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

- 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

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
- 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. Cyclin-dependent 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

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 [14]:

```
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.cross validation import StratifiedKFold
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

3.1.1. Reading Gene and Variation Data

```
In [2]:
```

```
data = pd.read_csv('training_variants')
print('Number of data points : ', data.shape[0])
print('Number of features : ', data.shape[1])
print('Features : ', data.columns.values)
data.head()

Number of data points : 3321
Number of features : 4
Features : ['ID' 'Gene' 'Variation' 'Class']
Out[2]:
```

	ID	Gene	Variation	Class
0	0	FAM58A	Truncating Mutations	1
1	1	CBL	W802*	2
2	2	CBL	Q249E	2
3	3	CBL	N454D	3
4	4	CBL	L399V	4

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 [3]:
```

```
# 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[3]:

	ID	ТЕХТ
0	0	Cyclin-dependent kinases (CDKs) regulate a var
1	1	Abstract Background Non-small cell lung canc
2	2	Abstract Background Non-small cell lung canc
3	3	Recent evidence has demonstrated that acquired
4	4	Oncogenic mutations in the monomeric Casitas B

3.1.3. Preprocessing of text

In [4]:

```
# loading stop words from nltk library
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
```

In [5]:

```
#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: 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
Time took for preprocessing the text: 236.8552379 seconds

In [6]:

```
#merging both gene_variations and text data based on ID
result = pd.merge(data, data_text,on='ID', how='left')
result.head()
```

Out[6]:

	ID	Gene	Variation	Class	TEXT
0	0	FAM58A	Truncating Mutations	1	cyclin dependent kinases cdks regulate variety
1	1	CBL	W802*	2	abstract background non small cell lung cancer
2	2	CBL	Q249E	2	abstract background non small cell lung cancer
3	3	CBL	N454D	3	recent evidence demonstrated acquired uniparen
4	4	CBL	L399V	4	oncogenic mutations monomeric casitas b lineag

In [7]:

```
result[result.isnull().any(axis=1)]
```

Out[7]:

	ID	Gene	Variation	Class	TEXT
1109	1109	FANCA	S1088F	1	NaN
1277	1277	ARID5B	Truncating Mutations	1	NaN
1407	1407	FGFR3	K508M	6	NaN
1639	1639	FLT1	Amplification	6	NaN
2755	2755	BRAF	G596C	7	NaN

In [8]:

```
result.loc[result['TEXT'].isnull(),'TEXT'] = result['Gene'] +' '+result['Variation']
```

In [9]:

```
result[result['ID']==1109]
```

	ID Gene		Variation	Class	TEXT		
1109	1109	FANCA	S1088F	1	FANCA 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 [10]:

```
y_true = result['Class'].values
result.Gene = result.Gene.str.replace('\s+', '_')
result.Variation = result.Variation.str.replace('\s+', '_')

# split the data into test and train by maintaining same distribution of output varaible 'y_true'
[stratify=y_true]
X_train, test_df, y_train, y_test = train_test_split(result, 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]
train_df, cv_df, y_train, y_cv = train_test_split(X_train, y_train, stratify=y_train, test_size=0.2)
```

We split the data into train, test and cross validation data sets, preserving the ratio of class distribution in the original data set

In [11]:

```
print('Number of data points in train data:', train_df.shape[0])
print('Number of data points in test data:', test_df.shape[0])
print('Number of data points in cross validation data:', cv_df.shape[0])

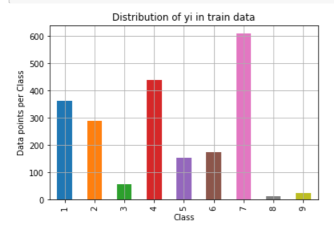
Number of data points in train data: 2124
Number of data points in test data: 665
Number of data points in cross validation data: 532
```

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

In [12]:

```
# it returns a dict, keys as class labels and values as the number of data points in that class
train_class_distribution = train_df['Class'].value_counts().sortlevel()
test_class_distribution = test_df['Class'].value_counts().sortlevel()
cv class distribution = cv df['Class'].value counts().sortlevel()
my colors = 'rgbkymc'
train_class_distribution.plot(kind='bar')
plt.xlabel('Class')
plt.ylabel('Data points per Class')
plt.title('Distribution of yi in train data')
plt.grid()
plt.show()
# 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.ro
und((train class distribution.values[i]/train df.shape[0]*100), 3), '%)')
print('-'*80)
my colors = 'rgbkymc'
test_class_distribution.plot(kind='bar')
plt.xlabel('Class')
plt.ylabel('Data points per Class')
plt.title('Distribution of yi in test data')
plt.grid()
```

```
plt.show()
# 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(-test_class_distribution.values)
for i in sorted yi:
    print('Number of data points in class', i+1, ':', test class distribution.values[i], '(', np.rou
nd((test_class_distribution.values[i]/test_df.shape[0]*100), 3), '%)')
print('-'*80)
my colors = 'rgbkymc'
cv class distribution.plot(kind='bar')
plt.xlabel('Class')
plt.ylabel('Data points per Class')
plt.title('Distribution of yi in cross validation data')
plt.grid()
plt.show()
# 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, ':',cv class distribution.values[i], '(', np.round
((cv class distribution.values[i]/cv df.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 %)

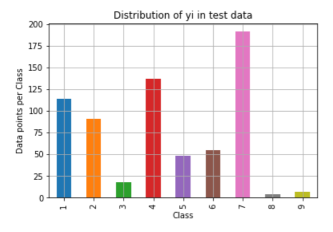
Number of data points in class 6 : 176 ( 8.286 %)

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 %)
```

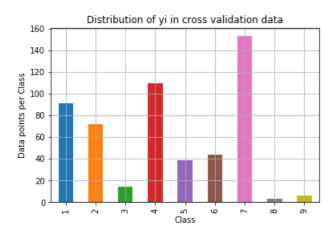
```
Number of data points in class 6: 55 (8.271 %)

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 %)
```

3.2 Prediction using a 'Random' Model

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

In [13]:

```
# This function plots the confusion matrices given y_i, y_i_hat.
def plot_confusion_matrix(test_y, predict_y):
   C = 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
   A = (((C.T)/(C.sum(axis=1))).T)
    #divid each element of the confusion matrix with the sum of elements in that column
    \# C = [[1, 2],
    \# 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
    B = (C/C.sum(axis=0))
    #divid 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]
```

```
# representing A in heatmap format
print("-"*20, "Confusion matrix", "-"*20)
plt.figure(figsize=(20,7))
sns.heatmap(C, annot=True, cmap="YlGnBu", fmt=".3f", xticklabels=labels, yticklabels=labels)
plt.xlabel('Predicted Class')
plt.ylabel('Original Class')
plt.show()
print("-"*20, "Precision matrix (Column Sum=1)", "-"*20)
plt.figure(figsize=(20,7))
sns.heatmap(B, annot=True, cmap="YlGnBu", fmt=".3f", xticklabels=labels, yticklabels=labels)
plt.xlabel('Predicted Class')
plt.ylabel('Original Class')
plt.show()
# representing B in heatmap format
print("-"*20, "Recall matrix (Row sum=1)", "-"*20)
plt.figure(figsize=(20,7))
sns.heatmap(A, annot=True, cmap="YlGnBu", fmt=".3f", xticklabels=labels, yticklabels=labels)
plt.xlabel('Predicted Class')
plt.ylabel('Original Class')
plt.show()
```

In [14]:

```
\# 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 = test_df.shape[0]
cv_data_len = cv_df.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)
```

Log loss on Cross Validation Data using Random Model 2.520718570871371 Log loss on Test Data using Random Model 2.4951496087584557 ------ Confusion matrix ------

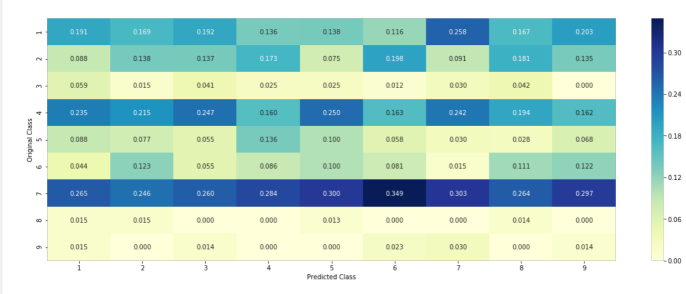
- ·	13.000	11.000	14.000	11.000	11.000	10.000	17.000	12.000	
7 -	6.000	9.000	10.000	14.000	6.000	17.000	6.000	13.000	10.000
m -	4.000	1.000	3.000	2.000	2.000	1.000	2.000	3.000	0.000
SS 4 -	16.000	14.000	18.000	13.000	20.000	14.000	16.000	14.000	12.000
Original Class 5	6.000	5.000	4.000	11.000	8.000	5.000	2.000	2.000	5.000
oric	3.000	8.000	4.000	7.000	8.000	7.000	1.000	8.000	9.000
۲.	18.000	16.000	19.000	23.000	24.000	30.000	20.000	19.000	22.000
∞ -	1.000	1.000	0.000	0.000	1.000	0.000	0.000	1.000	0.000
6 -	1.000	0.000	1.000	0.000	0.000	2.000	2.000	0.000	1.000
	i	2	3	4	5	6	7	8	9

- 15

- 10

- 5

----- Precision matrix (Columm Sum=1) -----



----- Recall matrix (Row sum=1) ------



3.3 Univariate Analysis

In [15]:

```
# code for response coding with Laplace smoothing.
# alpha : used for laplace smoothing
# feature: ['gene', 'variation']
# df: ['train_df', 'test_df', 'cv_df']
# 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 occured in class1 + 10*alpha / nu
mber of time it occurred 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):
    # walue count it contains a dist like
```

```
# print(train df['Gene'].value counts())
   # output:
           {BRCA1
                      174
            TP53
                      106
                      86
            EGFR
            BRCA2
                        7.5
                       69
            PTEN
            KTT
                       61
            BRAF
                       60
                       47
            ERBB2
            PDGFRA
   # print(train_df['Variation'].value counts())
   # output:
   # {
   # Truncating_Mutations
                                           63
   # Deletion
                                            43
   # Amplification
                                           43
   # Fusions
                                           22
                                            3
   # Overexpression
   # F.17K
                                            3
   # Q61L
   # S222D
   # P130S
                                            2
   # }
   value count = train df[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
           # 2486 2486 BRCA1
                                           S1841R
           # 2614 2614 BRCA1
                                             M1R
           # 2432 2432 BRCA1
# 2567 2567 BRCA1
# 2583 2583 BRCA1
                                           T.1657P
                                           T1685A
                                           E1660G
           # 2634 2634 BRCA1
                                           W1718L
           # cls cnt.shape[0] will return the number of rows
           cls cnt = train df.loc[(train df['Class']==k) & (train df[feature]==i)]
           # cls cnt.shape[0](numerator) will contain the number of time that particular feature (
ccured in whole data
          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, 0.068181818181818177,
0.13636363636363635,\ 0.25,\ 0.1931818181818181818,\ 0.03787878787878788,\ 0.03787878787878788,
0.03787878787878788],
         'TP53': [0.32142857142857145, 0.061224489795918366, 0.061224489795918366,
163265307, 0.056122448979591837],
  # 'EGFR': [0.05681818181818181816, 0.215909090909091, 0.0625, 0.068181818181818177,
0.068181818181818177, 0.0625, 0.34659090909090912, 0.0625, 0.05681818181818181816],
   # 'BRCA2': [0.1333333333333333, 0.0606060606060608, 0.0606060606060608,
0.07878787878787878782,\ 0.13939393939394,\ 0.345454545454546,\ 0.060606060606060608,
0.06060606060606060608, 0.060606060606060608],
  # 'PTEN': [0.069182389937106917, 0.062893081761006289, 0.069182389937106917,
761006289, 0.062893081761006289],
         'KIT': [0.066225165562913912, 0.25165562913907286, 0.072847682119205295,
0 070047600110005005 0 066005165560010010 0 066005165560010010 0 07150017000704700
```

value count. It contains a utcl like

```
U.U/204/0021132U0233, U.U00223103302312, U.U00223103302313, U.Z/13231/00U/34/UZ,
0.066225165562913912, 0.066225165562913912],
         'BRAF': [0.066666666666666666, 0.17999999999999, 0.073333333333333334,
gv dict = get gv fea dict(alpha, feature, df)
   # value_count is similar in get_gv_fea_dict
   value count = train df[feature].value counts()
   # gv fea: Gene variation feature, it will contain the feature for each feature value in the da
   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

s = sum(unique_genes.values);
h = unique_genes.values/s;

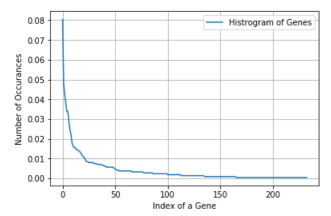
plt.xlabel('Index of a Gene')

plt.plot(h, label="Histrogram of Genes")

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

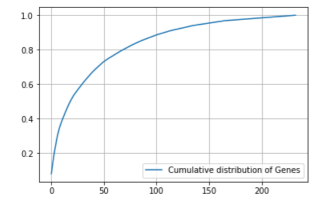
```
In [16]:
unique genes = train df['Gene'].value counts()
print('Number of Unique Genes :', unique_genes.shape[0])
# the top 10 genes that occured most
print(unique genes.head(10))
Number of Unique Genes: 233
BRCA1
          171
TP53
          102
           91
EGFR
BRCA2
           81
KTT
           72
PTEN
BRAF
           61
           52
ALK
ERBB2
           47
PDGFRA
          38
Name: Gene, dtype: int64
In [17]:
print("Ans: There are", unique_genes.shape[0], "different categories of genes in the train data, an
d they are distibuted as follows",)
                                                                                                  l Þ
Ans: There are 233 different categories of genes in the train data, and they are distibuted as fol
lows
In [18]:
```

```
plt.ylabel('Number of Occurances')
plt.legend()
plt.grid()
plt.show()
```



In [19]:

```
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 [20]:

```
# 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", train_df))
# test gene feature
test_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", test_df))
# cross validation gene feature
cv_gene_feature_responseCoding = np.array(get_gv_feature(alpha, "Gene", cv_df))
```

```
print ("train gene feature responseCoding is converted feature using respone coding method. The sha
pe of gene feature:", train_gene_feature_responseCoding.shape)
train gene feature responseCoding is converted feature using respone coding method. The shape of g
ene feature: (2124, 9)
In [22]:
# one-hot encoding of Gene feature.
gene vectorizer = TfidfVectorizer(max features = 1000)
train gene feature onehotCoding = gene vectorizer.fit transform(train df['Gene'])
test_gene_feature_onehotCoding = gene_vectorizer.transform(test_df['Gene'])
cv gene feature onehotCoding = gene vectorizer.transform(cv df['Gene'])
In [23]:
train_df['Gene'].head()
Out[23]:
1926
          SMO
2847
        BRCA2
3001
         KIT
1913
         SMO
1117
       FANCC
Name: Gene, dtype: object
In [24]:
gene_vectorizer.get_feature_names()
Out[24]:
['abl1',
 'acvr1',
 'ago2',
 'akt1',
 'akt2',
 'akt3',
 'alk',
 'apc',
 'ar',
 'araf',
 'arid1b',
 'arid2',
 'arid5b',
 'atm',
 'atr',
 'atrx',
 'aurka',
 'axl',
 'b2m',
 'bap1',
 'bard1',
 'bcl10',
 'bcl2',
 'bcl2111',
 'bcor',
 'braf',
 'brca1',
 'brca2',
 'brip1',
 'btk',
 'card11',
 'carm1',
 'casp8',
 'cbl',
 'ccnd1',
 'ccnd3',
 'ccne1',
 'cdh1',
 'cdk12',
 'cdk4',
```

```
'cdk6',
'cdk8',
'cdkn1a',
'cdkn1b',
'cdkn2a',
'cdkn2b',
'cdkn2c',
'cebpa',
'chek2',
'cic',
'crebbp',
'ctcf',
'ctla4',
'ctnnb1',
'ddr2',
'dicer1',
'dnmt3a',
'dnmt3b',
'egfr',
'elf3',
'ep300',
'epas1',
'epcam',
'erbb2',
'erbb3',
'erbb4',
'ercc2',
'ercc3',
'ercc4',
'erg',
'errfil',
'esr1',
'etv1',
'etv6',
'ewsr1',
'ezh2',
'fam58a',
'fanca',
'fancc',
'fat1',
'fbxw7',
'fgf19',
'fgf3',
'fgf4',
'fgfr1',
'fgfr2',
'fgfr3',
'fgfr4',
'flt3',
'foxa1',
'fox12',
'foxp1',
'gata3',
'gli1',
'gna11',
'gnas',
'h3f3a',
'hla',
'hnfla',
'hras',
'idh1',
'idh2',
'igf1r',
'ikbke',
'il7r',
'inpp4b',
'jak1',
'jak2',
'jun',
'kdm5a',
'kdm5c',
'kdm6a',
'kdr',
'keap1',
'kit',
'klf4',
'kmt2a',
```

```
'kmt2c',
'knstrn',
'kras',
'lats1',
'lats2',
'map2k1',
'map2k2',
'map2k4',
'map3k1',
'mapk1',
'mdm2',
'mdm4',
'med12',
'mef2b',
'met',
'mga',
'mlh1',
'msh2',
'msh6',
'mtor',
'myc',
'mycn',
'myd88',
'myod1',
'nf1',
'nf2',
'nfe212',
'nfkbia',
'nkx2',
'notch1',
'notch2',
'npm1',
'nras',
'nsd1',
'ntrk1',
'ntrk2',
'ntrk3',
'nup93',
'pbrm1',
'pdgfra',
'pdgfrb',
'pik3ca',
'pik3cb',
'pik3cd',
'pik3r1',
'pik3r2',
'pim1',
'pms1',
'pms2',
'pole',
'ppmld',
'ppp2r1a',
'prdm1',
'ptch1',
'pten',
'ptpn11',
'ptprd',
'ptprt',
'rac1',
'rad21',
'rad50',
'rad51c',
'rad51d',
'rad541',
'raf1',
'rasa1',
'rb1',
'rbm10',
'ret',
'rhoa',
'rictor',
'rit1',
'ros1',
'runx1',
'rxra',
'rybp',
'sdhb',
```

```
'setd2',
 'sf3b1',
 'shoc2',
 'shq1',
 'smad2',
 'smad3',
 'smad4',
 'smarca4',
 'smarcb1',
 'smo',
 'sos1',
 'sox9',
 'spop',
 'src',
 'srsf2',
 'stag2',
 'stat3',
 'stk11',
 'tcf3',
 'tcf712',
 'tert',
 'tet1',
 'tet2',
 'tgfbr1',
 'tgfbr2',
 'tmprss2',
 'tp53',
 'tp53bp1',
 'tsc1',
 'tsc2',
 'u2af1',
 'vhl',
 'whsc1'
 'whsc1l1'
 'xpol',
 'xrcc2',
 'yap1']
In [25]:
print("train gene feature onehotCoding is converted feature using one-hot encoding method. The sha
pe of gene feature:", train gene feature onehotCoding.shape)
```

train_gene_feature_onehotCoding is converted feature using one-hot encoding method. The shape of g ene feature: (2124, 232)

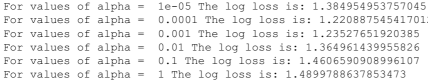
Q4. How good is this gene feature in predicting y_i?

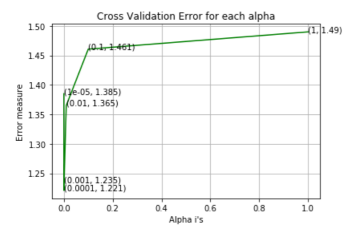
There are many ways to estimate how good a feature is, in predicting y_i. One of the good methods is to build a proper ML model using just this feature. In this case, we will build a logistic regression model using only Gene feature (one hot encoded) to predict y_i.

In [26]:

'sdhc',

```
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.clas
ses_, 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 lo
ss(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, p
redict y, labels=clf.classes , eps=1e-15))
For values of alpha = 1e-05 The log loss is: 1.384954953757045
For values of alpha = 0.0001 The log loss is: 1.220887545417012
For values of alpha = 0.001 The log loss is: 1.23527651920385
```





```
For values of best alpha = 0.0001 The train log loss is: 1.0526060099705743 For values of best alpha = 0.0001 The cross validation log loss is: 1.220887545417012 For values of best alpha = 0.0001 The test log loss is: 1.1843654625498663
```

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.

```
print("Q6. How many data points in Test and CV datasets are covered by the ", unique_genes.shape[0], " genes in train dataset?")

test_coverage=test_df[test_df['Gene'].isin(list(set(train_df['Gene'])))].shape[0]

cv_coverage=cv_df[cv_df['Gene'].isin(list(set(train_df['Gene'])))].shape[0]

print('Ans\n1. In test data',test_coverage, 'out of',test_df.shape[0], ":",(test_coverage/test_df.shape[0])*100)

print('2. In cross validation data',cv_coverage, 'out of ',cv_df.shape[0],":",(cv_coverage/cv_df.shape[0])*100)

Q6. How many data points in Test and CV datasets are covered by the 233 genes in train dataset?
Ans
1. In test data 643 out of 665 : 96.69172932330827
2. In cross validation data 512 out of 532 : 96.2406015037594
```

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 [28]:
```

```
unique_variations = train_df['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 : 1919
Truncating_Mutations 67
Deletion 52
Amplification 46
Fusions 26
M1R 2
EWSR1-ETV1_Fusion 2
Q61H 2
G12V 2
Promoter_Hypermethylation 2
T167A 2
Name: Variation, dtype: int64
```

In [29]:

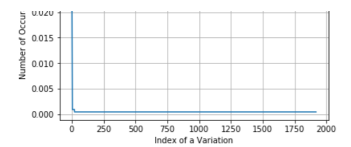
```
print("Ans: There are", unique_variations.shape[0] ,"different categories of variations in the
train data, and they are distibuted as follows",)
```

Ans: There are 1919 different categories of variations in the train data, and they are distibuted as follows

In [30]:

```
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()
```

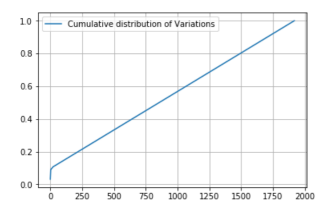
0.030		— Histrogram of Variations			
0.025 +					



In [31]:

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

```
[0.03154426 0.05602637 0.07768362 ... 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 [32]:

```
# alpha is used for laplace smoothing
alpha = 1
# train gene feature
train_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", train_df))
# test gene feature
test_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", test_df))
# cross validation gene feature
cv_variation_feature_responseCoding = np.array(get_gv_feature(alpha, "Variation", cv_df))
```

In [33]:

```
print("train_variation_feature_responseCoding is a converted feature using the response coding met
hod. The shape of Variation feature:", train_variation_feature_responseCoding.shape)
```

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

In [34]:

```
# one-hot encoding of variation feature.
variation_vectorizer = TfidfVectorizer(max_features = 1000)
train_variation_feature_onehotCoding = variation_vectorizer.fit_transform(train_df['Variation'])
test_variation_feature_onehotCoding = variation_vectorizer.transform(test_df['Variation'])
cv_variation_feature_onehotCoding = variation_vectorizer.transform(cv_df['Variation'])
```

In [35]:

```
print("train_variation_feature_onehotEncoded is converted feature using the onne-hot encoding meth
od. The shape of Variation feature:", train_variation_feature_onehotCoding.shape)
```

train_variation_feature_onehotEncoded is converted feature using the onne-hot encoding method. The shape of Variation feature: (2124, 1000)

Q10. How good is this Variation feature in predicting y i?

Let's build a model just like the earlier!

```
In [36]:
```

```
alpha = [10 ** x for x in range(-5, 1)]
# read more about SGDClassifier() at http://scikit-
learn.org/stable/modules/generated/sklearn.linear model.SGDClassifier.html
# default parameters
# SGDClassifier(loss='hinge', penalty='12', alpha=0.0001, 11 ratio=0.15, fit intercept=True, max i
ter=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.clas
ses_, 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.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=clf.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, p redict_y, labels=clf.classes_, eps=1e-15))
```

```
For values of alpha = 1e-05 The log loss is: 1.737476973365291

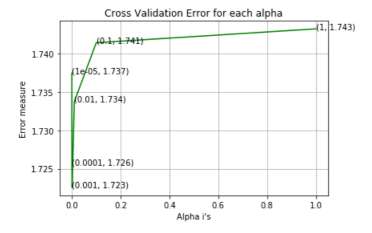
For values of alpha = 0.0001 The log loss is: 1.7255070794020986

For values of alpha = 0.001 The log loss is: 1.7225571047902302

For values of alpha = 0.01 The log loss is: 1.7337517466397754

For values of alpha = 0.1 The log loss is: 1.741443301227446

For values of alpha = 1 The log loss is: 1.7432509768377
```



```
For values of best alpha = 0.001 The train log loss is: 1.3611687786959856
For values of best alpha = 0.001 The cross validation log loss is: 1.7225571047902302
For values of best alpha = 0.001 The test log loss is: 1.74246364126082
```

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.

```
In [37]:
```

```
print("Q12. How many data points are covered by total ", unique_variations.shape[0], " variation i
n test and cross validation data sets?")
test_coverage=test_df[test_df['Variation'].isin(list(set(train_df['Variation'])))].shape[0]
cv_coverage=cv_df[cv_df['Variation'].isin(list(set(train_df['Variation'])))].shape[0]
print('Ans\n1. In test data',test_coverage, 'out of',test_df.shape[0], ":",(test_coverage/test_df.shape[0])*100)
print('2. In cross validation data',cv_coverage, 'out of ',cv_df.shape[0],":",(cv_coverage/cv_df.shape[0])*100)
```

Q12. How many data points are covered by total 1919 variation in test and cross validation data sets?

Ans

- 1. In test data 62 out of 665 : 9.323308270676693
- 2. In cross validation data 53 out of 532 : 9.962406015037594

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?

In [38]:

```
# cls_text is a data frame
# for every row in data fram consider the 'TEXT'
```

In [39]:

In [40]:

```
# building a CountVectorizer with all the words that occured minimum 3 times in train data
text_vectorizer = TfidfVectorizer(min_df=3,max_features=1000)
train_text_feature_onehotCoding = text_vectorizer.fit_transform(train_df['TEXT'])
# getting all the feature names (words)
train_text_features= text_vectorizer.get_feature_names()

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

# 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 [41]:

```
dict list = []
# dict list =[] contains 9 dictoinaries each corresponds to a class
for i in range(1,10):
    cls text = train df[train df['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(train_df)
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)
```

```
#response coding of text features
train_text_feature_responseCoding = get_text_responsecoding(train_df)
test_text_feature_responseCoding = get_text_responsecoding(test_df)
cv_text_feature_responseCoding = get_text_responsecoding(cv_df)
```

In [43]:

```
# 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 [44]:

```
# 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(test_df['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(cv_df['TEXT'])
# don't forget to normalize every feature
cv_text_feature_onehotCoding = normalize(cv_text_feature_onehotCoding, axis=0)
```

In [45]:

```
#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 [46]:

```
# Number of words for a given frequency.
print(Counter(sorted_text_occur))
```

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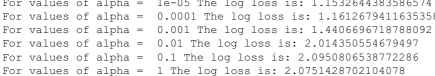
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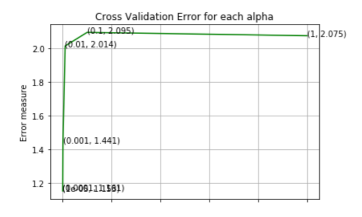
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1836078: 1, 6.299693823083686: 1})
4
```

In [47]:

```
π ασταυτι Ραταιισιστο
# SGDClassifier(loss='hinge', penalty='12', alpha=0.0001, 11_ratio=0.15, fit_intercept=True, max_i
ter=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.clas
ses_, 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=clf.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_lo
ss(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, p
redict_y, labels=clf.classes_, eps=1e-15))
For values of alpha = 1e-05 The log loss is: 1.1532644383586574
For values of alpha = 0.0001 The log loss is: 1.1612679411635358
For values of alpha = 0.001 The log loss is: 1.4406696718788092
For values of alpha = 0.01 The log loss is: 2.014350554679497
```





```
Alphai's

For values of best alpha = 1e-05 The train log loss is: 0.7412786504348623

For values of best alpha = 1e-05 The cross validation log loss is: 1.1532644383586574

For values of best alpha = 1e-05 The test log loss is: 1.1984396955548693
```

1.0

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

0.8

Ans. Yes, it seems like!

0.2

0.4

0.6

0.0

```
In [48]:
```

```
def get_intersec_text(df):
    df_text_vec = TfidfVectorizer(max_features = 1000, min_df = 3)
    df_text_fea = df_text_vec.fit_transform(df['TEXT'])
    df_text_features = df_text_vec.get_feature_names()

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 [49]:

```
len1,len2 = get_intersec_text(test_df)
print(np.round((len2/len1)*100, 3), "% of word of test data appeared in train data")
len1,len2 = get_intersec_text(cv_df)
print(np.round((len2/len1)*100, 3), "% of word of Cross Validation appeared in train data")
```

94.3 % of word of test data appeared in train data 93.6 % of word of Cross Validation appeared in train data

4. Machine Learning Models

In [63]:

```
#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 [64]:

```
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=le-15)
```

In [65]:

```
# 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 impleature names(indices, text, gene, var, no features):
    gene_count_vec = TfidfVectorizer(max_features = 1000)
    var count vec = TfidfVectorizer(max features = 1000)
    text count vec = TfidfVectorizer(min df=3, max features = 1000)
    gene vec = gene count vec.fit(train df['Gene'])
    var_vec = var_count_vec.fit(train_df['Variation'])
    text_vec = text_count_vec.fit(train_df['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_r
0))
        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")
4
```

Stacking the three types of features

In [66]:

```
# 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)).tocs
train_y = np.array(list(train_df['Class']))
test x onehotCoding = hstack((test gene var onehotCoding, test text feature onehotCoding)).tocsr()
test_y = np.array(list(test_df['Class']))
cv_x_onehotCoding = hstack((cv_gene_var_onehotCoding, cv_text_feature_onehotCoding)).tocsr()
cv_y = np.array(list(cv_df['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 [67]:
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, 2232)
(number of data points * number of features) in test data = (665, 2232)
(number of data points * number of features) in cross validation data = (532, 2232)
In [68]:
print(" Response encoding features :")
print("(number of data points * number of features) in train data = ", train x responseCoding.shap
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)
```

4.1. Base Line Model

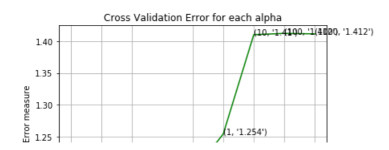
4.1.1. Naive Bayes

4.1.1.1. Hyper parameter tuning

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.CalibratedClassifier CV.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/
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))
    {\it \# 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=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 lo
ss(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, p
redict y, labels=clf.classes , eps=1e-15))
for alpha = 1e-05
Log Loss : 1.164111647557351
for alpha = 0.0001
Log Loss : 1.1636446127113975
for alpha = 0.001
Log Loss: 1.1627400999467232
for alpha = 0.1
Log Loss: 1.1924304270718766
for alpha = 1
Log Loss: 1.2541366464909083
for alpha = 10
Log Loss: 1.4103758317317077
for alpha = 100
Log Loss: 1.4124142354020242
for alpha = 1000
```



Log Loss : 1.4116634312711283

```
1.20 (1e-05. (0.064)(0.0064)1.163')

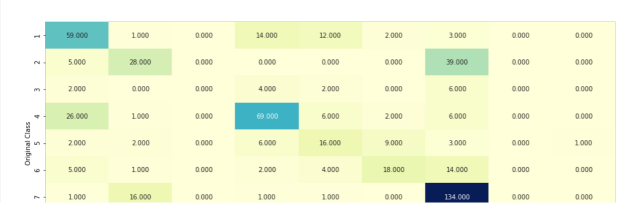
1.15 (-5 -4 -3 -1 0 1 2 3 Alpha i's
```

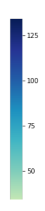
```
For values of best alpha = 0.001 The train log loss is: 0.7389550367162582
For values of best alpha = 0.001 The cross validation log loss is: 1.1627400999467232
For values of best alpha = 0.001 The test log loss is: 1.1886205094193842
```

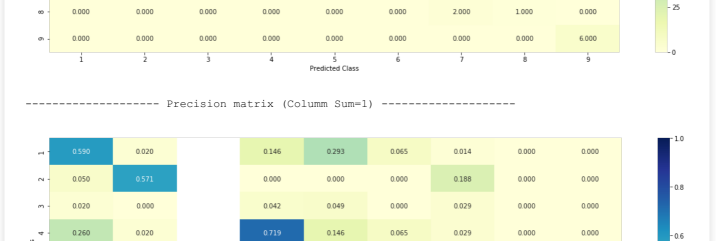
4.1.1.2. Testing the model with best hyper paramters

In [70]:

```
# 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()))
```







0.390

0.098

0.024

0.000

0.000

Predicted Class

0.290

0.000

0.000

0.000

0.014

0.068

0.010

0.000

0.000

0.000

0.000

0.000

0.143

0.000

0.000

0.000

- 0.2

----- Recall matrix (Row sum=1) ------

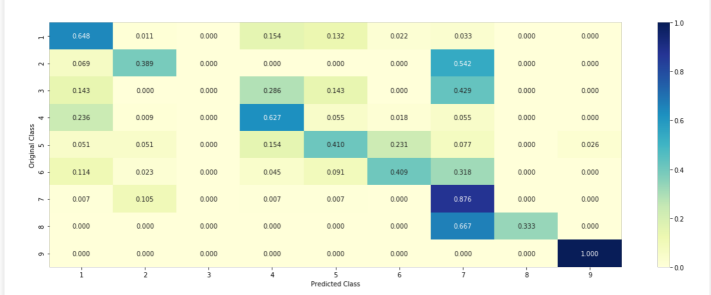
0.062

0.021

0.010

0.000

0.000



4.1.1.3. Feature Importance, Correctly classified point

```
In [71]:
```

Original C

0.020

0.050

0.010

0.000

0.000

0.041

0.020

0.327

0.000

0.000

```
test point index = 1
no feature = 100
predicted_cls = sig_clf.predict(test_x_onehotCoding[test_point_index])
print("Predicted Class :", predicted cls[0])
print("Predicted Class Probabilities:",
\verb"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],
test_df['TEXT'].iloc[test_point_index], test_df['Gene'].iloc[test_point_index], test_df['Variation']
.iloc[test point index], no feature)
```

Predicted Class : 4

```
10 Text feature [activity] present in test data point [True]
11 Text feature [protein] present in test data point [True]
13 Text feature [proteins] present in test data point [True]
14 Text feature [function] present in test data point [True]
15 Text feature [missense] present in test data point [True]
16 Text feature [results] present in test data point [True]
17 Text feature [whereas] present in test data point [True]
18 Text feature [experiments] present in test data point [True]
19 Text feature [acid] present in test data point [True]
20 Text feature [amino] present in test data point [True]
21 Text feature [suppressor] present in test data point [True]
24 Text feature [type] present in test data point [True]
25 Text feature [also] present in test data point [True]
26 Text feature [functional] present in test data point [True]
27 Text feature [shown] present in test data point [True]
28 Text feature [whether] present in test data point [True]
29 Text feature [described] present in test data point [True]
31 Text feature [wild] present in test data point [True]
32 Text feature [two] present in test data point [True]
33 Text feature [important] present in test data point [True]
34 Text feature [related] present in test data point [True]
35 Text feature [reduced] present in test data point [True]
36 Text feature [mutations] present in test data point [True]
37 Text feature [determined] present in test data point [True]
39 Text feature [either] present in test data point [True]
40 Text feature [may] present in test data point [True]
42 Text feature [ability] present in test data point [True]
43 Text feature [although] present in test data point [True]
44 Text feature [three] present in test data point [True]
45 Text feature [previously] present in test data point [True]
46 Text feature [determine] present in test data point [True]
47 Text feature [loss] present in test data point [True]
48 Text feature [containing] present in test data point [True]
49 Text feature [discussion] present in test data point [True]
50 Text feature [associated] present in test data point [True]
51 Text feature [therefore] present in test data point [True]
52 Text feature [analysis] present in test data point [True]
53 Text feature [levels] present in test data point [True]
54 Text feature [thus] present in test data point [True]
57 Text feature [mammalian] present in test data point [True]
58 Text feature [vitro] present in test data point [True]
63 Text feature [one] present in test data point [True]
65 Text feature [effect] present in test data point [True]
66 Text feature [lower] present in test data point [True]
67 Text feature [contribute] present in test data point [True]
68 Text feature [show] present in test data point [True]
69 Text feature [site] present in test data point [True]
70 Text feature [however] present in test data point [True]
71 Text feature [30] present in test data point [True]
72 Text feature [several] present in test data point [True]
74 Text feature [effects] present in test data point [True]
75 Text feature [indicated] present in test data point [True]
77 Text feature [similar] present in test data point [True]
78 Text feature [could] present in test data point [True]
80 Text feature [binding] present in test data point [True]
81 Text feature [introduction] present in test data point [True]
82 Text feature [possible] present in test data point [True]
83 Text feature [10] present in test data point [True]
85 Text feature [mutant] present in test data point [True]
86 Text feature [general] present in test data point [True]
87 Text feature [cells] present in test data point [True]
88 Text feature [addition] present in test data point [True]
89 Text feature [dna] present in test data point [True]
90 Text feature [bind] present in test data point [True]
91 Text feature [mutation] present in test data point [True]
92 Text feature [changes] present in test data point [True]
93 Text feature [found] present in test data point [True]
94 Text feature [see] present in test data point [True]
95 Text feature [substitutions] present in test data point [True]
96 Text feature [50] present in test data point [True]
97 Text feature [stability] present in test data point [True]
98 Text feature [mutants] present in test data point [True]
99 Text feature [terminal] present in test data point [True]
Out of the top 100 features 73 are present in query point
```

4.1.1.4. Feature Importance, Incorrectly classified point

In [72]: test_point_index = 100 no feature = 100 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], test df['TEXT'].iloc[test point index],test df['Gene'].iloc[test point index],test df['Variation'] .iloc[test point index], no feature) Predicted Class: 1 Predicted Class Probabilities: [[0.7247 0.0507 0.0131 0.0682 0.0339 0.0335 0.0694 0.0026 0.004]] Actual Class : 1 11 Text feature [one] present in test data point [True] 13 Text feature [results] present in test data point [True] 14 Text feature [therefore] present in test data point [True] 15 Text feature [protein] present in test data point [True] 16 Text feature [function] present in test data point [True] 17 Text feature [also] present in test data point [True] 18 Text feature [two] present in test data point [True] 19 Text feature [specific] present in test data point [True] 20 Text feature [however] present in test data point [True] 21 Text feature [role] present in test data point [True] 22 Text feature [type] present in test data point [True] 23 Text feature [region] present in test data point [True] 24 Text feature [table] present in test data point [True] 25 Text feature [whether] present in test data point [True] 26 Text feature [control] present in test data point [True] 27 Text feature [using] present in test data point [True] 28 Text feature [human] present in test data point [True] 29 Text feature [functions] present in test data point [True] 31 Text feature [may] present in test data point [True] 32 Text feature [either] present in test data point [True] 33 Text feature [shown] present in test data point [True] 34 Text feature [well] present in test data point [True] 35 Text feature [least] present in test data point [True] 36 Text feature [dna] present in test data point [True] 37 Text feature [essential] present in test data point [True] 38 Text feature [possible] present in test data point [True] 39 Text feature [discussion] present in test data point [True] 40 Text feature [deletion] present in test data point [True] 42 Text feature [wild] present in test data point [True] 43 Text feature [performed] present in test data point [True] 44 Text feature [effect] present in test data point [True] 45 Text feature [result] present in test data point [True] 46 Text feature [suggest] present in test data point [True] 48 Text feature [25] present in test data point [True] 49 Text feature [large] present in test data point [True] 50 Text feature [although] present in test data point [True] 52 Text feature [gene] present in test data point [True] 53 Text feature [proteins] present in test data point [True] 54 Text feature [expression] present in test data point [True] 55 Text feature [determined] present in test data point [True] 56 Text feature [containing] present in test data point [True] 57 Text feature [present] present in test data point [True] 58 Text feature [including] present in test data point [True] 59 Text feature [within] present in test data point [True] 60 Text feature [analysis] present in test data point [True] 61 Text feature [three] present in test data point [True] 62 Text feature [fig] present in test data point [True] 63 Text feature [similar] present in test data point [True] 64 Text feature [important] present in test data point [True] 65 Text feature [reduced] present in test data point [True] 66 Text feature [used] present in test data point [True]

67 Text feature [cell] present in test data point [True]
68 Text feature [mediated] present in test data point [True]

```
tene teacate [meatacea] probene in cobe aaca poinc
69 Text feature [another] present in test data point [True]
70 Text feature [data] present in test data point [True]
71 Text feature [different] present in test data point [True]
72 Text feature [together] present in test data point [True]
73 Text feature [previously] present in test data point [True]
75 Text feature [several] present in test data point [True]
76 Text feature [addition] present in test data point [True]
77 Text feature [indicated] present in test data point [True]
78 Text feature [example] present in test data point [True]
79 Text feature [additional] present in test data point [True]
80 Text feature [likely] present in test data point [True]
81 Text feature [described] present in test data point [True]
82 Text feature [30] present in test data point [True]
83 Text feature [whereas] present in test data point [True]
84 Text feature [15] present in test data point [True]
85 Text feature [significant] present in test data point [True]
86 Text feature [corresponding] present in test data point [True]
87 Text feature [observed] present in test data point [True]
88 Text feature [mutations] present in test data point [True]
89 Text feature [cancer] present in test data point [True]
90 Text feature [associated] present in test data point [True]
91 Text feature [previous] present in test data point [True]
92 Text feature [required] present in test data point [True]
93 Text feature [yet] present in test data point [True]
94 Text feature [furthermore] present in test data point [True]
95 Text feature [directly] present in test data point [True]
96 Text feature [indicating] present in test data point [True]
97 Text feature [first] present in test data point [True]
98 Text feature [respectively] present in test data point [True]
99 Text feature [ability] present in test data point [True]
Out of the top 100 features 83 are present in query point
```

4.2. K Nearest Neighbour Classification

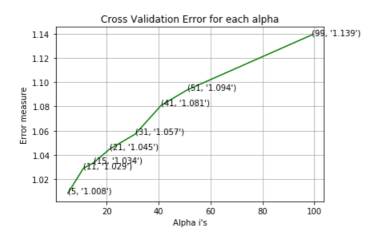
4.2.1. Hyper parameter tuning

In [73]:

```
# 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-ne
ighbors-geometric-intuition-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 arrav = []
```

```
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 lo
ss(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, p
redict y, labels=clf.classes , eps=1e-15))
for alpha = 5
Log Loss: 1.0082901840724636
```

```
Log Loss: 1.0082901840724636
for alpha = 11
Log Loss: 1.0290099490795332
for alpha = 15
Log Loss: 1.0336422613933147
for alpha = 21
Log Loss: 1.044671240525676
for alpha = 31
Log Loss: 1.0572406169994706
for alpha = 41
Log Loss: 1.080967997199732
for alpha = 51
Log Loss: 1.0936172427068644
for alpha = 99
Log Loss: 1.1389937619186208
```



```
For values of best alpha = 5 The train log loss is: 0.4755178649543858

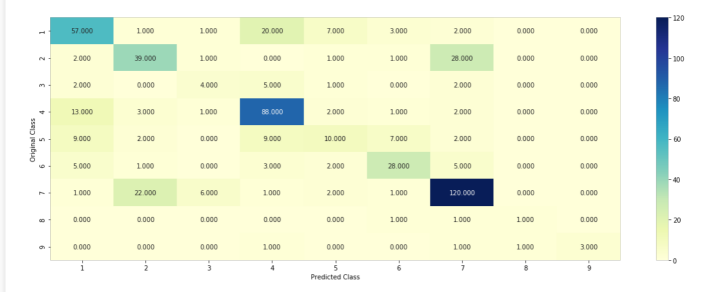
For values of best alpha = 5 The cross validation log loss is: 1.0082901840724636

For values of best alpha = 5 The test log loss is: 1.1184776899071618
```

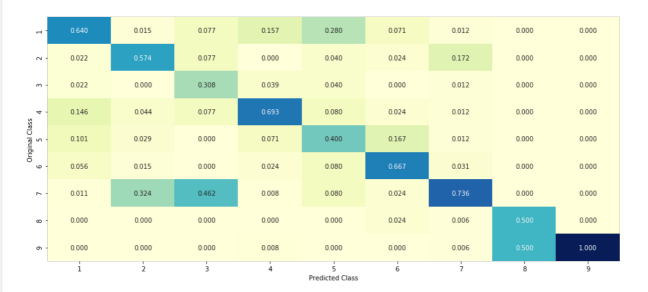
TOT VALUED OF DEDU AIPHA OF THE COOK TOG TOOD ID. I.IIO. 1, 100550, 1010

4.2.2. Testing the model with best hyper paramters

In [74]:



----- Precision matrix (Columm Sum=1) -----



- 0.8

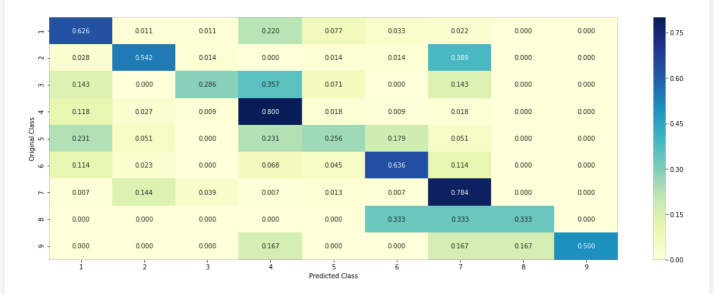
- 0.6

- 0.4

- 0.2

n 11 / /n 11

----- (kecall matrix (kow sum=1) ------



4.2.3. Sample Query point -1

```
In [75]:
```

11

```
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]])
print("Fequency of nearest points :",Counter(train_y[neighbors[1][0]]))

Predicted Class : 4
Actual Class : 4
```

The 5 nearest neighbours of the test points belongs to classes [4 4 4 4 4]

4.2.4. Sample Query Point-2

Fequency of nearest points : Counter({4: 5})

```
In [76]:
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
1)
print("the k value for knn is", alpha[best alpha], "and the nearest neighbours of the test points be
longs to classes",train y[neighbors[1][0]])
print("Fequency of nearest points :",Counter(train y[neighbors[1][0]]))
Predicted Class: 1
Actual Class : 1
the k value for knn is 5 and the nearest neighbours of the test points belongs to classes [1 1 1 1
```

4.3. Logistic Regression

4.3.1. With Class balancing

4.3.1.1. Hyper paramter tuning

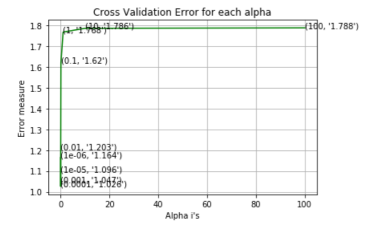
In [77]:

```
# read more about SGDClassifier() at http://scikit-
learn.org/stable/modules/generated/sklearn.linear model.SGDClassifier.html
# default parameters
# SGDClassifier(loss='hinge', penalty='12', alpha=0.0001, 11 ratio=0.15, fit intercept=True, max i
ter=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-in
tuition-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)
    \verb|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', ran
```

```
dom_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, p_redict_y, labels=clf.classes_, eps=1e-15))
```

```
for alpha = 1e-06
Log Loss: 1.1642713953663995
for alpha = 1e-05
Log Loss: 1.0958173677051277
for alpha = 0.0001
Log Loss: 1.0255817916912926
for alpha = 0.001
Log Loss : 1.0468027218518803
for alpha = 0.01
Log Loss: 1.2031750288141916
for alpha = 0.1
Log Loss: 1.6202351308931224
for alpha = 1
Log Loss : 1.7679919763730352
for alpha = 10
Log Loss: 1.7862074549485683
for alpha = 100
Log Loss: 1.7882999037580773
```



```
For values of best alpha = 0.0001 The train log loss is: 0.5861623344161264 For values of best alpha = 0.0001 The cross validation log loss is: 1.0255817916912926 For values of best alpha = 0.0001 The test log loss is: 1.044162743983839
```

4.3.1.2. Testing the model with best hyper paramters

In [78]:

125

- 100

- 75

- 50

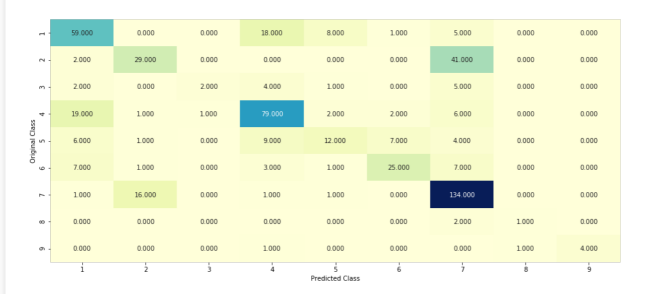
- 25

0.75

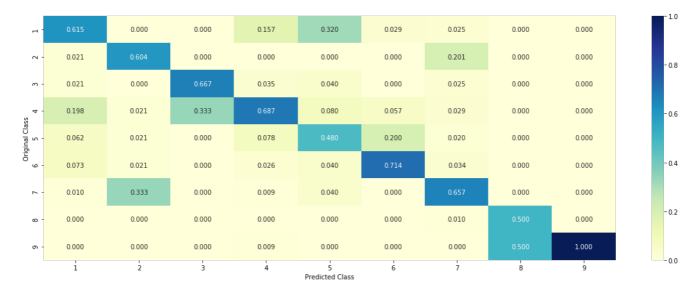
- 0.60

0.45

- 0.30

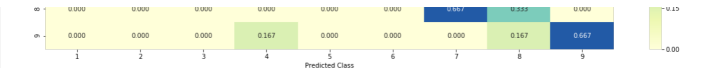


----- Precision matrix (Columm Sum=1) ------



----- Recall matrix (Row sum=1) ------





4.3.1.3. Feature Importance

```
In [79]:
```

```
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"])
        elif i< 18:
            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 [80]:
# from tabulate import tabulate
clf = SGDClassifier(class weight='balanced', alpha=alpha[best alpha], penalty='12', loss='log', ran
dom state=42)
clf.fit(train x onehotCoding,train y)
test_point_index = 1
no feature = 500
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],
test_df['TEXT'].iloc[test_point_index],test_df['Gene'].iloc[test_point_index],test_df['Variation']
.iloc[test point index], no feature)
Predicted Class: 4
Predicted Class Probabilities: [[0.0324 0.023 0.0067 0.8869 0.0195 0.0131 0.0148 0.0018 0.0017]]
Actual Class : 4
3 Text feature [suppressor] present in test data point [True]
17 Text feature [missense] present in test data point [True]
38 Text feature [families] present in test data point [True]
73 Text feature [mm] present in test data point [True]
80 Text feature [inactivation] present in test data point [True]
89 Text feature [mammalian] present in test data point [True]
92 Text feature [see] present in test data point [True]
99 Text feature [protein] present in test data point [True]
105 Text feature [yeast] present in test data point [True]
112 Text feature [null] present in test data point [True]
122 Text feature [proportion] present in test data point [True]
125 Text feature [homozygous] present in test data point [True]
133 Text feature [reduced] present in test data point [True]
147 Text feature [dna] present in test data point [True]
153 Text feature [consequences] present in test data point [True]
154 Text feature [family] present in test data point [True]
```

```
165 Text feature [localization] present in test data point [True]
168 Text feature [changes] present in test data point [True]
173 Text feature [germline] present in test data point [True]
189 Text feature [functional] present in test data point [True]
193 Text feature [appears] present in test data point [True]
194 Text feature [loss] present in test data point [True]
195 Text feature [require] present in test data point [True]
205 Text feature [represent] present in test data point [True]
219 Text feature [bind] present in test data point [True]
222 Text feature [cycle] present in test data point [True]
224 Text feature [kinases] present in test data point [True]
225 Text feature [show] present in test data point [True]
226 Text feature [38] present in test data point [True]
228 Text feature [specifically] present in test data point [True]
230 Text feature [plates] present in test data point [True]
232 Text feature [determine] present in test data point [True]
233 Text feature [1998] present in test data point [True]
234 Text feature [contribute] present in test data point [True]
237 Text feature [stability] present in test data point [True]
249 Text feature [plasmid] present in test data point [True]
255 Text feature [stained] present in test data point [True]
263 Text feature [2010] present in test data point [True]
266 Text feature [high] present in test data point [True]
271 Text feature [investigated] present in test data point [True]
279 Text feature [displayed] present in test data point [True]
280 Text feature [affected] present in test data point [True]
282 Text feature [defective] present in test data point [True]
291 Text feature [bound] present in test data point [True]
292 Text feature [risk] present in test data point [True]
293 Text feature [26] present in test data point [True]
294 Text feature [cannot] present in test data point [True]
299 Text feature [particular] present in test data point [True]
302 Text feature [several] present in test data point [True]
304 Text feature [western] present in test data point [True]
305 Text feature [cases] present in test data point [True]
309 Text feature [function] present in test data point [True]
314 Text feature [transfected] present in test data point [True]
315 Text feature [terminal] present in test data point [True]
321 Text feature [cellular] present in test data point [True]
324 Text feature [site] present in test data point [True]
333 Text feature [view] present in test data point [True]
334 Text feature [splice] present in test data point [True]
338 Text feature [early] present in test data point [True]
339 Text feature [observed] present in test data point [True]
340 Text feature [amino] present in test data point [True]
342 Text feature [relative] present in test data point [True]
343 Text feature [lower] present in test data point [True]
345 Text feature [comparison] present in test data point [True]
346 Text feature [red] present in test data point [True]
353 Text feature [37] present in test data point [True]
356 Text feature [associated] present in test data point [True]
362 Text feature [figure] present in test data point [True]
363 Text feature [mutants] present in test data point [True]
368 Text feature [described] present in test data point [True]
371 Text feature [altered] present in test data point [True]
375 Text feature [sds] present in test data point [True]
377 Text feature [involved] present in test data point [True]
380 Text feature [nuclear] present in test data point [True]
385 Text feature [inhibitory] present in test data point [True]
386 Text feature [groups] present in test data point [True]
392 Text feature [mouse] present in test data point [True]
394 Text feature [2a] present in test data point [True]
400 Text feature [low] present in test data point [True]
402 Text feature [transfection] present in test data point [True]
404 Text feature [cdk4] present in test data point [True]
405 Text feature [linked] present in test data point [True]
408 Text feature [antibodies] present in test data point [True]
409 Text feature [3b] present in test data point [True]
412 Text feature [similarly] present in test data point [True]
413 Text feature [times] present in test data point [True]
415 Text feature [isolated] present in test data point [True]
416 Text feature [2000] present in test data point [True]
418 Text feature [45] present in test data point [True]
421 Text feature [containing] present in test data point [True]
429 Text feature [genomic] present in test data point [True]
430 Text feature [buffer] present in test data point [True]
435 Text feature [indeed] present in test data point [True]
```

```
436 Text feature [40] present in test data point [True]
437 Text feature [pathogenic] present in test data point [True]
441 Text feature [key] present in test data point [True]
443 Text feature [phosphorylation] present in test data point [True]
444 Text feature [alterations] present in test data point [True]
445 Text feature [larger] present in test data point [True]
449 Text feature [decrease] present in test data point [True]
453 Text feature [partial] present in test data point [True]
454 Text feature [ca] present in test data point [True]
455 Text feature [flag] present in test data point [True]
457 Text feature [induced] present in test data point [True]
459 Text feature [genetic] present in test data point [True]
460 Text feature [29] present in test data point [True]
462 Text feature [via] present in test data point [True]
463 Text feature [gst] present in test data point [True]
465 Text feature [locus] present in test data point [True]
466 Text feature [individuals] present in test data point [True]
469 Text feature [proteins] present in test data point [True]
471 Text feature [distribution] present in test data point [True]
472 Text feature [reporter] present in test data point [True]
473 Text feature [assay] present in test data point [True]
474 Text feature [average] present in test data point [True]
481 Text feature [deletion] present in test data point [True]
483 Text feature [variant] present in test data point [True]
485 Text feature [whether] present in test data point [True]
488 Text feature [consistent] present in test data point [True]
490 Text feature [green] present in test data point [True]
496 Text feature [predicted] present in test data point [True]
499 Text feature [assays] present in test data point [True]
Out of the top 500 features 122 are present in query point
```

4.3.1.3.2. Incorrectly Classified point

```
In [81]:
test_point_index = 100
no feature = 500
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],
test df['TEXT'].iloc[test point index],test df['Gene'].iloc[test point index],test df['Variation']
.iloc[test point index], no feature)
Predicted Class: 1
Predicted Class Probabilities: [[9.648e-01 1.880e-02 1.000e-04 1.380e-02 1.300e-03 2.000e-04 9.000
 1.000e-04 0.000e+0011
Actual Class : 1
10 Text feature [panel] present in test data point [True]
52 Text feature [nuclear] present in test data point [True]
62 Text feature [deletion] present in test data point [True]
65 Text feature [repeats] present in test data point [True]
68 Text feature [rather] present in test data point [True]
99 Text feature [ability] present in test data point [True]
109 Text feature [therefore] present in test data point [True]
112 Text feature [defined] present in test data point [True]
117 Text feature [box] present in test data point [True]
121 Text feature [essential] present in test data point [True]
141 Text feature [functions] present in test data point [True]
144 Text feature [effect] present in test data point [True]
146 Text feature [21] present in test data point [True]
150 Text feature [deficient] present in test data point [True]
152 Text feature [corresponding] present in test data point [True]
154 Text feature [region] present in test data point [True]
155 Text feature [strong] present in test data point [True]
156 Text feature [analyses] present in test data point [True]
163 Text feature [mutational] present in test data point [True]
165 Text feature [17] present in test data point [True]
```

```
1/3 Text reature [page] present in test data point [True]
179 Text feature [calculated] present in test data point [True]
181 Text feature [indicated] present in test data point [True]
182 Text feature [notably] present in test data point [True]
187 Text feature [whole] present in test data point [True]
188 Text feature [gel] present in test data point [True]
191 Text feature [subjected] present in test data point [True]
192 Text feature [function] present in test data point [True]
194 Text feature [position] present in test data point [True]
195 Text feature [value] present in test data point [True]
196 Text feature [sequenced] present in test data point [True]
197 Text feature [change] present in test data point [True]
201 Text feature [impaired] present in test data point [True]
203 Text feature [identify] present in test data point [True]
204 Text feature [located] present in test data point [True]
206 Text feature [encoding] present in test data point [True]
218 Text feature [previous] present in test data point [True]
221 Text feature [percentage] present in test data point [True]
222 Text feature [showing] present in test data point [True]
223 Text feature [blood] present in test data point [True]
226 Text feature [splice] present in test data point [True]
227 Text feature [interact] present in test data point [True]
228 Text feature [splicing] present in test data point [True]
231 Text feature [set] present in test data point [True]
232 Text feature [another] present in test data point [True]
235 Text feature [next] present in test data point [True]
237 Text feature [sequences] present in test data point [True]
240 Text feature [least] present in test data point [True]
243 Text feature [localization] present in test data point [True]
244 Text feature [residues] present in test data point [True]
251 Text feature [performed] present in test data point [True]
252 Text feature [possibility] present in test data point [True]
254 Text feature [upon] present in test data point [True]
257 Text feature [induce] present in test data point [True]
258 Text feature [supplemental] present in test data point [True]
259 Text feature [signals] present in test data point [True]
260 Text feature [values] present in test data point [True]
262 Text feature [tested] present in test data point [True]
263 Text feature [conserved] present in test data point [True]
264 Text feature [primers] present in test data point [True]
266 Text feature [obtained] present in test data point [True]
268 Text feature [genome] present in test data point [True]
269 Text feature [strand] present in test data point [True]
270 Text feature [via] present in test data point [True]
271 Text feature [length] present in test data point [True]
274 Text feature [treated] present in test data point [True]
276 Text feature [signal] present in test data point [True]
277 Text feature [specific] present in test data point [True]
281 Text feature [possible] present in test data point [True]
282 Text feature [sample] present in test data point [True]
283 Text feature [defects] present in test data point [True]
285 Text feature [insertion] present in test data point [True]
286 Text feature [16] present in test data point [True]
287 Text feature [de] present in test data point [True]
289 Text feature [carrying] present in test data point [True]
290 Text feature [development] present in test data point [True]
291 Text feature [sequencing] present in test data point [True]
294 Text feature [medium] present in test data point [True]
295 Text feature [coding] present in test data point [True]
299 Text feature [size] present in test data point [True]
300 Text feature [remaining] present in test data point [True]
301 Text feature [taken] present in test data point [True]
303 Text feature [large] present in test data point [True]
304 Text feature [screening] present in test data point [True]
305 Text feature [chain] present in test data point [True]
306 Text feature [exon] present in test data point [True]
309 Text feature [wild] present in test data point [True]
310 Text feature [within] present in test data point [True]
311 Text feature [vitro] present in test data point [True]
317 Text feature [following] present in test data point [True]
321 Text feature [type] present in test data point [True]
322 Text feature [rt] present in test data point [True]
323 Text feature [frame] present in test data point [True]
327 Text feature [mean] present in test data point [True]
328 Text feature [rna] present in test data point [True]
329 Text feature [incubated] present in test data point [True]
332 Text feature [nucleotide] present in test data point [True]
```

```
333 Text feature [del] present in test data point [True]
334 Text feature [dependent] present in test data point [True]
336 Text feature [table] present in test data point [True]
339 Text feature [impact] present in test data point [True]
341 Text feature [selection] present in test data point [True]
342 Text feature [one] present in test data point [True]
347 Text feature [somatic] present in test data point [True]
348 Text feature [fold] present in test data point [True]
349 Text feature [heterozygous] present in test data point
350 Text feature [cell] present in test data point [True]
351 Text feature [manufacturer] present in test data point [True]
352 Text feature [additional] present in test data point [True]
353 Text feature [33] present in test data point [True]
357 Text feature [49] present in test data point [True]
360 Text feature [among] present in test data point [True]
361 Text feature [control] present in test data point [True]
362 Text feature [none] present in test data point [True]
363 Text feature [context] present in test data point [True]
371 Text feature [24] present in test data point [True]
373 Text feature [absence] present in test data point [True]
374 Text feature [exhibited] present in test data point [True]
375 Text feature [complex] present in test data point [True]
376 Text feature [established] present in test data point [True]
377 Text feature [deletions] present in test data point [True]
379 Text feature [stable] present in test data point [True]
380 Text feature [2001] present in test data point [True]
381 Text feature [sporadic] present in test data point [True]
385 Text feature [pcr] present in test data point [True]
387 Text feature [reverse] present in test data point [True]
388 Text feature [individual] present in test data point [True]
389 Text feature [domains] present in test data point [True]
396 Text feature [derived] present in test data point [True]
398 Text feature [reduced] present in test data point [True]
400 Text feature [analyzed] present in test data point [True]
401 Text feature [05] present in test data point [True]
402 Text feature [role] present in test data point [True]
403 Text feature [54] present in test data point [True]
404 Text feature [right] present in test data point [True]
405 Text feature [non] present in test data point [True]
406 Text feature [example] present in test data point [True]
407 Text feature [32] present in test data point [True]
409 Text feature [cause] present in test data point [True]
412 Text feature [allele] present in test data point [True]
413 Text feature [early] present in test data point [True]
414 Text feature [recently] present in test data point [True]
416 Text feature [evidence] present in test data point [True]
417 Text feature [complete] present in test data point [True]
419 Text feature [22] present in test data point [True]
420 Text feature [55] present in test data point [True]
421 Text feature [http] present in test data point [True]
422 Text feature [indeed] present in test data point [True]
423 Text feature [population] present in test data point [True]
425 Text feature [1997] present in test data point [True]
430 Text feature [assay] present in test data point [True]
432 Text feature [distinct] present in test data point [True]
435 Text feature [might] present in test data point [True]
436 Text feature [key] present in test data point [True]
438 Text feature [could] present in test data point [True]
439 Text feature [protein] present in test data point [True]
440 Text feature [results] present in test data point [True]
443 Text feature [revealed] present in test data point [True]
445 Text feature [residue] present in test data point [True]
446 Text feature [1b] present in test data point [True]
450 Text feature [25] present in test data point [True]
451 Text feature [lines] present in test data point [True]
452 Text feature [exons] present in test data point [True]
453 Text feature [generation] present in test data point [True]
454 Text feature [19] present in test data point [True]
455 Text feature [transcription] present in test data point [True]
458 Text feature [presence] present in test data point [True]
459 Text feature [furthermore] present in test data point [True]
466 Text feature [larger] present in test data point [True]
467 Text feature [finally] present in test data point [True]
468 Text feature [effects] present in test data point [True]
470 Text feature [limited] present in test data point [True]
471 Text feature [predicted] present in test data point [True]
473 Text feature [reported] present in test data point [True]
```

```
475 Text feature [nm] present in test data point [True]
476 Text feature [human] present in test data point [True]
478 Text feature [proteins] present in test data point [True]
480 Text feature [respectively] present in test data point [True]
481 Text feature [criteria] present in test data point [True]
484 Text feature [using] present in test data point [True]
485 Text feature [based] present in test data point [True]
480 Text feature [43] present in test data point [True]
490 Text feature [30] present in test data point [True]
491 Text feature [51] present in test data point [True]
492 Text feature [sensitivity] present in test data point [True]
496 Text feature [cohort] present in test data point [True]
497 Text feature [42] present in test data point [True]
498 Text feature [common] present in test data point [True]
499 Text feature [common] present in test data point [True]
Out of the top 500 features 188 are present in query point
```

4.3.2. Without Class balancing

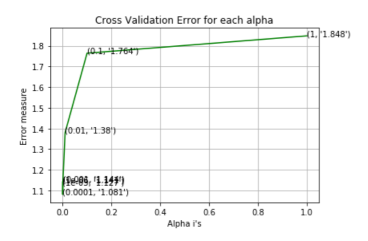
4.3.2.1. Hyper paramter tuning

In [82]:

```
# read more about SGDClassifier() at http://scikit-
learn.org/stable/modules/generated/sklearn.linear\ model.SGDClassifier.html
# default parameters
# SGDClassifier(loss='hinge', penalty='12', alpha=0.0001, 11 ratio=0.15, fit intercept=True, max i
# 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-in
tuition-1/
# find more about CalibratedClassifierCV here at http://scikit-
learn.org/stable/modules/generated/sklearn.calibration.CalibratedClassifier CV.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 arrav[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_lo
ss(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, p
redict y, labels=clf.classes , eps=1e-15))
```

```
for alpha = 1e-06
Log Loss : 1.1412991489533721
for alpha = 1e-05
Log Loss : 1.127222351318073
for alpha = 0.0001
Log Loss : 1.0809042565760936
for alpha = 0.001
Log Loss : 1.1442140220862396
for alpha = 0.01
Log Loss : 1.3801612857584817
for alpha = 0.1
Log Loss : 1.7642984247364517
for alpha = 1
Log Loss : 1.8480675582772192
```



```
For values of best alpha = 0.0001 The train log loss is: 0.5784459460818665
For values of best alpha = 0.0001 The cross validation log loss is: 1.0809042565760936
For values of best alpha = 0.0001 The test log loss is: 1.088411515127868
```

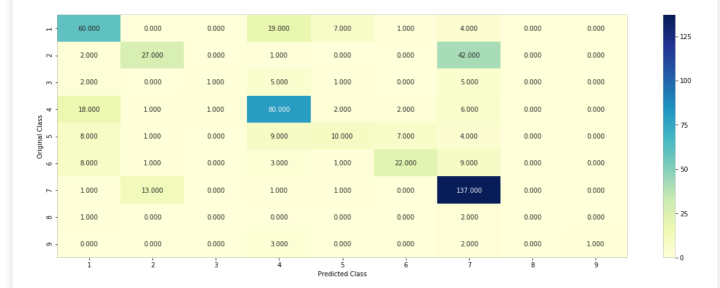
4.3.2.2. Testing model with best hyper parameters

In [83]:

```
# 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:
#-------

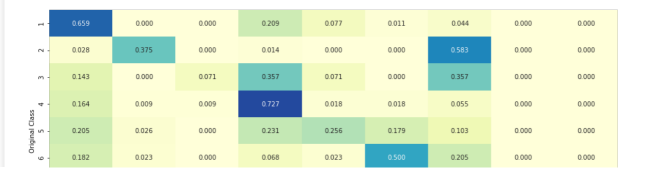
clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='log', random_state=42)
predict_and_plot_confusion_matrix(train_x_onehotCoding, train_y, cv_x_onehotCoding, cv_y, clf)
```



----- Precision matrix (Columm Sum=1) -----



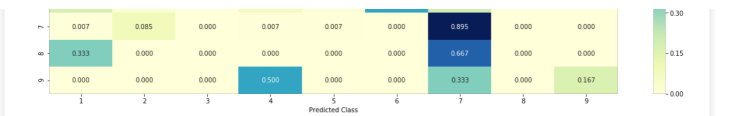
----- Recall matrix (Row sum=1) -----



0.75

- 0.60

0.45



4.3.2.3. Feature Importance, Correctly Classified point

```
In [84]:
clf = SGDClassifier(alpha=alpha[best alpha], penalty='12', loss='log', random state=42)
clf.fit(train_x_onehotCoding,train_y)
test point index = 1
no feature = 500
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],
test_df['TEXT'].iloc[test_point_index],test_df['Gene'].iloc[test_point_index],test_df['Variation']
.iloc[test point index], no feature)
Predicted Class: 4
Predicted Class Probabilities: [[0.032 0.0241 0.0053 0.8883 0.0182 0.0124 0.017 0.0012 0.0016]]
Actual Class : 4
______
O Text feature [suppressor] present in test data point [True]
27 Text feature [missense] present in test data point [True]
62 Text feature [families] present in test data point [True]
83 Text feature [protein] present in test data point [True]
89 Text feature [mammalian] present in test data point [True]
93 Text feature [inactivation] present in test data point [True]
99 Text feature [yeast] present in test data point [True]
100 Text feature [proportion] present in test data point [True]
104 Text feature [see] present in test data point [True]
108 Text feature [mm] present in test data point [True]
127 Text feature [bind] present in test data point [True]
128 Text feature [null] present in test data point [True]
150 Text feature [homozygous] present in test data point [True]
155 Text feature [reduced] present in test data point [True]
156 Text feature [family] present in test data point [True]
157 Text feature [specifically] present in test data point [True]
162 Text feature [dna] present in test data point [True]
163 Text feature [consequences] present in test data point [True]
164 Text feature [functional] present in test data point [True]
169 Text feature [require] present in test data point [True]
182 Text feature [represent] present in test data point [True]
186 Text feature [show] present in test data point [True]
191 Text feature [2010] present in test data point [True]
192 Text feature [germline] present in test data point [True]
194 Text feature [changes] present in test data point [True]
195 Text feature [determine] present in test data point [True]
196 Text feature [loss] present in test data point [True]
200 Text feature [plates] present in test data point [True]
201 Text feature [kinases] present in test data point [True]
210 Text feature [38] present in test data point [True]
221 Text feature [cannot] present in test data point [True]
231 Text feature [localization] present in test data point [True]
233 Text feature [defective] present in test data point [True]
235 Text feature [high] present in test data point [True]
238 Text feature [cycle] present in test data point [True]
243 Text feature [appears] present in test data point [True]
246 Text feature [contribute] present in test data point [True]
247 Text feature [plasmid] present in test data point [True]
253 Text feature [bound] present in test data point [True]
256 Text feature [1998] present in test data point [True]
```

262 Text feature [stained] present in test data point [True] 271 Text feature [displayed] present in test data point [True]

```
273 Text feature [several] present in test data point [True]
274 Text feature [risk] present in test data point [True]
275 Text feature [stability] present in test data point [True]
277 Text feature [function] present in test data point [True]
287 Text feature [26] present in test data point [True]
291 Text feature [cellular] present in test data point [True]
292 Text feature [investigated] present in test data point [True]
298 Text feature [western] present in test data point [True]
299 Text feature [early] present in test data point [True]
303 Text feature [site] present in test data point [True]
306 Text feature [cases] present in test data point [True]
310 Text feature [37] present in test data point [True]
312 Text feature [splice] present in test data point [True]
313 Text feature [observed] present in test data point [True]
316 Text feature [described] present in test data point [True]
317 Text feature [particular] present in test data point [True]
318 Text feature [affected] present in test data point [True]
320 Text feature [mouse] present in test data point [True]
321 Text feature [45] present in test data point [True]
325 Text feature [amino] present in test data point [True]
326 Text feature [view] present in test data point [True]
330 Text feature [relative] present in test data point [True]
332 Text feature [red] present in test data point [True]
339 Text feature [associated] present in test data point [True]
341 Text feature [transfected] present in test data point [True]
343 Text feature [inhibitory] present in test data point [True]
347 Text feature [lower] present in test data point [True]
350 Text feature [terminal] present in test data point [True]
351 Text feature [figure] present in test data point [True]
361 Text feature [3b] present in test data point [True]
368 Text feature [antibodies] present in test data point [True]
369 Text feature [partial] present in test data point [True]
370 Text feature [2a] present in test data point [True]
371 Text feature [40] present in test data point [True]
372 Text feature [mutants] present in test data point [True]
373 Text feature [induced] present in test data point [True]
380 Text feature [low] present in test data point [True]
384 Text feature [average] present in test data point [True]
392 Text feature [transfection] present in test data point [True]
393 Text feature [similarly] present in test data point [True]
395 Text feature [phosphorylation] present in test data point [True]
397 Text feature [medium] present in test data point [True]
399 Text feature [genomic] present in test data point [True]
402 Text feature [comparison] present in test data point [True]
403 Text feature [groups] present in test data point [True]
405 Text feature [times] present in test data point [True]
406 Text feature [key] present in test data point [True]
407 Text feature [larger] present in test data point [True]
410 Text feature [pathogenic] present in test data point [True]
418 Text feature [indeed] present in test data point [True]
422 Text feature [sds] present in test data point [True]
423 Text feature [29] present in test data point [True]
424 Text feature [assay] present in test data point [True]
425 Text feature [containing] present in test data point [True]
426 Text feature [variant] present in test data point [True]
427 Text feature [first] present in test data point [True]
439 Text feature [involved] present in test data point [True]
441 Text feature [breast] present in test data point [True]
445 Text feature [whether] present in test data point [True]
446 Text feature [correlated] present in test data point [True]
447 Text feature [isolated] present in test data point [True]
448 Text feature [2000] present in test data point [True]
449 Text feature [genetic] present in test data point [True]
451 Text feature [combined] present in test data point [True]
452 Text feature [via] present in test data point [True]
456 Text feature [predicted] present in test data point [True]
459 Text feature [assessment] present in test data point [True]
461 Text feature [decrease] present in test data point [True]
464 Text feature [cdk4] present in test data point [True]
466 Text feature [1999] present in test data point [True]
471 Text feature [ca] present in test data point [True]
472 Text feature [altered] present in test data point [True]
473 Text feature [nuclear] present in test data point [True]
477 Text feature [previously] present in test data point [True]
480 Text feature [assays] present in test data point [True]
481 Text feature [locus] present in test data point [True]
482 Text feature [proteins] present in test data point [True]
```

```
487 Text feature [considered] present in test data point [True]
489 Text feature [70] present in test data point [True]
494 Text feature [linked] present in test data point [True]
496 Text feature [bp] present in test data point [True]
497 Text feature [various] present in test data point [True]
498 Text feature [critical] present in test data point [True]
499 Text feature [eight] present in test data point [True]
Out of the top 500 features 126 are present in query point
```

4.3.2.4. Feature Importance, Inorrectly Classified point

```
In [85]:
test point index = 100
no feature = 500
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],
test_df['TEXT'].iloc[test_point_index],test_df['Gene'].iloc[test_point_index],test_df['Variation']
.iloc[test point index], no feature)
Predicted Class: 1
Predicted Class Probabilities: [[9.60e-01 2.38e-02 1.00e-04 1.35e-02 8.00e-04 2.00e-04 1.60e-03 0.
00e+00
 0.00e+0011
Actual Class : 1
12 Text feature [panel] present in test data point [True]
53 Text feature [nuclear] present in test data point [True]
60 Text feature [rather] present in test data point [True]
70 Text feature [repeats] present in test data point [True]
72 Text feature [ability] present in test data point [True]
73 Text feature [deletion] present in test data point [True]
88 Text feature [therefore] present in test data point [True]
97 Text feature [box] present in test data point [True]
112 Text feature [21] present in test data point [True]
121 Text feature [defined] present in test data point [True]
123 Text feature [effect] present in test data point [True]
130 Text feature [analyses] present in test data point [True]
138 Text feature [strong] present in test data point [True]
139 Text feature [essential] present in test data point [True]
141 Text feature [functions] present in test data point [True]
143 Text feature [corresponding] present in test data point [True]
153 Text feature [deficient] present in test data point [True]
155 Text feature [17] present in test data point [True]
158 Text feature [mutational] present in test data point [True]
162 Text feature [region] present in test data point [True]
163 Text feature [page] present in test data point [True]
173 Text feature [indicated] present in test data point [True]
176 Text feature [notably] present in test data point [True]
181 Text feature [calculated] present in test data point [True]
182 Text feature [position] present in test data point [True]
184 Text feature [whole] present in test data point [True]
189 Text feature [change] present in test data point [True]
193 Text feature [previous] present in test data point [True]
196 Text feature [another] present in test data point [True]
197 Text feature [function] present in test data point [True]
199 Text feature [gel] present in test data point [True]
200 Text feature [identify] present in test data point [True]
201 Text feature [impaired] present in test data point [True]
206 Text feature [encoding] present in test data point [True]
208 Text feature [value] present in test data point [True]
211 Text feature [subjected] present in test data point [True]
212 Text feature [blood] present in test data point [True]
213 Text feature [sequenced] present in test data point [True]
219 Text feature [supplemental] present in test data point [True]
220 Text feature [showing] present in test data point [True]
223 Text feature [induce] present in test data point [True]
229 Text feature [located] present in test data point [True]
220 Most footure [nost] procent in test data point [Mayor
```

```
ZOU TEXT TEATURE [MEXT] PRESENT IN LEST DATA POINT [TRUE]
232 Text feature [residues] present in test data point [True]
233 Text feature [least] present in test data point [True]
234 Text feature [set] present in test data point [True]
235 Text feature [signals] present in test data point [True]
236 Text feature [splicing] present in test data point [True]
238 Text feature [localization] present in test data point [True]
239 Text feature [sequences] present in test data point [True]
240 Text feature [possibility] present in test data point [True]
242 Text feature [values] present in test data point [True]
245 Text feature [splice] present in test data point [True]
246 Text feature [defects] present in test data point [True]
247 Text feature [interact] present in test data point [True]
250 Text feature [tested] present in test data point [True]
252 Text feature [via] present in test data point [True]
256 Text feature [selection] present in test data point [True]
257 Text feature [strand] present in test data point [True]
262 Text feature [screening] present in test data point [True]
264 Text feature [performed] present in test data point [True]
265 Text feature [insertion] present in test data point [True]
267 Text feature [percentage] present in test data point [True]
269 Text feature [obtained] present in test data point [True]
270 Text feature [upon] present in test data point [True]
272 Text feature [conserved] present in test data point [True]
273 Text feature [specific] present in test data point [True]
274 Text feature [medium] present in test data point [True]
278 Text feature [genome] present in test data point [True]
279 Text feature [signal] present in test data point [True]
281 Text feature [chain] present in test data point [True]
284 Text feature [sequencing] present in test data point [True]
286 Text feature [sample] present in test data point [True]
287 Text feature [length] present in test data point [True]
288 Text feature [16] present in test data point [True]
291 Text feature [within] present in test data point [True]
292 Text feature [wild] present in test data point [True]
295 Text feature [development] present in test data point [True]
296 Text feature [coding] present in test data point [True]
298 Text feature [treated] present in test data point [True]
300 Text feature [vitro] present in test data point [True]
301 Text feature [size] present in test data point [True]
304 Text feature [carrying] present in test data point [True]
305 Text feature [impact] present in test data point [True]
307 Text feature [taken] present in test data point [True]
309 Text feature [incubated] present in test data point [True]
311 Text feature [following] present in test data point [True]
313 Text feature [primers] present in test data point [True]
316 Text feature [del] present in test data point [True]
317 Text feature [possible] present in test data point [True]
318 Text feature [large] present in test data point [True]
319 Text feature [type] present in test data point [True]
322 Text feature [absence] present in test data point [True]
327 Text feature [fold] present in test data point [True]
328 Text feature [de] present in test data point [True]
329 Text feature [nucleotide] present in test data point [True]
331 Text feature [somatic] present in test data point [True]
336 Text feature [exon] present in test data point [True]
338 Text feature [dependent] present in test data point [True]
339 Text feature [rt] present in test data point [True]
340 Text feature [one] present in test data point [True]
343 Text feature [cell] present in test data point [True]
344 Text feature [remaining] present in test data point [True]
345 Text feature [none] present in test data point [True]
346 Text feature [table] present in test data point [True]
348 Text feature [mean] present in test data point [True]
350 Text feature [context] present in test data point [True]
353 Text feature [exhibited] present in test data point [True]
354 Text feature [additional] present in test data point [True]
357 Text feature [frame] present in test data point [True]
358 Text feature [among] present in test data point [True]
359 Text feature [33] present in test data point [True]
363 Text feature [rna] present in test data point [True]
365 Text feature [heterozygous] present in test data point [True]
369 Text feature [manufacturer] present in test data point [True]
372 Text feature [reduced] present in test data point [True]
374 Text feature [indeed] present in test data point [True]
375 Text feature [control] present in test data point [True]
377 Text feature [sporadic] present in test data point [True]
```

```
3/8 Text reature [evidence] present in test data point [True]
380 Text feature [analyzed] present in test data point [True]
381 Text feature [individual] present in test data point [True]
383 Text feature [05] present in test data point [True]
384 Text feature [reverse] present in test data point [True]
387 Text feature [cause] present in test data point [True]
389 Text feature [22] present in test data point [True]
390 Text feature [recently] present in test data point [True]
391 Text feature [established] present in test data point [True]
392 Text feature [stable] present in test data point [True]
394 Text feature [49] present in test data point [True]
396 Text feature [24] present in test data point [True]
401 Text feature [1b] present in test data point [True]
402 Text feature [allele] present in test data point [True]
403 Text feature [2001] present in test data point [True]
405 Text feature [example] present in test data point [True]
406 Text feature [pcr] present in test data point [True]
408 Text feature [could] present in test data point [True]
409 Text feature [lines] present in test data point [True]
410 Text feature [role] present in test data point [True]
412 Text feature [assay] present in test data point [True]
413 Text feature [non] present in test data point [True]
414 Text feature [1997] present in test data point [True]
415 Text feature [distinct] present in test data point [True]
417 Text feature [might] present in test data point [True]
418 Text feature [54] present in test data point [True]
420 Text feature [presence] present in test data point [True]
422 Text feature [32] present in test data point [True]
427 Text feature [deletions] present in test data point [True]
428 Text feature [http] present in test data point [True]
430 Text feature [sensitivity] present in test data point [True]
431 Text feature [population] present in test data point [True]
432 Text feature [complex] present in test data point [True]
434 Text feature [right] present in test data point [True]
435 Text feature [domains] present in test data point [True]
437 Text feature [43] present in test data point [True]
442 Text feature [key] present in test data point [True]
444 Text feature [derived] present in test data point [True]
445 Text feature [55] present in test data point [True]
446 Text feature [19] present in test data point [True]
448 Text feature [early] present in test data point [True]
449 Text feature [effects] present in test data point [True]
450 Text feature [results] present in test data point [True]
451 Text feature [limited] present in test data point [True]
452 Text feature [reported] present in test data point [True]
454 Text feature [larger] present in test data point [True]
455 Text feature [residue] present in test data point [True]
457 Text feature [predicted] present in test data point [True]
459 Text feature [protein] present in test data point [True]
463 Text feature [complete] present in test data point [True]
464 Text feature [generation] present in test data point [True]
467 Text feature [cohort] present in test data point [True]
470 Text feature [42] present in test data point [True]
473 Text feature [revealed] present in test data point [True]
474 Text feature [nm] present in test data point [True]
476 Text feature [furthermore] present in test data point [True]
479 Text feature [transcription] present in test data point [True]
480 Text feature [proteins] present in test data point [True]
484 Text feature [different] present in test data point [True]
486 Text feature [51] present in test data point [True]
487 Text feature [common] present in test data point [True]
488 Text feature [finally] present in test data point [True]
489 Text feature [based] present in test data point [True]
490 Text feature [exons] present in test data point [True]
492 Text feature [using] present in test data point [True]
493 Text feature [respectively] present in test data point [True]
494 Text feature [25] present in test data point [True]
Out of the top 500 features 186 are present in query point
```

4.4. Linear Support Vector Machines

4.4.1. Hyper paramter tuning

```
In [86]:
```

```
# read more about support vector machines with linear kernals here http://scikit-
learn.org/stable/modules/generated/sklearn.svm.SVC.html
# default parameters
# SVC(C=1.0, kernel='rbf', degree=3, gamma='auto', coef0=0.0, shrinking=True, probability=False, t
01=0.001.
# cache size=200, class weight=None, verbose=False, max iter=-1, decision function shape='ovr', ra
ndom 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', r
andom 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 lo
ss(y cv, predict y, labels=clf.classes_, eps=1e-15))
predict_y = sig_clf.predict_proba(test_x_onehotCoding)
                                                        Hmb- F-F 1-- 1-- 2-- 1-- 1-- 1-- 1-- 1-- 1--
```

```
print('For values of best alpha = ', alpha[best_alpha], "The test log loss is:",log_loss(y_test, p
redict_y, labels=clf.classes_, eps=1e-15))
for C = 1e-05
Log Loss: 1.1239671615818072
for C = 0.0001
Log Loss: 1.0760324189972004
```

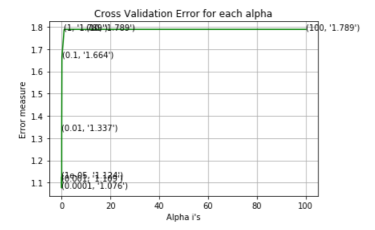
for C = 0.001Log Loss: 1.108906137026918 for C = 0.01Log Loss: 1.337427198724904 for C = 0.1Log Loss: 1.6642874744580503 Log Loss: 1.7887823762757156

for C = 10

Log Loss: 1.7887823316450397

for C = 100

Log Loss: 1.788782267348181



```
For values of best alpha = 0.0001 The train log loss is: 0.6797907549629029 For values of best alpha = 0.0001 The cross validation log loss is: 1.0760324189972004
For values of best alpha = 0.0001 The test log loss is: 1.110967409077816
```

4.4.2. Testing model with best hyper parameters

In [87]:

```
# read more about support vector machines with linear kernals here http://scikit-
learn.org/stable/modules/generated/sklearn.svm.SVC.html
# default parameters
# SVC(C=1.0, kernel='rbf', degree=3, gamma='auto', coef0=0.0, shrinking=True, probability=False, t
01=0.001.
# cache size=200, class weight=None, verbose=False, max iter=-1, decision function shape='ovr', ra
ndom 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/
# clf = SVC(C=alpha[best_alpha],kernel='linear',probability=True, class_weight='balanced')
clf = SGDClassifier(alpha=alpha[best_alpha], penalty='12', loss='hinge',
random state=42,class weight='balanced')
predict_and_plot_confusion_matrix(train_x_onehotCoding, train_y,cv_x_onehotCoding,cv_y, clf)
```

```
Log loss: 1.0760324189972004
Number of mis-classified points : 0.34962406015037595
----- Confusion matrix -----
```



- 125

- 25

- 0.8

-0.6

- 0.4

- 0.2

- 0.75

- 0.60

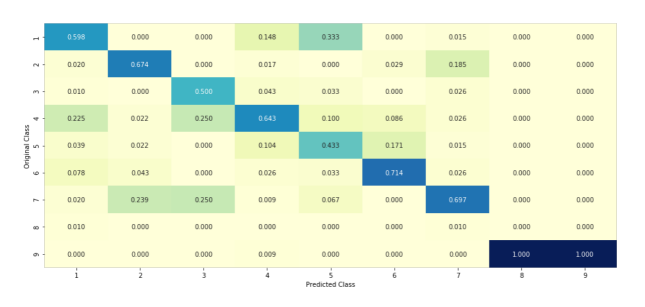
- 0.45

- 0.30

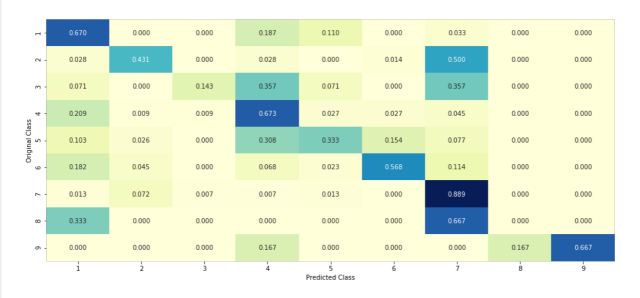
-0.15

- 0.00

----- Precision matrix (Columm Sum=1) -----



----- Recall matrix (Row sum=1) ------



4.3.3. Feature Importance

4.3.3.1. For Correctly classified point

```
In [88]:
clf = SGDClassifier(alpha=alpha[best alpha], penalty='12', loss='hinge', random state=42)
clf.fit(train x onehotCoding,train y)
test point index = 1
# test point index = 100
no feature = 500
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],
test df['TEXT'].iloc[test point index],test_df['Gene'].iloc[test_point_index],test_df['Variation']
.iloc[test point index], no feature)
Predicted Class: 4
Actual Class: 4
O Text feature [suppressor] present in test data point [True]
59 Text feature [missense] present in test data point [True]
61 Text feature [families] present in test data point [True]
64 Text feature [proportion] present in test data point [True]
142 Text feature [mammalian] present in test data point [True]
143 Text feature [bind] present in test data point [True]
145 Text feature [mm] present in test data point [True]
147 Text feature [yeast] present in test data point [True]
150 Text feature [family] present in test data point [True]
151 Text feature [defective] present in test data point [True]
160 Text feature [specifically] present in test data point [True]
162 Text feature [1998] present in test data point [True]
163 Text feature [plates] present in test data point [True]
164 Text feature [see] present in test data point [True]
167 Text feature [require] present in test data point [True]
168 Text feature [inactivation] present in test data point [True]
169 Text feature [protein] present in test data point [True]
170 Text feature [bound] present in test data point [True]
171 Text feature [homozygous] present in test data point [True]
173 Text feature [dna] present in test data point [True]
174 Text feature [determine] present in test data point [True]
175 Text feature [38] present in test data point [True]
179 Text feature [high] present in test data point [True]
182 Text feature [consequences] present in test data point [True]
185 Text feature [changes] present in test data point [True]
187 Text feature [functional] present in test data point [True]
188 Text feature [show] present in test data point [True]
191 Text feature [site] present in test data point [True]
197 Text feature [2010] present in test data point [True]
198 Text feature [induced] present in test data point [True]
199 Text feature [medium] present in test data point [True]
200 Text feature [average] present in test data point [True]
201 Text feature [reduced] present in test data point [True]
202 Text feature [several] present in test data point [True]
204 Text feature [represent] present in test data point [True]
205 Text feature [plasmid] present in test data point [True]
208 Text feature [cannot] present in test data point [True]
209 Text feature [partial] present in test data point [True]
210 Text feature [loss] present in test data point [True]
211 Text feature [37] present in test data point [True]
214 Text feature [figure] present in test data point [True]
215 Text feature [null] present in test data point [True]
216 Text feature [mutants] present in test data point [True]
217 Text feature [phosphorylation] present in test data point [True]
219 Text feature [view] present in test data point [True]
221 Text feature [inhibitory] present in test data point [True]
222 Text feature [correlated] present in test data point [True]
224 Text feature [function] present in test data point [True]
225 Text feature [cases] present in test data point [True]
227 Text feature [29] present in test data point [True]
229 Text feature [described] present in test data point [True]
230 Text feature [similarly] present in test data point [True]
231 Text feature [observed] present in test data point [True]
```

```
235 Text reature [kinases] present in test data point [True]
236 Text feature [line] present in test data point [True]
237 Text feature [localization] present in test data point [True]
358 Text feature [critical] present in test data point [True]
360 Text feature [sds] present in test data point [True]
361 Text feature [contribute] present in test data point [True]
362 Text feature [2000] present in test data point [True]
365 Text feature [germline] present in test data point [True]
367 Text feature [cellular] present in test data point [True]
368 Text feature [early] present in test data point [True]
369 Text feature [affected] present in test data point [True]
371 Text feature [comparison] present in test data point [True]
373 Text feature [displayed] present in test data point [True]
375 Text feature [mouse] present in test data point [True]
376 Text feature [amino] present in test data point [True]
378 Text feature [1999] present in test data point [True]
379 Text feature [stained] present in test data point [True]
381 Text feature [growth] present in test data point [True]
382 Text feature [first] present in test data point [True]
383 Text feature [times] present in test data point [True]
384 Text feature [investigated] present in test data point [True]
385 Text feature [groups] present in test data point [True]
386 Text feature [26] present in test data point [True]
387 Text feature [associated] present in test data point [True]
388 Text feature [absence] present in test data point [True]
389 Text feature [particular] present in test data point [True]
390 Text feature [affinity] present in test data point [True]
391 Text feature [isolated] present in test data point [True]
393 Text feature [primary] present in test data point [True]
396 Text feature [western] present in test data point [True]
397 Text feature [suggested] present in test data point [True]
398 Text feature [70] present in test data point [True]
399 Text feature [relative] present in test data point [True]
402 Text feature [splice] present in test data point [True]
403 Text feature [decrease] present in test data point [True]
405 Text feature [containing] present in test data point [True]
406 Text feature [considered] present in test data point [True]
408 Text feature [materials] present in test data point [True]
411 Text feature [low] present in test data point [True]
415 Text feature [45] present in test data point [True]
417 Text feature [transfected] present in test data point [True]
420 Text feature [effective] present in test data point [True]
421 Text feature [impact] present in test data point [True]
422 Text feature [23] present in test data point [True]
423 Text feature [combined] present in test data point [True]
425 Text feature [3b] present in test data point [True]
426 Text feature [following] present in test data point [True]
428 Text feature [reports] present in test data point [True]
431 Text feature [indeed] present in test data point [True]
433 Text feature [40] present in test data point [True]
434 Text feature [locus] present in test data point [True]
435 Text feature [significantly] present in test data point [True]
436 Text feature [various] present in test data point [True]
437 Text feature [cycle] present in test data point [True]
438 Text feature [2a] present in test data point [True]
439 Text feature [appears] present in test data point [True]
440 Text feature [via] present in test data point [True]
442 Text feature [variant] present in test data point [True]
444 Text feature [university] present in test data point [True]
447 Text feature [antibodies] present in test data point [True]
452 Text feature [risk] present in test data point [True]
455 Text feature [breast] present in test data point [True]
456 Text feature [ca] present in test data point [True]
457 Text feature [48] present in test data point [True]
460 Text feature [lines] present in test data point [True]
463 Text feature [http] present in test data point [True]
464 Text feature [larger] present in test data point [True]
466 Text feature [pathogenic] present in test data point [True]
467 Text feature [stability] present in test data point [True]
468 Text feature [likely] present in test data point [True]
469 Text feature [consistent] present in test data point [True]
470 Text feature [kinase] present in test data point [True]
477 Text feature [genomic] present in test data point [True]
478 Text feature [number] present in test data point [True]
480 Text feature [followed] present in test data point [True]
485 Text feature [large] present in test data point [True]
486 Text feature [gst] present in test data point [True]
```

```
48/ Text reature [based] present in test data point [True]
491 Text feature [terminal] present in test data point [True]
494 Text feature [identified] present in test data point [True]
495 Text feature [buffer] present in test data point [True]
496 Text feature [key] present in test data point [True]
498 Text feature [history] present in test data point [True]
499 Text feature [genetic] present in test data point [True]
Out of the top 500 features 137 are present in query point
```

4.3.3.2. For Incorrectly classified point

```
In [89]:
test point index = 100
no feature = 500
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],
test df['TEXT'].iloc[test point index],test df['Gene'].iloc[test point index],test df['Variation']
.iloc[test point index], no feature)
Predicted Class: 1
Predicted Class Probabilities: [[9.003e-01 7.040e-02 3.000e-04 1.790e-02 4.300e-03 4.000e-04 6.000
e - 0.3
  2.000e-04 2.000e-0411
Actual Class : 1
10 Text feature [panel] present in test data point [True]
55 Text feature [21] present in test data point [True]
56 Text feature [ability] present in test data point [True]
57 Text feature [rather] present in test data point [True]
58 Text feature [analyses] present in test data point [True]
59 Text feature [repeats] present in test data point [True]
127 Text feature [box] present in test data point [True]
129 Text feature [deletion] present in test data point [True]
131 Text feature [nuclear] present in test data point [True]
132 Text feature [therefore] present in test data point [True]
134 Text feature [previous] present in test data point [True]
136 Text feature [selection] present in test data point [True]
138 Text feature [identify] present in test data point [True]
139 Text feature [supplemental] present in test data point [True]
140 Text feature [strong] present in test data point [True]
142 Text feature [page] present in test data point [True]
143 Text feature [whole] present in test data point [True]
144 Text feature [another] present in test data point [True]
147 Text feature [mutational] present in test data point [True]
149 Text feature [showing] present in test data point [True]
151 Text feature [defined] present in test data point [True]
153 Text feature [impaired] present in test data point [True]
154 Text feature [values] present in test data point [True]
156 Text feature [effect] present in test data point [True]
158 Text feature [17] present in test data point [True]
159 Text feature [essential] present in test data point [True]
161 Text feature [sequenced] present in test data point [True]
162 Text feature [calculated] present in test data point [True]
164 Text feature [gel] present in test data point [True]
166 Text feature [signals] present in test data point [True]
167 Text feature [corresponding] present in test data point [True]
168 Text feature [taken] present in test data point [True]
170 Text feature [indicated] present in test data point [True]
172 Text feature [position] present in test data point [True]
175 Text feature [screening] present in test data point [True]
176 Text feature [signal] present in test data point [True]
178 Text feature [blood] present in test data point [True]
181 Text feature [wild] present in test data point [True]
182 Text feature [functions] present in test data point [True]
183 Text feature [deficient] present in test data point [True]
184 Text feature [induce] present in test data point [True]
185 Text feature [located] present in test data point [True]
186 Text feature [value] present in test data point [True]
```

```
TOO TEAC TEACUTE [VALUE] PLESENC IN CESC GACA POINC [ITGE]
187 Text feature [subjected] present in test data point [True]
190 Text feature [fold] present in test data point [True]
191 Text feature [least] present in test data point [True]
192 Text feature [notably] present in test data point [True]
193 Text feature [residues] present in test data point [True]
194 Text feature [16] present in test data point [True]
196 Text feature [next] present in test data point [True]
197 Text feature [percentage] present in test data point [True]
199 Text feature [development] present in test data point [True]
201 Text feature [somatic] present in test data point [True]
204 Text feature [encoding] present in test data point [True]
206 Text feature [type] present in test data point [True]
208 Text feature [sequencing] present in test data point [True]
209 Text feature [medium] present in test data point [True]
210 Text feature [none] present in test data point [True]
211 Text feature [obtained] present in test data point [True]
212 Text feature [performed] present in test data point [True]
214 Text feature [localization] present in test data point [True]
215 Text feature [possibility] present in test data point [True]
216 Text feature [defects] present in test data point [True]
217 Text feature [sample] present in test data point [True]
296 Text feature [sporadic] present in test data point [True]
299 Text feature [carrying] present in test data point [True]
300 Text feature [incubated] present in test data point [True]
301 Text feature [sequences] present in test data point [True]
302 Text feature [conditions] present in test data point [True]
303 Text feature [heterozygous] present in test data point [True]
304 Text feature [22] present in test data point [True]
306 Text feature [change] present in test data point [True]
308 Text feature [exon] present in test data point [True]
309 Text feature [specific] present in test data point [True]
312 Text feature [upon] present in test data point [True]
313 Text feature [exhibited] present in test data point [True]
314 Text feature [genome] present in test data point [True]
315 Text feature [del] present in test data point [True]
316 Text feature [evidence] present in test data point [True]
317 Text feature [rt] present in test data point [True]
318 Text feature [mean] present in test data point [True]
319 Text feature [stable] present in test data point [True]
320 Text feature [splice] present in test data point [True]
324 Text feature [indeed] present in test data point [True]
325 Text feature [analyzed] present in test data point [True]
326 Text feature [region] present in test data point [True]
328 Text feature [05] present in test data point [True]
331 Text feature [55] present in test data point [True]
332 Text feature [table] present in test data point [True]
333 Text feature [chain] present in test data point [True]
334 Text feature [coding] present in test data point [True]
337 Text feature [set] present in test data point [True]
339 Text feature [cell] present in test data point [True]
340 Text feature [1997] present in test data point [True]
341 Text feature [tested] present in test data point [True]
342 Text feature [length] present in test data point [True]
345 Text feature [within] present in test data point [True]
346 Text feature [right] present in test data point [True]
347 Text feature [54] present in test data point [True]
348 Text feature [presence] present in test data point [True]
349 Text feature [population] present in test data point [True]
350 Text feature [absence] present in test data point [True]
351 Text feature [signaling] present in test data point [True]
352 Text feature [2001] present in test data point [True]
354 Text feature [reduced] present in test data point [True]
355 Text feature [reported] present in test data point [True]
356 Text feature [cause] present in test data point [True]
357 Text feature [sensitivity] present in test data point [True]
359 Text feature [allele] present in test data point [True]
362 Text feature [limited] present in test data point [True]
367 Text feature [primers] present in test data point [True]
368 Text feature [51] present in test data point [True]
369 Text feature [splicing] present in test data point [True]
370 Text feature [could] present in test data point [True]
371 Text feature [one] present in test data point [True]
373 Text feature [distinct] present in test data point [True]
374 Text feature [impact] present in test data point [True]
375 Text feature [insertion] present in test data point [True]
376 Text feature [treated] present in test data point [True]
377 Tayt fasture [might] present in test data point [True]
```

```
Jii tevr tearnte [mtdur] breseur tu rest nara botur [true]
378 Text feature [via] present in test data point [True]
380 Text feature [recently] present in test data point [True]
381 Text feature [key] present in test data point [True]
382 Text feature [function] present in test data point [True]
384 Text feature [context] present in test data point [True]
385 Text feature [cultured] present in test data point [True]
386 Text feature [insertions] present in test data point [True]
387 Text feature [possible] present in test data point [True]
389 Text feature [non] present in test data point [True]
390 Text feature [lines] present in test data point [True]
391 Text feature [pcr] present in test data point [True]
392 Text feature [interact] present in test data point [True]
394 Text feature [studied] present in test data point [True]
395 Text feature [exons] present in test data point [True]
396 Text feature [among] present in test data point [True]
397 Text feature [predicted] present in test data point [True]
399 Text feature [remaining] present in test data point [True]
401 Text feature [1b] present in test data point [True]
402 Text feature [nucleotide] present in test data point [True]
404 Text feature [conserved] present in test data point [True]
406 Text feature [even] present in test data point [True]
407 Text feature [additional] present in test data point [True]
408 Text feature [particular] present in test data point
409 Text feature [43] present in test data point [True]
412 Text feature [patients] present in test data point [True]
413 Text feature [plasmid] present in test data point [True]
414 Text feature [de] present in test data point [True]
415 Text feature [common] present in test data point [True]
416 Text feature [related] present in test data point [True]
418 Text feature [strand] present in test data point [True]
419 Text feature [30] present in test data point [True]
420 Text feature [provided] present in test data point [True]
421 Text feature [different] present in test data point [True]
423 Text feature [33] present in test data point [True]
424 Text feature [52] present in test data point [True]
425 Text feature [reverse] present in test data point [True]
427 Text feature [selected] present in test data point [True]
429 Text feature [gene] present in test data point [True]
430 Text feature [combination] present in test data point [True]
431 Text feature [following] present in test data point [True]
433 Text feature [cohort] present in test data point [True]
434 Text feature [rna] present in test data point [True]
436 Text feature [min] present in test data point [True]
437 Text feature [finally] present in test data point [True]
438 Text feature [24] present in test data point [True]
439 Text feature [frame] present in test data point [True]
440 Text feature [mutagenesis] present in test data point [True]
441 Text feature [inhibitors] present in test data point [True]
443 Text feature [part] present in test data point [True]
444 Text feature [control] present in test data point [True]
445 Text feature [method] present in test data point [True]
447 Text feature [frequently] present in test data point [True]
448 Text feature [independent] present in test data point [True]
449 Text feature [phenotype] present in test data point [True]
450 Text feature [vitro] present in test data point [True]
451 Text feature [large] present in test data point [True]
454 Text feature [mutant] present in test data point [True]
455 Text feature [100] present in test data point [True]
457 Text feature [effects] present in test data point [True]
458 Text feature [42] present in test data point [True]
459 Text feature [assay] present in test data point [True]
460 Text feature [current] present in test data point [True]
462 Text feature [involving] present in test data point [True]
468 Text feature [diagnosis] present in test data point [True]
471 Text feature [association] present in test data point [True]
472 Text feature [2005] present in test data point [True]
473 Text feature [using] present in test data point [True]
474 Text feature [homozygous] present in test data point [True]
476 Text feature [study] present in test data point [True]
478 Text feature [first] present in test data point [True]
479 Text feature [manufacturer] present in test data point [True]
481 Text feature [established] present in test data point [True]
482 Text feature [ng] present in test data point [True]
483 Text feature [role] present in test data point [True]
484 Text feature [provide] present in test data point [True]
486 Text feature [example] present in test data point [True]
```

```
488 Text feature [majority] present in test data point [True]
489 Text feature [13] present in test data point [True]
491 Text feature [49] present in test data point [True]
492 Text feature [compared] present in test data point [True]
493 Text feature [codon] present in test data point [True]
494 Text feature [blot] present in test data point [True]
495 Text feature [41] present in test data point [True]
497 Text feature [1a] present in test data point [True]
498 Text feature [2004] present in test data point [True]
498 Text feature [2004] present in test data point [True]
Out of the top 500 features 206 are present in query point
```

4.5 Random Forest Classifier

4.5.1. Hyper paramter tuning (With One hot Encoding)

In [90]:

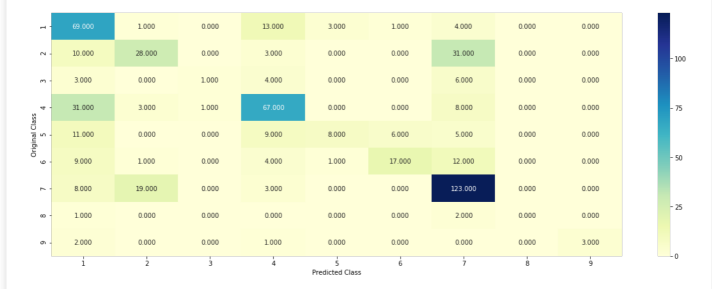
```
# default parameters
# sklearn.ensemble.RandomForestClassifier(n estimators=10, criterion='qini', max depth=None, min s
# min samples leaf=1, min weight fraction leaf=0.0, max features='auto', max leaf nodes=None, min
impurity decrease=0.0,
# min impurity split=None, bootstrap=True, oob score=False, 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-fores
t-and-their-construction-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
       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))
Illfia av = nlt euhnlate()
```

```
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 alpha%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 : 1.2104650476104701
for n estimators = 100 and max depth = 10
Log Loss : 1.229839981617805
for n estimators = 200 and max depth = 5
Log Loss : 1.194056714239944
for n estimators = 200 and max depth = 10
Log Loss : 1.2169872464100477
for n estimators = 500 and max depth = 5
Log Loss: 1.181379091988994
for n estimators = 500 and max depth = 10
Log Loss : 1.2134859279603696
for n estimators = 1000 and max depth = 5
Log Loss: 1.182933407723163
for n estimators = 1000 and max depth = 10
Log Loss: 1.2089507869204237
for n_{estimators} = 2000 and max depth = 5
Log Loss: 1.1830596828378452
for n estimators = 2000 and max depth = 10
Log Loss : 1.2095645950553566
For values of best estimator = 500 The train log loss is: 0.8409143472123352
For values of best estimator = 500 The cross validation log loss is: 1.181379091988994
For values of best estimator = 500 The test log loss is: 1.2216369378538312
```

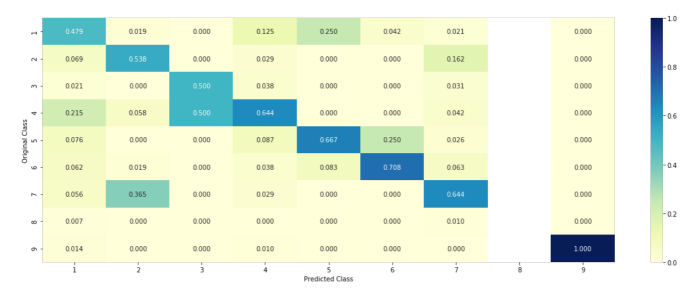
4.5.2. Testing model with best hyper parameters (One Hot Encoding)

```
In [91]:
```

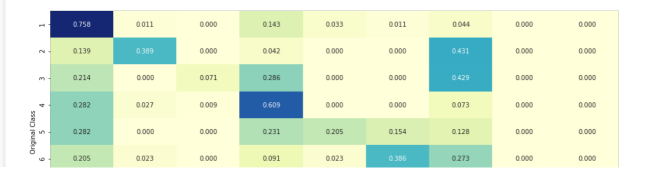
IIY, ax - pic.suppicts()



----- Precision matrix (Columm Sum=1) ------



----- Recall matrix (Row sum=1) ------

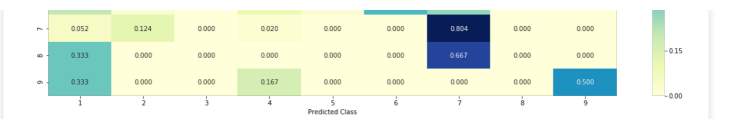


0.75

- 0 60

-045

0.30



4.5.3. Feature Importance

4.5.3.1. Correctly Classified point

```
In [92]:
# test point index = 10
clf = RandomForestClassifier(n estimators=alpha[int(best alpha/2)], criterion='gini', max depth=max
_depth[int(best_alpha%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 = 1
no feature = 100
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], test df['TEXT'].iloc[test point index],test df['Gene'].
iloc[test point index], test df['Variation'].iloc[test point index], no feature)
Predicted Class: 4
Predicted Class Probabilities: [[0.1574 0.0133 0.0113 0.6882 0.0552 0.0456 0.0225 0.0032 0.0033]]
Actual Class : 4
O Text feature [kinase] present in test data point [True]
2 Text feature [suppressor] present in test data point [True]
4 Text feature [inhibitors] present in test data point [True]
5 Text feature [function] present in test data point [True]
6 Text feature [missense] present in test data point [True]
9 Text feature [phosphorylation] present in test data point [True]
10 Text feature [loss] present in test data point [True]
12 Text feature [deleterious] present in test data point [True]
13 Text feature [protein] present in test data point [True]
15 Text feature [stability] present in test data point [True]
19 Text feature [pathogenic] present in test data point [True]
20 Text feature [neutral] present in test data point [True]
25 Text feature [functional] present in test data point [True]
27 Text feature [cell] present in test data point [True]
33 Text feature [classified] present in test data point [True]
34 Text feature [proteins] present in test data point [True]
36 Text feature [kinases] present in test data point [True]
40 Text feature [downstream] present in test data point [True]
41 Text feature [cells] present in test data point [True]
46 Text feature [expected] present in test data point [True]
48 Text feature [growth] present in test data point [True]
52 Text feature [predicted] present in test data point [True]
55 Text feature [functions] present in test data point [True]
56 Text feature [defective] present in test data point [True]
61 Text feature [variants] present in test data point [True]
63 Text feature [splice] present in test data point [True]
65 Text feature [months] present in test data point [True]
66 Text feature [57] present in test data point [True]
69 Text feature [proliferation] present in test data point [True]
70 Text feature [expression] present in test data point [True]
71 Text feature [yeast] present in test data point [True]
74 Text feature [nuclear] present in test data point [True]
75 Text feature [inactivation] present in test data point [True]
76 Text feature [phosphorylated] present in test data point [True]
80 Text feature [inhibition] present in test data point [True]
```

```
81 Text feature [potential] present in test data point [True]
82 Text feature [breast] present in test data point [True]
84 Text feature [use] present in test data point [True]
85 Text feature [expressing] present in test data point [True]
86 Text feature [sensitivity] present in test data point [True]
88 Text feature [variant] present in test data point [True]
89 Text feature [null] present in test data point [True]
92 Text feature [ability] present in test data point [True]
94 Text feature [clinical] present in test data point [True]
98 Text feature [type] present in test data point [True]
99 Text feature [21] present in test data point [True]
Out of the top 100 features 46 are present in query point
```

4.5.3.2. Inorrectly Classified point

```
In [93]:
test point index = 100
no feature = 100
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("Actuall Class :", test y[test point index])
indices = np.argsort(-clf.feature importances )
print("-"*50)
get_impfeature_names(indices[:no_feature], test_df['TEXT'].iloc[test_point_index],test_df['Gene'].
iloc[test_point_index], test_df['Variation'].iloc[test_point_index], no_feature)
Predicted Class: 1
Predicted Class Probabilities: [[0.6281 0.0386 0.0059 0.1477 0.0504 0.0502 0.0324 0.017 0.0298]]
Actuall Class : 1
_____
4 Text feature [inhibitors] present in test data point [True]
5 Text feature [function] present in test data point [True]
6 Text feature [missense] present in test data point [True]
7 Text feature [activation] present in test data point [True]
9 Text feature [phosphorylation] present in test data point [True]
13 Text feature [protein] present in test data point [True]
15 Text feature [stability] present in test data point [True]
17 Text feature [treatment] present in test data point [True]
18 Text feature [brca1] present in test data point [True]
19 Text feature [pathogenic] present in test data point [True]
25 Text feature [functional] present in test data point [True]
26 Text feature [therapeutic] present in test data point [True]
27 Text feature [cell] present in test data point [True]
28 Text feature [activate] present in test data point [True]
30 Text feature [signaling] present in test data point [True]
31 Text feature [therapy] present in test data point [True]
32 Text feature [brca2] present in test data point [True]
33 Text feature [classified] present in test data point [True]
34 Text feature [proteins] present in test data point [True]
35 Text feature [treated] present in test data point [True]
40 Text feature [downstream] present in test data point [True]
41 Text feature [cells] present in test data point [True]
46 Text feature [expected] present in test data point [True]
50 Text feature [survival] present in test data point [True]
52 Text feature [predicted] present in test data point [True]
53 Text feature [patients] present in test data point [True]
55 Text feature [functions] present in test data point [True]
60 Text feature [brca] present in test data point [True]
61 Text feature [variants] present in test data point [True]
63 Text feature [splice] present in test data point [True]
64 Text feature [response] present in test data point [True]
68 Text feature [repair] present in test data point [True]
70 Text feature [expression] present in test data point [True]
74 Text feature [nuclear] present in test data point [True]
81 Text feature [potential] present in test data point [True]
82 Text feature [breast] present in test data point [True]
84 Text feature [use] present in test data point [True]
85 Text feature [expressing] present in test data point [True]
86 Text feature [sensitivity] present in test data point [True]
87 Text feature [pathway] present in test data point [True]
88 Text feature [variant] present in test data point [True]
```

```
89 Text feature [null] present in test data point [True]
92 Text feature [ability] present in test data point [True]
94 Text feature [clinical] present in test data point [True]
98 Text feature [type] present in test data point [True]
99 Text feature [21] present in test data point [True]
Out of the top 100 features 46 are present in query point
```

4.5.3. Hyper paramter tuning (With Response Coding)

In [94]:

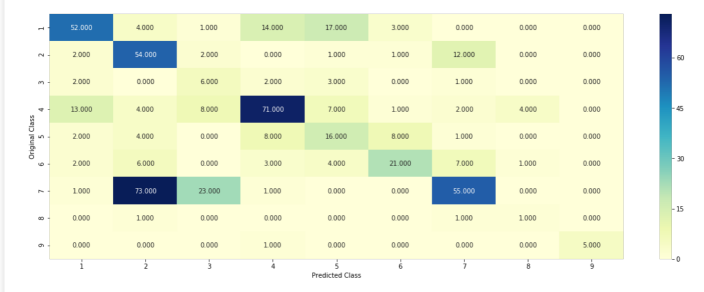
```
# default parameters
# sklearn.ensemble.RandomForestClassifier(n estimators=10, criterion='gini', max depth=None, min s
amples split=2,
# min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_
impurity decrease=0.0,
# min impurity split=None, bootstrap=True, 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-fores
t-and-their-construction-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 alpha%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 : 2.201308963715666
for n estimators = 10 and max depth = 3
Log Loss : 1.715576402352545
for n estimators = 10 and max depth = 5
Log Loss: 1.4611230896289988
for n estimators = 10 and max depth = 10
Log Loss: 1.7468155715605362
for n_{estimators} = 50 and max depth = 2
Log Loss: 1.7190133197710857
for n estimators = 50 and max depth = 3
Log Loss: 1.4837863479799054
for n estimators = 50 and max depth = 5
Log Loss : 1.413886352151792
for n estimators = 50 and max depth = 10
Log Loss: 1.5606902745576023
for n_{estimators} = 100 and max depth = 2
Log Loss: 1.557195317852617
for n estimators = 100 and max depth = 3
Log Loss : 1.4832220897946087
for n estimators = 100 and max depth = 5
Log Loss: 1.2537573027728035
for n estimators = 100 and max depth = 10
Log Loss: 1.5128031426075377
for n estimators = 200 and max depth = 2
Log Loss: 1.624233611986707
for n_{estimators} = 200 and max depth = 3
Log Loss: 1.5255797829657924
for n estimators = 200 and max depth = 5
Log Loss : 1.2654943694462788
for n_{estimators} = 200 and max depth = 10
Log Loss : 1.5317255553062172
for n estimators = 500 and max depth = 2
Log Loss : 1.6791549372006445
for n estimators = 500 and max depth = 3
Log Loss: 1.565164538412316
for n estimators = 500 and max depth = 5
Log Loss: 1.293726277866204
for n estimators = 500 and max depth = 10
Log Loss: 1.5660587192387527
for n estimators = 1000 and max depth = 2
Log Loss : 1.6482182778866807
for n estimators = 1000 and max depth = 3
Log Loss : 1.5569256071386133
for n estimators = 1000 and max depth = 5
Log Loss: 1.2778922308136347
for n estimators = 1000 and max depth = 10
Log Loss: 1.5484922596041149
For values of best alpha = 100 The train log loss is: 0.054526937802532705
For values of best alpha = 100 The cross validation log loss is: 1.2537573027728035
For values of best alpha = 100 The test log loss is: 1.324091335741171
```

In [95]:

```
# default parameters
# sklearn.ensemble.RandomForestClassifier(n estimators=10, criterion='gini', max depth=None, min s
amples split=2,
# min samples leaf=1, min weight fraction leaf=0.0, max features='auto', max leaf nodes=None, min
impurity decrease=0.0,
# 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-fores
t-and-their-construction-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)
```

Log loss: 1.2537573027728035 Number of mis-classified points: 0.4718045112781955 ------ Confusion matrix ------



1.0

- 0.8

- 0.6

- 0.4

-02

----- Precision matrix (Columm Sum=1) -----

r	0.703	0.027	0.025	0.140	0.354	0.088	0.000	0.000	0.000
- 2	0.027	0.370	0.050	0.000	0.021	0.029	0.152	0.000	0.000
m -	0.027	0.000	0.150	0.020	0.062	0.000	0.013	0.000	0.000
Original Class 5 4	0.176	0.027	0.200	0.710	0.146	0.029	0.025	0.667	0.000
	0.027	0.027	0.000	0.080	0.333	0.235	0.013	0.000	0.000
Ori	0.027	0.041	0.000	0.030	0.083	0.618	0.089	0.167	0.000
۲ -	0.014			0.010	0.000	0.000	0.696	0.000	0.000
ω -	0.000	0.007	0.000	0.000	0.000	0.000	0.013	0.167	0.000



4.5.5. Feature Importance

4.5.5.1. Correctly Classified point

```
In [96]:
```

```
clf = RandomForestClassifier(n estimators=alpha[int(best alpha/4)], criterion='gini', max depth=max
depth[int(best alpha%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 = 1
no_feature = 27
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: [[0.1094 0.0174 0.2285 0.5241 0.0206 0.0293 0.0074 0.0259 0.0374]]
Actual Class : 4
Variation is important feature
Variation is important feature
Variation is important feature
Variation is important feature
Gene is important feature
Variation is important feature
Variation is important feature
Text is important feature
Text is important feature
Gene is important feature
```

```
Text is important feature
Text is important feature
Text is important feature
Gene is important feature
Variation is important feature
Gene is important feature
Gene is important feature
Text is important feature
Gene is important feature
Variation is important feature
Variation is important feature
Text is important feature
Text is important feature
Gene is important feature
Text is important feature
Gene is important feature
Gene is important feature
```

4.5.5.2. Incorrectly Classified point

```
In [97]:
```

```
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("Predicted Class Probabilities:",
\verb"np.round(sig_clf.predict_proba(test_x_responseCoding[test_point_index].reshape(1,-1)), 4))" in the property of the probability of the probabil
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")
                 print("Text is important feature")
Predicted Class: 1
Predicted Class Probabilities: [[0.9829 0.0014 0.0018 0.0046 0.001 0.0023 0.0015 0.0018 0.0027]]
Actual Class : 1
 ______
Variation is important feature
Variation is important feature
Variation is important feature
Variation is important feature
Gene is important feature
Variation is important feature
Variation is important feature
Text is important feature
Text is important feature
Gene is important feature
Text is important feature
Text is important feature
Text is important feature
Gene is important feature
Variation is important feature
Gene is important feature
Gene is important feature
Text is important feature
Gene is important feature
Variation is important feature
Variation is important feature
Text is important feature
Text is important feature
Gene is important feature
Text is important feature
Gene is important feature
Gene is important feature
```

4.7 Stack the models

4.7.1 testing with hyper parameter tuning

In [98]:

```
# read more about SGDClassifier() at http://scikit-
learn.org/stable/modules/generated/sklearn.linear model.SGDClassifier.html
# default parameters
# SGDClassifier(loss='hinge', penalty='12', alpha=0.0001, 11 ratio=0.15, fit intercept=True, max i
# 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-in
tuition-1/
# read more about support vector machines with linear kernals here http://scikit-
learn.org/stable/modules/generated/sklearn.svm.SVC.html
# default parameters
# SVC(C=1.0, kernel='rbf', degree=3, gamma='auto', coef0=0.0, shrinking=True, probability=False, t
# cache size=200, class weight=None, verbose=False, max iter=-1, decision function shape='ovr', ra
ndom 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/sklearn.ensemble.RandomForestClassifier.html \\
# default parameters
# sklearn.ensemble.RandomForestClassifier(n estimators=10, criterion='gini', max depth=None, min s
amples split=2,
# min_samples_leaf=1, min_weight_fraction_leaf=0.0, max_features='auto', max_leaf_nodes=None, min_
impurity decrease=0.0,
# 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-fores
t-and-their-construction-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=1, 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=0.001)
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_onehot
Coding())))
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 p
robas=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, sc
lf.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
                                                                                                 I
Logistic Regression: Log Loss: 1.04
Support vector machines : Log Loss: 1.79
Naive Bayes : Log Loss: 1.16
Stacking Classifer: for the value of alpha: 0.000100 Log Loss: 2.178
Stacking Classifer: for the value of alpha: 0.001000 Log Loss: 2.033
Stacking Classifer: for the value of alpha: 0.010000 Log Loss: 1.505
Stacking Classifer: for the value of alpha: 0.100000 Log Loss: 1.124
Stacking Classifer: for the value of alpha: 1.000000 Log Loss: 1.134
Stacking Classifer: for the value of alpha: 10.000000 Log Loss: 1.253
```

4.7.2 testing the model with the best hyper parameters

0.000

0.000

28.000

13.000

5.000

4.000

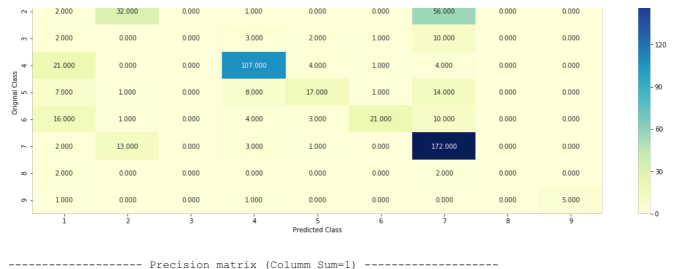
0.000

0.000

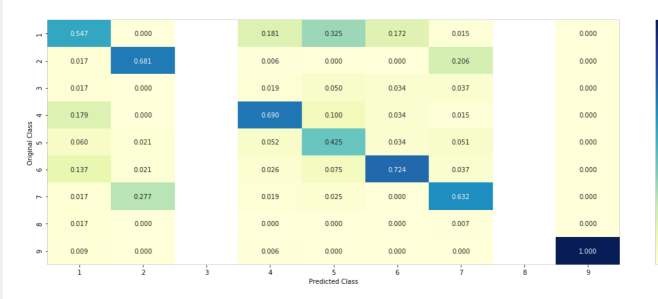
64.000

```
In [99]:
```

```
lr = LogisticRegression(C=0.1)
sclf = StackingClassifier(classifiers=[sig_clf1, sig_clf2, sig_clf3], meta_classifier=lr, use_proba
sclf.fit(train x onehotCoding, train y)
log error = log loss(train y, sclf.predict proba(train x onehotCoding))
print("Log loss (train) on the stacking classifier :",log_error)
log error = log loss(cv y, sclf.predict proba(cv x onehotCoding))
print("Log loss (CV) on the stacking classifier :",log error)
log error = log loss(test y, sclf.predict proba(test x onehotCoding))
print("Log loss (test) on the stacking classifier :",log error)
print("Number of missclassified point :", np.count_nonzero((sclf.predict(test_x_onehotCoding)-
test y))/test y.shape[0])
plot confusion matrix(test y=test y, predict y=sclf.predict(test x onehotCoding))
Log loss (train) on the stacking classifier : 0.7988623633997219
Log loss (CV) on the stacking classifier: 1.1236622548128592
Log loss (test) on the stacking classifier: 1.1440981600121238
Number of missclassified point : 0.37142857142857144
----- Confusion matrix -----
```



----- Precision matrix (Columm Sum=1) ------



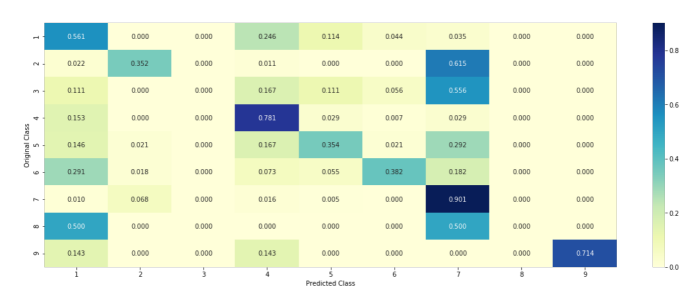
0.6

- 0.4

- 0.2

0.0

----- Recall matrix (Row sum=1) ------



4.7.3 Maximum Voting classifier

In [100]:

#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=

```
vclr.fit(train_x_onenotCoding, 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))
```

- 150

- 120

- 90

60

- 30

- 0.8

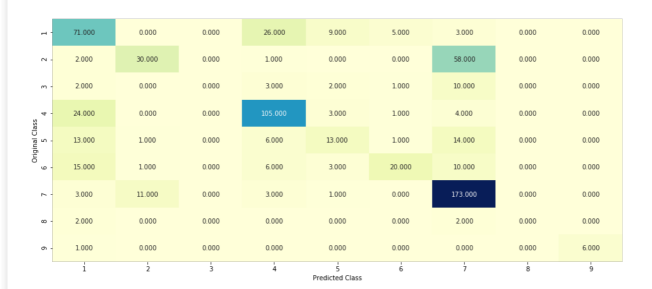
-0.6

- 0.4

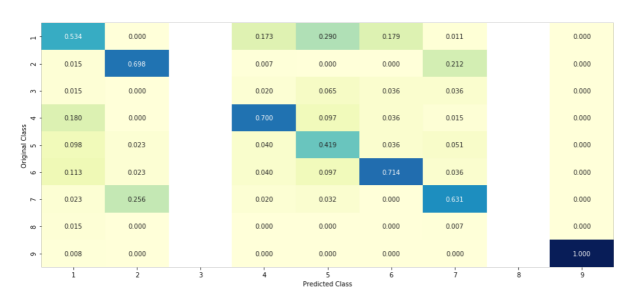
- 0.2

- 0 0

- 0.6

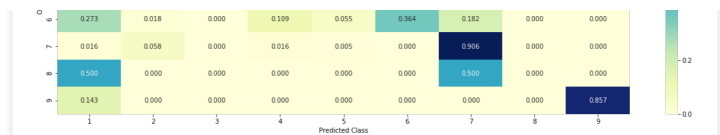






----- Recall matrix (Row sum=1) ------

0.000
0.000
0.000
0.000
0.000



Conclusion

- According to our ploblem statment: We need to predict the probability of each data-point belonging to each of the nine classes. i.e Classify the given genetic variations/mutations based on evidence from text-based clinical literature
- · Lets Start ->
 - 1. As we know we have dataset which contains ID,Gene,Variation,Test,Class and we have given class lables i.e 1-9 and genetic mutation has been classified on this.
 - 2. So lets start with Exploratory Data Analysis but before that we will do some Preprocessing of text so that we will able to rectify that is there any row which has null value and after doing this we will split our whole data into Train,Cv and test and then we will do some EDA on these we will able to visialuze the distribution of data or the class in the train test and cv.
 - 3. And after that lets apply some random models to that we will able to recify the worst performance of the models, and then we will start working on the models
 - 4. Now after getting performance of the random models, lets do some featurization on our data set as we know we have dataset which contains ID,Gene,Variation,Test,Class from which we have Gene, variance and test as a features. So lets start with univariate Analysis on each features one by one and try to get the performance of each. Here we will work with two type of featurization i.e
 - Response coding
 - One hot coding

Note: 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. And at the same time we will try to get how good is this feature in predicting y_i, and Is the feature stable across all the data sets (Test, Train, Cross validation), the distribution if the features and lot more.

- 5. After doing all above now we will Start with our Machine learning models but before that we will combine our all featurized features into one i.e Stacking the three types of features and then will apply different machine learning models on it and try to get the best out of it by apply hyperparameter tuning on it each of the models.
- 6. The most imp this is to visualise the confusion matrix, precison and recall using heat map which help us to visualize the performance of our models so that we will able to pick best out of these.

Note: In this we will Apply All the models with tf-idf features (Replace CountVectorizer with tfidfVectorizer) and Instead of using all the words in the dataset, use only the top 1000 words based of tf-idf values

Performance

In [19]:

```
from prettytable import PrettyTable
from termcolor import colored
print(colored('Performance Table','green'))
x = PrettyTable()
x.field_names =["Models","Train","CV","Test","Misclassified(%)"]

x.add_row(["Naive Bayes (One hot coding)",0.73,1.16,1.18,0.377])
x.add_row(["KNN (Response)",0.47,1.008,1.11,0.3421])
x.add_row(["LR(Class balanced) one hot coding",0.58,1.02,1.044,0.3515])
x.add_row(["LR(Class unbalanced) one hot coding",0.57,1.08,1.088,0.3646])
x.add_row(["Lr SVM one hot encoding",0.67,1.07,1.11,0.34])
x.add_row(["Random Forest one hot coding",0.84,1.18,1.22,0.40])
x.add_row(["Random Forest Response coding",0.05,1.25,1.32,0.47])
x.add_row(["Stacking classifier",0.79,1.12,1.14,0.37])
x.add_row(["Maximum Voting Classifier",0.94,1.16,1.19,0.37])
```

print(x)

Performance Table

+	Models	İ	Train		CV	İ	Test	İ	Misclassified(%)	+
	Naive Bayes (One hot coding) KNN (Response) (Class balanced) one hot coding (Class unbalanced) one hot coding Lr SVM one hot encoding Random Forest one hot coding	 	0.73 0.47 0.58 0.57 0.67	 	1.16 1.008 1.02 1.08 1.07		1.18 1.11 1.044 1.088 1.11 1.22	1 1 1 1 1	0.377 0.3421 0.3515 0.3646 0.34 0.4	+
	Random Forest Response coding Stacking classifier Maximum Voting Classifier		0.05 0.79 0.94	 	1.25 1.12 1.16		1.32 1.14 1.19	İ	0.47 0.37 0.37	1 1 1