in test data point [True]
904 Text feature [1c] present in test data point [True]
907 Text feature [designed] present in test data point [True]
908 Text feature [instability] present in test data point [True]
911 Text feature [addressed] present in test data point [True]
913 Text feature [difficult] present in test data point [True]
920 Text feature [mouse] present in test data point [True]
959 Text feature [research] present in test data point [True]
962 Text feature [heterozygous] present in test data point [True]
963 Text feature [next] present in test data point [True]
964 Text feature [version] present in test data point [True]
965 Text feature [allowed] present in test data point [True]
967 Text feature [recognized] present in test data point [True]
969 Text feature [subtle] present in test data point [True]
971 Text feature [culture] present in test data point [True]
973 Text feature [200] present in test data point [True]
974 Text feature [six] present in test data point [True]
975 Text feature [assays] present in test data point [True]
976 Text feature [members] present in test data point [True]
977 Text feature [promega] present in test data point [True]
978 Text feature [rabbit] present in test data point [True]
980 Text feature [model] present in test data point [True]
981 Text feature [provides] present in test data point [True]
982 Text feature [form] present in test data point [True]
983 Text feature [transferred] present in test data point [True]
984 Text feature [plates] present in test data point [True]
985 Text feature [80] present in test data point [True]
986 Text feature [resulted] present in test data point [True]
989 Text feature [lipofectamine] present in test data point [True]
991 Text feature [frame] present in test data point [True]
Out of the top 1000 features 479 are present in query point
```

4.1.1.4. Feature Importance, Incorrectly classified point

```
Predicted Class : 7
Predicted Class Probabilities: [[3.750e-02 9.570e-02 4.400e-03 7.190e-02 1.790e-02 1.500e-02 7.557e-01
  1.600e-03 3.000e-04]]
Actual Class: 1
______
16 Text feature [cells] present in test data point [True]
19 Text feature [kinase] present in test data point [True]
20 Text feature [activated] present in test data point [True]
21 Text feature [cell] present in test data point [True]
22 Text feature [factor] present in test data point [True]
23 Text feature [activation] present in test data point [True]
24 Text feature [phosphorylation] present in test data point [True]
25 Text feature [downstream] present in test data point [True]
26 Text feature [contrast] present in test data point [True]
27 Text feature [presence] present in test data point [True]
28 Text feature [shown] present in test data point [True]
29 Text feature [expressing] present in test data point [True]
30 Text feature [inhibitor] present in test data point [True]
31 Text feature [growth] present in test data point [True]
32 Text feature [also] present in test data point [True]
33 Text feature [however] present in test data point [True]
34 Text feature [signaling] present in test data point [True]
36 Text feature [addition] present in test data point [True]
37 Text feature [suggest] present in test data point [True]
38 Text feature [independent] present in test data point [True]
39 Text feature [treated] present in test data point [True]
41 Text feature [10] present in test data point [True]
42 Text feature [compared] present in test data point [True]
43 Text feature [previously] present in test data point [True]
44 Text feature [1a] present in test data point [True]
45 Text feature [mechanism] present in test data point [True]
46 Text feature [constitutively] present in test data point [True]
47 Text feature [potential] present in test data point [True]
48 Text feature [similar] present in test data point [True]
49 Text feature [increased] present in test data point [True]
50 Text feature [figure] present in test data point [True]
51 Text feature [tyrosine] present in test data point [True]
52 Text feature [3b] present in test data point [True]
54 Text feature [interestingly] present in test data point [True]
56 Text feature [showed] present in test data point [True]
57 Text feature [followed] present in test data point [True]
58 Text feature [mutations] present in test data point [True]
59 Text feature [treatment] present in test data point [True]
63 Text feature [mutant] present in test data point [True]
64 Text feature [well] present in test data point [True]
66 Text feature [using] present in test data point [True]
67 Text feature [inhibition] present in test data point [True]
68 Text feature [activating] present in test data point [True]
69 Text feature [3a] present in test data point [True]
70 Text feature [consistent] present in test data point [True]
71 Text feature [enhanced] present in test data point [True]
72 Text feature [may] present in test data point [True]
73 Text feature [described] present in test data point [True]
74 Text feature [fig] present in test data point [True]
75 Text feature [antibodies] present in test data point [True]
76 Text feature [confirmed] present in test data point [True]
77 Text feature [reported] present in test data point [True]
78 Text feature [including] present in test data point [True]
79 Text feature [total] present in test data point [True]
80 Text feature [absence] present in test data point [True]
81 Text feature [furthermore] present in test data point [True]
83 Text feature [proliferation] present in test data point [True]
84 Text feature [sensitive] present in test data point [True]
85 Text feature [obtained] present in test data point [True]
86 Text feature [observed] present in test data point [True]
87 Text feature [survival] present in test data point [True]
88 Text feature [mutation] present in test data point [True]
89 Text feature [inhibitors] present in test data point [True]
90 Text feature [induced] present in test data point [True]
91 Text feature [expression] present in test data point [True]
94 Text feature [inhibited] present in test data point [True]
95 Text feature [culture] present in test data point [True]
96 Text feature [small] present in test data point [True]
97 Text feature [expressed] present in test data point [True]
98 Text feature [respectively] present in test data point [True]
100 Text feature [approximately] present in test data point [True]
102 Text feature [together] present in test data point [True]
104 Text feature [concentrations] present in test data point [True]
105 Text feature [western] present in test data point [True]
106 Text feature [suggesting] present in test data point [True]
107 Text feature [recent] present in test data point [True]
108 Text feature [either] present in test data point [True]
109 Text feature [hours] present in test data point [True]
111 Text feature [increase] present in test data point [True]
112 Text feature [molecular] present in test data point [True]
113 Text feature [cultured] present in test data point [True]
114 Text feature [4a] present in test data point [True]
120 Text feature [discussion] present in test data point [True]
```

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121 Text feature [study] present in test data point [True]
122 Text feature [identified] present in test data point [True]
124 Text feature [domain] present in test data point [True]
125 Text feature [12] present in test data point [True]
126 Text feature [show] present in test data point [True]
128 Text feature [receptor] present in test data point [True]
129 Text feature [studies] present in test data point [True]
131 Text feature [15] present in test data point [True]
132 Text feature [approved] present in test data point [True]
133 Text feature [findings] present in test data point [True]
134 Text feature [common] present in test data point [True]
136 Text feature [activate] present in test data point [True]
137 Text feature [1b] present in test data point [True]
138 Text feature [thus] present in test data point [True]
139 Text feature [whether] present in test data point [True]
140 Text feature [therapeutic] present in test data point [True]
142 Text feature [high] present in test data point [True]
143 Text feature [resulting] present in test data point [True]
147 Text feature [due] present in test data point [True]
148 Text feature [development] present in test data point [True]
151 Text feature [13] present in test data point [True]
152 Text feature [4b] present in test data point [True]
153 Text feature [tumor] present in test data point [True]
154 Text feature [different] present in test data point [True]
155 Text feature [assessed] present in test data point [True]
156 Text feature [3c] present in test data point [True]
157 Text feature [lines] present in test data point [True]
158 Text feature [oncogenic] present in test data point [True]
160 Text feature [revealed] present in test data point [True]
161 Text feature [positive] present in test data point [True]
162 Text feature [although] present in test data point [True]
163 Text feature [results] present in test data point [True]
164 Text feature [within] present in test data point [True]
166 Text feature [detected] present in test data point [True]
167 Text feature [trials] present in test data point [True]
168 Text feature [next] present in test data point [True]
169 Text feature [anti] present in test data point [True]
170 Text feature [additional] present in test data point [True]
171 Text feature [phospho] present in test data point [True]
172 Text feature [2b] present in test data point [True]
173 Text feature [specific] present in test data point [True]
174 Text feature [result] present in test data point [True]
175 Text feature [20] present in test data point [True]
176 Text feature [determine] present in test data point [True]
177 Text feature [conditions] present in test data point [True]
178 Text feature [single] present in test data point [True]
179 Text feature [could] present in test data point [True]
180 Text feature [active] present in test data point [True]
182 Text feature [whereas] present in test data point [True]
183 Text feature [measured] present in test data point [True]
184 Text feature [dependent] present in test data point [True]
185 Text feature [several] present in test data point [True]
186 Text feature [targeted] present in test data point [True]
187 Text feature [lysates] present in test data point [True]
189 Text feature [mediated] present in test data point [True]
190 Text feature [analysis] present in test data point [True]
191 Text feature [1c] present in test data point [True]
192 Text feature [medium] present in test data point [True]
193 Text feature [patients] present in test data point [True]
194 Text feature [genomic] present in test data point [True]
196 Text feature [malignant] present in test data point [True]
198 Text feature [target] present in test data point [True]
200 Text feature [relative] present in test data point [True]
201 Text feature [effects] present in test data point [True]
202 Text feature [new] present in test data point [True]
203 Text feature [three] present in test data point [True]
204 Text feature [transformation] present in test data point [True]
206 Text feature [pathway] present in test data point [True]
208 Text feature [one] present in test data point [True]
209 Text feature [analyzed] present in test data point [True]
210 Text feature [time] present in test data point [True]
211 Text feature [promote] present in test data point [True]
212 Text feature [major] present in test data point [True]
213 Text feature [progression] present in test data point [True]
214 Text feature [human] present in test data point [True]
216 Text feature [indicating] present in test data point [True]
219 Text feature [2a] present in test data point [True]
220 Text feature [clinical] present in test data point [True]
222 Text feature [mechanisms] present in test data point [True]
223 Text feature [27] present in test data point [True]
224 Text feature [18] present in test data point [True]
225 Text feature [point] present in test data point [True]
226 Text feature [mitogen] present in test data point [True]
227 Text feature [free] present in test data point [True]
228 Text feature [might] present in test data point [True]
229 Text feature [supplemented] present in test data point [True]
230 Text feature [significantly] present in test data point [True]
231 Text feature [kinases] present in test data point [True]
```

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232 Text feature [indicated] present in test data point [True]
233 Text feature [lysed] present in test data point [True]
234 Text feature [table] present in test data point [True]
236 Text feature [confirm] present in test data point [True]
237 Text feature [overexpression] present in test data point [True]
238 Text feature [present] present in test data point [True]
239 Text feature [gene] present in test data point [True]
240 Text feature [somatic] present in test data point [True]
241 Text feature [tumors] present in test data point [True]
244 Text feature [sequencing] present in test data point [True]
247 Text feature [non] present in test data point [True]
248 Text feature [therapies] present in test data point [True]
252 Text feature [sigma] present in test data point [True]
254 Text feature [seeded] present in test data point [True]
255 Text feature [test] present in test data point [True]
256 Text feature [drug] present in test data point [True]
258 Text feature [therapy] present in test data point [True]
259 Text feature [first] present in test data point [True]
260 Text feature [17] present in test data point [True]
262 Text feature [vitro] present in test data point [True]
264 Text feature [suggested] present in test data point [True]
265 Text feature [days] present in test data point [True]
268 Text feature [standard] present in test data point [True]
269 Text feature [identification] present in test data point [True]
270 Text feature [lung] present in test data point [True]
271 Text feature [wt] present in test data point [True]
272 Text feature [less] present in test data point [True]
273 Text feature [generated] present in test data point [True]
274 Text feature [per] present in test data point [True]
276 Text feature [cancers] present in test data point [True]
278 Text feature [following] present in test data point [True]
279 Text feature [institutional] present in test data point [True]
280 Text feature [download] present in test data point [True]
281 Text feature [resulted] present in test data point [True]
282 Text feature [novel] present in test data point [True]
283 Text feature [30] present in test data point [True]
284 Text feature [determined] present in test data point [True]
285 Text feature [patient] present in test data point [True]
288 Text feature [negative] present in test data point [True]
289 Text feature [manufacturer] present in test data point [True]
290 Text feature [24] present in test data point [True]
291 Text feature [collected] present in test data point [True]
292 Text feature [distinct] present in test data point [True]
296 Text feature [level] present in test data point [True]
298 Text feature [sequenced] present in test data point [True]
299 Text feature [according] present in test data point [True]
302 Text feature [biotechnology] present in test data point [True]
303 Text feature [25] present in test data point [True]
304 Text feature [transduction] present in test data point [True]
305 Text feature [protocol] present in test data point [True]
306 Text feature [molecule] present in test data point [True]
308 Text feature [finding] present in test data point [True]
310 Text feature [normal] present in test data point [True]
311 Text feature [selected] present in test data point [True]
313 Text feature [significant] present in test data point [True]
314 Text feature [stimulation] present in test data point [True]
316 Text feature [provided] present in test data point [True]
317 Text feature [santa] present in test data point [True]
318 Text feature [incubated] present in test data point [True]
319 Text feature [levels] present in test data point [True]
320 Text feature [pcr] present in test data point [True]
321 Text feature [mutated] present in test data point [True]
322 Text feature [samples] present in test data point [True]
323 Text feature [harbor] present in test data point [True]
324 Text feature [48] present in test data point [True]
325 Text feature [cancer] present in test data point [True]
326 Text feature [added] present in test data point [True]
328 Text feature [cruz] present in test data point [True]
329 Text feature [remains] present in test data point [True]
330 Text feature [line] present in test data point [True]
331 Text feature [40] present in test data point [True]
332 Text feature [later] present in test data point [True]
333 Text feature [containing] present in test data point [True]
334 Text feature [regulated] present in test data point [True]
337 Text feature [disease] present in test data point [True]
339 Text feature [key] present in test data point [True]
340 Text feature [harboring] present in test data point [True]
342 Text feature [primary] present in test data point [True]
343 Text feature [indeed] present in test data point [True]
344 Text feature [experiments] present in test data point [True]
345 Text feature [50] present in test data point [True]
346 Text feature [96] present in test data point [True]
347 Text feature [mapk] present in test data point [True]
348 Text feature [collection] present in test data point [True]
349 Text feature [19] present in test data point [True]
350 Text feature [even] present in test data point [True]
351 Text feature [caused] present in test data point [True]
352 Text feature [represent] present in test data point [True]
```

```
353 Text feature [introduced] present in test data point [True]
354 Text feature [support] present in test data point [True]
355 Text feature [open] present in test data point [True]
356 Text feature [activity] present in test data point [True]
357 Text feature [form] present in test data point [True]
360 Text feature [100] present in test data point [True]
361 Text feature [cellular] present in test data point [True]
362 Text feature [22] present in test data point [True]
364 Text feature [low] present in test data point [True]
366 Text feature [still] present in test data point [True]
371 Text feature [highly] present in test data point [True]
372 Text feature [software] present in test data point [True]
373 Text feature [occurred] present in test data point [True]
374 Text feature [14] present in test data point [True]
383 Text feature [agents] present in test data point [True]
384 Text feature [completely] present in test data point [True]
385 Text feature [remain] present in test data point [True]
388 Text feature [doses] present in test data point [True]
389 Text feature [regulation] present in test data point [True]
390 Text feature [greater] present in test data point [True]
392 Text feature [therefore] present in test data point [True]
393 Text feature [3d] present in test data point [True]
395 Text feature [fold] present in test data point [True]
396 Text feature [type] present in test data point [True]
397 Text feature [minutes] present in test data point [True]
398 Text feature [manner] present in test data point [True]
401 Text feature [moreover] present in test data point [True]
403 Text feature [11] present in test data point [True]
404 Text feature [response] present in test data point [True]
405 Text feature [green] present in test data point [True]
406 Text feature [agent] present in test data point [True]
407 Text feature [concentration] present in test data point [True]
408 Text feature [driven] present in test data point [True]
409 Text feature [would] present in test data point [True]
411 Text feature [stained] present in test data point [True]
412 Text feature [4c] present in test data point [True]
413 Text feature [benefit] present in test data point [True]
416 Text feature [institute] present in test data point [True]
418 Text feature [ml] present in test data point [True]
420 Text feature [combination] present in test data point [True]
423 Text feature [complete] present in test data point [True]
424 Text feature [established] present in test data point [True]
425 Text feature [contained] present in test data point [True]
426 Text feature [possibility] present in test data point [True]
429 Text feature [observation] present in test data point [True]
430 Text feature [identify] present in test data point [True]
431 Text feature [upon] present in test data point [True]
434 Text feature [16] present in test data point [True]
438 Text feature [effectively] present in test data point [True]
439 Text feature [understanding] present in test data point [True]
442 Text feature [event] present in test data point [True]
443 Text feature [used] present in test data point [True]
444 Text feature [powerpoint] present in test data point [True]
446 Text feature [cases] present in test data point [True]
448 Text feature [whose] present in test data point [True]
451 Text feature [s1] present in test data point [True]
452 Text feature [oncogene] present in test data point [True]
455 Text feature [conformation] present in test data point [True]
456 Text feature [epidermal] present in test data point [True]
459 Text feature [amplification] present in test data point [True]
460 Text feature [activates] present in test data point [True]
461 Text feature [plates] present in test data point [True]
462 Text feature [sensitivity] present in test data point [True]
465 Text feature [signal] present in test data point [True]
468 Text feature [potent] present in test data point [True]
473 Text feature [particularly] present in test data point [True]
474 Text feature [4d] present in test data point [True]
479 Text feature [32] present in test data point [True]
480 Text feature [located] present in test data point [True]
482 Text feature [status] present in test data point [True]
483 Text feature [day] present in test data point [True]
484 Text feature [observations] present in test data point [True]
487 Text feature [selective] present in test data point [True]
488 Text feature [endogenous] present in test data point [True]
491 Text feature [extracted] present in test data point [True]
493 Text feature [threonine] present in test data point [True]
494 Text feature [metastatic] present in test data point [True]
495 Text feature [generation] present in test data point [True]
496 Text feature [control] present in test data point [True]
499 Text feature [blotting] present in test data point [True]
500 Text feature [prism] present in test data point [True]
503 Text feature [targeting] present in test data point [True]
507 Text feature [sample] present in test data point [True]
508 Text feature [28] present in test data point [True]
515 Text feature [required] present in test data point [True]
518 Text feature [80] present in test data point [True]
521 Text feature [33] present in test data point [True]
523 Text feature [alone] present in test data point [True]
```

```
524 Text feature [extent] present in test data point [True]
525 Text feature [erk] present in test data point [True]
530 Text feature [data] present in test data point [True]
532 Text feature [wild] present in test data point [True]
536 Text feature [transfection] present in test data point [True]
539 Text feature [resistant] present in test data point [True]
540 Text feature [exon] present in test data point [True]
542 Text feature [respective] present in test data point [True]
544 Text feature [modest] present in test data point [True]
546 Text feature [amplified] present in test data point [True]
547 Text feature [purchased] present in test data point [True]
551 Text feature [whole] present in test data point [True]
553 Text feature [provide] present in test data point [True]
554 Text feature [forms] present in test data point [True]
555 Text feature [reagent] present in test data point [True]
558 Text feature [72] present in test data point [True]
559 Text feature [respond] present in test data point [True]
565 Text feature [number] present in test data point [True]
566 Text feature [sites] present in test data point [True]
567 Text feature [constructs] present in test data point [True]
568 Text feature [dmso] present in test data point [True]
570 Text feature [associated] present in test data point [True]
575 Text feature [preclinical] present in test data point [True]
580 Text feature [mouse] present in test data point [True]
581 Text feature [transformed] present in test data point [True]
582 Text feature [atcc] present in test data point [True]
583 Text feature [200] present in test data point [True]
586 Text feature [90] present in test data point [True]
587 Text feature [multiple] present in test data point [True]
588 Text feature [early] present in test data point [True]
589 Text feature [reverse] present in test data point [True]
590 Text feature [exposure] present in test data point [True]
591 Text feature [density] present in test data point [True]
596 Text feature [future] present in test data point [True]
605 Text feature [blocked] present in test data point [True]
606 Text feature [strategies] present in test data point [True]
611 Text feature [block] present in test data point [True]
613 Text feature [promotes] present in test data point [True]
614 Text feature [reduced] present in test data point [True]
615 Text feature [needed] present in test data point [True]
618 Text feature [confer] present in test data point [True]
619 Text feature [effect] present in test data point [True]
620 Text feature [include] present in test data point [True]
621 Text feature [virus] present in test data point [True]
622 Text feature [generate] present in test data point [True]
624 Text feature [sufficient] present in test data point [True]
627 Text feature [exhibited] present in test data point [True]
628 Text feature [involving] present in test data point [True]
630 Text feature [drive] present in test data point [True]
631 Text feature [remained] present in test data point [True]
632 Text feature [majority] present in test data point [True]
633 Text feature [board] present in test data point [True]
635 Text feature [23] present in test data point [True]
640 Text feature [extracellular] present in test data point [True]
641 Text feature [numerous] present in test data point [True]
645 Text feature [short] present in test data point [True]
647 Text feature [harvested] present in test data point [True]
648 Text feature [pbs] present in test data point [True]
651 Text feature [protein] present in test data point [True]
655 Text feature [fbs] present in test data point [True]
658 Text feature [many] present in test data point [True]
659 Text feature [months] present in test data point [True]
660 Text feature [develop] present in test data point [True]
662 Text feature [review] present in test data point [True]
668 Text feature [red] present in test data point [True]
671 Text feature [washed] present in test data point [True]
679 Text feature [millipore] present in test data point [True]
681 Text feature [importantly] present in test data point [True]
682 Text feature [least] present in test data point [True]
686 Text feature [lysis] present in test data point [True]
692 Text feature [hypothesized] present in test data point [True]
693 Text feature [nucleotide] present in test data point [True]
697 Text feature [every] present in test data point [True]
700 Text feature [overnight] present in test data point [True]
701 Text feature [second] present in test data point [True]
702 Text feature [wide] present in test data point [True]
703 Text feature [obvious] present in test data point [True]
704 Text feature [formation] present in test data point [True]
706 Text feature [model] present in test data point [True]
707 Text feature [capable] present in test data point [True]
710 Text feature [expected] present in test data point [True]
711 Text feature [near] present in test data point [True]
713 Text feature [1e] present in test data point [True]
716 Text feature [context] present in test data point [True]
718 Text feature [alterations] present in test data point [True]
719 Text feature [events] present in test data point [True]
721 Text feature [leukemia] present in test data point [True]
722 Text feature [notably] present in test data point [True]
```

```
723 Text feature [mean] present in test data point [True]
728 Text feature [inhibit] present in test data point [True]
729 Text feature [insensitive] present in test data point [True]
730 Text feature [egfr] present in test data point [True]
731 Text feature [proteins] present in test data point [True]
732 Text feature [reports] present in test data point [True]
733 Text feature [21] present in test data point [True]
736 Text feature [5a] present in test data point [True]
737 Text feature [sds] present in test data point [True]
739 Text feature [phenotype] present in test data point [True]
740 Text feature [reaction] present in test data point [True]
741 Text feature [buffer] present in test data point [True]
742 Text feature [corresponding] present in test data point [True]
746 Text feature [maintain] present in test data point [True]
747 Text feature [types] present in test data point [True]
753 Text feature [hour] present in test data point [True]
755 Text feature [see] present in test data point [True]
760 Text feature [equivalent] present in test data point [True]
761 Text feature [genes] present in test data point [True]
765 Text feature [administered] present in test data point [True]
766 Text feature [case] present in test data point [True]
768 Text feature [biology] present in test data point [True]
769 Text feature [viral] present in test data point [True]
770 Text feature [characterization] present in test data point [True]
772 Text feature [biosciences] present in test data point [True]
773 Text feature [transferred] present in test data point [True]
776 Text feature [end] present in test data point [True]
781 Text feature [500] present in test data point [True]
782 Text feature [known] present in test data point [True]
784 Text feature [1d] present in test data point [True]
785 Text feature [understand] present in test data point [True]
788 Text feature [site] present in test data point [True]
789 Text feature [viable] present in test data point [True]
793 Text feature [reagents] present in test data point [True]
794 Text feature [pattern] present in test data point [True]
795 Text feature [potentially] present in test data point [True]
797 Text feature [44] present in test data point [True]
798 Text feature [gain] present in test data point [True]
805 Text feature [exhibit] present in test data point [True]
810 Text feature [matched] present in test data point [True]
815 Text feature [technologies] present in test data point [True]
818 Text feature [rates] present in test data point [True]
819 Text feature [evaluate] present in test data point [True]
821 Text feature [fluorescent] present in test data point [True]
825 Text feature [regulate] present in test data point [True]
826 Text feature [60] present in test data point [True]
828 Text feature [immediately] present in test data point [True]
829 Text feature [overcome] present in test data point [True]
831 Text feature [top] present in test data point [True]
834 Text feature [36] present in test data point [True]
837 Text feature [listed] present in test data point [True]
840 Text feature [toward] present in test data point [True]
843 Text feature [selection] present in test data point [True]
844 Text feature [tested] present in test data point [True]
848 Text feature [5c] present in test data point [True]
858 Text feature [hand] present in test data point [True]
859 Text feature [29] present in test data point [True]
860 Text feature [use] present in test data point [True]
864 Text feature [blue] present in test data point [True]
865 Text feature [fixed] present in test data point [True]
870 Text feature [sought] present in test data point [True]
871 Text feature [comparison] present in test data point [True]
873 Text feature [system] present in test data point [True]
874 Text feature [encodes] present in test data point [True]
876 Text feature [except] present in test data point [True]
880 Text feature [extraction] present in test data point [True]
882 Text feature [crystal] present in test data point [True]
890 Text feature [modified] present in test data point [True]
891 Text feature [clearly] present in test data point [True]
900 Text feature [life] present in test data point [True]
904 Text feature [recurrent] present in test data point [True]
906 Text feature [blocking] present in test data point [True]
914 Text feature [essential] present in test data point [True]
919 Text feature [notion] present in test data point [True]
921 Text feature [particular] present in test data point [True]
923 Text feature [methods] present in test data point [True]
926 Text feature [s4] present in test data point [True]
930 Text feature [synthesized] present in test data point [True]
933 Text feature [transduced] present in test data point [True]
935 Text feature [colony] present in test data point [True]
938 Text feature [like] present in test data point [True]
940 Text feature [implicated] present in test data point [True]
941 Text feature [values] present in test data point [True]
950 Text feature [thereby] present in test data point [True]
951 Text feature [characteristics] present in test data point [True]
952 Text feature [lentiviral] present in test data point [True]
955 Text feature [supplementary] present in test data point [True]
958 Text feature [period] present in test data point [True]
```

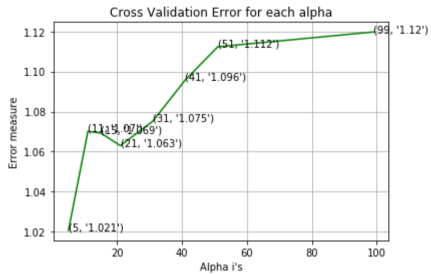
```
961 Text feature [probed] present in test data point [True]
963 Text feature [6b] present in test data point [True]
970 Text feature [2000] present in test data point [True]
971 Text feature [nm] present in test data point [True]
974 Text feature [times] present in test data point [True]
977 Text feature [occurring] present in test data point [True]
978 Text feature [frame] present in test data point [True]
980 Text feature [plus] present in test data point [True]
982 Text feature [association] present in test data point [True]
986 Text feature [alter] present in test data point [True]
987 Text feature [indicates] present in test data point [True]
990 Text feature [inactive] present in test data point [True]
993 Text feature [provides] present in test data point [True]
995 Text feature [acquired] present in test data point [True]
996 Text feature [striking] present in test data point [True]
Out of the top 1000 features 538 are present in query point
```

4.2. K Nearest Neighbour Classification

4.2.1. Hyper parameter tuning

```
In [75]: # find more about KNeighborsClassifier()
         # here http://scikit-learn.org/stable/modules/generated/sklearn.neighbors.KNeighborsClassifier.html
         # ______
         # default parameter
         # KNeighborsClassifier(n_neighbors=5, weights='uniform', algorithm='auto', leaf_size=30, p=2,
         # metric='minkowski', metric_params=None, n_jobs=1, **kwargs)
         # methods of
         \# fit(X, y) : Fit the model using X as training data and y as target values
         # predict(X):Predict the class labels for the provided data
         # predict_proba(X):Return probability estimates for the test data X.
         #-----
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/k-nearest-neighbors-geometric-in
         tuition-with-a-toy-example-1/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # -----
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         #-----
         # video link:
         #-----
         alpha = [5, 11, 15, 21, 31, 41, 51, 99]
         cv_log_error_array = []
         for i in alpha:
             print("for alpha =", i)
             clf = KNeighborsClassifier(n_neighbors=i)
             clf.fit(train_x_responseCoding, train_y)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_x_responseCoding, train_y)
             sig_clf_probs = sig_clf.predict_proba(cv_x_responseCoding)
             cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
             # to avoid rounding error while multiplying probabilites we use log-probability estimates
             print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = KNeighborsClassifier(n neighbors=alpha[best alpha])
         clf.fit(train_x_responseCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_responseCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_responseCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_,eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_responseCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_x_responseCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

```
for alpha = 5
Log Loss : 1.0207102110793702
for alpha = 11
Log Loss : 1.0701898518708892
for alpha = 15
Log Loss : 1.069260952087364
for alpha = 21
Log Loss: 1.0630030649170528
for alpha = 31
Log Loss: 1.0752161883257838
for alpha = 41
Log Loss : 1.0957851900888886
for alpha = 51
Log Loss: 1.1123202098355227
for alpha = 99
Log Loss : 1.1197198931974635
```



For values of best alpha = 5 The train log loss is: 0.5088538403861076

For values of best alpha = 5 The cross validation log loss is: 1.0207102110793702

For values of best alpha = 5 The test log loss is: 1.042859714523589

4.2.2. Testing the model with best hyper paramters

Number of mis-classified points : 0.34210526315789475

----- Confusion matrix -----55.000 1.000 0.000 23.000 4.000 7.000 1.000 0.000 0.000 - 100 1.000 39.000 1.000 1.000 0.000 1.000 29.000 0.000 0.000 4.000 0.000 0.000 1.000 7.000 0.000 2.000 0.000 0.000 - 80 14.000 2.000 1.000 4.000 1.000 2.000 1.000 0.000 Original Class 6.000 3.000 0.000 8.000 12.000 4.000 6.000 0.000 0.000 - 60 29.000 6.000 3.000 0.000 4.000 1.000 1.000 0.000 0.000 - 40 0.000 26.000 5.000 2.000 2.000 0.000 118.000 0.000 0.000 0.000 1.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 - 20 0.000 0.000 0.000 0.000 0.000 1.000 1.000 4.000 Predicted Class ----- Precision matrix (Columm Sum=1) -----1.0 0.013 0.000 0.181 0.174 0.167 0.006 0.000 0.000 0.071 0.008 0.000 0.012 0.024 0.180 0.000 0.000 - 0.8 0.000 0.013 0.031 0.000 0.000 0.000 0.012 0.000 0.171 0.026 0.071 0.174 0.024 0.012 0.333 0.000 - 0.6 0.073 0.039 0.000 0.063 0.095 0.037 0.000 0.000 - 0.4 0.039 0.000 0.031 0.073 0.043 0.006 0.000 0.000 0.342 0.357 0.000 0.016 0.087 0.000 0.000 0.000 - 0.2 0.000 0.013 0.000 0.000 0.000 0.000 0.006 0.333 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.006 0.333 1.000 - 0.0 8 Predicted Class ----- Recall matrix (Row sum=1) ------- 0.75 0.011 0.000 0.253 0.044 0.077 0.011 0.000 0.000 0.014 0.014 0.014 0.000 0.014 0.000 0.000 - 0.60 0.286 0.000 0.071 0.000 0.000 0.143 0.000 0.000 0.127 0.018 0.009 0.036 0.009 0.018 0.009 0.000 - 0.45 0.154 0.077 0.000 0.205 0.308 0.103 0.154 0.000 0.000 0.136 0.068 0.023 0.000 0.091 0.023 0.000 0.000 - 0.30 0.000 0.170 0.033 0.013 0.013 0.000 0.000 0.000 -0.15 0.333 0.333 0.333 0.000 0.000 0.000 0.000 0.000 0.000 0.167 0.000 0.000 0.000 0.000 0.000 0.000 0.167 - 0.00 2 'n 8 9

Predicted Class

```
In [77]: clf = KNeighborsClassifier(n_neighbors=alpha[best_alpha])
    clf.fit(train_x_responseCoding, train_y)
    sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
    sig_clf.fit(train_x_responseCoding, train_y)

    test_point_index = 1
    predicted_cls = sig_clf.predict(test_x_responseCoding[0].reshape(1,-1))
    print("Predicted Class :", predicted_cls[0])
    print("Actual Class :", test_y[test_point_index])
    neighbors = clf.kneighbors(test_x_responseCoding[test_point_index].reshape(1, -1), alpha[best_alpha])
    print("The ",alpha[best_alpha]," nearest neighbours of the test points belongs to classes",train_y[neighbors[1][0]])

    Predicted Class : 1
    Actual Class : 5
    The 5 nearest neighbours of the test points belongs to classes [1 5 4 1 1]
    Fequency of nearest points : Counter({1: 3, 5: 1, 4: 1})
```

4.2.4. Sample Query Point-2

```
In [78]: | clf = KNeighborsClassifier(n_neighbors=alpha[best_alpha])
         clf.fit(train_x_responseCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_responseCoding, train_y)
         test_point_index = 100
         predicted_cls = sig_clf.predict(test_x_responseCoding[test_point_index].reshape(1,-1))
         print("Predicted Class :", predicted_cls[0])
         print("Actual Class :", test_y[test_point_index])
         neighbors = clf.kneighbors(test_x_responseCoding[test_point_index].reshape(1, -1), alpha[best_alpha])
         print("the k value for knn is",alpha[best_alpha],"and the nearest neighbours of the test points belongs to classes",tr
         ain_y[neighbors[1][0]])
         print("Fequency of nearest points :",Counter(train_y[neighbors[1][0]]))
         Predicted Class: 4
         Actual Class : 4
         the k value for knn is 5 and the nearest neighbours of the test points belongs to classes [4 4 4 3 7]
         Fequency of nearest points : Counter({4: 3, 3: 1, 7: 1})
```

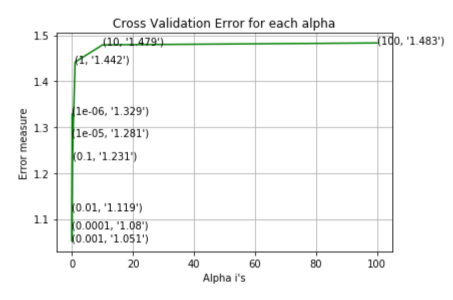
4.3. Logistic Regression

4.3.1. With Class balancing

4.3.1.1. Hyper paramter tuning

```
In [79]: | # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear_model.SGDClassifi
         er.html
         # -----
         # default parameters
         # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
         # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
         # class weight=None, warm_start=False, average=False, n_iter=None)
         # some of methods
         # fit(X, y[, coef_init, intercept_init, ...]) Fit linear model with Stochastic Gradient Descent.
         \# predict(X) Predict class labels for samples in X.
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/geometric-intuition-1/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # ------
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         #-----
         # video Link:
         #-----
         alpha = [10 ** x for x in range(-6, 3)]
         cv_log_error_array = []
         for i in alpha:
             print("for alpha =", i)
             clf = SGDClassifier(class_weight='balanced', alpha=i, penalty='12', loss='log', random_state=42)
             clf.fit(train_x_onehotCoding, train_y)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_x_onehotCoding, train_y)
             sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
             cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
             # to avoid rounding error while multiplying probabilites we use log-probability estimates
             print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train_x_onehotCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_onehotCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best alpha],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict y = sig clf.predict proba(test x onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

for alpha = 1e-06Log Loss: 1.3286245067921225 for alpha = 1e-05 Log Loss : 1.281251066396459 for alpha = 0.0001Log Loss : 1.079926912296904 for alpha = 0.001Log Loss : 1.0511161987442945 for alpha = 0.01Log Loss: 1.1194022062599829 for alpha = 0.1Log Loss: 1.230877272340499 for alpha = 1Log Loss : 1.441711651755352 for alpha = 10Log Loss : 1.4792202960456888 for alpha = 100Log Loss: 1.4833463424991187



For values of best alpha = 0.001 The train log loss is: 0.5227411978926324

For values of best alpha = 0.001 The cross validation log loss is: 1.0511161987442945

For values of best alpha = 0.001 The test log loss is: 1.014271573198189

4.3.1.2. Testing the model with best hyper paramters

----- Confusion matrix ------ 125 53.000 1.000 1.000 24.000 5.000 3.000 4.000 0.000 0.000 2.000 36.000 0.000 1.000 0.000 1.000 32.000 0.000 0.000 - 100 4.000 0.000 0.000 0.000 6.000 0.000 4.000 0.000 0.000 18.000 2.000 1.000 5.000 1.000 13.000 0.000 0.000 Original Class - 75 7.000 2.000 0.000 6.000 13.000 4.000 7.000 0.000 0.000 6.000 0.000 0.000 4.000 3.000 28.000 3.000 0.000 0.000 - 50 1.000 16.000 1.000 2.000 2.000 0.000 131.000 0.000 0.000 - 25 0.000 1.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 1.000 1.000 4.000 Predicted Class ----- Precision matrix (Columm Sum=1) -----1.0 0.017 0.111 0.216 0.179 0.081 0.020 0.000 0.000 0.000 0.009 0.000 0.023 0.027 0.163 0.000 0.000 - 0.8 0.000 0.000 0.036 0.000 0.000 0.020 0.000 0.000 0.207 0.034 0.111 0.179 0.027 0.066 0.000 0.000 - 0.6 Original Class 0.080 0.034 0.000 0.054 0.464 0.108 0.036 0.000 0.000 - 0.4 0.000 0.036 0.069 0.000 0.107 0.015 0.000 0.000 0.276 0.011 0.111 0.018 0.071 0.000 0.000 0.000 - 0.2 0.000 0.017 0.000 0.000 0.000 0.000 0.005 0.000 0.005 0.000 0.000 0.000 0.000 0.000 0.000 1.000 თ -- 0.0 2 8 9 Predicted Class ----- Recall matrix (Row sum=1) ------0.011 0.011 0.264 0.055 0.033 0.044 0.000 0.000 - 0.75 0.000 0.014 0.028 0.000 0.014 0.000 0.000 0.000 0.286 0.000 0.000 0.000 0.286 0.000 0.000 - 0.60 0.045 0.164 0.018 0.009 0.009 0.118 0.000 0.000 - 0.45 0.179 0.051 0.000 0.154 0.333 0.103 0.179 0.000 0.000 0.136 0.000 0.000 0.091 0.068 0.068 0.000 0.000 0.30 0.007 0.105 0.007 0.013 0.013 0.000 0.856 0.000 0.000 -0.15 0.333 0.000 0.333 0.333 0.000 0.000 0.000 0.000 0.000

0.000

Predicted Class

0.000

0.167

7

0.000

4

0.167

8

9

- 0.00

0.000

2

0.000

0.000

i

```
In [0]: | def get_imp_feature_names(text, indices, removed_ind = []):
            word_present = 0
            tabulte_list = []
            incresingorder_ind = 0
            for i in indices:
                if i < train_gene_feature_onehotCoding.shape[1]:</pre>
                     tabulte_list.append([incresingorder_ind, "Gene", "Yes"])
                     tabulte_list.append([incresingorder_ind, "Variation", "Yes"])
                if ((i > 17) & (i not in removed_ind)) :
                    word = train_text_features[i]
                     yes_no = True if word in text.split() else False
                    if yes_no:
                         word_present += 1
                     tabulte_list.append([incresingorder_ind,train_text_features[i], yes_no])
                incresingorder_ind += 1
            print(word_present, "most importent features are present in our query point")
            print("-"*50)
            print("The features that are most importent of the ",predicted_cls[0]," class:")
            print (tabulate(tabulte_list, headers=["Index",'Feature name', 'Present or Not']))
```

4.3.1.3.1. Correctly Classified point

```
In [84]: # from tabulate import tabulate
         clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train_x_onehotCoding,train_y)
         test_point_index = 12
         no feature = 1000
         predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
         print("Predicted Class :", predicted_cls[0])
         print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
         print("Actual Class :", test_y[test_point_index])
         indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
         print("-"*50)
         get_impfeature_names(indices[0],
                              x_test['TEXT'].iloc[test_point_index],
                              x_test['Gene'].iloc[test_point_index],
                              x_test['Variation'].iloc[test_point_index],
                              no_feature)
         Predicted Class: 4
         Predicted Class Probabilities: [[4.900e-03 7.900e-03 2.600e-03 6.221e-01 3.163e-01 3.920e-02 4.700e-03
           1.800e-03 5.000e-04]]
         Actual Class : 4
         263 Text feature [germline] present in test data point [True]
         306 Text feature [y371h] present in test data point [True]
         341 Text feature [tagged] present in test data point [True]
         355 Text feature [suppressor] present in test data point [True]
         613 Text feature [popmusic] present in test data point [True]
         924 Text feature [hcl] present in test data point [True]
         Out of the top 1000 features 6 are present in query point
```

4.3.1.3.2. Incorrectly Classified point

```
In [86]: | test point index = 5
         no feature = 1000
         predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
         print("Predicted Class :", predicted_cls[0])
         print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
         print("Actual Class :", test_y[test_point_index])
         indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
         print("-"*50)
         get_impfeature_names(indices[0],
                              x_test['TEXT'].iloc[test_point_index],
                              x_test['Gene'].iloc[test_point_index],
                              x_test['Variation'].iloc[test_point_index],
                              no_feature)
         Predicted Class : 7
         Predicted Class Probabilities: [[0.0335 0.126 0.0073 0.0537 0.0241 0.0195 0.7289 0.006 0.0011]]
         Actual Class : 1
         25 Text feature [constitutively] present in test data point [True]
         32 Text feature [activated] present in test data point [True]
         35 Text feature [oncoprotein] present in test data point [True]
         46 Text feature [oncogene] present in test data point [True]
         89 Text feature [elicited] present in test data point [True]
         152 Text feature [11e] present in test data point [True]
         153 Text feature [mitogen] present in test data point [True]
         156 Text feature [transformation] present in test data point [True]
         158 Text feature [activation] present in test data point [True]
         195 Text feature [downstream] present in test data point [True]
         278 Text feature [skmel28] present in test data point [True]
         286 Text feature [phosphorylation] present in test data point [True]
         326 Text feature [hgf] present in test data point [True]
         352 Text feature [erk] present in test data point [True]
         384 Text feature [phospho] present in test data point [True]
         386 Text feature [mapk] present in test data point [True]
         431 Text feature [biopsy] present in test data point [True]
         451 Text feature [overexpression] present in test data point [True]
         525 Text feature [activating] present in test data point [True]
         538 Text feature [tyrosine] present in test data point [True]
         542 Text feature [expressing] present in test data point [True]
         576 Text feature [melanocytes] present in test data point [True]
         583 Text feature [receptor] present in test data point [True]
         619 Text feature [activate] present in test data point [True]
         634 Text feature [doses] present in test data point [True]
         792 Text feature [cultured] present in test data point [True]
         815 Text feature [malignant] present in test data point [True]
         903 Text feature [plx4032] present in test data point [True]
         906 Text feature [remain] present in test data point [True]
         910 Text feature [inhibitor] present in test data point [True]
         938 Text feature [transformed] present in test data point [True]
         958 Text feature [signaling] present in test data point [True]
         Out of the top 1000 features 32 are present in query point
```

4.3.2. Without Class balancing

4.3.2.1. Hyper paramter tuning

```
In [87]: # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear_model.SGDClassifi
         er.htmL
         # -----
         # default parameters
         # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
         # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
         # class weight=None, warm_start=False, average=False, n_iter=None)
         # some of methods
         # fit(X, y[, coef_init, intercept_init, ...]) Fit linear model with Stochastic Gradient Descent.
         \# predict(X) Predict class labels for samples in X.
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/geometric-intuition-1/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         \# predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         #-----
         # video link:
         alpha = [10 ** x for x in range(-6, 1)]
         cv_log_error_array = []
         for i in alpha:
             print("for alpha =", i)
             clf = SGDClassifier(alpha=i, penalty='12', loss='log', random_state=42)
             clf.fit(train x onehotCoding, train y)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_x_onehotCoding, train_y)
             sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
             cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
             print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train_x_onehotCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_onehotCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict y = sig clf.predict proba(test x onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

for alpha = 1e-06
Log Loss : 1.2804729730683213
for alpha = 1e-05

Log Loss : 1.2239806482265947

for alpha = 0.0001

Log Loss: 1.0963867561753107

for alpha = 0.001 Log Loss : 1.0715726841165094

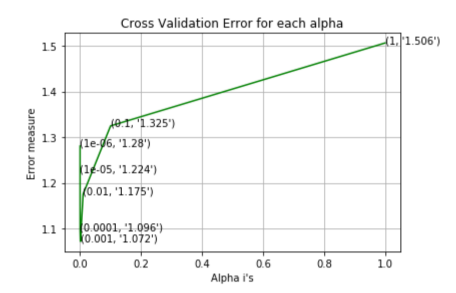
for alpha = 0.01

Log Loss : 1.1751276000460735

for alpha = 0.1 Log Loss : 1.3246934516703148

for alpha = 1

Log Loss : 1.506323364941991



For values of best alpha = 0.001 The train log loss is: 0.5205306499237352

For values of best alpha = 0.001 The cross validation log loss is: 1.0715726841165094

For values of best alpha = 0.001 The test log loss is: 1.0406989550967904

4.3.2.2. Testing model with best hyper parameters

----- Confusion matrix -----53.000 0.000 1.000 25.000 5.000 3.000 4.000 0.000 0.000 - 125 2.000 37.000 0.000 1.000 0.000 1.000 31.000 0.000 0.000 - 100 0.000 6.000 0.000 0.000 4.000 0.000 4.000 0.000 0.000 13.000 1.000 1.000 5.000 1.000 15.000 0.000 0.000 Original Class - 75 8.000 2.000 0.000 7.000 12.000 3.000 7.000 0.000 0.000 5.000 0.000 0.000 6.000 2.000 28.000 3.000 0.000 0.000 - 50 1.000 13.000 0.000 1.000 2.000 0.000 136.000 0.000 0.000 - 25 0.000 1.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 2.000 1.000 3.000 Predicted Class ----- Precision matrix (Columm Sum=1) ------1.0 0.000 0.167 0.208 0.192 0.083 0.020 0.000 0.000 0.000 0.008 0.000 0.024 0.028 0.153 0.000 0.000 - 0.8 0.000 0.000 0.050 0.000 0.000 0.020 0.000 0.000 0.159 0.019 0.167 0.192 0.028 0.074 0.000 0.000 - 0.6 0.098 0.037 0.000 0.058 0.462 0.083 0.034 0.000 0.000 - 0.4 0.000 0.000 0.050 0.015 0.061 0.077 0.000 0.000 0.241 0.012 0.000 0.008 0.077 0.000 0.000 0.000 - 0.2 0.000 0.019 0.000 0.000 0.000 0.000 0.005 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.010 1.000 - 0.0 2 8 Predicted Class ----- Recall matrix (Row sum=1) ------0.000 0.011 0.275 0.055 0.033 0.044 0.000 0.000 - 0.75 0.000 0.014 0.028 0.000 0.014 0.000 0.000 0.000 0.286 0.000 0.000 0.000 0.286 0.000 0.000 - 0.60 0.118 0.045 0.009 0.009 0.009 0.136 0.000 0.000 - 0.45 0.205 0.051 0.000 0.179 0.308 0.077 0.179 0.000 0.000 0.114 0.000 0.000 0.136 0.045 0.068 0.000 0.000 0.30 0.007 0.085 0.000 0.007 0.013 0.000 0.889 0.000 0.000 0.333 0.333 0.333 -0.15 0.000 0.000 0.000 0.000 0.000 0.000

0.000

Predicted Class

0.000

0.000

4

0.333

7

0.167

8

9

- 0.00

0.000

0.000

2

0.000

```
In [89]: | clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
         clf.fit(train x onehotCoding,train y)
         test_point_index = 100
         no_feature = 1000
         predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
         print("Predicted Class :", predicted_cls[0])
         print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
         print("Actual Class :", test y[test point index])
         indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
         print("-"*50)
         get_impfeature_names(indices[0],
                              x_test['TEXT'].iloc[test_point_index],
                              x_test['Gene'].iloc[test_point_index],
                              x_test['Variation'].iloc[test_point_index],
                              no_feature)
         Predicted Class: 4
         Predicted Class Probabilities: [[2.000e-02 3.220e-02 5.300e-03 8.965e-01 9.800e-03 9.900e-03 2.350e-02
           2.200e-03 7.000e-04]]
         Actual Class : 4
         231 Text feature [biallelic] present in test data point [True]
         246 Text feature [impairing] present in test data point [True]
         377 Text feature [synergic] present in test data point [True]
         402 Text feature [704] present in test data point [True]
         427 Text feature [families] present in test data point [True]
         524 Text feature [donor] present in test data point [True]
         719 Text feature [grandmother] present in test data point [True]
         847 Text feature [severely] present in test data point [True]
         877 Text feature [5th] present in test data point [True]
         893 Text feature [anticipation] present in test data point [True]
         Out of the top 1000 features 10 are present in query point
```

4.3.2.4. Feature Importance, Inorrectly Classified point

```
In [90]: | test point index = 1
         no feature = 1000
         predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
         print("Predicted Class :", predicted_cls[0])
         print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
         print("Actual Class :", test_y[test_point_index])
         indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
         print("-"*50)
         get_impfeature_names(indices[0],
                               x_test['TEXT'].iloc[test_point_index],
                               x_test['Gene'].iloc[test_point_index],
                               x_test['Variation'].iloc[test_point_index],
                               no_feature)
         Predicted Class : 1
         Predicted Class Probabilities: [[7.270e-01 1.890e-02 3.000e-03 1.383e-01 8.670e-02 1.270e-02 1.290e-02
           5.000e-04 0.000e+00]]
         Actual Class : 5
         36 Text feature [misinterpretation] present in test data point [True]
         143 Text feature [rationalization] present in test data point [True]
         157 Text feature [s1613c] present in test data point [True]
         260 Text feature [v1833m] present in test data point [True]
         283 Text feature [unstructured] present in test data point [True]
         547 Text feature [dbd] present in test data point [True]
         552 Text feature [repeats] present in test data point [True]
         554 Text feature [rendered] present in test data point [True]
         569 Text feature [chemoprevention] present in test data point [True]
         694 Text feature [folding] present in test data point [True]
         864 Text feature [ese] present in test data point [True]
         893 Text feature [rodent] present in test data point [True]
         909 Text feature [truncating] present in test data point [True]
         990 Text feature [calorimetric] present in test data point [True]
```

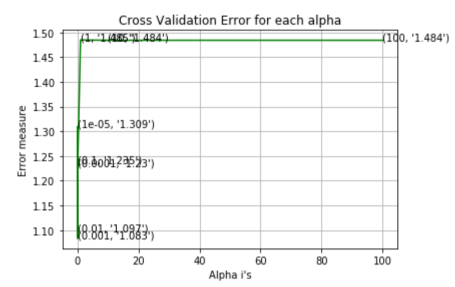
4.4. Linear Support Vector Machines

Out of the top 1000 features 14 are present in query point

4.4.1. Hyper paramter tuning

```
In [91]: # read more about support vector machines with linear kernals here http://scikit-learn.org/stable/modules/generated/sk
         Learn.svm.SVC.htmL
         # -----
         # default parameters
         # SVC(C=1.0, kernel='rbf', degree=3, gamma='auto', coef0=0.0, shrinking=True, probability=False, tol=0.001,
         # cache_size=200, class_weight=None, verbose=False, max_iter=-1, decision_function_shape='ovr', random_state=None)
         # Some of methods of SVM()
         \# fit(X, y, [sample\_weight]) Fit the SVM model according to the given training data.
         \# predict(X) Perform classification on samples in X.
         # -----
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/mathematical-derivation-copy-8/
         # -----
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # -----
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         #-----
         # video link:
         #-----
         alpha = [10 ** x for x in range(-5, 3)]
         cv_log_error_array = []
         for i in alpha:
            print("for C =", i)
             clf = SVC(C=i,kernel='linear',probability=True, class_weight='balanced')
            clf = SGDClassifier( class_weight='balanced', alpha=i, penalty='12', loss='hinge', random_state=42)
             clf.fit(train_x_onehotCoding, train_y)
             sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
             sig_clf.fit(train_x_onehotCoding, train_y)
             sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
             cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
             print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         fig, ax = plt.subplots()
         ax.plot(alpha, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         # clf = SVC(C=i,kernel='linear',probability=True, class_weight='balanced')
         clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='hinge', random_state=42)
         clf.fit(train x onehotCoding, train y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_onehotCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_x_onehotCoding)
         print('For values of best alpha = ',
               alpha[best_alpha],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

```
for C = 1e-05
Log Loss: 1.3091203579003237
for C = 0.0001
Log Loss : 1.2295636226048856
for C = 0.001
Log Loss : 1.0832675057609182
for C = 0.01
Log Loss : 1.0971099861721214
for C = 0.1
Log Loss: 1.2353906667767993
for C = 1
Log Loss : 1.4846582194755762
for C = 10
Log Loss: 1.4840503434177905
for C = 100
Log Loss: 1.4840503526488262
```



```
For values of best alpha = 0.001 The train log loss is: 0.5598621871187457

For values of best alpha = 0.001 The cross validation log loss is: 1.0832675057609182

For values of best alpha = 0.001 The test log loss is: 1.074936412241439
```

4.4.2. Testing model with best hyper parameters

----- Confusion matrix -----125 52.000 0.000 1.000 25.000 7.000 4.000 2.000 0.000 0.000 2.000 36.000 0.000 1.000 0.000 32.000 1.000 0.000 0.000 - 100 4.000 0.000 0.000 6.000 0.000 0.000 4.000 0.000 0.000 13.000 0.000 2.000 5.000 2.000 17.000 0.000 0.000 Original Class 5.000 3.000 0.000 8.000 16.000 3.000 4.000 0.000 0.000 - 50 6.000 0.000 0.000 4.000 2.000 28.000 4.000 0.000 0.000 3.000 18.000 1.000 2.000 4.000 0.000 125.000 0.000 0.000 - 25 0.000 1.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 0.000 1.000 0.000 0.000 0.000 0.000 3.000 0.000 2.000 Predicted Class ----- Precision matrix (Columm Sum=1) ------1.0 0.000 0.100 0.216 0.206 0.105 0.010 0.000 0.000 0.000 0.009 0.000 0.025 0.026 0.167 0.000 0.000 - 0.8 0.000 0.034 0.000 0.000 0.000 0.021 0.000 0.000 0.160 0.000 0.200 0.147 0.053 0.089 0.000 0.000 - 0.6 0.062 0.052 0.000 0.069 0.471 0.079 0.021 0.000 0.000 - 0.4 0.000 0.034 0.074 0.000 0.059 0.021 0.000 0.000 0.310 0.037 0.100 0.017 0.118 0.000 0.000 0.000 - 0.2 0.000 0.017 0.000 0.000 0.000 0.000 0.005 1.000 0.000 0.000 0.000 0.000 0.009 0.000 0.000 0.016 0.000 1.000 - 0.0 2 9 Predicted Class ----- Recall matrix (Row sum=1) ------0.000 0.011 0.275 0.077 0.044 0.022 0.000 0.000 - 0.75 0.000 0.014 0.028 0.000 0.014 0.000 0.000 - 0.60 0.286 0.000 0.000 0.000 0.000 0.286 0.000 0.000 0.118 0.645 0.045 0.000 0.018 0.018 0.155 0.000 0.000 - 0.45 0.128 0.077 0.000 0.205 0.077 0.103 0.000 0.000 0.136 0.000 0.000 0.091 0.045 0.091 0.000 0.000 - 0.30 0.020 0.118 0.007 0.013 0.026 0.000 0.817 0.000 0.000 -0.15 0.333 0.000 0.333 0.333 0.000 0.000 0.000 0.000 0.000 0.000 0.167 0.000 0.000 0.000 0.000 0.000 0.333 - 0.00 7 9 4

Predicted Class

4.3.3. Feature Importance

4.3.3.1. For Correctly classified point

```
In [94]: | clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='hinge', random_state=42)
            clf.fit(train_x_onehotCoding,train_y)
            test_point_index = 12
            # test_point_index = 100
            no_feature = 1000
            predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
            print("Predicted Class :", predicted_cls[0])
            print("Predicted Class Probabilities:", np.round(sig clf.predict proba(test x onehotCoding[test point index]),4))
            print("Actual Class :", test_y[test_point_index])
            indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
            print("-"*50)
            get_impfeature_names(indices[0],
                                  x_test['TEXT'].iloc[test_point_index],
                                 x_test['Gene'].iloc[test_point_index],
                                  x_test['Variation'].iloc[test_point_index],
                                  no_feature)
            Predicted Class : 4
            Predicted Class Probabilities: [[0.0365 0.0115 0.0108 0.5439 0.3064 0.0334 0.048 0.004 0.0055]]
            Actual Class : 4
            268 Text feature [germline] present in test data point [True]
            375 Text feature [endogenous] present in test data point [True]
            401 Text feature [y371h] present in test data point [True]
            488 Text feature [tagged] present in test data point [True]
            569 Text feature [attributed] present in test data point [True]
            591 Text feature [linked] present in test data point [True]
            634 Text feature [coordinating] present in test data point [True]
            661 Text feature [popmusic] present in test data point [True]
            751 Text feature [sodium] present in test data point [True]
            832 Text feature [lipofectamine] present in test data point [True]
            951 Text feature [consequences] present in test data point [True]
            952 Text feature [suppressor] present in test data point [True]
            998 Text feature [place] present in test data point [True]
            Out of the top 1000 features 13 are present in query point
4.3.3.2. For Incorrectly classified point
   In [96]: | test_point_index = 1
            no_feature = 1000
            predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
            print("Predicted Class :", predicted_cls[0])
            print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
            print("Actual Class :", test_y[test_point_index])
            indices = np.argsort(-clf.coef_)[predicted_cls-1][:,:no_feature]
            print("-"*50)
            get_impfeature_names(indices[0],
                                  x_test['TEXT'].iloc[test_point_index],
                                  x_test['Gene'].iloc[test_point_index],
                                  x_test['Variation'].iloc[test_point_index],
                                  no_feature)
            Predicted Class : 1
            Predicted Class Probabilities: [[0.637  0.0728  0.0114  0.0926  0.0999  0.0136  0.0615  0.0054  0.0057]]
            Actual Class : 5
            108 Text feature [misinterpretation] present in test data point [True]
            139 Text feature [rationalization] present in test data point [True]
            146 Text feature [s1613c] present in test data point [True]
            235 Text feature [valuable] present in test data point [True]
            241 Text feature [rendered] present in test data point [True]
            266 Text feature [v1833m] present in test data point [True]
            273 Text feature [unstructured] present in test data point [True]
            334 Text feature [m1652i] present in test data point [True]
            337 Text feature [24h] present in test data point [True]
            491 Text feature [chemoprevention] present in test data point [True]
            560 Text feature [arrived] present in test data point [True]
```

564 Text feature [calorimetric] present in test data point [True] 580 Text feature [shortage] present in test data point [True]

695 Text feature [computation] present in test data point [True]

596 Text feature [dbd] present in test data point [True]

727 Text feature [m1652] present in test data point [True]
779 Text feature [reagents] present in test data point [True]
784 Text feature [frameshift] present in test data point [True]
825 Text feature [inactivation] present in test data point [True]

863 Text feature [tools] present in test data point [True]
876 Text feature [v1833] present in test data point [True]
877 Text feature [i1855] present in test data point [True]
878 Text feature [11854] present in test data point [True]
879 Text feature [y1853] present in test data point [True]
892 Text feature [retaining] present in test data point [True]
Out of the top 1000 features 25 are present in query point

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4.5.1. Hyper paramter tuning (With One hot Encoding)

```
In [97]: | # ------
         # default parameters
         # sklearn.ensemble.RandomForestClassifier(n_estimators=10, criterion='gini', max_depth=None, min_samples_split=2,
         # min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_impurity_decrease=0.
         # min_impurity_split=None, bootstrap=True, oob_score=False, n_jobs=1, random_state=None, verbose=0, warm_start=False,
         # class_weight=None)
         # Some of methods of RandomForestClassifier()
         \# fit(X, y, [sample\_weight]) Fit the SVM model according to the given training data.
         \# predict(X) Perform classification on samples in X.
                              Perform classification on samples in X.
         # predict_proba (X)
         # some of attributes of RandomForestClassifier()
         # feature_importances_ : array of shape = [n_features]
         # The feature importances (the higher, the more important the feature).
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/random-forest-and-their-construc
         tion-2/
         # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
         CalibratedClassifierCV.html
         # ------
         # default paramters
         # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
         # some of the methods of CalibratedClassifierCV()
         # fit(X, y[, sample_weight])
Fit the calibrated model
         # get_params([deep]) Get parameters for this estimator.
         # predict(X) Predict the target of new samples.
         # predict_proba(X) Posterior probabilities of classification
         #-----
         # video link:
         #-----
         alpha = [100,200,500,1000,2000]
         max_depth = [5, 10]
         cv_log_error_array = []
         for i in alpha:
             for j in max_depth:
                 print("for n_estimators =", i,"and max depth = ", j)
                 clf = RandomForestClassifier(n_estimators=i, criterion='gini', max_depth=j, random_state=42, n_jobs=-1)
                 clf.fit(train_x_onehotCoding, train_y)
                 sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
                 sig_clf.fit(train_x_onehotCoding, train_y)
                 sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
                 cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
                 print("Log Loss :",log_loss(cv_y, sig_clf_probs))
         '''fig, ax = plt.subplots()
         features = np.dot(np.array(alpha)[:,None],np.array(max_depth)[None]).ravel()
         ax.plot(features, cv_log_error_array,c='g')
         for i, txt in enumerate(np.round(cv_log_error_array,3)):
             ax.annotate((alpha[int(i/2)],max_depth[int(i%2)],str(txt)), (features[i],cv_log_error_array[i]))
         plt.grid()
         plt.title("Cross Validation Error for each alpha")
         plt.xlabel("Alpha i's")
         plt.ylabel("Error measure")
         plt.show()
         best_alpha = np.argmin(cv_log_error_array)
         clf = RandomForestClassifier(n_estimators=alpha[int(best_alpha/2)], criterion='gini', max_depth=max_depth[int(best_alp
         ha%2)], random state=42, n jobs=-1)
         clf.fit(train_x_onehotCoding, train_y)
         sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
         sig_clf.fit(train_x_onehotCoding, train_y)
         predict_y = sig_clf.predict_proba(train_x_onehotCoding)
         print('For values of best estimator = ',
               alpha[int(best_alpha/2)],
               "The train log loss is:",
               log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
         print('For values of best estimator = ',
               alpha[int(best_alpha/2)],
               "The cross validation log loss is:",
               log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
         predict_y = sig_clf.predict_proba(test_x_onehotCoding)
         print('For values of best estimator = ',
               alpha[int(best_alpha/2)],
               "The test log loss is:",
               log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

```
for n_{estimators} = 100 and max depth = 5
Log Loss: 1.2503658683902228
for n_{estimators} = 100 and max depth = 10
Log Loss : 1.1869730488857573
for n_estimators = 200 and max depth = 5
Log Loss: 1.2429350726830202
for n_{estimators} = 200 and max depth = 10
Log Loss: 1.1806222819028016
for n_{estimators} = 500 and max depth = 5
Log Loss: 1.2328980304958208
for n_{estimators} = 500 and max depth = 10
Log Loss : 1.176527882241779
for n_estimators = 1000 and max depth = 5
Log Loss: 1.2300760069362329
for n_{estimators} = 1000 and max depth = 10
Log Loss : 1.1760573969739745
for n_{estimators} = 2000 and max depth = 5
Log Loss: 1.2307665089414377
for n_estimators = 2000 and max depth = 10
Log Loss : 1.1785316711423004
For values of best estimator = 1000 The train log loss is: 0.6720057995286121
For values of best estimator = 1000 The cross validation log loss is: 1.1760573969739745
For values of best estimator = 1000 The test log loss is: 1.1625737611100833
```

4.5.2. Testing model with best hyper parameters (One Hot Encoding)

```
In [98]: # -----
         # default parameters
         # sklearn.ensemble.RandomForestClassifier(n_estimators=10, criterion='gini', max_depth=None, min_samples_split=2,
         # min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_impurity_decrease=0.
         # min_impurity_split=None, bootstrap=True, oob_score=False, n_jobs=1, random_state=None, verbose=0, warm_start=False,
         # class_weight=None)
         # Some of methods of RandomForestClassifier()
         # fit(X, y, [sample_weight])
                                       Fit the SVM model according to the given training data.
         \# predict(X) Perform classification on samples in X.
         # predict_proba (X)
                               Perform classification on samples in X.
         # some of attributes of RandomForestClassifier()
         # feature_importances_ : array of shape = [n_features]
         # The feature importances (the higher, the more important the feature).
         # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/random-forest-and-their-construc
         tion-2/
         clf = RandomForestClassifier(n_estimators=alpha[int(best_alpha/2)], criterion='gini', max_depth=max_depth[int(best_alpha/2)]
         ha%2)], random_state=42, n_jobs=-1)
         predict_and_plot_confusion_matrix(train_x_onehotCoding, train_y,cv_x_onehotCoding,cv_y, clf)
```

----- Confusion matrix -----37.000 8.000 0.000 35.000 2.000 3.000 6.000 0.000 0.000 125 0.000 39.000 0.000 6.000 0.000 1.000 26.000 0.000 0.000 - 100 4.000 2.000 0.000 1.000 0.000 0.000 7.000 0.000 0.000 11.000 3.000 0.000 3.000 1.000 20.000 0.000 0.000 Original Class - 75 5.000 4.000 0.000 9.000 6.000 6.000 9.000 0.000 0.000 6.000 2.000 0.000 8.000 1.000 25.000 2.000 0.000 0.000 - 50 1.000 18.000 0.000 1.000 0.000 0.000 133.000 0.000 0.000 - 25 0.000 1.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 4.000 0.000 2.000 Predicted Class ----- Precision matrix (Columm Sum=1) -----1.0 0.105 0.259 0.143 0.083 0.029 0.000 0.000 0.000 0.000 0.044 0.028 0.125 0.000 0.000 - 0.8 0.000 0.030 0.143 0.000 0.034 0.013 0.000 0.000 0.183 0.039 0.214 0.028 0.096 0.000 0.000 - 0.6 Original Class 0.083 0.053 0.067 0.429 0.167 0.043 0.000 0.000 - 0.4 0.026 0.100 0.059 0.071 0.010 0.000 0.000 0.017 0.237 0.007 0.000 0.000 0.000 0.000 - 0.2 0.000 0.013 0.000 0.000 0.000 0.005 1.000 0.000 0.000 0.000 0.000 0.000 0.000 0.019 0.000 1.000 - 0.0 9 Predicted Class ----- Recall matrix (Row sum=1) ------0.407 0.088 0.000 0.385 0.022 0.033 0.066 0.000 0.000 - 0.75 0.000 0.083 0.361 0.000 0.000 0.014 0.000 0.000 0.000 0.286 0.143 0.000 0.071 0.000 0.000 0.000 - 0.60 0.027 0.000 0.027 0.100 0.009 0.182 0.000 0.000 - 0.45 0.103 0.000 0.231 0.154 0.154 0.231 0.000 0.136 0.045 0.000 0.182 0.023 0.045 0.000 0.000 0.007 0.118 0.000 0.007 0.000 0.000 0.869 0.000 0.000 0.333 0.333 0.333 -0.15 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.333 - 0.00 7 9

Predicted Class

4.5.3. Feature Importance

4.5.3.1. Correctly Classified point

```
In [100]: # test_point_index = 10
          clf = RandomForestClassifier(n_estimators=alpha[int(best_alpha/2)], criterion='gini', max_depth=max_depth[int(best_alpha/2)]
          ha%2)], random_state=42, n_jobs=-1)
          clf.fit(train_x_onehotCoding, train_y)
          sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
          sig_clf.fit(train_x_onehotCoding, train_y)
          test_point_index = 12
          no_feature = 1000
          predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
          print("Predicted Class :", predicted_cls[0])
          print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_onehotCoding[test_point_index]),4))
          print("Actual Class :", test_y[test_point_index])
          indices = np.argsort(-clf.feature_importances_)
          print("-"*50)
          get_impfeature_names(indices[:no_feature],
                               x_test['TEXT'].iloc[test_point_index],
                               x_test['Gene'].iloc[test_point_index],
                               x_test['Variation'].iloc[test_point_index],
                               no_feature)
```

```
Predicted Class : 4
Predicted Class Probabilities: [[0.1259 0.0881 0.0259 0.4699 0.096 0.0665 0.1134 0.0067 0.0076]]
0 Text feature [kinase] present in test data point [True]
2 Text feature [phosphorylation] present in test data point [True]
3 Text feature [activation] present in test data point [True]
4 Text feature [inhibitors] present in test data point [True]
5 Text feature [tyrosine] present in test data point [True]
6 Text feature [signaling] present in test data point [True]
7 Text feature [suppressor] present in test data point [True]
8 Text feature [function] present in test data point [True]
9 Text feature [activated] present in test data point [True]
10 Text feature [oncogenic] present in test data point [True]
14 Text feature [loss] present in test data point [True]
15 Text feature [growth] present in test data point [True]
16 Text feature [treated] present in test data point [True]
24 Text feature [drug] present in test data point [True]
25 Text feature [activate] present in test data point [True]
27 Text feature [missense] present in test data point [True]
28 Text feature [downstream] present in test data point [True]
34 Text feature [receptor] present in test data point [True]
36 Text feature [protein] present in test data point [True]
38 Text feature [oncogene] present in test data point [True]
40 Text feature [cells] present in test data point [True]
42 Text feature [variants] present in test data point [True]
47 Text feature [egfr] present in test data point [True]
51 Text feature [kinases] present in test data point [True]
54 Text feature [functional] present in test data point [True]
55 Text feature [effective] present in test data point [True]
60 Text feature [stability] present in test data point [True]
63 Text feature [patients] present in test data point [True]
73 Text feature [potential] present in test data point [True]
75 Text feature [classified] present in test data point [True]
77 Text feature [lines] present in test data point [True]
80 Text feature [cell] present in test data point [True]
81 Text feature [neutral] present in test data point [True]
84 Text feature [predicted] present in test data point [True]
86 Text feature [active] present in test data point [True]
90 Text feature [patient] present in test data point [True]
93 Text feature [dna] present in test data point [True]
103 Text feature [functions] present in test data point [True]
104 Text feature [progression] present in test data point [True]
112 Text feature [conserved] present in test data point [True]
116 Text feature [proteins] present in test data point [True]
117 Text feature [starved] present in test data point [True]
118 Text feature [tumors] present in test data point [True]
123 Text feature [activity] present in test data point [True]
130 Text feature [likely] present in test data point [True]
132 Text feature [independent] present in test data point [True]
133 Text feature [drugs] present in test data point [True]
135 Text feature [kit] present in test data point [True]
136 Text feature [affected] present in test data point [True]
142 Text feature [retained] present in test data point [True]
144 Text feature [use] present in test data point [True]
148 Text feature [expression] present in test data point [True]
151 Text feature [ring] present in test data point [True]
154 Text feature [assays] present in test data point [True]
159 Text feature [therapies] present in test data point [True]
164 Text feature [phosphorylated] present in test data point [True]
165 Text feature [concentrations] present in test data point [True]
174 Text feature [disrupt] present in test data point [True]
176 Text feature [enhanced] present in test data point [True]
183 Text feature [database] present in test data point [True]
189 Text feature [3b] present in test data point [True]
193 Text feature [lung] present in test data point [True]
198 Text feature [2b] present in test data point [True]
207 Text feature [oncogenes] present in test data point [True]
208 Text feature [transfected] present in test data point [True]
211 Text feature [predictions] present in test data point [True]
213 Text feature [structure] present in test data point [True]
214 Text feature [presence] present in test data point [True]
219 Text feature [affect] present in test data point [True]
222 Text feature [tagged] present in test data point [True]
223 Text feature [mechanism] present in test data point [True]
224 Text feature [small] present in test data point [True]
226 Text feature [hcl] present in test data point [True]
227 Text feature [information] present in test data point [True]
229 Text feature [model] present in test data point [True]
232 Text feature [interaction] present in test data point [True]
235 Text feature [well] present in test data point [True]
236 Text feature [assay] present in test data point [True]
238 Text feature [experiments] present in test data point [True]
241 Text feature [il] present in test data point [True]
244 Text feature [type] present in test data point [True]
245 Text feature [leukemia] present in test data point [True]
248 Text feature [family] present in test data point [True]
251 Text feature [effect] present in test data point [True]
```

```
254 Text feature [effects] present in test data point [True]
256 Text feature [binding] present in test data point [True]
258 Text feature [mutant] present in test data point [True]
259 Text feature [genes] present in test data point [True]
260 Text feature [results] present in test data point [True]
261 Text feature [3a] present in test data point [True]
263 Text feature [study] present in test data point [True]
266 Text feature [introduction] present in test data point [True]
267 Text feature [expressed] present in test data point [True]
268 Text feature [contrast] present in test data point [True]
274 Text feature [mutants] present in test data point [True]
277 Text feature [19] present in test data point [True]
281 Text feature [amino] present in test data point [True]
282 Text feature [21] present in test data point [True]
283 Text feature [developed] present in test data point [True]
290 Text feature [27] present in test data point [True]
291 Text feature [based] present in test data point [True]
292 Text feature [case] present in test data point [True]
293 Text feature [residues] present in test data point [True]
294 Text feature [structural] present in test data point [True]
296 Text feature [probability] present in test data point [True]
298 Text feature [specific] present in test data point [True]
299 Text feature [evolutionary] present in test data point [True]
300 Text feature [used] present in test data point [True]
303 Text feature [known] present in test data point [True]
304 Text feature [constructs] present in test data point [True]
307 Text feature [majority] present in test data point [True]
313 Text feature [substitution] present in test data point [True]
314 Text feature [ha] present in test data point [True]
316 Text feature [identified] present in test data point [True]
317 Text feature [defined] present in test data point [True]
318 Text feature [available] present in test data point [True]
321 Text feature [mutation] present in test data point [True]
325 Text feature [bind] present in test data point [True]
326 Text feature [showed] present in test data point [True]
329 Text feature [sequence] present in test data point [True]
331 Text feature [significantly] present in test data point [True]
332 Text feature [rearrangements] present in test data point [True]
334 Text feature [large] present in test data point [True]
336 Text feature [figure] present in test data point [True]
337 Text feature [shown] present in test data point [True]
339 Text feature [molecular] present in test data point [True]
341 Text feature [01] present in test data point [True]
342 Text feature [system] present in test data point [True]
343 Text feature [detected] present in test data point [True]
345 Text feature [recently] present in test data point [True]
346 Text feature [levels] present in test data point [True]
347 Text feature [ability] present in test data point [True]
349 Text feature [positive] present in test data point [True]
350 Text feature [11] present in test data point [True]
352 Text feature [sequencing] present in test data point [True]
356 Text feature [line] present in test data point [True]
357 Text feature [impact] present in test data point [True]
358 Text feature [gene] present in test data point [True]
359 Text feature [domain] present in test data point [True]
360 Text feature [induced] present in test data point [True]
363 Text feature [tumor] present in test data point [True]
365 Text feature [received] present in test data point [True]
366 Text feature [43] present in test data point [True]
367 Text feature [approach] present in test data point [True]
371 Text feature [mm] present in test data point [True]
372 Text feature [wild] present in test data point [True]
376 Text feature [data] present in test data point [True]
377 Text feature [pathways] present in test data point [True]
379 Text feature [compared] present in test data point [True]
382 Text feature [targeting] present in test data point [True]
383 Text feature [times] present in test data point [True]
384 Text feature [multiple] present in test data point [True]
385 Text feature [similar] present in test data point [True]
388 Text feature [another] present in test data point [True]
392 Text feature [complex] present in test data point [True]
393 Text feature [directed] present in test data point [True]
396 Text feature [least] present in test data point [True]
398 Text feature [anti] present in test data point [True]
399 Text feature [within] present in test data point [True]
400 Text feature [determined] present in test data point [True]
402 Text feature [although] present in test data point [True]
403 Text feature [37] present in test data point [True]
405 Text feature [hours] present in test data point [True]
412 Text feature [18] present in test data point [True]
416 Text feature [observed] present in test data point [True]
417 Text feature [rather] present in test data point [True]
418 Text feature [higher] present in test data point [True]
420 Text feature [important] present in test data point [True]
422 Text feature [mapping] present in test data point [True]
424 Text feature [17] present in test data point [True]
426 Text feature [12] present in test data point [True]
428 Text feature [containing] present in test data point [True]
```

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431 Text feature [acids] present in test data point [True]
432 Text feature [studies] present in test data point [True]
433 Text feature [previously] present in test data point [True]
434 Text feature [time] present in test data point [True]
436 Text feature [catalytic] present in test data point [True]
438 Text feature [studied] present in test data point [True]
439 Text feature [would] present in test data point [True]
440 Text feature [could] present in test data point [True]
444 Text feature [methods] present in test data point [True]
445 Text feature [targeted] present in test data point [True]
446 Text feature [e3] present in test data point [True]
447 Text feature [substitutions] present in test data point [True]
451 Text feature [four] present in test data point [True]
453 Text feature [reported] present in test data point [True]
455 Text feature [obtained] present in test data point [True]
457 Text feature [cancer] present in test data point [True]
458 Text feature [various] present in test data point [True]
460 Text feature [transfection] present in test data point [True]
462 Text feature [vector] present in test data point [True]
463 Text feature [whether] present in test data point [True]
464 Text feature [findings] present in test data point [True]
470 Text feature [heterozygosity] present in test data point [True]
471 Text feature [signal] present in test data point [True]
482 Text feature [less] present in test data point [True]
484 Text feature [interactions] present in test data point [True]
485 Text feature [mg] present in test data point [True]
486 Text feature [20] present in test data point [True]
489 Text feature [16] present in test data point [True]
490 Text feature [increase] present in test data point [True]
491 Text feature [detection] present in test data point [True]
492 Text feature [vivo] present in test data point [True]
494 Text feature [confer] present in test data point [True]
502 Text feature [47] present in test data point [True]
503 Text feature [confirmed] present in test data point [True]
507 Text feature [found] present in test data point [True]
511 Text feature [somatic] present in test data point [True]
512 Text feature [regulatory] present in test data point [True]
514 Text feature [culture] present in test data point [True]
515 Text feature [consistent] present in test data point [True]
516 Text feature [13] present in test data point [True]
517 Text feature [fold] present in test data point [True]
518 Text feature [table] present in test data point [True]
522 Text feature [32] present in test data point [True]
523 Text feature [included] present in test data point [True]
525 Text feature [antibodies] present in test data point [True]
526 Text feature [39] present in test data point [True]
527 Text feature [combined] present in test data point [True]
528 Text feature [residue] present in test data point [True]
529 Text feature [human] present in test data point [True]
533 Text feature [changes] present in test data point [True]
534 Text feature [page] present in test data point [True]
535 Text feature [cancers] present in test data point [True]
537 Text feature [general] present in test data point [True]
541 Text feature [whereas] present in test data point [True]
543 Text feature [s1] present in test data point [True]
544 Text feature [region] present in test data point [True]
546 Text feature [repeat] present in test data point [True]
548 Text feature [samples] present in test data point [True]
551 Text feature [applied] present in test data point [True]
552 Text feature [involved] present in test data point [True]
554 Text feature [relative] present in test data point [True]
560 Text feature [2a] present in test data point [True]
561 Text feature [resulting] present in test data point [True]
563 Text feature [destabilized] present in test data point [True]
564 Text feature [analysis] present in test data point [True]
566 Text feature [final] present in test data point [True]
570 Text feature [associated] present in test data point [True]
571 Text feature [several] present in test data point [True]
573 Text feature [tested] present in test data point [True]
575 Text feature [addition] present in test data point [True]
576 Text feature [transduction] present in test data point [True]
577 Text feature [significant] present in test data point [True]
580 Text feature [method] present in test data point [True]
581 Text feature [form] present in test data point [True]
583 Text feature [bridge] present in test data point [True]
586 Text feature [either] present in test data point [True]
588 Text feature [ubiquitin] present in test data point [True]
591 Text feature [selection] present in test data point [True]
592 Text feature [80] present in test data point [True]
594 Text feature [additional] present in test data point [True]
596 Text feature [also] present in test data point [True]
597 Text feature [lesser] present in test data point [True]
602 Text feature [site] present in test data point [True]
603 Text feature [gain] present in test data point [True]
606 Text feature [interest] present in test data point [True]
609 Text feature [fully] present in test data point [True]
610 Text feature [previous] present in test data point [True]
614 Text feature [three] present in test data point [True]
```

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617 Text feature [mean] present in test data point [True]
619 Text feature [conformation] present in test data point [True]
625 Text feature [first] present in test data point [True]
627 Text feature [15] present in test data point [True]
628 Text feature [significance] present in test data point [True]
631 Text feature [five] present in test data point [True]
634 Text feature [increased] present in test data point [True]
638 Text feature [parallel] present in test data point [True]
639 Text feature [value] present in test data point [True]
640 Text feature [ligase] present in test data point [True]
641 Text feature [group] present in test data point [True]
642 Text feature [mechanisms] present in test data point [True]
643 Text feature [concentration] present in test data point [True]
648 Text feature [values] present in test data point [True]
649 Text feature [non] present in test data point [True]
650 Text feature [41] present in test data point [True]
651 Text feature [assessed] present in test data point [True]
652 Text feature [detailed] present in test data point [True]
654 Text feature [single] present in test data point [True]
655 Text feature [stimulated] present in test data point [True]
657 Text feature [manufacturer] present in test data point [True]
658 Text feature [analyzed] present in test data point [True]
661 Text feature [led] present in test data point [True]
662 Text feature [23] present in test data point [True]
664 Text feature [inactive] present in test data point [True]
665 Text feature [complexes] present in test data point [True]
666 Text feature [lost] present in test data point [True]
669 Text feature [related] present in test data point [True]
670 Text feature [example] present in test data point [True]
671 Text feature [control] present in test data point [True]
674 Text feature [however] present in test data point [True]
676 Text feature [comparison] present in test data point [True]
678 Text feature [examined] present in test data point [True]
681 Text feature [49] present in test data point [True]
687 Text feature [report] present in test data point [True]
688 Text feature [present] present in test data point [True]
696 Text feature [certain] present in test data point [True]
697 Text feature [possible] present in test data point [True]
700 Text feature [membranes] present in test data point [True]
701 Text feature [mutations] present in test data point [True]
703 Text feature [tris] present in test data point [True]
704 Text feature [substrate] present in test data point [True]
710 Text feature [especially] present in test data point [True]
711 Text feature [24] present in test data point [True]
712 Text feature [syndrome] present in test data point [True]
713 Text feature [years] present in test data point [True]
714 Text feature [pdb] present in test data point [True]
717 Text feature [two] present in test data point [True]
719 Text feature [33] present in test data point [True]
721 Text feature [cases] present in test data point [True]
725 Text feature [28] present in test data point [True]
728 Text feature [incubated] present in test data point [True]
732 Text feature [negative] present in test data point [True]
733 Text feature [mutagenesis] present in test data point [True]
734 Text feature [show] present in test data point [True]
735 Text feature [change] present in test data point [True]
736 Text feature [discussion] present in test data point [True]
737 Text feature [helix] present in test data point [True]
738 Text feature [sites] present in test data point [True]
739 Text feature [may] present in test data point [True]
741 Text feature [many] present in test data point [True]
742 Text feature [area] present in test data point [True]
743 Text feature [using] present in test data point [True]
744 Text feature [distribution] present in test data point [True]
748 Text feature [specificity] present in test data point [True]
750 Text feature [disease] present in test data point [True]
751 Text feature [alter] present in test data point [True]
752 Text feature [even] present in test data point [True]
753 Text feature [30] present in test data point [True]
755 Text feature [finally] present in test data point [True]
756 Text feature [correlation] present in test data point [True]
758 Text feature [strong] present in test data point [True]
759 Text feature [allele] present in test data point [True]
761 Text feature [plasmid] present in test data point [True]
762 Text feature [unknown] present in test data point [True]
765 Text feature [critical] present in test data point [True]
769 Text feature [high] present in test data point [True]
770 Text feature [egf] present in test data point [True]
772 Text feature [agreement] present in test data point [True]
773 Text feature [located] present in test data point [True]
777 Text feature [10] present in test data point [True]
778 Text feature [homozygous] present in test data point [True]
780 Text feature [acid] present in test data point [True]
781 Text feature [one] present in test data point [True]
783 Text feature [types] present in test data point [True]
786 Text feature [inactivate] present in test data point [True]
789 Text feature [26] present in test data point [True]
790 Text feature [required] present in test data point [True]
```

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794 Text feature [statistical] present in test data point [True]
796 Text feature [underlying] present in test data point [True]
799 Text feature [despite] present in test data point [True]
806 Text feature [42] present in test data point [True]
809 Text feature [22] present in test data point [True]
810 Text feature [respectively] present in test data point [True]
814 Text feature [test] present in test data point [True]
817 Text feature [relatively] present in test data point [True]
823 Text feature [development] present in test data point [True]
828 Text feature [see] present in test data point [True]
836 Text feature [become] present in test data point [True]
839 Text feature [nucleotide] present in test data point [True]
844 Text feature [intermediate] present in test data point [True]
845 Text feature [25] present in test data point [True]
852 Text feature [35] present in test data point [True]
856 Text feature [nacl] present in test data point [True]
860 Text feature [performed] present in test data point [True]
866 Text feature [plasmids] present in test data point [True]
876 Text feature [full] present in test data point [True]
878 Text feature [identify] present in test data point [True]
881 Text feature [e2] present in test data point [True]
885 Text feature [fig] present in test data point [True]
886 Text feature [potentially] present in test data point [True]
887 Text feature [features] present in test data point [True]
888 Text feature [strongly] present in test data point [True]
889 Text feature [limited] present in test data point [True]
890 Text feature [sequences] present in test data point [True]
897 Text feature [predict] present in test data point [True]
900 Text feature [population] present in test data point [True]
901 Text feature [noted] present in test data point [True]
902 Text feature [mutated] present in test data point [True]
905 Text feature [promega] present in test data point [True]
909 Text feature [new] present in test data point [True]
910 Text feature [100] present in test data point [True]
911 Text feature [peptide] present in test data point [True]
914 Text feature [rtk] present in test data point [True]
915 Text feature [set] present in test data point [True]
919 Text feature [define] present in test data point [True]
921 Text feature [assess] present in test data point [True]
922 Text feature [result] present in test data point [True]
923 Text feature [thus] present in test data point [True]
926 Text feature [29] present in test data point [True]
927 Text feature [interestingly] present in test data point [True]
936 Text feature [shows] present in test data point [True]
937 Text feature [score] present in test data point [True]
939 Text feature [prior] present in test data point [True]
940 Text feature [due] present in test data point [True]
941 Text feature [approaches] present in test data point [True]
942 Text feature [salt] present in test data point [True]
943 Text feature [partially] present in test data point [True]
944 Text feature [complete] present in test data point [True]
945 Text feature [suppressors] present in test data point [True]
946 Text feature [reduced] present in test data point [True]
949 Text feature [formation] present in test data point [True]
950 Text feature [specifically] present in test data point [True]
952 Text feature [resulted] present in test data point [True]
956 Text feature [frequency] present in test data point [True]
957 Text feature [calculated] present in test data point [True]
958 Text feature [34] present in test data point [True]
959 Text feature [highly] present in test data point [True]
962 Text feature [46] present in test data point [True]
963 Text feature [respect] present in test data point [True]
964 Text feature [selective] present in test data point [True]
965 Text feature [second] present in test data point [True]
968 Text feature [underwent] present in test data point [True]
973 Text feature [characterized] present in test data point [True]
982 Text feature [buffer] present in test data point [True]
988 Text feature [might] present in test data point [True]
989 Text feature [sigma] present in test data point [True]
990 Text feature [whole] present in test data point [True]
991 Text feature [referred] present in test data point [True]
994 Text feature [added] present in test data point [True]
999 Text feature [status] present in test data point [True]
Out of the top 1000 features 421 are present in query point
```

```
Predicted Class : 1
Predicted Class Probabilities: [[0.3221 0.0448 0.0253 0.2688 0.1451 0.1407 0.0411 0.0059 0.0061]]
0 Text feature [kinase] present in test data point [True]
3 Text feature [activation] present in test data point [True]
7 Text feature [suppressor] present in test data point [True]
8 Text feature [function] present in test data point [True]
9 Text feature [activated] present in test data point [True]
12 Text feature [constitutive] present in test data point [True]
14 Text feature [loss] present in test data point [True]
15 Text feature [growth] present in test data point [True]
19 Text feature [nonsense] present in test data point [True]
25 Text feature [activate] present in test data point [True]
26 Text feature [brca1] present in test data point [True]
27 Text feature [missense] present in test data point [True]
36 Text feature [protein] present in test data point [True]
40 Text feature [cells] present in test data point [True]
42 Text feature [variants] present in test data point [True]
43 Text feature [yeast] present in test data point [True]
44 Text feature [clinical] present in test data point [True]
54 Text feature [functional] present in test data point [True]
57 Text feature [deleterious] present in test data point [True]
60 Text feature [stability] present in test data point [True]
62 Text feature [f3] present in test data point [True]
63 Text feature [patients] present in test data point [True]
73 Text feature [potential] present in test data point [True]
74 Text feature [factor] present in test data point [True]
75 Text feature [classified] present in test data point [True]
76 Text feature [frameshift] present in test data point [True]
77 Text feature [lines] present in test data point [True]
79 Text feature [unclassified] present in test data point [True]
80 Text feature [cell] present in test data point [True]
81 Text feature [neutral] present in test data point [True]
84 Text feature [predicted] present in test data point [True]
87 Text feature [expressing] present in test data point [True]
88 Text feature [brca2] present in test data point [True]
91 Text feature [variant] present in test data point [True]
93 Text feature [dna] present in test data point [True]
97 Text feature [carriers] present in test data point [True]
99 Text feature [likelihood] present in test data point [True]
103 Text feature [functions] present in test data point [True]
106 Text feature [sensitivity] present in test data point [True]
108 Text feature [repair] present in test data point [True]
112 Text feature [conserved] present in test data point [True]
115 Text feature [core] present in test data point [True]
116 Text feature [proteins] present in test data point [True]
120 Text feature [mammalian] present in test data point [True]
123 Text feature [activity] present in test data point [True]
125 Text feature [inactivation] present in test data point [True]
127 Text feature [sensitive] present in test data point [True]
130 Text feature [likely] present in test data point [True]
131 Text feature [transform] present in test data point [True]
132 Text feature [independent] present in test data point [True]
136 Text feature [affected] present in test data point [True]
137 Text feature [truncating] present in test data point [True]
141 Text feature [damage] present in test data point [True]
144 Text feature [use] present in test data point [True]
145 Text feature [classify] present in test data point [True]
147 Text feature [neutrality] present in test data point [True]
148 Text feature [expression] present in test data point [True]
150 Text feature [brct] present in test data point [True]
154 Text feature [assays] present in test data point [True]
156 Text feature [nuclear] present in test data point [True]
168 Text feature [terminal] present in test data point [True]
171 Text feature [conservation] present in test data point [True]
172 Text feature [classification] present in test data point [True]
173 Text feature [bic] present in test data point [True]
174 Text feature [disrupt] present in test data point [True]
175 Text feature [ovarian] present in test data point [True]
178 Text feature [odds] present in test data point [True]
183 Text feature [database] present in test data point [True]
189 Text feature [3b] present in test data point [True]
190 Text feature [putative] present in test data point [True]
191 Text feature [favor] present in test data point [True]
196 Text feature [79] present in test data point [True]
198 Text feature [2b] present in test data point [True]
202 Text feature [individuals] present in test data point [True]
208 Text feature [transfected] present in test data point [True]
211 Text feature [predictions] present in test data point [True]
213 Text feature [structure] present in test data point [True]
214 Text feature [presence] present in test data point [True]
216 Text feature [reporter] present in test data point [True]
219 Text feature [affect] present in test data point [True]
221 Text feature [absence] present in test data point [True]
224 Text feature [small] present in test data point [True]
227 Text feature [information] present in test data point [True]
228 Text feature [1863] present in test data point [True]
```

```
229 Text feature [model] present in test data point [True]
232 Text feature [interaction] present in test data point [True]
234 Text feature [predisposition] present in test data point [True]
235 Text feature [well] present in test data point [True]
236 Text feature [assay] present in test data point [True]
238 Text feature [experiments] present in test data point [True]
242 Text feature [folding] present in test data point [True]
244 Text feature [type] present in test data point [True]
247 Text feature [68] present in test data point [True]
248 Text feature [family] present in test data point [True]
249 Text feature [pedigree] present in test data point [True]
251 Text feature [effect] present in test data point [True]
256 Text feature [binding] present in test data point [True]
257 Text feature [role] present in test data point [True]
258 Text feature [mutant] present in test data point [True]
259 Text feature [genes] present in test data point [True]
260 Text feature [results] present in test data point [True]
261 Text feature [3a] present in test data point [True]
263 Text feature [study] present in test data point [True]
266 Text feature [introduction] present in test data point [True]
267 Text feature [expressed] present in test data point [True]
274 Text feature [mutants] present in test data point [True]
275 Text feature [expected] present in test data point [True]
277 Text feature [19] present in test data point [True]
281 Text feature [amino] present in test data point [True]
282 Text feature [21] present in test data point [True]
284 Text feature [currently] present in test data point [True]
286 Text feature [deletion] present in test data point [True]
287 Text feature [splicing] present in test data point [True]
289 Text feature [multifactorial] present in test data point [True]
290 Text feature [27] present in test data point [True]
291 Text feature [based] present in test data point [True]
292 Text feature [case] present in test data point [True]
293 Text feature [residues] present in test data point [True]
294 Text feature [structural] present in test data point [True]
298 Text feature [specific] present in test data point [True]
299 Text feature [evolutionary] present in test data point [True]
300 Text feature [used] present in test data point [True]
303 Text feature [known] present in test data point [True]
304 Text feature [constructs] present in test data point [True]
305 Text feature [controls] present in test data point [True]
308 Text feature [occurrence] present in test data point [True]
311 Text feature [breast] present in test data point [True]
313 Text feature [substitution] present in test data point [True]
316 Text feature [identified] present in test data point [True]
317 Text feature [defined] present in test data point [True]
318 Text feature [available] present in test data point [True]
319 Text feature [transcription] present in test data point [True]
320 Text feature [published] present in test data point [True]
321 Text feature [mutation] present in test data point [True]
322 Text feature [nhgri] present in test data point [True]
325 Text feature [bind] present in test data point [True]
326 Text feature [showed] present in test data point [True]
327 Text feature [50] present in test data point [True]
329 Text feature [sequence] present in test data point [True]
330 Text feature [co] present in test data point [True]
331 Text feature [significantly] present in test data point [True]
334 Text feature [large] present in test data point [True]
336 Text feature [figure] present in test data point [True]
337 Text feature [shown] present in test data point [True]
342 Text feature [system] present in test data point [True]
343 Text feature [detected] present in test data point [True]
345 Text feature [recently] present in test data point [True]
346 Text feature [levels] present in test data point [True]
347 Text feature [ability] present in test data point [True]
349 Text feature [positive] present in test data point [True]
350 Text feature [11] present in test data point [True]
352 Text feature [sequencing] present in test data point [True]
353 Text feature [personal] present in test data point [True]
354 Text feature [western] present in test data point [True]
356 Text feature [line] present in test data point [True]
357 Text feature [impact] present in test data point [True]
358 Text feature [gene] present in test data point [True]
359 Text feature [domain] present in test data point [True]
363 Text feature [tumor] present in test data point [True]
366 Text feature [43] present in test data point [True]
367 Text feature [approach] present in test data point [True]
372 Text feature [wild] present in test data point [True]
376 Text feature [data] present in test data point [True]
378 Text feature [risk] present in test data point [True]
379 Text feature [compared] present in test data point [True]
380 Text feature [promoter] present in test data point [True]
384 Text feature [multiple] present in test data point [True]
385 Text feature [similar] present in test data point [True]
387 Text feature [research] present in test data point [True]
396 Text feature [least] present in test data point [True]
397 Text feature [outside] present in test data point [True]
400 Text feature [determined] present in test data point [True]
```

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402 Text feature [although] present in test data point [True]
403 Text feature [37] present in test data point [True]
404 Text feature [stress] present in test data point [True]
410 Text feature [determine] present in test data point [True]
412 Text feature [18] present in test data point [True]
414 Text feature [number] present in test data point [True]
417 Text feature [rather] present in test data point [True]
418 Text feature [higher] present in test data point [True]
420 Text feature [important] present in test data point [True]
421 Text feature [53] present in test data point [True]
424 Text feature [17] present in test data point [True]
425 Text feature [myriad] present in test data point [True]
426 Text feature [12] present in test data point [True]
427 Text feature [measured] present in test data point [True]
428 Text feature [containing] present in test data point [True]
431 Text feature [acids] present in test data point [True]
432 Text feature [studies] present in test data point [True]
433 Text feature [previously] present in test data point [True]
438 Text feature [studied] present in test data point [True]
440 Text feature [could] present in test data point [True]
442 Text feature [wt] present in test data point [True]
444 Text feature [methods] present in test data point [True]
447 Text feature [substitutions] present in test data point [True]
450 Text feature [leading] present in test data point [True]
451 Text feature [four] present in test data point [True]
452 Text feature [predictive] present in test data point [True]
453 Text feature [reported] present in test data point [True]
455 Text feature [obtained] present in test data point [True]
457 Text feature [cancer] present in test data point [True]
459 Text feature [vitro] present in test data point [True]
460 Text feature [transfection] present in test data point [True]
462 Text feature [vector] present in test data point [True]
463 Text feature [whether] present in test data point [True]
469 Text feature [uncertain] present in test data point [True]
470 Text feature [heterozygosity] present in test data point [True]
472 Text feature [transcriptional] present in test data point [True]
478 Text feature [http] present in test data point [True]
481 Text feature [orthologs] present in test data point [True]
482 Text feature [less] present in test data point [True]
483 Text feature [basis] present in test data point [True]
484 Text feature [interactions] present in test data point [True]
486 Text feature [20] present in test data point [True]
489 Text feature [16] present in test data point [True]
490 Text feature [increase] present in test data point [True]
492 Text feature [vivo] present in test data point [True]
494 Text feature [confer] present in test data point [True]
495 Text feature [medium] present in test data point [True]
497 Text feature [occur] present in test data point [True]
499 Text feature [90] present in test data point [True]
500 Text feature [unable] present in test data point [True]
501 Text feature [including] present in test data point [True]
502 Text feature [47] present in test data point [True]
503 Text feature [confirmed] present in test data point [True]
504 Text feature [predispose] present in test data point [True]
505 Text feature [conservative] present in test data point [True]
507 Text feature [found] present in test data point [True]
510 Text feature [counseling] present in test data point [True]
513 Text feature [genetic] present in test data point [True]
515 Text feature [consistent] present in test data point [True]
516 Text feature [13] present in test data point [True]
517 Text feature [fold] present in test data point [True]
518 Text feature [table] present in test data point [True]
519 Text feature [impaired] present in test data point [True]
520 Text feature [pocket] present in test data point [True]
522 Text feature [32] present in test data point [True]
523 Text feature [included] present in test data point [True]
525 Text feature [antibodies] present in test data point [True]
526 Text feature [39] present in test data point [True]
527 Text feature [combined] present in test data point [True]
528 Text feature [residue] present in test data point [True]
529 Text feature [human] present in test data point [True]
531 Text feature [contribute] present in test data point [True]
533 Text feature [changes] present in test data point [True]
537 Text feature [general] present in test data point [True]
540 Text feature [together] present in test data point [True]
541 Text feature [whereas] present in test data point [True]
542 Text feature [integrity] present in test data point [True]
544 Text feature [region] present in test data point [True]
548 Text feature [samples] present in test data point [True]
551 Text feature [applied] present in test data point [True]
555 Text feature [cause] present in test data point [True]
560 Text feature [2a] present in test data point [True]
561 Text feature [resulting] present in test data point [True]
564 Text feature [analysis] present in test data point [True]
568 Text feature [dependent] present in test data point [True]
570 Text feature [associated] present in test data point [True]
573 Text feature [tested] present in test data point [True]
575 Text feature [addition] present in test data point [True]
```

```
577 Text feature [significant] present in test data point [True]
579 Text feature [novel] present in test data point [True]
580 Text feature [method] present in test data point [True]
584 Text feature [prediction] present in test data point [True]
585 Text feature [suppression] present in test data point [True]
586 Text feature [either] present in test data point [True]
592 Text feature [80] present in test data point [True]
594 Text feature [additional] present in test data point [True]
596 Text feature [also] present in test data point [True]
600 Text feature [antibody] present in test data point [True]
602 Text feature [site] present in test data point [True]
604 Text feature [et] present in test data point [True]
605 Text feature [evaluated] present in test data point [True]
610 Text feature [previous] present in test data point [True]
611 Text feature [susceptibility] present in test data point [True]
613 Text feature [exon] present in test data point [True]
614 Text feature [three] present in test data point [True]
621 Text feature [prophylactic] present in test data point [True]
625 Text feature [first] present in test data point [True]
626 Text feature [contains] present in test data point [True]
627 Text feature [15] present in test data point [True]
628 Text feature [significance] present in test data point [True]
630 Text feature [suggests] present in test data point [True]
631 Text feature [five] present in test data point [True]
634 Text feature [increased] present in test data point [True]
636 Text feature [surface] present in test data point [True]
638 Text feature [parallel] present in test data point [True]
649 Text feature [non] present in test data point [True]
650 Text feature [41] present in test data point [True]
651 Text feature [assessed] present in test data point [True]
652 Text feature [detailed] present in test data point [True]
654 Text feature [single] present in test data point [True]
658 Text feature [analyzed] present in test data point [True]
660 Text feature [basal] present in test data point [True]
662 Text feature [23] present in test data point [True]
663 Text feature [remaining] present in test data point [True]
669 Text feature [related] present in test data point [True]
670 Text feature [example] present in test data point [True]
671 Text feature [control] present in test data point [True]
672 Text feature [evaluate] present in test data point [True]
674 Text feature [however] present in test data point [True]
675 Text feature [polymorphisms] present in test data point [True]
678 Text feature [examined] present in test data point [True]
679 Text feature [need] present in test data point [True]
681 Text feature [49] present in test data point [True]
682 Text feature [amplified] present in test data point [True]
684 Text feature [domains] present in test data point [True]
686 Text feature [al] present in test data point [True]
688 Text feature [present] present in test data point [True]
690 Text feature [revealed] present in test data point [True]
696 Text feature [certain] present in test data point [True]
697 Text feature [possible] present in test data point [True]
698 Text feature [1b] present in test data point [True]
699 Text feature [1a] present in test data point [True]
701 Text feature [mutations] present in test data point [True]
702 Text feature [repeats] present in test data point [True]
704 Text feature [substrate] present in test data point [True]
705 Text feature [lack] present in test data point [True]
706 Text feature [threshold] present in test data point [True]
711 Text feature [24] present in test data point [True]
717 Text feature [two] present in test data point [True]
719 Text feature [33] present in test data point [True]
721 Text feature [cases] present in test data point [True]
725 Text feature [28] present in test data point [True]
729 Text feature [3c] present in test data point [True]
732 Text feature [negative] present in test data point [True]
735 Text feature [change] present in test data point [True]
736 Text feature [discussion] present in test data point [True]
737 Text feature [helix] present in test data point [True]
738 Text feature [sites] present in test data point [True]
739 Text feature [may] present in test data point [True]
741 Text feature [many] present in test data point [True]
743 Text feature [using] present in test data point [True]
746 Text feature [longer] present in test data point [True]
747 Text feature [4b] present in test data point [True]
748 Text feature [specificity] present in test data point [True]
750 Text feature [disease] present in test data point [True]
752 Text feature [even] present in test data point [True]
753 Text feature [30] present in test data point [True]
755 Text feature [finally] present in test data point [True]
759 Text feature [allele] present in test data point [True]
761 Text feature [plasmid] present in test data point [True]
762 Text feature [unknown] present in test data point [True]
769 Text feature [high] present in test data point [True]
772 Text feature [agreement] present in test data point [True]
773 Text feature [located] present in test data point [True]
777 Text feature [10] present in test data point [True]
780 Text feature [acid] present in test data point [True]
```

```
781 Text feature [one] present in test data point [True]
782 Text feature [numbering] present in test data point [True]
783 Text feature [types] present in test data point [True]
786 Text feature [inactivate] present in test data point [True]
787 Text feature [v1736a] present in test data point [True]
788 Text feature [recent] present in test data point [True]
790 Text feature [required] present in test data point [True]
793 Text feature [fusion] present in test data point [True]
798 Text feature [bach1] present in test data point [True]
799 Text feature [despite] present in test data point [True]
802 Text feature [stable] present in test data point [True]
803 Text feature [week] present in test data point [True]
806 Text feature [42] present in test data point [True]
807 Text feature [evidence] present in test data point [True]
809 Text feature [22] present in test data point [True]
810 Text feature [respectively] present in test data point [True]
812 Text feature [causing] present in test data point [True]
813 Text feature [common] present in test data point [True]
814 Text feature [test] present in test data point [True]
816 Text feature [six] present in test data point [True]
817 Text feature [relatively] present in test data point [True]
818 Text feature [pcr] present in test data point [True]
822 Text feature [direct] present in test data point [True]
824 Text feature [suggest] present in test data point [True]
825 Text feature [differences] present in test data point [True]
826 Text feature [mrna] present in test data point [True]
828 Text feature [see] present in test data point [True]
833 Text feature [cannot] present in test data point [True]
835 Text feature [sd] present in test data point [True]
839 Text feature [nucleotide] present in test data point [True]
844 Text feature [intermediate] present in test data point [True]
845 Text feature [25] present in test data point [True]
851 Text feature [indicate] present in test data point [True]
852 Text feature [35] present in test data point [True]
853 Text feature [hydrophobic] present in test data point [True]
860 Text feature [performed] present in test data point [True]
861 Text feature [transcript] present in test data point [True]
864 Text feature [sources] present in test data point [True]
865 Text feature [comparable] present in test data point [True]
867 Text feature [corresponding] present in test data point [True]
874 Text feature [combination] present in test data point [True]
875 Text feature [measure] present in test data point [True]
877 Text feature [fusions] present in test data point [True]
884 Text feature [activities] present in test data point [True]
885 Text feature [fig] present in test data point [True]
887 Text feature [features] present in test data point [True]
896 Text feature [follows] present in test data point [True]
897 Text feature [predict] present in test data point [True]
898 Text feature [standard] present in test data point [True]
900 Text feature [population] present in test data point [True]
901 Text feature [noted] present in test data point [True]
905 Text feature [promega] present in test data point [True]
908 Text feature [consent] present in test data point [True]
910 Text feature [100] present in test data point [True]
911 Text feature [peptide] present in test data point [True]
915 Text feature [set] present in test data point [True]
918 Text feature [conclude] present in test data point [True]
921 Text feature [assess] present in test data point [True]
922 Text feature [result] present in test data point [True]
923 Text feature [thus] present in test data point [True]
926 Text feature [29] present in test data point [True]
927 Text feature [interestingly] present in test data point [True]
928 Text feature [luciferase] present in test data point [True]
929 Text feature [generated] present in test data point [True]
935 Text feature [interpretation] present in test data point [True]
936 Text feature [shows] present in test data point [True]
937 Text feature [score] present in test data point [True]
940 Text feature [due] present in test data point [True]
943 Text feature [partially] present in test data point [True]
944 Text feature [complete] present in test data point [True]
946 Text feature [reduced] present in test data point [True]
952 Text feature [resulted] present in test data point [True]
956 Text feature [frequency] present in test data point [True]
958 Text feature [34] present in test data point [True]
959 Text feature [highly] present in test data point [True]
962 Text feature [46] present in test data point [True]
963 Text feature [respect] present in test data point [True]
964 Text feature [selective] present in test data point [True]
965 Text feature [second] present in test data point [True]
969 Text feature [dual] present in test data point [True]
972 Text feature [proven] present in test data point [True]
975 Text feature [seven] present in test data point [True]
981 Text feature [still] present in test data point [True]
987 Text feature [3d] present in test data point [True]
988 Text feature [might] present in test data point [True]
991 Text feature [referred] present in test data point [True]
Out of the top 1000 features 434 are present in query point
```

4.5.3. Hyper paramter tuning (With Response Coding)	

```
In [103]: # ------
          # default parameters
          # sklearn.ensemble.RandomForestClassifier(n_estimators=10, criterion='gini', max_depth=None, min_samples_split=2,
          # min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_impurity_decrease=0.
          # min_impurity_split=None, bootstrap=True, oob_score=False, n_jobs=1, random_state=None, verbose=0, warm_start=False,
          # class_weight=None)
          # Some of methods of RandomForestClassifier()
          \# fit(X, y, [sample\_weight]) Fit the SVM model according to the given training data.
          \# predict(X) Perform classification on samples in X.
                               Perform classification on samples in X.
          # predict_proba (X)
          # some of attributes of RandomForestClassifier()
          # feature_importances_ : array of shape = [n_features]
          # The feature importances (the higher, the more important the feature).
          # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/random-forest-and-their-construc
          tion-2/
          # find more about CalibratedClassifierCV here at http://scikit-learn.org/stable/modules/generated/sklearn.calibration.
          CalibratedClassifierCV.html
          # ------
          # default paramters
          # sklearn.calibration.CalibratedClassifierCV(base_estimator=None, method='sigmoid', cv=3)
          # some of the methods of CalibratedClassifierCV()
          # fit(X, y[, sample_weight])
Fit the calibrated model
          # get_params([deep]) Get parameters for this estimator.
          # predict(X) Predict the target of new samples.
          # predict_proba(X) Posterior probabilities of classification
          #-----
          # video link:
          #-----
          alpha = [10,50,100,200,500,1000]
          max_depth = [2,3,5,10]
          cv_log_error_array = []
          for i in alpha:
              for j in max_depth:
                  print("for n_estimators =", i,"and max depth = ", j)
                  clf = RandomForestClassifier(n_estimators=i, criterion='gini', max_depth=j, random_state=42, n_jobs=-1)
                  clf.fit(train_x_responseCoding, train_y)
                  sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
                  sig_clf.fit(train_x_responseCoding, train_y)
                  sig_clf_probs = sig_clf.predict_proba(cv_x_responseCoding)
                  cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
                  print("Log Loss :",log_loss(cv_y, sig_clf_probs))
          fig, ax = plt.subplots()
          features = np.dot(np.array(alpha)[:,None],np.array(max_depth)[None]).ravel()
          ax.plot(features, cv_log_error_array,c='g')
          for i, txt in enumerate(np.round(cv_log_error_array,3)):
              ax.annotate((alpha[int(i/4)],max_depth[int(i%4)],str(txt)), (features[i],cv_log_error_array[i]))
          plt.grid()
          plt.title("Cross Validation Error for each alpha")
          plt.xlabel("Alpha i's")
          plt.ylabel("Error measure")
          plt.show()
          best_alpha = np.argmin(cv_log_error_array)
          clf = RandomForestClassifier(n_estimators=alpha[int(best_alpha/4)], criterion='gini', max_depth=max_depth[int(best_alp
          ha%4)], random state=42, n jobs=-1)
          clf.fit(train_x_responseCoding, train_y)
          sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
          sig_clf.fit(train_x_responseCoding, train_y)
          predict_y = sig_clf.predict_proba(train_x_responseCoding)
          print('For values of best alpha = ',
                alpha[int(best_alpha/4)],
                "The train log loss is:",
                log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(cv_x_responseCoding)
          print('For values of best alpha = ',
                alpha[int(best_alpha/4)],
                "The cross validation log loss is:",
                log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(test_x_responseCoding)
          print('For values of best alpha = ',
                alpha[int(best alpha/4)],
                "The test log loss is:",
                log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
```

```
for n_estimators = 10 and max depth = 2
Log Loss: 2.028729151890328
for n_estimators = 10 and max depth = 3
Log Loss: 1.7785418239759783
for n_estimators = 10 and max depth = 5
Log Loss: 1.7227725500903117
for n_estimators = 10 and max depth = 10
Log Loss: 1.7758739721968217
for n_estimators = 50 and max depth = 2
Log Loss: 1.551911707052268
for n_{estimators} = 50 and max depth = 3
Log Loss: 1.3879542300015055
for n_estimators = 50 and max depth = 5
Log Loss: 1.326427418987433
for n_estimators = 50 and max depth = 10
Log Loss: 1.7012899190898199
for n_estimators = 100 and max depth = 2
Log Loss: 1.4999410953211652
for n_estimators = 100 and max depth = 3
Log Loss: 1.441355346035283
for n_estimators = 100 and max depth = 5
Log Loss: 1.3252757573526668
for n_estimators = 100 and max depth = 10
Log Loss: 1.6197637694335536
for n_estimators = 200 and max depth = 2
Log Loss: 1.5724756257010042
for n_estimators = 200 and max depth = 3
Log Loss: 1.4575507254276303
for n_estimators = 200 and max depth = 5
Log Loss: 1.3717281530237928
for n_estimators = 200 and max depth = 10
Log Loss: 1.5809442783361776
for n_{estimators} = 500 and max depth = 2
Log Loss: 1.6334876757608925
for n_{estimators} = 500 and max depth = 3
Log Loss: 1.5014970272237314
for n_{estimators} = 500 and max depth = 5
Log Loss: 1.3498893006607293
for n_{estimators} = 500 and max depth = 10
Log Loss: 1.6338271124612158
for n estimators = 1000 and max depth = 2
Log Loss: 1.6166343912659635
for n_{estimators} = 1000 and max depth = 3
Log Loss: 1.5147897117238742
for n_estimators = 1000 and max depth = 5
Log Loss: 1.3404535776145738
for n_{estimators} = 1000 and max depth = 10
Log Loss: 1.5950803869403798
For values of best alpha = 100 The train log loss is: 0.0586846034926599
For values of best alpha = 100 The cross validation log loss is: 1.3252757573526668
For values of best alpha = 100 The test log loss is: 1.3461728638769181
```

4.5.4. Testing model with best hyper parameters (Response Coding)

```
In [104]: | # ------
          # default parameters
          # sklearn.ensemble.RandomForestClassifier(n_estimators=10, criterion='gini', max_depth=None, min_samples_split=2,
          # min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_impurity_decrease=0.
          # min_impurity_split=None, bootstrap=True, oob_score=False, n_jobs=1, random_state=None, verbose=0, warm_start=False,
          # class_weight=None)
          # Some of methods of RandomForestClassifier()
          # fit(X, y, [sample_weight])
                                       Fit the SVM model according to the given training data.
          \# predict(X) Perform classification on samples in X.
          # predict_proba (X)
                                Perform classification on samples in X.
          # some of attributes of RandomForestClassifier()
          # feature_importances_ : array of shape = [n_features]
          # The feature importances (the higher, the more important the feature).
          # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/random-forest-and-their-construc
          tion-2/
          clf = RandomForestClassifier(max_depth=max_depth[int(best_alpha%4)], n_estimators=alpha[int(best_alpha/4)], criterion
          ='gini', max_features='auto',random_state=42)
          predict_and_plot_confusion_matrix(train_x_responseCoding, train_y,cv_x_responseCoding,cv_y, clf)
```

----- Confusion matrix -----3.000 2.000 16.000 12.000 5.000 0.000 5.000 0.000 - 75 1.000 4.000 2.000 0.000 0.000 6.000 2.000 0.000 2.000 1.000 6.000 0.000 4.000 0.000 1.000 0.000 0.000 - 60 17.000 3.000 9.000 12.000 1.000 1.000 3.000 1.000 Original Class - 45 1.000 7.000 1.000 5.000 15.000 5.000 3.000 1.000 1.000 2.000 7.000 0.000 3.000 6.000 26.000 0.000 0.000 0.000 - 30 0.000 84.000 21.000 0.000 0.000 0.000 1.000 0.000 - 15 0.000 1.000 0.000 1.000 0.000 0.000 0.000 1.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 1.000 2.000 3.000 -0 Predicted Class ----- Precision matrix (Columm Sum=1) -------0.018 0.047 0.178 0.245 0.135 0.333 0.000 0.000 - 0.75 0.350 0.014 0.093 0.022 0.000 0.000 0.102 0.133 0.000 - 0.60 0.028 0.006 0.140 0.000 0.082 0.000 0.017 0.000 0.000 0.239 0.018 0.209 0.700 0.245 0.027 0.200 0.200 0.017 - 0.45 0.014 0.043 0.023 0.056 0.306 0.135 0.051 0.067 0.200 0.043 0.000 0.028 0.033 0.122 0.703 0.000 0.000 0.000 - 0.30 0.000 0.000 0.000 0.000 0.067 0.000 -0.15 0.000 0.006 0.000 0.011 0.000 0.000 0.000 0.067 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.017 0.133 - 0.00 Predicted Class ----- Recall matrix (Row sum=1) ------0.033 0.022 0.176 0.132 0.055 0.000 0.055 0.000 - 0.75 0.056 0.028 0.000 0.014 0.792 0.000 0.083 0.028 0.000 - 0.60 0.071 0.000 0.286 0.143 0.000 0.071 0.000 0.000 0.027 0.109 0.155 0.082 0.009 0.009 0.027 0.009 - 0.45 0.179 0.026 0.128 0.128 0.077 0.026 0.026 0.045 0.159 0.000 0.068 0.136 0.000 0.000 0.000 - 0.30 0.000 0.137 0.000 0.000 0.000 0.307 0.007 0.000 - 0.15 0.333 0.333 0.333 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.333 0.000 0.000 0.000 0.000 0.167 - 0.00 4 'n 8 9

Predicted Class

4.5.5. Feature Importance

4.5.5.1. Correctly Classified point

```
In [105]: | clf = RandomForestClassifier(n_estimators=alpha[int(best_alpha/4)], criterion='gini', max_depth=max_depth[int(best_alpha/4)]
          ha%4)], random_state=42, n_jobs=-1)
          clf.fit(train_x_responseCoding, train_y)
          sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
          sig_clf.fit(train_x_responseCoding, train_y)
          test_point_index = 100
          no_feature = 1000
          predicted_cls = sig_clf.predict(test_x_responseCoding[test_point_index].reshape(1,-1))
          print("Predicted Class :", predicted_cls[0])
          print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_responseCoding[test_point_index].reshape
          (1,-1)),4))
          print("Actual Class :", test_y[test_point_index])
          # indices = np.argsort(-clf.feature_importances_)
          # print("-"*50)
          # for i in indices:
                if i<9:
                    print("Gene is important feature")
          #
                elif i<18:
                    print("Variation is important feature")
          #
          #
                else:
          #
                    print("Text is important feature")
```

Predicted Class : 4
Predicted Class Probabilities: [[0.0766 0.1183 0.2165 0.3039 0.0436 0.0757 0.0571 0.0629 0.0454]]
Actual Class : 4

4.5.5.2. Incorrectly Classified point

```
In [106]: test_point_index = 31
          predicted_cls = sig_clf.predict(test_x_responseCoding[test_point_index].reshape(1,-1))
          print("Predicted Class :", predicted_cls[0])
          print("Predicted Class Probabilities:", np.round(sig_clf.predict_proba(test_x_responseCoding[test_point_index].reshape
          (1,-1),4))
          print("Actual Class :", test_y[test_point_index])
          # indices = np.argsort(-clf.feature_importances_)
          # print("-"*50)
          # for i in indices:
                if i<9:
                    print("Gene is important feature")
                elif i<18:
                    print("Variation is important feature")
          #
          #
                else:
                    print("Text is important feature")
```

Predicted Class : 2
Predicted Class Probabilities: [[0.0069 0.7685 0.0586 0.0107 0.0103 0.0258 0.1004 0.0125 0.0062]]
Actual Class : 7

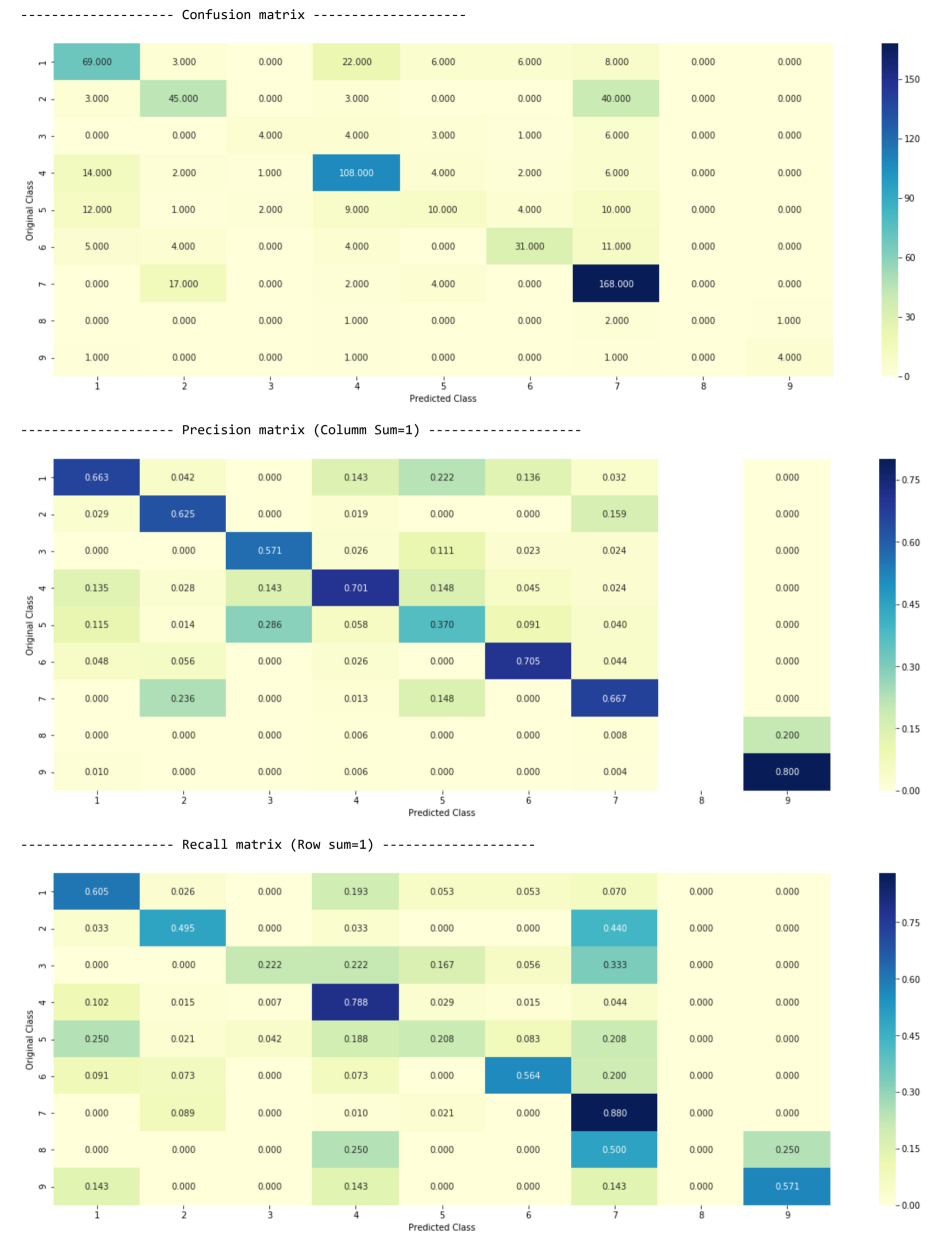
4.7 Stack the models

4.7.1 testing with hyper parameter tuning

```
In [107]: # read more about SGDClassifier() at http://scikit-learn.org/stable/modules/generated/sklearn.linear_model.SGDClassifi
          er.html
          # -----
          # default parameters
          # SGDClassifier(loss='hinge', penalty='l2', alpha=0.0001, l1_ratio=0.15, fit_intercept=True, max_iter=None, tol=None,
          # shuffle=True, verbose=0, epsilon=0.1, n_jobs=1, random_state=None, learning_rate='optimal', eta0=0.0, power_t=0.5,
          # class_weight=None, warm_start=False, average=False, n_iter=None)
          # some of methods
          \# fit(X, y[, coef_init, intercept_init, ...]) Fit linear model with Stochastic Gradient Descent.
          # predict(X) Predict class labels for samples in X.
          # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/geometric-intuition-1/
          # read more about support vector machines with linear kernals here http://scikit-learn.org/stable/modules/generated/sk
          Learn.svm.SVC.html
          # default parameters
          # SVC(C=1.0, kernel='rbf', degree=3, gamma='auto', coef0=0.0, shrinking=True, probability=False, tol=0.001,
          # cache_size=200, class_weight=None, verbose=False, max_iter=-1, decision_function_shape='ovr', random_state=None)
          # Some of methods of SVM()
          \# fit(X, y, [sample\_weight]) Fit the SVM model according to the given training data.
          \# predict(X) Perform classification on samples in X.
          # -----
          # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/mathematical-derivation-copy-8/
          # read more about support vector machines with linear kernals here http://scikit-learn.org/stable/modules/generated/sk
          learn.ensemble.RandomForestClassifier.html
          # -----
          # default parameters
          # sklearn.ensemble.RandomForestClassifier(n_estimators=10, criterion='gini', max_depth=None, min_samples_split=2,
          # min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_impurity_decrease=0.
          # min_impurity_split=None, bootstrap=True, oob_score=False, n_jobs=1, random_state=None, verbose=0, warm_start=False,
          # class_weight=None)
          # Some of methods of RandomForestClassifier()
          # fit(X, y, [sample_weight]) Fit the SVM model according to the given training data.
          \# predict(X) Perform classification on samples in X.
          # predict_proba (X) Perform classification on samples in X.
          # some of attributes of RandomForestClassifier()
          # feature_importances_ : array of shape = [n_features]
          # The feature importances (the higher, the more important the feature).
          # video link: https://www.appliedaicourse.com/course/applied-ai-course-online/lessons/random-forest-and-their-construc
          tion-2/
          # -----
          clf1 = SGDClassifier(alpha=0.001, penalty='12', loss='log', class_weight='balanced', random_state=0)
          clf1.fit(train_x_onehotCoding, train_y)
          sig clf1 = CalibratedClassifierCV(clf1, method="sigmoid")
          clf2 = SGDClassifier(alpha=0.01, penalty='12', loss='hinge', class_weight='balanced', random_state=0)
          clf2.fit(train_x_onehotCoding, train_y)
          sig_clf2 = CalibratedClassifierCV(clf2, method="sigmoid")
          clf3 = MultinomialNB(alpha=1000)
          clf3.fit(train_x_onehotCoding, train_y)
          sig_clf3 = CalibratedClassifierCV(clf3, method="sigmoid")
          sig_clf1.fit(train_x_onehotCoding, train_y)
          print("Logistic Regression : Log Loss: %0.2f" % (log_loss(cv_y, sig_clf1.predict_proba(cv_x_onehotCoding))))
          sig_clf2.fit(train_x_onehotCoding, train_y)
          print("Support vector machines : Log Loss: %0.2f" % (log_loss(cv_y, sig_clf2.predict_proba(cv_x_onehotCoding))))
          sig clf3.fit(train_x_onehotCoding, train_y)
          print("Naive Bayes : Log Loss: %0.2f" % (log loss(cv y, sig clf3.predict proba(cv x onehotCoding))))
          print("-"*50)
          alpha = [0.0001,0.001,0.01,0.1,1,10]
          best_alpha = 999
          for i in alpha:
              lr = LogisticRegression(C=i)
              sclf = StackingClassifier(classifiers=[sig_clf1, sig_clf2, sig_clf3], meta_classifier=lr, use_probas=True)
              sclf.fit(train x onehotCoding, train y)
              print("Stacking Classifer: for the value of alpha: %f Log Loss: %0.3f" % (i, log_loss(cv_y, sclf.predict_proba(cv
          _x_onehotCoding))))
              log error =log loss(cv y, sclf.predict proba(cv x onehotCoding))
              if best_alpha > log_error:
                  best alpha = log error
```

4.7.2 testing the model with the best hyper parameters

Log loss (train) on the stacking classifier: 0.6015459146688893 Log loss (CV) on the stacking classifier: 1.0959530429401525 Log loss (test) on the stacking classifier: 1.0572334308842748 Number of missclassified point: 0.3398496240601504



```
In [109]: #Refer:http://scikit-learn.org/stable/modules/generated/sklearn.ensemble.VotingClassifier.html
    from sklearn.ensemble import VotingClassifier
    vclf = VotingClassifier(estimators=[('lr', sig_clf1), ('svc', sig_clf2), ('rf', sig_clf3)], voting='soft')
    vclf.fit(train_x_onehotCoding, train_y)
    print("Log loss (train) on the VotingClassifier :", log_loss(train_y, vclf.predict_proba(train_x_onehotCoding)))
    print("Log loss (CV) on the VotingClassifier :", log_loss(cv_y, vclf.predict_proba(cv_x_onehotCoding)))
    print("Log loss (test) on the VotingClassifier :", log_loss(test_y, vclf.predict_proba(test_x_onehotCoding)))
    print("Number of missclassified point :", np.count_nonzero((vclf.predict(test_x_onehotCoding) - test_y))/test_y.shape[0]
    ])
    plot_confusion_matrix(test_y=test_y, predict_y=vclf.predict(test_x_onehotCoding))
```

Log loss (train) on the VotingClassifier: 0.6843786340991019 Log loss (CV) on the VotingClassifier: 1.0156508836071645 Log loss (test) on the VotingClassifier: 1.019685657541371 Number of missclassified point: 0.3458646616541353

----- Confusion matrix ------66.000 2.000 0.000 21.000 8.000 6.000 11.000 0.000 0.000 - 150 3.000 39.000 0.000 3.000 0.000 0.000 46.000 0.000 0.000 0.000 6.000 3.000 3.000 0.000 0.000 1.000 5.000 0.000 - 120 14.000 1.000 1.000 4.000 2.000 9.000 0.000 0.000 - 90 9.000 0.000 3.000 8.000 12.000 4.000 12.000 0.000 0.000 5.000 2.000 0.000 4.000 0.000 30.000 14.000 0.000 0.000 60 0.000 13.000 2.000 2.000 4.000 0.000 170.000 0.000 0.000 0.000 0.000 - 30 0.000 0.000 0.000 0.000 2.000 1.000 1.000 0.000 0.000 1.000 0.000 1.000 0.000 0.000 0.000 5.000 -0 Predicted Class ----- Precision matrix (Columm Sum=1) ------1.0 0.035 0.258 0.000 0.142 0.140 0.041 0.000 0.000 0.031 0.000 0.020 0.000 0.000 0.171 0.000 - 0.8 0.000 0.000 0.020 0.097 0.023 0.019 0.000 0.000 0.018 0.083 0.129 0.047 0.033 0.143 0.000 0.000 - 0.6 0.250 0.054 0.387 0.092 0.000 0.093 0.045 0.000 0.000 - 0.4 0.051 0.035 0.000 0.027 0.000 0.052 0.000 0.228 0.167 0.014 0.129 0.000 0.000 0.000 - 0.2 0.167 0.000 0.000 0.000 0.000 0.000 0.000 0.007 1.000 0.010 0.000 0.000 0.007 0.000 0.000 0.000 0.000 0.833 - 0.0 Predicted Class ----- Recall matrix (Row sum=1) ------0.184 0.070 0.018 0.000 0.053 0.096 0.000 0.000 - 0.75 0.033 0.033 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.333 0.167 0.167 0.056 0.000 0.000 - 0.60 0.102 0.007 0.007 0.774 0.029 0.015 0.066 0.000 0.000 0.167 0.250 0.083 0.250 - 0.45 0.188 0.000 0.062 0.000 0.000 0.091 0.036 0.000 0.073 0.000 0.545 0.255 0.000 0.000 - 0.30 0.010 0.010 0.021 0.000 0.890 0.000 0.000 0.000 0.000 0.000 0.250 - 0.15 0.000 0.000 0.143 0.000 0.000 0.000 0.000 0.143 - 0.00 9 i Predicted Class

From above models we can evaluate that 'Logistic Regression With Class balancing' is far better choice than others. So we will try countVectorizer features with both unigrams and bigrams to see whether it will reduce the log loss further or not.

Logistic Regression With Class Balancing using countVectorizer features with both unigrams and bigrams

Gene Feature

```
In [0]: #response-coding of the Gene feature
        # alpha is used for laplace smoothing
        alpha = 1
        # train gene feature
        train_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_train))
        # test gene feature
        test_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_test))
        # cross validation gene feature
        cv_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_cv))
In [0]: # one-hot encoding of Gene feature.
        gene_vectorizer = CountVectorizer(ngram_range=(1, 2))
        train_gene_feature_onehotCoding = gene_vectorizer.fit_transform(x_train['Gene'])
        test_gene_feature_onehotCoding = gene_vectorizer.transform(x_test['Gene'])
        cv_gene_feature_onehotCoding = gene_vectorizer.transform(x_cv['Gene'])
        # don't forget to normalize every feature
        train_gene_feature_onehotCoding = normalize(train_gene_feature_onehotCoding, axis=0)
        test_gene_feature_onehotCoding = normalize(test_gene_feature_onehotCoding, axis=0)
        cv_gene_feature_onehotCoding = normalize(cv_gene_feature_onehotCoding, axis=0)
```

Variation Feature

```
In [0]: # alpha is used for laplace smoothing
alpha = 1

# train gene feature
train_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_train))

# test gene feature
test_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_test))

# cross validation gene feature
cv_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_cv))

In [0]: # one-hot encoding of variation feature.
variation_vectorizer = CountVectorizer(ngram_range=(1, 2))
train_variation_feature_onehotCoding = variation_vectorizer.fit_transform(x_train['Variation'])
test_variation_feature_onehotCoding = variation_vectorizer.transform(x_test['Variation'])

# don't forget to normalize every feature
train_variation_feature_onehotCoding = normalize(train_variation_feature_onehotCoding, axis=0)
```

test_variation_feature_onehotCoding = normalize(test_variation_feature_onehotCoding, axis=0)
cv_variation_feature_onehotCoding = normalize(cv_variation_feature_onehotCoding, axis=0)

Text Feature

```
In [115]: # building a CountVectorizer with all the words that occured minimum 3 times in train data
    text_vectorizer = CountVectorizer(min_df=3,ngram_range=(1, 2))
    train_text_feature_onehotCoding = text_vectorizer.fit_transform(x_train['TEXT'])

# getting all the feature names (words)
    train_text_features= text_vectorizer.get_feature_names()

# train_text_feature_onehotCoding.sum(axis=0).A1 will sum every row and returns (1*number of features) vector
    train_text_fea_counts = train_text_feature_onehotCoding.sum(axis=0).A1

# zip(list(text_features),text_fea_counts) will zip a word with its number of times it occured
    text_fea_dict = dict(zip(list(train_text_features),train_text_fea_counts))

print("Total number of unique words in train data :", len(train_text_features))
```

Total number of unique words in train data : 769276

```
In [0]: #response coding of text features
            train_text_feature_responseCoding = get_text_responsecoding(x_train)
            test_text_feature_responseCoding = get_text_responsecoding(x_test)
            cv_text_feature_responseCoding = get_text_responsecoding(x_cv)
            # https://stackoverflow.com/a/16202486
            # we convert each row values such that they sum to 1
            train_text_feature_responseCoding = (train_text_feature_responseCoding.T/train_text_feature_responseCoding.sum(axis=1
            )).T
            test_text_feature_responseCoding = (test_text_feature_responseCoding.T/test_text_feature_responseCoding.sum(axis=1)).
            cv_text_feature_responseCoding = (cv_text_feature_responseCoding.T/cv_text_feature_responseCoding.sum(axis=1)).T
    In [0]: | # don't forget to normalize every feature
            train_text_feature_onehotCoding = normalize(train_text_feature_onehotCoding, axis=0)
             # we use the same vectorizer that was trained on train data
            test_text_feature_onehotCoding = text_vectorizer.transform(x_test['TEXT'])
             # don't forget to normalize every feature
            test_text_feature_onehotCoding = normalize(test_text_feature_onehotCoding, axis=0)
            # we use the same vectorizer that was trained on train data
            cv_text_feature_onehotCoding = text_vectorizer.transform(x_cv['TEXT'])
            # don't forget to normalize every feature
            cv_text_feature_onehotCoding = normalize(cv_text_feature_onehotCoding, axis=0)
Stack above three features
    In [0]: # merging gene, variance and text features
            # building train, test and cross validation data sets
             \# \ a = [[1, 2],
                   [3, 4]]
            # b = [[4, 5],
                  [6, 7]]
            # hstack(a, b) = [[1, 2, 4, 5],
                             [ 3, 4, 6, 7]]
            train_gene_var_onehotCoding = hstack((train_gene_feature_onehotCoding,train_variation_feature_onehotCoding))
             test_gene_var_onehotCoding = hstack((test_gene_feature_onehotCoding,test_variation_feature_onehotCoding))
            cv_gene_var_onehotCoding = hstack((cv_gene_feature_onehotCoding,cv_variation_feature_onehotCoding))
            train_x_onehotCoding = hstack((train_gene_var_onehotCoding, train_text_feature_onehotCoding)).tocsr()
            train_y = np.array(list(y_train['Class']))
            test_x_onehotCoding = hstack((test_gene_var_onehotCoding, test_text_feature_onehotCoding)).tocsr()
            test_y = np.array(list(y_test['Class']))
            cv_x_onehotCoding = hstack((cv_gene_var_onehotCoding, cv_text_feature_onehotCoding)).tocsr()
```

```
train_gene_var_responseCoding = np.hstack((train_gene_feature_responseCoding,train_variation_feature_responseCoding))
          test_gene_var_responseCoding = np.hstack((test_gene_feature_responseCoding,test_variation_feature_responseCoding))
          cv_gene_var_responseCoding = np.hstack((cv_gene_feature_responseCoding,cv_variation_feature_responseCoding))
          train_x_responseCoding = np.hstack((train_gene_var_responseCoding, train_text_feature_responseCoding))
          test_x_responseCoding = np.hstack((test_gene_var_responseCoding, test_text_feature_responseCoding))
          cv_x_responseCoding = np.hstack((cv_gene_var_responseCoding, cv_text_feature_responseCoding))
In [119]: print("One hot encoding features :")
          print("(number of data points * number of features) in train data = ", train_x_onehotCoding.shape)
          print("(number of data points * number of features) in test data = ", test_x_onehotCoding.shape)
          print("(number of data points * number of features) in cross validation data =", cv_x_onehotCoding.shape)
          One hot encoding features :
          (number of data points * number of features) in train data = (2124, 771564)
          (number of data points * number of features) in test data = (665, 771564)
          (number of data points * number of features) in cross validation data = (532, 771564)
In [120]: print(" Response encoding features :")
          print("(number of data points * number of features) in train data = ", train_x_responseCoding.shape)
```

print("(number of data points * number of features) in test data = ", test_x_responseCoding.shape)

(number of data points * number of features) in train data = (2124, 27) (number of data points * number of features) in test data = (665, 27)

(number of data points * number of features) in cross validation data = (532, 27)

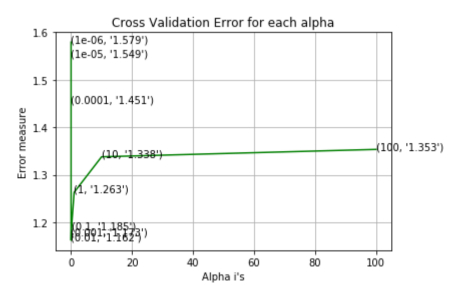
print("(number of data points * number of features) in cross validation data =", cv_x_responseCoding.shape)

cv_y = np.array(list(y_cv['Class']))

Response encoding features :

```
In [121]: | alpha = [10 ** x for x in range(-6, 3)]
          cv_log_error_array = []
          for i in alpha:
              print("for alpha =", i)
              clf = SGDClassifier(class_weight='balanced', alpha=i, penalty='12', loss='log', random_state=42)
              clf.fit(train_x_onehotCoding, train_y)
              sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
              sig clf.fit(train x onehotCoding, train y)
              sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
              cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
              # to avoid rounding error while multiplying probabilites we use log-probability estimates
              print("Log Loss :",log_loss(cv_y, sig_clf_probs))
          fig, ax = plt.subplots()
          ax.plot(alpha, cv_log_error_array,c='g')
          for i, txt in enumerate(np.round(cv_log_error_array,3)):
              ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
          plt.grid()
          plt.title("Cross Validation Error for each alpha")
          plt.xlabel("Alpha i's")
          plt.ylabel("Error measure")
          plt.show()
          best_alpha = np.argmin(cv_log_error_array)
          clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
          clf.fit(train_x_onehotCoding, train_y)
          sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
          sig_clf.fit(train_x_onehotCoding, train_y)
          predict_y = sig_clf.predict_proba(train_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best_alpha],
                "The train log loss is:",
                log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best alpha],
                "The cross validation log loss is:",
                log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(test_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best_alpha], "The test log loss is:",
                log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
          for alpha = 1e-06
          Log Loss: 1.5792285866014613
          for alpha = 1e-05
          Log Loss: 1.549337116347172
          for alpha = 0.0001
          Log Loss: 1.4507715783792976
          for alpha = 0.001
```

Log Loss: 1.5792285866014613
for alpha = 1e-05
Log Loss: 1.549337116347172
for alpha = 0.0001
Log Loss: 1.4507715783792976
for alpha = 0.001
Log Loss: 1.172640110999891
for alpha = 0.01
Log Loss: 1.1620017733274437
for alpha = 0.1
Log Loss: 1.184997700369832
for alpha = 1
Log Loss: 1.2629453423714603
for alpha = 10
Log Loss: 1.3377980615323617
for alpha = 100
Log Loss: 1.3533760588405503



```
For values of best alpha = 0.01 The train log loss is: 0.715249194715444

For values of best alpha = 0.01 The cross validation log loss is: 1.1620017733274437

For values of best alpha = 0.01 The test log loss is: 1.1469722139871348
```

Predicted Class

Still model does not decreases log loss values after using unigram and bigram features

Lets apply feature engineering on the data and then apply logistic regression again

Lets merge gene and variation data into one list and apply TFidfVectorizer on top of it.

Gene Feature

```
In [0]: result = pd.merge(data_variants, data_text,on='ID', how='left')
        result.loc[result['TEXT'].isnull(),'TEXT'] = result['Gene'] +' '+result['Variation']
        y_true = result['Class'].values
        result.Gene = result.Gene.str.replace('\s+', '_')
        result.Variation = result.Variation.str.replace('\s+', '_')
        x_train, x_test, y_train, y_test = train_test_split(result, y_true, stratify=y_true, test_size=0.2)
        x_train, x_cv, y_train, y_cv = train_test_split(x_train, y_train, stratify=y_train, test_size=0.2)
In [0]: # get_gv_fea_dict: Get Gene varaition Feature Dict
        def get_gv_fea_dict(alpha, feature, df):
            value_count = x_train[feature].value_counts()
            gv_dict = dict()
            for i, denominator in value_count.items():
                vec = []
                for k in range(1,10):
                    cls_cnt = x_train.loc[(x_train['Class']==k) & (x_train[feature]==i)]
                    vec.append((cls_cnt.shape[0] + alpha*10)/ (denominator + 90*alpha))
                gv_dict[i]=vec
            return gv_dict
        # Get Gene variation feature
        def get_gv_feature(alpha, feature, df):
            gv_dict = get_gv_fea_dict(alpha, feature, df)
            value_count = x_train[feature].value_counts()
            gv_fea = []
            for index, row in df.iterrows():
                if row[feature] in dict(value_count).keys():
                    gv fea.append(gv dict[row[feature]])
                    gv_fea.append([1/9,1/9,1/9,1/9,1/9,1/9,1/9,1/9])
            return gv_fea
In [0]: #response-coding of the Gene feature
        # alpha is used for laplace smoothing
        alpha = 1
        # train gene feature
        train_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_train))
        # test gene feature
        test_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_test))
        # cross validation gene feature
        cv_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", x_cv))
In [0]: | # one-hot encoding of Gene feature.
        gene_vectorizer = TfidfVectorizer()
        train_gene_feature_onehotCoding = gene_vectorizer.fit_transform(x_train['Gene'])
        test_gene_feature_onehotCoding = gene_vectorizer.transform(x_test['Gene'])
        cv_gene_feature_onehotCoding = gene_vectorizer.transform(x_cv['Gene'])
```

Variation Feature

```
In [0]: # alpha is used for laplace smoothing
    alpha = 1

# train gene feature
    train_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_train))

# test gene feature
    test_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_test))

# cross validation gene feature
    cv_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", x_cv))
In [0]: # one-hot encoding of variation feature.
    variation vectorizer = TfidfVectorizer()
```

train_variation_feature_onehotCoding = variation_vectorizer.fit_transform(x_train['Variation'])

test_variation_feature_onehotCoding = variation_vectorizer.transform(x_test['Variation'])
cv variation feature onehotCoding = variation vectorizer.transform(x cv['Variation'])

In [0]: | def extract_dictionary_paddle(cls_text):

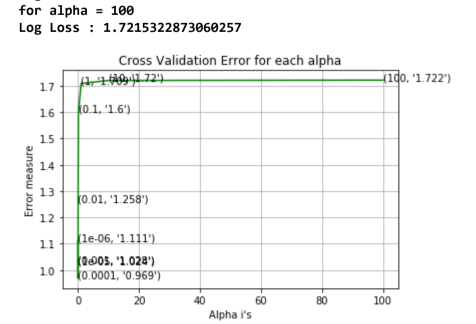
dictionary = defaultdict(int)

```
for index, row in cls_text.iterrows():
                  for word in row['TEXT'].split():
                      dictionary[word] +=1
              return dictionary
          import math
          #https://stackoverflow.com/a/1602964
          def get_text_responsecoding(df):
              text_feature_responseCoding = np.zeros((df.shape[0],9))
              for i in range(0,9):
                  row_index = 0
                  for index, row in df.iterrows():
                      sum_prob = 0
                      for word in row['TEXT'].split():
                          sum_prob += math.log(((dict_list[i].get(word,0)+10 )/(total_dict.get(word,0)+90)))
                      text_feature_responseCoding[row_index][i] = math.exp(sum_prob/len(row['TEXT'].split()))
                      row_index += 1
              return text_feature_responseCoding
In [130]: # building a CountVectorizer with all the words that occured minimum 3 times in train data
          text_vectorizer = TfidfVectorizer()
          train_text_feature_onehotCoding = text_vectorizer.fit_transform(x_train['TEXT'])
          # getting all the feature names (words)
          train_text_features= text_vectorizer.get_feature_names()
          # train_text_feature_onehotCoding.sum(axis=0).A1 will sum every row and returns (1*number of features) vector
          train_text_fea_counts = train_text_feature_onehotCoding.sum(axis=0).A1
          # zip(list(text_features),text_fea_counts) will zip a word with its number of times it occured
          text_fea_dict = dict(zip(list(train_text_features),train_text_fea_counts))
          print("Total number of unique words in train data :", len(train_text_features))
          Total number of unique words in train data : 126830
 In [0]: | dict_list = []
          # dict_list =[] contains 9 dictoinaries each corresponds to a class
          for i in range(1,10):
              cls_text = x_train[x_train['Class']==i]
              # build a word dict based on the words in that class
              dict_list.append(extract_dictionary_paddle(cls_text))
              # append it to dict_list
          # dict_list[i] is build on i'th class text data
          # total_dict is buid on whole training text data
          total_dict = extract_dictionary_paddle(x_train)
          confuse_array = []
          for i in train_text_features:
              ratios = []
              max_val = -1
              for j in range(0,9):
                  ratios.append((dict_list[j][i]+10 )/(total_dict[i]+90))
              confuse_array.append(ratios)
          confuse_array = np.array(confuse_array)
 In [0]: #response coding of text features
          train_text_feature_responseCoding = get_text_responsecoding(x_train)
          test_text_feature_responseCoding = get_text_responsecoding(x_test)
          cv_text_feature_responseCoding = get_text_responsecoding(x_cv)
          # https://stackoverflow.com/a/16202486
          # we convert each row values such that they sum to 1
          train_text_feature_responseCoding = (train_text_feature_responseCoding.T/train_text_feature_responseCoding.sum(axis=1)
          test text feature responseCoding = (test text feature responseCoding.T/test text feature responseCoding.sum(axis=1)).
          cv_text_feature_responseCoding = (cv_text_feature_responseCoding.T/cv_text_feature_responseCoding.sum(axis=1)).T
 In [0]: | test_text_feature_onehotCoding = text_vectorizer.transform(x_test['TEXT'])
          cv_text_feature_onehotCoding = text_vectorizer.transform(x_cv['TEXT'])
```

```
In [0]: # Collecting all the genes and variations data into a single list
            gene_variation = []
             for gene in data_variants['Gene'].values:
                 gene_variation.append(gene)
             for variation in data_variants['Variation'].values:
                 gene_variation.append(variation)
    In [0]: | tfidfVectorizer = TfidfVectorizer(max_features=1000)
             text2 = tfidfVectorizer.fit_transform(gene_variation)
            gene_variation_features = tfidfVectorizer.get_feature_names()
             train_text = tfidfVectorizer.transform(x_train['TEXT'])
            test text = tfidfVectorizer.transform(x test['TEXT'])
             cv_text = tfidfVectorizer.transform(x_cv['TEXT'])
Stack above three features
```

```
In [0]: | train_gene_var_onehotCoding = hstack((train_gene_feature_onehotCoding, train_variation_feature_onehotCoding))
          test_gene_var_onehotCoding = hstack((test_gene_feature_onehotCoding,test_variation_feature_onehotCoding))
          cv_gene_var_onehotCoding = hstack((cv_gene_feature_onehotCoding,cv_variation_feature_onehotCoding))
          # Adding the train_text feature
          train_x_onehotCoding = hstack((train_gene_var_onehotCoding, train_text))
          train_x_onehotCoding = hstack((train_x_onehotCoding, train_text_feature_onehotCoding)).tocsr()
          train_y = np.array(list(x_train['Class']))
          # Adding the test_text feature
          test_x_onehotCoding = hstack((test_gene_var_onehotCoding, test_text))
          test_x_onehotCoding = hstack((test_x_onehotCoding, test_text_feature_onehotCoding)).tocsr()
          test_y = np.array(list(x_test['Class']))
          # Adding the cv_text feature
          cv_x_onehotCoding = hstack((cv_gene_var_onehotCoding, cv_text))
          cv_x_onehotCoding = hstack((cv_x_onehotCoding, cv_text_feature_onehotCoding)).tocsr()
          cv_y = np.array(list(x_cv['Class']))
          train_gene_var_responseCoding = np.hstack((train_gene_feature_responseCoding,train_variation_feature_responseCoding))
          test_gene_var_responseCoding = np.hstack((test_gene_feature_responseCoding,test_variation_feature_responseCoding))
          cv_gene_var_responseCoding = np.hstack((cv_gene_feature_responseCoding,cv_variation_feature_responseCoding))
          train_x_responseCoding = np.hstack((train_gene_var_responseCoding, train_text_feature_responseCoding))
          test_x_responseCoding = np.hstack((test_gene_var_responseCoding, test_text_feature_responseCoding))
          cv_x_responseCoding = np.hstack((cv_gene_var_responseCoding, cv_text_feature_responseCoding))
In [141]: | print("One hot encoding features :")
          print("(number of data points * number of features) in train data = ", train_x_onehotCoding.shape)
          print("(number of data points * number of features) in test data = ", test_x_onehotCoding.shape)
          print("(number of data points * number of features) in cross validation data =", cv_x_onehotCoding.shape)
          One hot encoding features :
          (number of data points * number of features) in train data = (2124, 130016)
          (number of data points * number of features) in test data = (665, 130016)
          (number of data points * number of features) in cross validation data = (532, 130016)
In [142]: print(" Response encoding features :")
          print("(number of data points * number of features) in train data = ", train_x_responseCoding.shape)
          print("(number of data points * number of features) in test data = ", test_x_responseCoding.shape)
          print("(number of data points * number of features) in cross validation data =", cv_x_responseCoding.shape)
           Response encoding features :
          (number of data points * number of features) in train data = (2124, 27)
           (number of data points * number of features) in test data = (665, 27)
          (number of data points * number of features) in cross validation data = (532, 27)
```

```
In [143]: | alpha = [10 ** x for x in range(-6, 3)]
          cv_log_error_array = []
          for i in alpha:
              print("for alpha =", i)
              clf = SGDClassifier(class_weight='balanced', alpha=i, penalty='12', loss='log', random_state=42)
              clf.fit(train_x_onehotCoding, train_y)
              sig_clf = CalibratedClassifierCV(clf, method="sigmoid")
              sig_clf.fit(train_x_onehotCoding, train_y)
              sig_clf_probs = sig_clf.predict_proba(cv_x_onehotCoding)
              cv_log_error_array.append(log_loss(cv_y, sig_clf_probs, labels=clf.classes_, eps=1e-15))
              # to avoid rounding error while multiplying probabilites we use log-probability estimates
              print("Log Loss :",log_loss(cv_y, sig_clf_probs))
          fig, ax = plt.subplots()
          ax.plot(alpha, cv_log_error_array,c='g')
          for i, txt in enumerate(np.round(cv_log_error_array,3)):
              ax.annotate((alpha[i],str(txt)), (alpha[i],cv_log_error_array[i]))
          plt.grid()
          plt.title("Cross Validation Error for each alpha")
          plt.xlabel("Alpha i's")
          plt.ylabel("Error measure")
          plt.show()
          best_alpha = np.argmin(cv_log_error_array)
          clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
          clf.fit(train_x_onehotCoding, train_y)
          sig clf = CalibratedClassifierCV(clf, method="sigmoid")
          sig_clf.fit(train_x_onehotCoding, train_y)
          predict_y = sig_clf.predict_proba(train_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best_alpha],
                "The train log loss is:",
                log_loss(y_train, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(cv_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best_alpha],
                "The cross validation log loss is:",
                log_loss(y_cv, predict_y, labels=clf.classes_, eps=1e-15))
          predict_y = sig_clf.predict_proba(test_x_onehotCoding)
          print('For values of best alpha = ',
                alpha[best_alpha], "The test log loss is:",
                log_loss(y_test, predict_y, labels=clf.classes_, eps=1e-15))
          for alpha = 1e-06
          Log Loss: 1.1109656126983949
          for alpha = 1e-05
          Log Loss: 1.0241360362529663
```



for alpha = 0.0001

for alpha = 0.001

for alpha = 0.01

for alpha = 0.1

for alpha = 1

for alpha = 10

Log Loss: 0.9688562180861012

Log Loss: 1.027737900802265

Log Loss: 1.2576779587627898

Log Loss: 1.60016682723826

Log Loss: 1.709106939445485

Log Loss: 1.720344494189662

```
For values of best alpha = 0.0001 The train log loss is: 0.4278060174196659

For values of best alpha = 0.0001 The cross validation log loss is: 0.9688562180861012

For values of best alpha = 0.0001 The test log loss is: 0.9854267985727088
```

In [144]: | clf = SGDClassifier(class_weight='balanced', alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42) predict_and_plot_confusion_matrix(train_x_onehotCoding, train_y, cv_x_onehotCoding, cv_y, clf) Log loss: 0.9688562180861012 Number of mis-classified points: 0.34210526315789475 ----- Confusion matrix -----53.000 22.000 5.000 5.000 3.000 0.000 3.000 0.000 0.000 - 125 1.000 35.000 0.000 1.000 0.000 3.000 32.000 0.000 0.000 1.000 0.000 0.000 3.000 2.000 1.000 7.000 0.000 0.000 - 100 0.000 1.000 6.000 1.000 0.000 Original Class 5 - 75 4.000 2.000 1.000 6.000 16.000 5.000 5.000 0.000 0.000 10.000 3.000 0.000 3.000 1.000 22.000 5.000 0.000 0.000 - 50 0.000 10.000 0.000 3.000 1.000 0.000 139.000 0.000 0.000 - 25 0.000 0.000 0.000 0.000 0.000 1.000 1.000 0.000 0.000 0.000 1.000 0.000 0.000 0.000 5.000 0.000 -0 Predicted Class ----- Precision matrix (Columm Sum=1) ------0.053 0.000 0.192 0.075 0.026 0.186 0.000 0.000 0.011 0.000 0.008 0.000 0.075 0.164 0.000 0.000 - 0.8 0.011 0.000 0.000 0.025 0.077 0.025 0.036 0.000 0.000 0.213 0.070 0.000 0.038 0.150 0.005 0.000 0.000 - 0.6 0.045 0.035 1.000 0.051 0.125 0.026 0.000 0.000 - 0.4 0.112 0.053 0.000 0.025 0.038 0.026 0.000 0.000 0.000 0.175 0.000 0.025 0.038 0.000 0.000 0.000 - 0.2 0.000 0.000 0.000 0.000 0.011 0.000 0.000 0.005 1.000 0.000 0.000 0.000 0.008 0.000 0.000 0.000 0.000 1.000 -0.0 i 3 8 6 Predicted Class ----- Recall matrix (Row sum=1) ------0.033 0.000 0.242 0.055 0.033 0.055 0.000 0.000 0.000 0.014 0.000 0.014 0.042 0.000 0.000 0.071 0.000 0.000 0.214 0.143 0.071 0.000 - 0.6 0.009 0.173 0.036 0.000 0.055 0.009 0.000 0.000 0.103 0.051 0.026 0.154 0.410 0.128 0.128 0.000 Origin - 0.4 0.114 0.227 0.068 0.000 0.068 0.023 0.000 0.000 0.000 0.065 0.000 0.020 0.007 0.000 0.908 0.000 0.000 - 0.2 0.333 0.000 0.000 0.000 0.000 0.000 0.333 0.333 0.000 0.000 0.000 0.000 0.167 0.000 0.000 0.000 0.000 0.833 - 0.0 3 4 ģ 6

Predicted Class

Late elimmariza all above models including the feature engineering approach

```
In [147]: print()
          from prettytable import PrettyTable
          ptable = PrettyTable()
          ptable.title = "*** Model Summary *** [Performance Metric: Log-Loss]"
          ptable.field_names=["Model Name","Train","CV","Test","% Misclassified Points"]
          ptable.add_row(["Naive Bayes","0.94","1.21","1.19","43"])
          ptable.add_row(["KNN","0.50","1.02","1.04","34"])
          ptable.add_row(["Logistic Regression With Class balancing","0.52","1.05","1.01","35"])
          ptable.add_row(["Logistic Regression Without Class balancing","0.52","1.07","1.04","34"])
          ptable.add_row(["Linear SVM","0.55","1.08","1.07","36"])
          ptable.add_row(["Random Forest Classifier With One hot Encoding","0.67","1.17","1.16","40"])
          ptable.add_row(["Random Forest Classifier With Response Coding","0.05","1.32","1.34","50"])
          ptable.add_row(["Stack Models:LR+NB+SVM","0.60","1.09","1.05","33"])
          ptable.add_row(["Maximum Voting classifier","0.68","1.01","1.01","34"])
          ptable.add_row(["Logistic Regression with class balancing using unigram and bigram","0.71","1.16","1.14","41"])
          ptable.add_row(["Logistic Regression with class balancing after feature engineering","0.42","0.96","0.98","34"])
          print(ptable)
          print()
```

Model Name	Train	cv	Test	% Misclassified Points
Naive Bayes	0.94	1.21	1.19	43
KNN	0.50	1.02	1.04	34
Logistic Regression With Class balancing	0.52	1.05	1.01	35
Logistic Regression Without Class balancing	0.52	1.07	1.04	34
Linear SVM	0.55	1.08	1.07	36
Random Forest Classifier With One hot Encoding	0.67	1.17	1.16	40
Random Forest Classifier With Response Coding	0.05	1.32	1.34	50
Stack Models:LR+NB+SVM	0.60	1.09	1.05	33
Maximum Voting classifier	0.68	1.01	1.01	34
Logistic Regression with class balancing using unigram and bigram	0.71	1.16	1.14	41
Logistic Regression with class balancing after feature engineering	0.42	0.96	0.98	34

In [0]: