Bicycle Crash analysis NEISS data 1999 to 2018

National Electronic Injury Surveillance System

File: app_neiss_hospital_injuries.ipynb

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"CPSC's National Electronic Injury Surveillance System (NEISS) is a national probability sample of hospitals in the U.S. and its territories. Patient information is collected from each NEISS hospital for every emergency visit involving an injury associated with consumer products."

https://catalog.data.gov/dataset/cpscs-national-electronic-injury-surveillance-system-neiss (https://catalog.data.gov/dataset/cpscs-national-electronic-injury-surveillance-system-neiss)

https://www.cpsc.gov/cqibin/NEISSQuery/home.aspx (https://www.cpsc.gov/cqibin/NEISSQuery/home.aspx)

```
In [1]: # Imports
        import calendar
        from code_id_translator import *
        from datetime import datetime
        import graphviz
        from IPython.display import display
        import ipywidgets as widgets
        from ipywidgets import interact, fixed
        import matplotlib.dates as mdates
        from matplotlib.lines import Line2D
        import matplotlib.pyplot as plt
        from neiss backend import *
        import numpy as np
        import os
        import pandas as pd
        import pickle
        import re
        from scipy.stats import chi2 contingency
        from scipy.stats import chi2
        from sklearn import metrics
        from sklearn import preprocessing
        from sklearn.discriminant_analysis import LinearDiscriminantAnalysis
        from sklearn.feature selection import SelectKBest, f classif, mutual inf
        o classif, chi2, f regression, mutual info regression
        from sklearn.feature_selection import SelectPercentile, SelectFpr, Selec
        tFdr, SelectFwe, GenericUnivariateSelect
        from sklearn.linear model import LinearRegression, LogisticRegression
        from sklearn.metrics import mean squared error, r2 score, scorer
        from sklearn.model selection import train test split, GridSearchCV
        from sklearn.naive_bayes import GaussianNB
        from sklearn.pipeline import make pipeline
        from sklearn.pipeline import Pipeline
        from sklearn.preprocessing import StandardScaler
        from sklearn.svm import SVC
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.tree import export_graphviz, plot_tree
        import seaborn as sns
        import sys
        from time based graphs import *
        import xlrd
```

```
In [2]: %matplotlib inline
  plt.style.use('seaborn-whitegrid')
```

```
In [3]: class Includes():
    TimeBasedGraphs = True
    PearsonChiSquared = True
    PythonCorr = True
    LinearRegressionChi2 = True
    LogisticRegression = True
    GaussianNB = True
    DecisionTree = True
    LdaSvmPipeline = True
```

• Selection of geographic areas called primary sampling units (PSU) that are defined within sampling strata.

https://www.cdc.gov/nchs/nhis/singleton_psu.htm (https://www.cdc.gov/nchs/nhis/singleton_psu.htm)

Read the pickled neissYYYY.xlsx file

convert_neiss_original_data_to_pckl.ipynb is used to create the pickled file

- If the neiss_data.pckl file exists read it as the data file. - Otherwise, raise an exception. See convert_neiss_original_data_to_pckl.ipynb for the creation of the file

Reading /Users/mcorbett/Boston University/MET_CS677_DataScienceWithPyth on/Project/data/NEISS/neiss data.pckl ... done!

```
In [5]: dfNeiss.shape
Out[5]: (7352927, 19)
```

```
In [6]:
           dfNeiss.head()
Out[6]:
               CPSC Case Number Treatment Date Age Sex Race Other Race Body Part Diagnosis Ot
           0
                                                                              0
                                                                                                   71
                           100001
                                                           2
                                                                1.0
                                                                                        31
                                        1999-12-24
                                                    41
                                                                              0
           1
                           100002
                                        1999-12-27
                                                    80
                                                                2.0
                                                                                        31
                                                                                                   57
           2
                           100003
                                                                              0
                                                                                        75
                                                                                                   53
                                        1999-12-27
                                                                1.0
           3
                           100005
                                        1999-12-28
                                                                0.0
                                                                           NaN
                                                                                        94
                                                                                                   53
                                                    18
                                                           1
           4
                           100009
                                        1999-12-28
                                                    19
                                                                0.0
                                                                           NaN
                                                                                        92
                                                                                                   64
In [ ]:
```

Update Neiss with column code dictionary from Excel file

```
In [7]: column_codes_fname = neiss_pathname + '/column_codes.xlsx'
    column_dictionary = getColumnCodeDictionary(column_codes_fname)
    Neiss.setColumnCodeDictionary(column_dictionary)
```

Code to take the Neiss dictionaries for column codes and write them out to the column codes.xlsx file.

```
In [8]: neiss = Neiss(dfNeiss)
In [9]: dfNeiss.shape
Out[9]: (7352927, 19)
```

Build a categorical dataframe (with a subset of the overall data 3000 random rows)

The dataframe is built of columns that are only categorical in nature.

```
In [10]: class DataFramePruner():
             def init (self, dict prune=dict(), max output rows=None, keep end
         =True):
                 Prune a dataframe based on a dictionary and the maximum number o
         f output rows
                 Args:
                   dict prune (dict): Of the form {col name : max output catego
         ries, ...}
                   max output rows (int): If none, all rows are kept. Otherwis
         e, the dataframe is clipped to this number of rows maximum.
                   keep end (bool): True, max output rows is relative to the end
         of the dataframe. False, from beginning.
                 self.dict_prune = dict_prune
                 self.max_output_rows = max_output_rows
                 self.keep end = keep end
             def limitColumnCategoriesTo(df, column name, num categories):
                  Get the top 'num categories' most frequent names in self.df[col
         umn name]
                  Args:
                      column name (str): The name of the column to limit the cat
         egories on
                      num categories (int): The maximum number of unique values
          to retain in 'column name'
                 selected = df[column name].value counts()[:num categories].index
         .tolist()
                 return df[df[column name].isin(selected)]
             def prune(self, df):
                 for column name in self.dict prune.keys():
                     df = DataFramePruner. limitColumnCategoriesTo(df, column nam
         e, self.dict prune[column name])
                 if None != self.max output rows:
                     if True == self.keep end:
                         df = df[-self.max output rows : ]
                     else:
                         df = df[0 : self.max output rows]
                 return df
```

```
In [11]: class NeissSubset():
             def __init__(self, df, categories, dataframe_pruner=None):
                 self.df = df.copy()
                 self.categories = categories.copy()
                 self.df = self.df.xs(self.categories, axis=1)
                 self.df.dropna(inplace=True)
                 self.df['Race'] = [int(x) for x in self.df['Race']]
                 if None != dataframe pruner:
                     self.df = dataframe pruner.prune(self.df)
                 self.updateCodeIdVariables()
             def updateCodeIdVariables(self):
                 # Get the code ID translator for the dataframe
                 self.codeIdTranslator = CodeIdTranslatorDataFrame(self.df, self.
         categories)
                 self.codeIdTranslator.transformColumns()
                 # Get the code ID translators for the dataframe
                 currentState = self.codeIdTranslator.getState()
                 self.codeIdTranslator.setState('id')
                 self.dfIdToCode = self.codeIdTranslator.getDataFrame().copy()
                 self.codeIdTranslator.setState('code')
                 self.dfCodeToId = self.codeIdTranslator.getDataFrame().copy()
                 self.codeIdTranslator.setState(currentState)
             def limitColumnCategoriesTo(self, column name, num categories):
                  Get the top 'num categories' most frequent names in self.df[col
         umn name]
                  Args:
                      column name (str): The name of the column to limit the cat
         egories on
                      num categories (int): The maximum number of unique values
          to retain in 'column name'
                 selected = self.df[column name].value counts()[:num categories].
         index.tolist()
                 self.df = self.df[self.df[column name].isin(selected)]
                 #self.updateCodeIdVariables()
             def limitMaxRowsTo(self, num rows):
                 self.df = self.df.sample(num rows)
                 #self.updateCodeIdVariables()
             def getDataFrame(self):
                 return self.df
             def getCategories(self):
                 return self.categories
```

```
def getCodeIdTranslator(self):
    return self.codeIdTranslator

def getIdToCodeDataframe(self):
    return self.dfIdToCode

def getCodeToIdDataframe(self):
    return self.dfCodeToId
]: output_choice = 'Diagnosis'
```

```
In [13]: neissSubset.getDataFrame().shape
```

Out[13]: (1000000, 11)

```
In [14]: neissSubset.getDataFrame().head()
```

Out[14]:

	Sex	Race	Body_Part	Diagnosis	Disposition	Location	Fire_Involvement	Product_1	Prc
33328	3	2	16	10	1	9	1	396	
33329	3	2	5	5	1	2	1	580	
33331	3	2	9	10	1	2	1	165	
33332	2	2	6	2	1	2	1	79	
33333	2	2	11	4	1	2	1	519	

Show the IdToCode and CodeToId translators for the neissSubset dataframe

In [15]: neissSubset.getIdToCodeDataframe().head()

Out[15]:

	Sex	Race	Body_Part	Diagnosis	Disposition	Location	Fire_Involvement	Product_1	Prc
33328	3	2	16	10	1	9	1	396	
33329	3	2	5	5	1	2	1	580	
33331	3	2	9	10	1	2	1	165	
33332	2	2	6	2	1	2	1	79	
33333	2	2	11	4	1	2	1	519	

In [16]: neissSubset.getCodeToIdDataframe().head()

Out[16]:

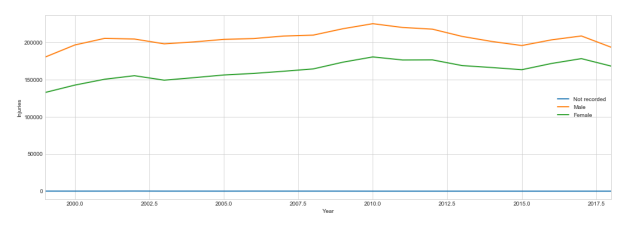
	Sex	Race	Body_Part	Diagnosis	Disposition	Location	Fire_Involvement	Product_1	Pro
33328	2	1	82	71	1	9	0	1290	
33329	2	1	34	57	1	1	0	1807	
33331	2	1	38	71	1	1	0	610	
33332	1	1	35	53	1	1	0	374	
33333	1	1	76	56	1	1	0	1616	

Show time based graphs of male/female injuries - whole dataframe used

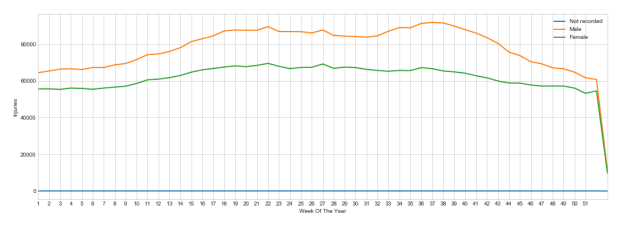
```
In [17]: if True == Includes.TimeBasedGraphs:
    stat_name = 'Sex'
    date_name = 'Treatment_Date'

    TimeBasedGraphs(dfNeiss, Neiss.getColumnDictionary(stat_name), date_name, stat_name).show()
```

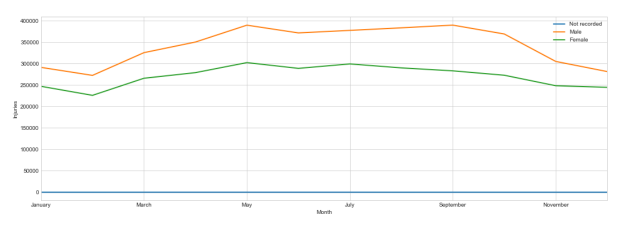
Injuries Per Year



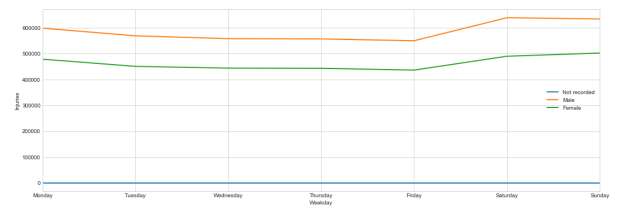
Injuries per week (1 ... 52) of the year



Injuries Per Month of the Year



Injuries Per Weekday of the Year



Functions/Classes used for the python correlations and pearson chi squared correlations

```
def getHighCorrelations(df, dfCategoricalCorrMatrix, minValue):
In [18]:
             For each column in the dataframe determine which rows equal or excee
         d the minimum value
             Args:
                                      The original dataframe
                 df (pd.DataFrame):
                 dfCategoricalCorrMatrix (list of (row name, col name) tuples):
          The categorical matrix
                 minValue (int): The value the row/column cell must equal or exc
         eed
             Returns:
               A list of tuples containing (row, column) where the value equalled
         or exceeded the minimum value
             high correlations = []
             for yIndex, y in enumerate(dfCategoricalCorrMatrix.index):
                 for xIndex, x in enumerate(dfCategoricalCorrMatrix.columns):
                     #if xIndex >= yIndex:
                          break
                     if (x != y) and (dfCategoricalCorrMatrix[y][x] > minValue):
                         y = y
                         X = x
                         if len(df[X].unique()) > len(df[Y].unique()):
                             # Keep the smallest item on the X axis
                             Y, X = X, Y
                         if (Y, X) not in high correlations:
                             high correlations.insert(-1, (Y, X))
             high correlations.sort()
             return high correlations
```

```
In [19]: class UserSelectableSwarmScatterPlots():
             def init (self, high correlations, code id translator):
                 self.button = widgets.Button(description="Click Me!")
                 self.output = widgets.Output()
                 self.high_correlations = high_correlations
                 self.code_id_translator = code_id_translator
             def show(self):
                 button = widgets.Button(description="Click Me!")
                 output = widgets.Output()
                 #print(self.high correlations.values)
                 select = self.high_correlations[0]
                 #print(select)
                 lCorrelations = ['{}, {}'.format(y, x) for y, x in self.high_cor
         relations]
                 correlationDropDownSel = widgets.Dropdown(
                     options=1Correlations,
                     value=lCorrelations[0],
                     description='correlations',
                     disabled=False,
                 )
                 typeDropDownSel = widgets.Dropdown(
                     options=['swarm', 'scatter'],
                     value='swarm',
                     description='plot_type',
                     disabled=False,
                 )
                 wHBox = widgets.HBox([correlationDropDownSel, typeDropDownSel])
                 wVBox = widgets.VBox([wHBox, button, output])
                 display(wVBox)
                 def on button clicked(b):
                     with output:
                         sel = correlationDropDownSel.value
                          (xSel, ySel) = [x.strip() for x in sel.split(',')]
                         print('-{}-, -{}-'.format(xSel, ySel))
                         correlations(self.code id translator, xSel, ySel, typeDr
         opDownSel.value)
                         #sns.pairplot(dfSel, hue=xSel)
                 button.on click(on button clicked)
```

```
In [20]: class CategoricalMatrixHeatMap:
    def __init__(self, title, dfCategoricalMatrix):
        self.title = title
        self.dfCategoricalMatrix = dfCategoricalMatrix

def show(self, figsize=(10, 10)):
        fig, ax = plt.subplots(figsize=figsize)

        g = sns.heatmap(self.dfCategoricalMatrix, annot=True, linewidths = 0.4, ax=ax)
        g.set_title(self.title)

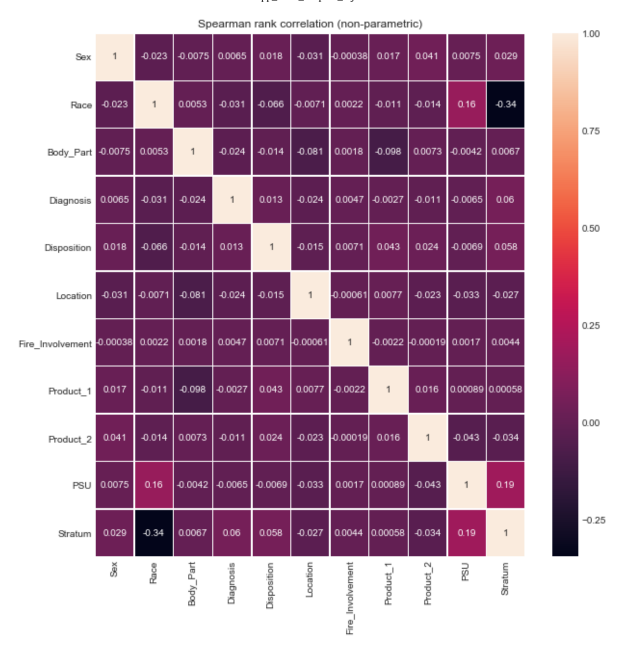
# Fix the top and bottom margins of the heatmap
        bottom_y, top_y = plt.ylim()
        bottom_y += 0.5
        top_y -= 0.5
        plt.ylim(bottom_y, top_y)
        plt.show()
```

Python (pandas.DataFrame.corr) - dataframe subset used

```
In [21]: if True == Includes.PythonCorr:
             methods = {
                  'pearson' : 'Pearson R Correlation (parametric)', # not good fo
         r categorical data
                     # For pearson:
                     # both variables should be normally distributed
                     # There should be no significant outliers
                     # Each variable should be continuous
                     # The two variables have a linear relationship
                     # The observations are paired observations.
                     # Should support homoscedascity. Homoscedascity simply refe
         rs to 'equal variances'.
                  'kendall' : 'Kendall Tau-b rank correlation (non-parametric)',
                     # The variables are measured on an ordinal or continuous sca
         le.
                     # Desirable if your data appears to follow a monotonic relat
         ionship.
                  'spearman': 'Spearman rank correlation (non-parametric)'
                     # Does not assume that both datasets are normally distribute
         d
                 }
             dfCategoricalMatrices = {}
             for key in methods.keys():
                 dfCategoricalMatrices[key] = neissSubset.getDataFrame().corr(met
         hod = key)
                 heatMap = CategoricalMatrixHeatMap(methods[key], dfCategoricalMa
         trices[key])
                 heatMap.show()
```

Pearson R Correlation (parametric) 1.00 -0.026 -0.0091 0.0055 0.012 -0.082 -0.00062 0.032 0.039 0.0081 0.032 -0.026 0.0084 -0.023 -0.049 0.018 0.0018 -0.0092 -0.013 0.14 -0.31 Race 0.75 -0.089 0.0084 0.0026 -0.0091 1 -0.0094 -0.025 0.0011 -0.12-0.0096 -0.0046 Body_Part Diagnosis 0.0055 -0.023 -0.0094 0.043 0.0056 0.0038 0.00098 -0.021 -0.0054 0.056 0.50 0.019 0.047 0.012 -0.049 -0.025 0.043 1 -0.061 0.0066 0.045 -0.011 Disposition -0.082 0.018 -0.089 0.0056 -0.061 1 -0.0052 0.008 -0.083 -0.031 -0.078 Location 0.25 Fire_Involvement -0.00062 0.0018 0.0011 0.0038 0.0066 -0.0052 -0.0032 -0.0015 0.0015 0.0043 0.032 -0.0092 -0.12 0.00098 0.045 0.008 -0.0032 0.012 -0.0022 -0.0003 Product_1 0.00 0.039 -0.013 -0.0096 -0.021 0.019 -0.083 -0.0015 0.012 -0.038 -0.034 Product_2 PSU 0.0081 0.14 -0.0046 -0.0054 -0.011 -0.031 0.0015 -0.0022 -0.038 0.19 Stratum 0.032 -0.31 0.0026 0.056 0.047 -0.078 0.0043 -0.0003 -0.034 -0.25 Product_1 Š Location Stratum Product_2 28 Race Diagnosis Fire_Involvement Body_Part Disposition

Kendall Tau-b rank correlation (non-parametric) 1.00 -0.021 -0.0063 0.018 -0.029 -0.00038 0.0062 0.026 0.0056 0.014 0.04 -0.021 0.0042 -0.025 -0.061 -0.0062 0.002 -0.0081 -0.013 0.12 -0.3 Race 0.75 0.0042 -0.065 0.0051 -0.0063 1 -0.032 -0.011 0.0015 -0.068 0.0058 -0.003 Body_Part Diagnosis 0.0056 -0.025 -0.032 1 0.011 -0.019 0.0041 -0.0022 -0.0089 -0.0048 0.047 0.50 -0.014 0.052 0.018 -0.061 -0.011 1 0.0071 0.035 0.023 -0.0056 Disposition -0.029 -0.0062 -0.065 -0.019 -0.014 1 -0.00056 0.0046 -0.02 -0.025 -0.022 Location 0.25 Fire_Involvement -0.00038 0.002 0.0015 0.0041 0.0071 -0.00056 -0.0018 | 0.00018 | 0.0014 0.004 0.014 -0.0081 -0.068 -0.0022 0.035 0.0046 -0.0018 0.013 0.00065 0.00056 Product_1 0.00 0.04 -0.013 0.0058 -0.0089 0.023 -0.02 -0.00018 0.013 -0.035 -0.029 Product_2 PSU 0.0062 0.12 -0.003 -0.0048 -0.0056 -0.025 0.0014 0.00065 -0.035 0.052 Stratum -0.3 0.0051 0.047 -0.022 0.004 0.00056 -0.029 -0.25 Product_1 Š Location Stratum Fire_Involvement Product_2 28 Race Body_Part Diagnosis Disposition



```
In [22]:
         #if True == Includes.PythonCorr:
               for key in dfCategoricalMatrices.keys():
         #
                   print('{}:\n{}'.format(key, dfCategoricalMatrices))
         #
                   print()
         if True == Includes.PythonCorr:
In [23]:
             highCorrelationsPythonCorr = getHighCorrelations(
                  neissSubset.getDataFrame(),
                  dfCategoricalMatrices['spearman'],
                  0.1)
         #if True == Includes.PythonCorr:
         #
               for n in highCorrelationsPythonCorr:
```

print(n)

#

```
In [25]: if True == Includes.PythonCorr:
    plots = UserSelectableSwarmScatterPlots(highCorrelationsPythonCorr,
    neissSubset.getCodeIdTranslator())
    plots.show()
```

 Analysis: The pandas.DataFrame.corr method is not that great when used with categorical data given an output with many categories.

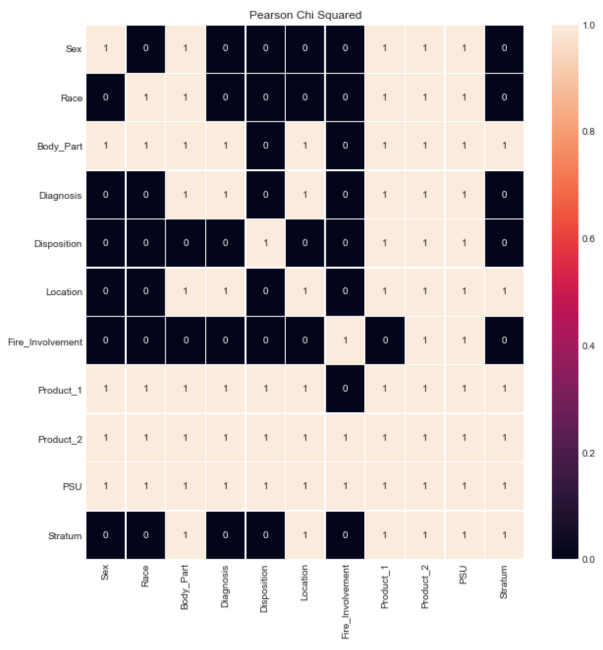
PearsonChiSquared - dataframe subset used

https://machinelearningmastery.com/chi-squared-test-for-machine-learning/ (https://machinelearningmastery.com/chi-squared-test-for-machine-learning/)

'The Pearson's chi-squared statistical hypothesis is an example of a test for independence between categorical variables.'

```
In [26]: if True == Includes.PearsonChiSquared:
    pearsonChiSquared = PearsonChiSquared(neissSubset.getDataFrame())
    dfPersonChiSquaredCategoricalCorrMatrix = pearsonChiSquared.getCorrM
    atrixDataframe(neissSubset.getCategories())
    #print(dfCategoricalCorrMatrix.head())

    heatMap = CategoricalMatrixHeatMap('Pearson Chi Squared', dfPersonChiSquaredCategoricalCorrMatrix)
    heatMap.show()
```



```
In [28]: #if True == Includes.PearsonChiSquared:
    # for n in highCorrelationsPearsonChiSquared:
    # print(n)

In [29]: if True == Includes.PearsonChiSquared:
    plots = UserSelectableSwarmScatterPlots(highCorrelationsPearsonChiSquared, neissSubset.getCodeIdTranslator())
    plots.show()
```

 Analysis: Pearson ChiSquared is a better measure of correlation between categorical data than the three python functions shown below (used with pandas.DataFrame.corr):

'pearson': 'Pearson R Correlation (parametric)', # not good for categorical data

'kendall': 'Kendall Tau-b rank correlation (non-parametric)'

'spearman': 'Spearman rank correlation (non-parametric)'

Linear regression (using chi2 & KBest) - dataframe subset used

https://scikit-learn.org/stable/modules/generated/sklearn.feature_selection.SelectKBest.html (https://scikit-learn.org/stable/modules/generated/sklearn.feature_selection.SelectKBest.html)

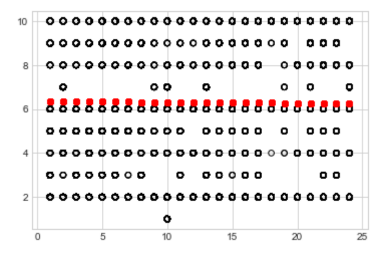
- f classif: ANOVA F-value between label/feature for classification tasks.
- mutual_info_classif: Mutual information for a discrete target.
- chi2: Chi-squared stats of non-negative features for classification tasks.
- f_regression: F-value between label/feature for regression tasks.
- mutual_info_regression: Mutual information for a continuous target.
- SelectPercentile: Select features based on percentile of the highest scores.
- SelectFpr: Select features based on a false positive rate test.
- SelectFdr: Select features based on an estimated false discovery rate.
- SelectFwe: Select features based on family-wise error rate.
- GenericUnivariateSelect: Univariate feature selector with configurable mode.

```
In [30]: if True == Includes.LinearRegressionChi2:
             class WorkingLinearRegressionChi2():
                 def __init__(self, outputFeature, categories, dfIdToCode, k_feat
         ures=2):
                      self.outputFeature = outputFeature
                      self.categories = categories.copy()
                      self.dfKBest = dfIdToCode.copy()
                      self.categories.remove(self.outputFeature)
                      self.y = self.dfKBest[self.outputFeature]
                      self.selector = SelectKBest(chi2, k=k features)
                      self.selector.fit(self.dfKBest[self.categories].values, self
         .y)
                      self.selector.get support()
                      self.selected columns = np.asarray(self.categories)[self.sel
         ector.get_support()]
                      self.X = self.dfKBest[self.selected columns]
                 def plot_scatter(X,Y,R=None):
                     plt.scatter(X, Y, s=32, marker='o', facecolors='none', edgec
         olors='k')
                      if R is not None:
                          plt.scatter(X, R, color='red', linewidth=0.5)
                     plt.show()
                 def showShape(self):
                     print('X.shape={}'.format(self.X.shape))
                     print()
                 def showSelectedColumns(self):
                      print('selected columns={}'.format(self.selected columns))
                     print()
                 def showSelectorScores(self):
                      print('selector.scores ={}'.format(self.selector.scores ))
                     print()
                 def showSelectorSupport(self):
                     print('selector.get support()={}'.format(self.selector.get s
         upport()))
                     print()
                 def showPlots(self):
                      for category in self.X:
                          print('x=', category)
                          x = np.asarray(self.dfKBest[category]).reshape(-1, 1)
                          regressor = LinearRegression(normalize=True).fit(x, self
         ·y)
                                    = regressor.predict(x)
                          y pred
                          WorkingLinearRegressionChi2.plot scatter(x, self.y, y pr
         ed)
                          print("R-squared score: {:.4f}".format(r2 score(self.y,
         y pred)))
```

```
print()
                         print()
In [31]:
        if True == Includes.LinearRegressionChi2:
             linearRegressionChi2 = WorkingLinearRegressionChi2(
                 output choice,
                 neissSubset.getCategories(),
                 neissSubset.getIdToCodeDataframe(),
                 k features=5)
In [32]: if True == Includes.LinearRegressionChi2:
             linearRegressionChi2.showShape()
             linearRegressionChi2.showSelectedColumns()
             linearRegressionChi2.showSelectorScores()
             linearRegressionChi2.showSelectorSupport()
         X.shape=(1000000, 5)
         selected_columns=['Body_Part' 'Location' 'Product_1' 'Product_2' 'PSU']
         selector.scores_=[8.72381044e+02 6.76128427e+02 3.46631214e+05 2.491715
         16e+04
          8.74721628e+04 8.11917568e-02 2.10697710e+06 9.77344423e+06
          9.79400124e+04 6.12615193e+03]
         selector.get support()=[False False True False True False True True
```

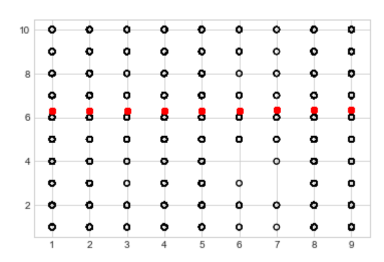
True False]

x= Body_Part



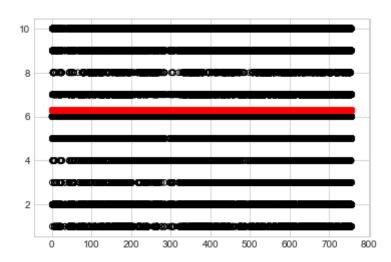
R-squared score: 0.0001

x= Location

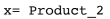


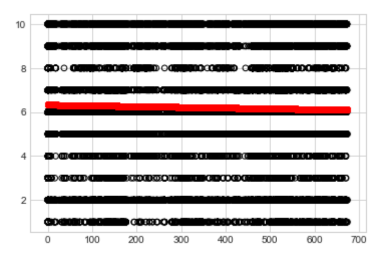
R-squared score: 0.0000

x= Product_1



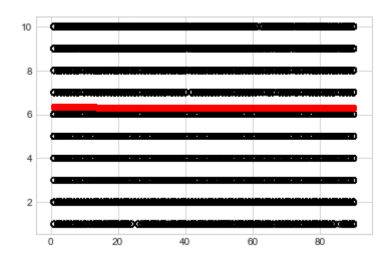
R-squared score: 0.0000





R-squared score: 0.0004

x= PSU



R-squared score: 0.0000

• Analysis: The data does not follow a linear regression model

Logistic Regression

```
In [34]: class WorkingHeatmap():
             Plots a seaborn fixing the edges of the plot
             def __init__(self, x_label, y_label,
                          title='WorkingHeatmap',
                          y bottom adjust=-1.5, y top adjust=-0.5,
                          x_left_adjust=None, x_right_adjust=None,
                          normalize_columns=False):
                 self.x label
                                        = x label
                 self.y_label
                                        = y_label
                 self.title
                                        = title
                 self.normalize columns = normalize columns
             def plot(self,
                 data, vmin=None, vmax=None, cmap=None, center=None, robust=False
                 annot=None, fmt='.2g', annot_kws=None, linewidths=0, linecolor=
         'white',
                 cbar=True, cbar kws=None, cbar ax=None, square=False, xticklabel
         s='auto',
                 yticklabels='auto', mask=None, ax=None, **kwargs):
                 if type(data) != pd.DataFrame:
                     data = pd.DataFrame(data)
                 dN = data
                 if True == self.normalize columns:
                     dT = data.copy()
                     dS = dT.sum(axis=0)[:, np.newaxis]
                     dN = dT.astype('float') / dS.T.astype('float')
                 q = sns.heatmap(
                     dN, vmin, vmax, cmap, center, robust,
                     annot, fmt, annot_kws, linewidths, linecolor,
                     cbar, cbar kws, cbar ax, square, xticklabels,
                     yticklabels, mask, ax, **kwargs)
                 g.set title(self.title)
                 ax.set xlabel(self.x label)
                 ax.set ylabel(self.y label)
                 if self.y bottom adjust or self.y top adjust or self.x left adju
         st or self.x right adjust:
                     if self.y bottom adjust or self.y top adjust:
                         bottom y, top y = ax.get ylim()
                         if self.y bottom adjust:
                             bottom y += self.y bottom adjust
```

```
if self.y_top_adjust:
    top_y += self.y_top_adjust

ax.set_ylim(bottom=bottom_y, top=top_y)

if self.x_left_adjust or self.x_right_adjust:
    left_x, right_x = ax.get_xlim()

if self.x_left_adjust:
    left_x += self.x_left_adjust

if self.x_right_adjust:
    right_x += self.x_right_adjust

ax.set_xlim(left=left_x, right=right_x)
```

```
In [35]: if True == Includes.LogisticRegression:
             class WorkingLogisticRegression():
                 Based loosly on: https://acadgild.com/blog/logistic-regression-
         multiclass-classification
                 def __init__(self, outputFeature, categories, dfIdToCode, codeId
         Translator, max iter=5000):
                     self.outputFeature = outputFeature
                     self.categories = categories.copy()
                     self.df = dfIdToCode.copy()
                     self.codeIdTranslator = codeIdTranslator
                     self.categories.remove(self.outputFeature)
                     self.inputs = self.df[self.categories]
                     self.output = self.df[self.outputFeature]
                     self.x train, self.x test, self.y train, self.y test = train
         _test_split(
                         self.inputs, self.output, test size=1/7.0, random state=
         122)
                     # Standardize the data
                     scaler = StandardScaler()
                     # Fit on training set only.
                     scaler.fit(self.x train)
                     # Apply transform to both the training set and the test set.
                     self.x train = scaler.transform(self.x train)
                     self.x_test = scaler.transform(self.x_test)
                     # Fit the model
                     # For multiclass problems, only 'newton-cg', 'sag', 'saga' a
         nd 'lbfgs' handle multinomial loss.
                     self.model = LogisticRegression(solver = 'newton-cg', multi
         class='multinomial', max iter=max iter)
                     self.model.fit(self.x train, self.y train)
                     # Validate the fitting
                     # use the model to make predictions with the test data
                     self.y pred = self.model.predict(self.x test)
                     self.probs = self.model.predict proba(self.x test)
                     test score = self.model.score(self.x test, self.y test)
                     print('test_score =', test_score)
                     # how did our model perform?
                     self.count misclassified = (self.y test != self.y pred).sum
         ()
                     self.accuracy = metrics.accuracy score(self.y test, self.y p
         red)
                     self.confusion matrix = metrics.confusion matrix(self.y test
         , self.y pred)
```

```
# Create predicted versus actual dataframe
            target names = self.output.unique()
            target dict = column dictionary[self.outputFeature]
            dfTest = pd.DataFrame(self.y_test, columns=[self.outputFeatu
re])
            self.codeIdTranslator._transform(dfTest, 'idToCode', self.ou
tputFeature)
            y test = [target dict[x] for x in dfTest[self.outputFeature
]]
            self.y test = y test
            #print('y test={}'.format( np.sort(np.unique(y test)) ))
            dfPred = pd.DataFrame(self.y pred, columns=[self.outputFeatu
re])
            self.codeIdTranslator._transform(dfPred, 'idToCode', self.ou
tputFeature)
            y pred = [target dict[x] for x in dfPred[self.outputFeature
11
            #print('y pred={}'.format( np.sort(np.unique(y pred)) ))
            dfTargetNames = pd.DataFrame(target_names, columns=[self.out
putFeature])
            self.codeIdTranslator. transform(dfTargetNames, 'idToCode',
self.outputFeature)
            target_names = [target_dict[x] for x in dfTargetNames[self.
outputFeature]]
            #print('target names={}'.format( np.sort(np.unique(target na
mes))))
            self.dfPredictedVersusActual = pd.DataFrame(self.probs, colu
mns=target names).round(4)
            self.dfPredictedVersusActual.insert(0, 'target class',
                                                                       У
test)
            self.dfPredictedVersusActual.insert(1, 'predicted_class', y_
pred)
        def showAccuracy(self):
            print('Accuracy: {:.2f}'.format(self.accuracy))
        def showMissclassifiedSamples(self):
            print('Misclassified samples: {} out of {}'.format(self.coun
t misclassified, len(self.y test)))
        def showConfusionMatrix(self):
            labels=np.unique(self.y test)
            fig, ax = plt.subplots(figsize=(14, 14))
            heatmap = WorkingHeatmap(title='WorkingLogisticRegression',
                x_label='actual', y_label='predicted',
                y_bottom_adjust=0.5, y_top_adjust=-0.5,
                normalize columns=True)
            # This sets the yticks "upright" with 0, as opposed to sidew
ays with 90.
```

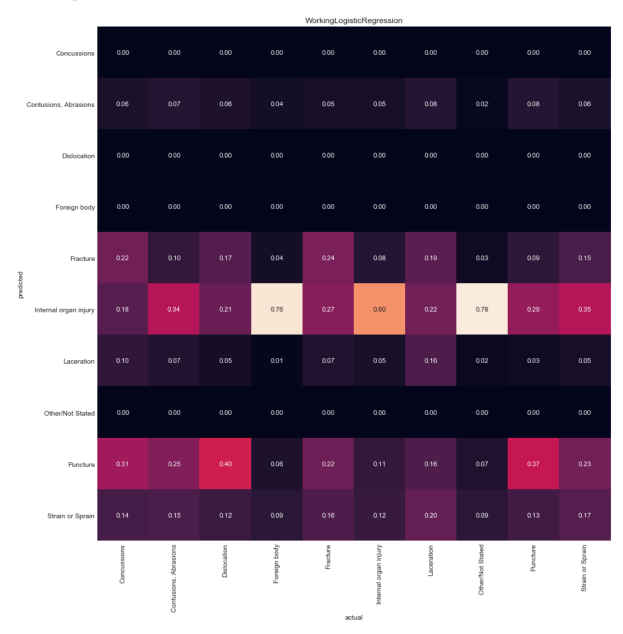
```
In [36]: if True == Includes.LogisticRegression:
    logisticRegression = WorkingLogisticRegression(
        output_choice,
        neissSubset.getCategories(),
        neissSubset.getIdToCodeDataframe(),  # neissSubset.getIdToCode
    Dataframe()[0:1500]
    neissSubset.getCodeIdTranslator(),
        max_iter=100)
```

/Users/mcorbett/anaconda3/lib/python3.6/site-packages/scipy/optimize/li nesearch.py:314: LineSearchWarning: The line search algorithm did not c onverge

warn('The line search algorithm did not converge', LineSearchWarning)
/Users/mcorbett/anaconda3/lib/python3.6/site-packages/sklearn/utils/opt
imize.py:195: UserWarning: Line Search failed
 warnings.warn('Line Search failed')

test score = 0.26411541530750815

Misclassified samples: 105127 out of 142858 Accuracy: 0.26



	target_class	predicted_class	Other/Not Stated	Fracture	Contusions, Abrasions	Foreign body	Laceration	Inte oı in
0	Fracture	Strain or Sprain	0.0141	0.1824	0.0289	0.0043	0.1487	0.1
1	Laceration	Laceration	0.0127	0.1608	0.0023	0.1164	0.1158	0.2
2	Internal organ injury	Laceration	0.0128	0.1769	0.0158	0.0130	0.1332	0.1
3	Fracture	Strain or Sprain	0.0158	0.1908	0.0353	0.0023	0.1622	0.0
4	Contusions, Abrasions	Strain or Sprain	0.0291	0.1867	0.0424	0.0021	0.1974	0.0
142853	Fracture	Strain or Sprain	0.0214	0.1948	0.0253	0.0030	0.1690	0.0
142854	Strain or Sprain	Strain or Sprain	0.0446	0.1323	0.0304	0.0021	0.1646	0.0
142855	Laceration	Contusions, Abrasions	0.0243	0.1844	0.0098	0.0184	0.1687	0.1
142856	Strain or Sprain	Contusions, Abrasions	0.0176	0.1925	0.0093	0.0203	0.1636	0.1
142857	Other/Not Stated	Fracture	0.0135	0.0778	0.0191	0.0048	0.3560	0.0

142858 rows × 12 columns

• Analysis: The data does not follow a logistic regression model

Gaussian Naive Bayes (GaussianNB) - dataframe subset used

Can perform online updates to model parameters via partial_fit method.

For details on algorithm used to update feature means and variance online,

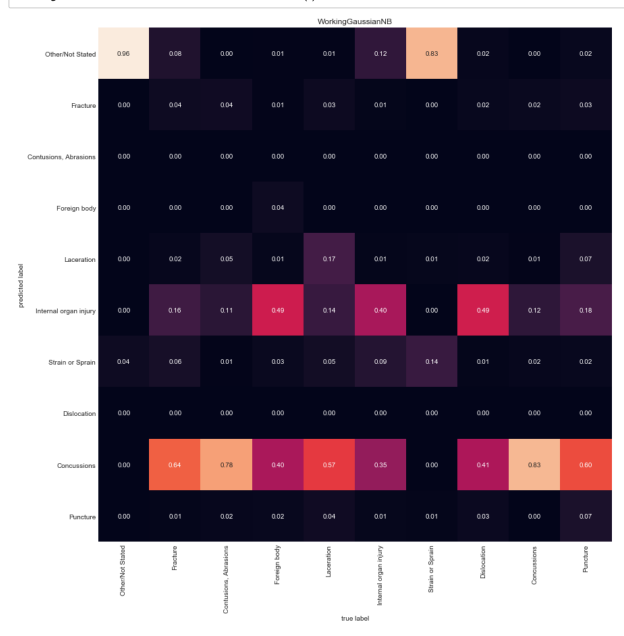
see Stanford CS tech report STAN-CS-79-773 by Chan, Golub, and LeVeque:

http://i.stanford.edu/pub/cstr/reports/cs/tr/79/773/CS-TR-79-773.pdf (http://i.stanford.edu/pub/cstr/reports/cs/tr/79/773/CS-TR-79-773.pdf)

```
In [38]: if True == Includes.GaussianNB:
             class WorkingGaussianNB():
                 def __init__(self, outputFeature, categories, dfIdToCode, codeId
         Translator, verbose=False):
                     self.outputFeature = outputFeature
                     self.categories = categories.copy()
                     self.df = dfIdToCode.copy()
                     self.codeIdTranslator = codeIdTranslator
                     self.categories.remove(outputFeature)
                     # print('categories=', self.categories)
                     # Build the Label encoder
                     self.le = {}
                     for col in self.df.columns:
                         self.le[col] = preprocessing.LabelEncoder()
                         self.le[col].fit(self.df[col].unique())
                         if True == verbose:
                             print('{0:12s} => {1}'.format(col, self.le[col].clas
         ses_))
                     self.y_test = self.df[self.outputFeature]
                     self.y labels = self.columnOutputValueIdToString(self.y test
         .unique(), self.outputFeature)
                 def columnOutputValueIdToString(self, y values, column name):
                     target dict = column dictionary[column name]
                     dfValues = pd.DataFrame(y values, columns=[column name])
                     self.codeIdTranslator. transform(dfValues, 'idToCode', colum
         n name)
                     return [target dict[x] for x in dfValues[column name]]
                 def showPredicted(self):
                     print("Number of mislabeled points out of a total {} points
          : {}, performance {:05.2f}%"
                            .format(
                                self.df.shape[0],
                                (self.y_test != self.y_pred).sum(),
                                100*(1-(self.y_test != self.y_pred).sum()/self.df.
         shape[0])
                     ))
                 def showConfusionMatrix(self):
                     fig, ax = plt.subplots(figsize=(14, 14))
                     heatmap = WorkingHeatmap(title='WorkingGaussianNB',
                         x label='true label', y label='predicted label',
                         y bottom adjust=0.5, y top adjust=-0.5,
                         normalize columns=True)
                     # This sets the yticks "upright" with 0, as opposed to sidew
         ays with 90.
                     #plt.yticks(rotation=0) # does not appear to be working...
```

```
heatmap.plot(
                self.confusion matrix.T, square=True, annot=True, fmt='.
2f', cbar=False,
                ax=ax, xticklabels=self.y labels, yticklabels=self.y lab
els)
            plt.show()
        def trainClassifier(self, show predicted versus actual=True, sho
w confusion matrices=True):
            # Drop categories with low scores
            categories = self.categories
            df = self.df
            # Train classifier
            gnb = GaussianNB()
            gnb.fit(
                df[categories].values,
                df[self.outputFeature])
            y_pred = gnb.predict(self.df[categories])
            self.y pred = y pred
            self.confusion matrix = metrics.confusion matrix(self.y test
, self.y pred,)
```

Number of mislabeled points out of a total 1000000 points: 719923, per formance 28.01%



In []:

Decision Tree - dataframe subset used

http://benalexkeen.com/decision-tree-classifier-in-python-using-scikit-learn/ (http://benalexkeen.com/decision-tree-classifier-in-python-using-scikit-learn/)

DecisionTreeClassifier(

```
class_weight=None, criterion='gini', max_depth=None,
max_features=None, max_leaf_nodes=None,
min_impurity_split=1e-07, min_samples_leaf=1,
min_samples_split=2, min_weight_fraction_leaf=0.0,
presort=False, random_state=None, splitter='best')
```

• Analysis: The GausianNB works better then either the Linear or Logistic Regressions.

```
In [43]: if True == Includes.DecisionTree:
             class WorkingDecisionTree():
                 def __init__(self, df, output_choice, dataframe pruner=None):
                     self.df = df.copy()
                     self.output_choice = output_choice
                     if None != dataframe_pruner:
                         self.df = dataframe pruner.prune(self.df)
                     # Remove columns containing NaN or columns where the number
          of unique items is greater than
                     toBeDropped = []
                     for col in self.df.columns:
                         if self.df[col].isnull().values.any():
                             toBeDropped.append(col)
                     # Also remove the case number and the narrative
                     toBeDropped.extend(['CPSC_Case_Number', 'Narrative_1'])
                     # The following dates (1999 ... 2013) contain codes that do
          not match the column codes table.
                     self.df = self.df['Treatment Date'].dt.year.isin(li
         st(range(1999, 2013)))]
                     self.df.drop(toBeDropped, axis=1, inplace=True)
                     # Remove values from dates (2014 ... 2018) that do not have
          column codes for them
                     self.df = self.df[~self.df['Product 1'].isin([1841, 1903])]
                     self.df = self.df[~self.df['Product 2'].isin([1841, 1903])]
                     self.df['mDate'] = mdates.date2num(self.df['Treatment Date'
         ])
                     self.df.drop(['Treatment Date'], axis=1, inplace=True)
                     self.dfDecisionTree = Neiss.translateCodesToIds(self.df)
                     self.replaceValueWithStringInColumn('Product 1', 0, 'Zero')
         # Keep the zero.
                     self.replaceValueWithStringInColumn('Product 2', 0, 'Zero')
         # Especially for Product 2.
                     self.replaceValueWithStringInColumn('Fire Involvement', 4,
         'InvalidCode')
                     #self.checkForNumericValueInColumn('Disposition')
                     self.checkForNumericValueInColumn('Fire Involvement')
                     self.checkForNumericValueInColumn('Product 1')
                     self.checkForNumericValueInColumn('Product 2')
                     self.dfOneHot = WorkingDecisionTree.transformToOneHotEncoded
         Dataframe(self.dfDecisionTree)
                     #print('dfOneHot.shape =', self.dfOneHot.shape)
                     #print('dfOneHot.columns =', self.dfOneHot.columns)
```

```
self.y labels org = self.columnOutputValueCodeToString(
                self.df[self.output_choice].unique(),
                self.output_choice)
            self.y labels = []
            for col_value in self.dfOneHot.columns:
                if col value[0] == self.output choice:
                    self.y_labels.append(col_value)
            self.x labels = [value for value in self.dfOneHot.columns if
value not in self.y labels]
            self.x = self.dfOneHot[self.x labels]
            self.y = self.dfOneHot[self.y labels]
            #print('x labels =', self.x labels)
            #print('x.shape =', self.x.shape)
            #print('y.shape =', self.y.shape)
            #print()
            #for col in self.dfDecisionTree.columns:
                 print('{:20s}\tnunique={}\tnum nulls={}\ttype={}'.forma
t(
            #
                     col,
            #
                     self.dfDecisionTree[col].nunique(),
            #
                     self.dfDecisionTree[col].isnull().sum(),
                     self.dfDecisionTree(col).dtype))
            print('Done')
        def testNull(obj, name):
            if pd.isnull(obj).any():
                raise Exception('Exception: {} is null'.format(name))
        def showConfusionMatrix(self):
            fig, ax = plt.subplots(figsize=(14, 14))
            heatmap = WorkingHeatmap(title='WorkingDecisionTree',
                x_label='true label', y_label='predicted label',
                y bottom adjust=0.5, y top adjust=-0.5,
                normalize columns=True)
            # This sets the yticks "upright" with 0, as opposed to sidew
ays with 90.
            #plt.yticks(rotation=0) # does not appear to be working...
            heatmap.plot(
                self.confusion matrix.T, square=True, annot=True, fmt='.
2f', cbar=False,
                ax=ax, xticklabels=self.y labels, yticklabels=self.y lab
els)
            plt.show()
        def columnOutputValueCodeToString(self, y_values, column_name):
            target dict = column dictionary[column name]
            dfValues = pd.DataFrame(y values, columns=[column name])
            return [target dict[x] for x in dfValues[column name]]
```

```
def fullTestTrainAccuracy(self, criterion='gini', graph_viz=Fals
e, out file=None, render name=None):
             # gini is the default criterion
            clfDecisionTree = DecisionTreeClassifier(criterion=criterion
)
            clfDecisionTreeFit = clfDecisionTree.fit(self.x, self.y)
            y pred = clfDecisionTreeFit.predict(self.x)
            self.y labels = self.y labels org
            self.y_test = self.columnOutputValueCodeToString(
                self.df[self.output choice], self.output choice)
            dfPredY = pd.DataFrame(y pred, columns=self.y.columns)
            dfPredY = WorkingDecisionTree.transformFromOneHotEncodedData
frame(dfPredY)
            self.y pred = dfPredY[self.output_choice]
            WorkingDecisionTree.testNull(self.y test, 'self.y test')
            WorkingDecisionTree.testNull(self.y_pred, 'self.y_pred')
            self.confusion matrix = metrics.confusion matrix(self.y test
, self.y_pred)
            # Model accuracy
            accuracy = metrics.accuracy_score(self.y, y_pred)
            print('Accuracy =', accuracy)
            # This code generates the following error for some reason:
                 Error: neiss 2013 2018: syntax error in line 743 near
            if True == graph viz:
                dot data = export graphviz(
                    clfDecisionTreeFit, out file=out file,
                    feature names = self.x labels, # inputs
                    class_names = self.y_labels,
                                                     # outputs
                    filled=True, rounded=True,
                    special characters=True)
                graph = graphviz.Source(dot data)
                if None != render name:
                    graph.render(render name)
        def runClassifier(self, classifier, X test, y test):
            y pred = classifier.predict(X test)
            dfTestY = WorkingDecisionTree.transformFromOneHotEncodedData
frame(self.y_test)
            dfPredY = pd.DataFrame(y pred, columns=self.y test.columns)
            dfPredY = WorkingDecisionTree.transformFromOneHotEncodedData
frame(dfPredY)
            self.y test = dfTestY[self.output choice]
            self.y pred = dfPredY[self.output choice]
```

```
self.confusion matrix = metrics.confusion matrix(self.y test
, self.y pred)
            self.y labels = self.y test.unique()
            #print('self.y test')
            #display(self.y test)
            #print('self.y pred')
            #display(self.y pred)
            # Model accuracy
            accuracy = metrics.accuracy_score(y_test, y_pred)
           print('Accuracy =', accuracy)
            # This code generates the following error for some reason:
                 Error: neiss 2013 2018: syntax error in line 743 near
            if True == graph_viz:
                dot_data = export_graphviz(
                    clfDecisionTreeFit, out file=out file,
                    feature_names = self.x_labels,
                                                     # inputs
                    class_names = self.y_labels,
                                                     # outputs
                    filled=True, rounded=True,
                    special_characters=True)
                graph = graphviz.Source(dot_data)
                if None != render name:
                    graph.render(render name)
        def splitTestTrainAccuracy(self,
            criterion='gini', random_state=1, test_size=0.3,
            graph viz=False, out file=None, render name=None):
            X train, X test, y train, y test = train test split(
                self.x, self.y, test size=0.3, random state=bu id)
            self.y test = y test
            clfDecisionTree = DecisionTreeClassifier(criterion=criterion
)
            clfDecisionTreeFit = clfDecisionTree.fit(X train, y train)
            print(clfDecisionTreeFit)
            self. runClassifier(clfDecisionTreeFit, X test, y test)
        def gridSearch(self, random state=1, test size=0.3,
            graph viz=False, out file=None, render name=None):
            # http://benalexkeen.com/decision-tree-classifier-in-python-
using-scikit-learn/
            #DecisionTreeClassifier(class weight=None, criterion='gini',
max depth=None,
                        max features=None, max leaf nodes=None,
            #
                         min impurity split=1e-07, min samples leaf=1,
            #
                         min samples split=2, min weight fraction leaf=
0.0,
                         presort=False, random state=None, splitter='bes
```

```
t')
            tree parameters = {
                'criterion' : ['gini', 'entropy'],
                'min samples_split' : [2, 4, 8, 16, 32, 64, 128],
                'max_depth' : [16, 32, 64, 128, 256]
            }
            X_train, X_test, y_train, y_test = train_test_split(
                self.x, self.y, test size=0.3, random state=bu id)
            self.y test = y test
            #print('X train')
            #display(X train)
            #print('y train')
            #display(y train)
            clfDecisionTree = GridSearchCV(DecisionTreeClassifier(), tre
e_parameters, cv=5)
            clfDecisionTreeFit = clfDecisionTree.fit(X train, y train)
            print(clfDecisionTreeFit)
            self. runClassifier(clfDecisionTreeFit, X test, y test)
            # num leafs = [1, 5, 10, 20, 50, 100]
        def replaceValueWithStringInColumn(self, column name, replace va
lue, with_string):
            self.dfDecisionTree[column name] = [
                with string if str(x) == '{}'.format(replace value) else
x
                for x in self.dfDecisionTree[column name]] # Replace ze
ros in col='disposition' with 'Unknown'
        def checkForNumericValueInColumn(self, columnName):
            for index, value in enumerate(self.dfDecisionTree[columnName
]):
                if type(value) == int:
                    if 'Treatment Date' in self.dfDecisionTree.columns:
                        print('{}: Found int={} at {} - {}'.format(
                            columnName, value, index, dfDecisionTree['Tr
eatment_Date'].iloc[index]))
                    else:
                        print('{}: Found int={} at {}'.format(columnName
, value, index))
        def transformFromOneHotEncodedDataframe(df):
            df1 = df.copy()
            columns = df.columns
            for col in df.columns:
                (column name, value) = col
                if column name not in df1.columns:
                    df1[column name] = np.nan
                is set = (1 == df[col])
                df1[column name][is set] = value
```

```
df1.drop(columns, axis=1, inplace=True)
            return df1
        def transformToOneHotEncodedDataframe(df):
            Take columns that are objects and turn them into multiple on
e-hot columns.
            Args:
                df
                     (pd.DataFrame): The dataframe to convert to a one
-hot dataframe
            Returns:
                dfOneHot (pd.DataFrame): A dataframe with the original
 objects replaced with one-hot versions.
            Example dataframe:
                Cost(dollars) Item
                2
                                'Baseball'
                5
                                'Baseball Glove'
                7
                                'Helmet'
            Will become a dataframe similar to:
                Cost(dollars) ('Item', 'Baseball') ('Item', 'Basebal
1 Glove') ('Item', 'Helmet)
0
                5
                                                        1
0
                7
                               0
                                                        0
1
            By making a tuple for the column header it is easy to determ
ine the variable that the one-hot
            column represents.
            dfOneHot = df.copy()
            updateFreq = 100
            print('Dots are printed every {} translations during one-hot
transformation'.format(updateFreq))
            for col in df.columns:
                if df[col].dtype == object:
                    print('One-hot transforming {:20s}'.format(col), end
='\t')
                    unique = pd.unique(df[col].sort values(ascending=Tru
e))
                    nUnique = len(unique)
                    print('nUnique={}'.format(nUnique), end='\t')
```

```
In [44]: if True == Includes.DecisionTree:
    df = neiss.getDataFrame()
    # max_output_rows = 1000000
    # max_output_rows = 20000
    # max_output_rows = 10000
    max_output_rows = 5000
    # max_output_rows = 100
    # max_output_rows = len(df) // 4
    pruner = DataFramePruner(dict_prune={output_choice : 10}, max_output_rows=max_output_rows)

    decisionTree = WorkingDecisionTree(df, output_choice, pruner)
```

```
Dots are printed every 100 translations during one-hot transformation
One-hot transforming Sex
                                                nUnique=2
complete
One-hot transforming Race
                                                nUnique=7
complete
One-hot transforming Body Part
                                                nUnique=24
complete
One-hot transforming Diagnosis
                                                nUnique=10
complete
One-hot transforming Disposition
                                                nUnique=6
complete
One-hot transforming Location
                                                nUnique=8
complete
One-hot transforming Fire Involvement
                                                nUnique=3
complete
One-hot transforming Product 1
                                                nUnique=355
complete
One-hot transforming Product 2
                                                nUnique=166
complete
One-hot transforming Stratum
                                                nUnique=5
complete
Done
```

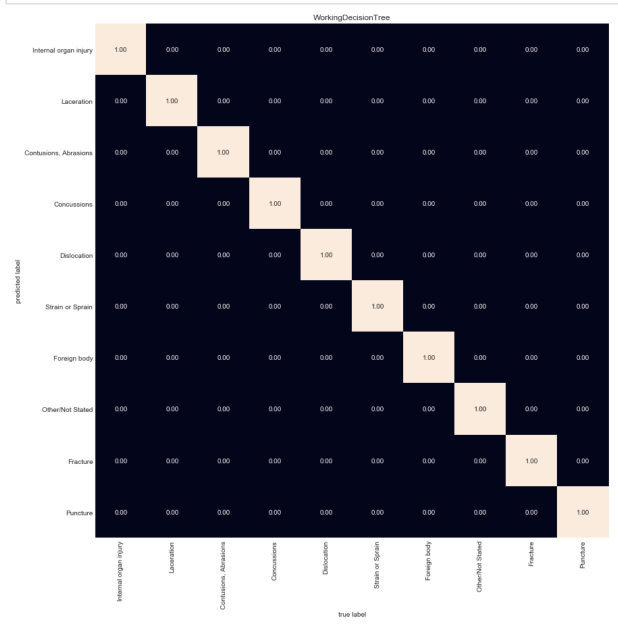
• Full with 100% training set, 100% testing set

Accuracy = 1.0

/Users/mcorbett/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:249: SettingWithCopyWarning:

A value is trying to be set on a copy of a slice from a DataFrame

See the caveats in the documentation: http://pandas.pydata.org/pandas-docs/stable/user_guide/indexing.html#returning-a-view-versus-a-copy



```
In [47]: if True == Includes.DecisionTree:
    if True == graph_viz:
    !open '{}.pdf'.format(render_name)
```

• Now with 70% training set, 30% testing set

```
In [48]: if True == Includes.DecisionTree:
    bu_id = 7286
    render_name='neiss_70_30_split'
    graph_viz=False
    decisionTree.splitTestTrainAccuracy(
        criterion='gini', graph_viz=graph_viz, render_name=render_name,
        test_size=0.3, random_state=bu_id)
```

DecisionTreeClassifier(class_weight=None, criterion='gini', max_depth=None,

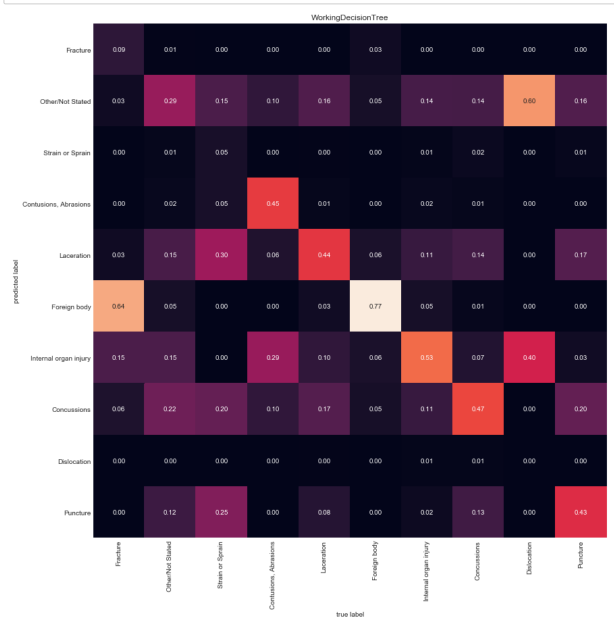
max_features=None, max_leaf_nodes=None,
min_impurity_decrease=0.0, min_impurity_split=No
min_samples_leaf=1, min_samples_split=2,
min_weight_fraction_leaf=0.0, presort=False,
random_state=None, splitter='best')

ne,

/Users/mcorbett/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:249: SettingWithCopyWarning:

A value is trying to be set on a copy of a slice from a DataFrame

See the caveats in the documentation: http://pandas.pydata.org/pandas-docs/stable/user_guide/indexing.html#returning-a-view-versus-a-copy

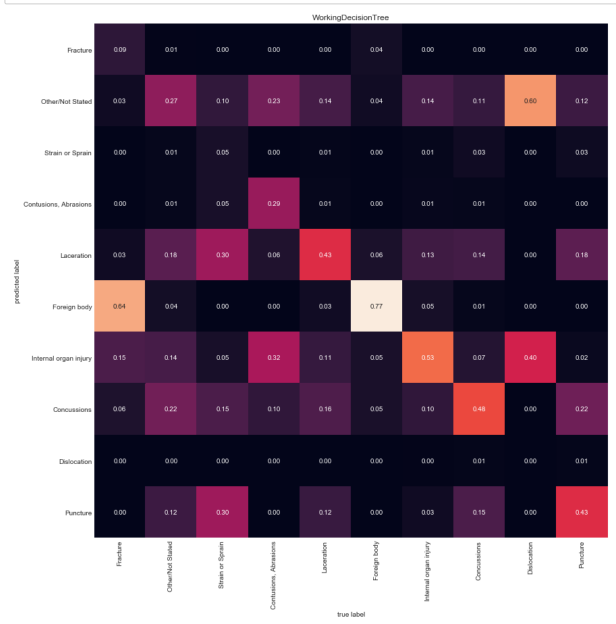


```
In [50]: if True == Includes.DecisionTree:
    if True == graph_viz:
    !open '{}.pdf'.format(render_name)
```

Now using GridSearchCV

```
In [51]: if True == Includes.DecisionTree:
             bu id = 7286
             render_name='neiss_grid_search_cv'
             graph_viz=False
             decisionTree.gridSearch(graph viz=graph viz, render name=render name
         , test_size=0.3, random_state=bu id)
         GridSearchCV(cv=5, error_score='raise-deprecating',
                      estimator=DecisionTreeClassifier(class_weight=None,
                                                        criterion='gini', max dep
         th=None,
                                                        max features=None,
                                                        max leaf nodes=None,
                                                        min_impurity_decrease=0.
         0,
                                                        min_impurity_split=None,
                                                        min samples leaf=1,
                                                        min_samples_split=2,
                                                        min weight fraction leaf=
         0.0,
                                                        presort=False, random_sta
         te=None,
                                                        splitter='best'),
                      iid='warn', n_jobs=None,
                      param grid={'criterion': ['gini', 'entropy'],
                                   'max_depth': [16, 32, 64, 128, 256],
                                   'min samples split': [2, 4, 8, 16, 32, 64, 12
         8]},
                      pre dispatch='2*n jobs', refit=True, return train score=Fa
         lse,
                      scoring=None, verbose=0)
         Accuracy = 0.4626666666666666667
         /Users/mcorbett/anaconda3/lib/python3.6/site-packages/ipykernel launche
         r.py:249: SettingWithCopyWarning:
         A value is trying to be set on a copy of a slice from a DataFrame
         See the caveats in the documentation: http://pandas.pydata.org/pandas-d
         ocs/stable/user guide/indexing.html#returning-a-view-versus-a-copy
```

```
In [52]: if True == Includes.DecisionTree:
    decisionTree.showConfusionMatrix()
```



```
In [53]: if True == Includes.DecisionTree:
    if True == graph_viz:
       !open '{}.pdf'.format(render_name)
```

• Analysis: The DecisionTree looks like it is taking shape. The accurracy is upto 50%. Unfortunately, feeding more rows into the dataframe causes a NaN error. Running out of memory?

LDA (fit based on class) / Support Vector Machines (SVM) / Pipeline

```
In [54]: if True == Includes.LdaSvmPipeline:
             class WorkingLdaSvmPipeline:
                 def __init__(self,
                              outputFeature, categories, dfIdToCode,
                              codeIdTranslator, max_iter=5000,
                              dataframe pruner=None):
                     self.outputFeature = outputFeature
                     self.categories = categories.copy()
                     self.df = dfIdToCode.copy()
                     self.codeIdTranslator = codeIdTranslator
                     self.categories.remove(self.outputFeature)
                     if None != dataframe pruner:
                         self.df = dataframe_pruner.prune(self.df)
                     self.inputs = self.df[self.categories]
                     self.output = self.df[self.outputFeature]
                     self.x train, self.x test, self.y train, self.y test = train
         _test_split(
                         self.inputs, self.output, test_size=1/7.0, random_state=
         122)
                     # Standardize the data
                     self.scaler = StandardScaler()
                     # --- ---
                     n components=self.y test.nunique()
                     print('n_components=', n_components)
                     lda tree parameters = {
                         'n components' : [n components],
                     self.clfLdaGridCV = GridSearchCV(LinearDiscriminantAnalysis
         (), lda tree parameters, cv=5)
                     # --- ---
                     svc_tree_parameters = {
                         'kernel' : ['rbf'],
                         'C'
                                       : [1000, 1E6],
                                   : ['auto'],
                         'qamma'
                         'class weight' : ['balanced']
                     self.clfSvcGridCV = GridSearchCV(SVC(), svc tree parameters,
         cv=5)
                     # --- ---
                     self.model = Pipeline([
                         ('scaler', self.scaler),
                         ('lda', self.clfLdaGridCV),
                         ('svc', self.clfSvcGridCV)
```

```
])
            print('Model built')
        def columnOutputValueIdToString(self, y_values, column_name):
            target_dict = column_dictionary[column_name]
            dfValues = pd.DataFrame(y_values, columns=[column_name])
            self.codeIdTranslator. transform(dfValues, 'idToCode', colum
n name)
            return [target_dict[x] for x in dfValues[column_name]]
        def compute(self):
            self.model.fit(self.x_train, self.y_train)
            self.y pred = self.model.predict(self.x test)
            self.confusion matrix = metrics.confusion matrix(self.y test
, self.y pred)
            self.y_labels = self.y_test.unique()
            self.y labels = self.columnOutputValueIdToString(self.y labe
ls, self.outputFeature)
            # Model accuracy
            accuracy = metrics.accuracy score(self.y test, self.y pred)
            print('Accuracy =', accuracy)
        def testNull(obj, name):
            if pd.isnull(obj).any():
                raise Exception('Exception: {} is null'.format(name))
        def showConfusionMatrix(self):
            fig, ax = plt.subplots(figsize=(14, 14))
            heatmap = WorkingHeatmap(title='WorkingDecisionTree',
                x_label='true label', y_label='predicted label',
                y_bottom_adjust=0.5, y_top_adjust=-0.5,
                normalize columns=True)
            # This sets the yticks "upright" with 0, as opposed to sidew
ays with 90.
            #plt.yticks(rotation=0) # does not appear to be working...
            heatmap.plot(
                self.confusion matrix.T, square=True, annot=True, fmt='.
2f', cbar=False,
                ax=ax, xticklabels=self.y_labels, yticklabels=self.y_lab
els)
            plt.show()
```

```
In [55]: if True == Includes.LdaSvmPipeline:
    max_output_rows = 5000
    # max_output_rows = len(df) // 4

pruner = DataFramePruner(
    dict_prune={output_choice : 10},
    max_output_rows=max_output_rows)

ldaSvmPipeline = WorkingLdaSvmPipeline(
    output_choice,
    neissSubset.getCategories(),
    neissSubset.getIdToCodeDataframe(),
    neissSubset.getCodeIdTranslator(),
    max_iter=100,
    dataframe_pruner=pruner)
```

n_components= 10
Model built

/Users/mcorbett/anaconda3/lib/python3.6/site-packages/sklearn/discrimin ant_analysis.py:466: ChangedBehaviorWarning: n_components cannot be lar ger than $min(n_features, n_classes - 1)$. Using $min(n_features, n_classes - 1) = min(10, 10 - 1) = 9$ components.

ChangedBehaviorWarning)

/Users/mcorbett/anaconda3/lib/python3.6/site-packages/sklearn/discrimin ant_analysis.py:472: FutureWarning: In version 0.23, setting n_componen ts > min(n_features, n_classes - 1) will raise a ValueError. You should set n_components to None (default), or a value smaller or equal to min (n features, n classes - 1).

warnings.warn(future_msg, FutureWarning)

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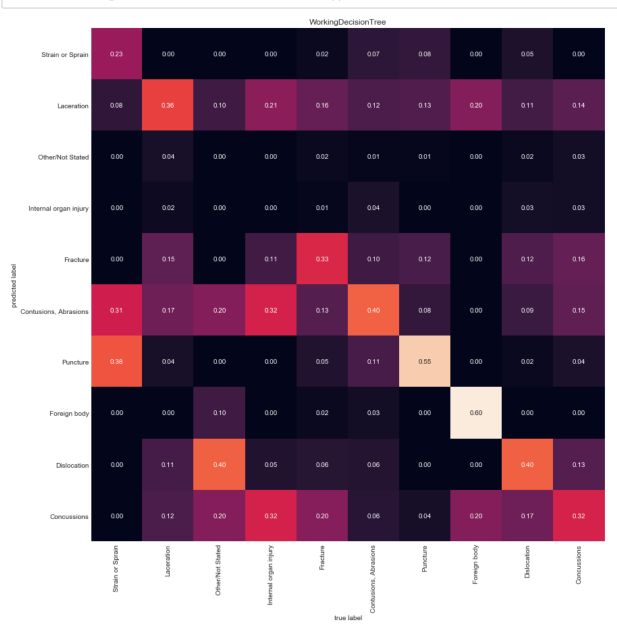
ger than $min(n_features, n_classes - 1)$. Using $min(n_features, n_classes - 1) = min(10, 10 - 1) = 9$ components.

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warnings.warn(future_msg, FutureWarning)

Accuracy = 0.3734265734265734



Analysis: For the same size dataset the DecisionTree appears to be faaster than the Scaler, LDA,SVM pipeline and provides better classification.