## chronic\_disease

## April 2, 2019

Chronic Diseases Data Analysis Cleaning Process Following are the steps we followed for data analysis

1. Import the libraries

```
In [1]: #Import the Libraries
        import numpy as np
        import pandas as pd
        import matplotlib.pyplot as plt
        %matplotlib inline
        # Algorithms
        from sklearn import linear_model
        from sklearn.linear_model import LinearRegression
        import seaborn as sns
        import warnings
        warnings.filterwarnings('ignore')
  2.Load the dataset.
In [2]: dataset=pd.read_csv('data/data.csv')
In [3]: dataset.head(3)
Out[3]:
                          Indicator Category \
        O Behavioral Health/Substance Abuse
        1 Behavioral Health/Substance Abuse
        2 Behavioral Health/Substance Abuse
                                                   Indicator Year
                                                                    Sex \
        O Opioid-Related Unintentional Drug Overdose Mor...
                                                              2010 Both
        1 Opioid-Related Unintentional Drug Overdose Mor...
                                                              2010 Both
        2 Opioid-Related Unintentional Drug Overdose Mor...
                                                              2010 Both
         Race/Ethnicity Value
                                                           Place \
        0
                    All
                            1.7
                                                  Washington, DC
```

```
1
             All
                    2.2 Fort Worth (Tarrant County), TX
2
                            Oakland (Alameda County), CA
             All
                    2.3
                          BCHC Requested Methodology \
  Age-Adjusted rate of opioid-related mortality ...
  Age-adjusted rate of opioid-related mortality ...
2 Age-adjusted rate of opioid-related mortality ...
0
  D.C. Department of Health, Center for Policy, ...
               National Center for Health Statistics
1
2
                                           CDC Wonder
                                              Methods
0
                                                  NaN
1
                                                  NaN
  Age-adjusted rate of opioid-related mortality ...
                                                Notes \
  This indicator is not exclusive of other drugs...
  This indicator is not exclusive of other drugs...
  Data is for Alameda County. This indicator is ...
   90% Confidence Level - Low 90% Confidence Level - High \
0
                          NaN
                                                        NaN
1
                          NaN
                                                        NaN
2
                                                        NaN
                          {\tt NaN}
   95% Confidence Level - Low
                               95% Confidence Level - High
0
                          NaN
                                                        3.0
1
                          1.5
2
                          1.6
                                                        3.2
```

Above we saw the column names and we might need to fix the spaces in the column names. In order to change that we need to first know what are the actual names of the columns.

We do that using the pandas function columns to list all the columns

Now we rename the columns

```
'Notes':'notes', '90% Confidence Level - Low':'90pc_con_lvl-low', '90% Confidence '95% Confidence Level - Low':'95pc_con_lvl-low','95% Confidence Level - High':'
```

3. Now we need to filter the data according to the indicator category. We use one of the values "Cancer".

```
In [6]: chronic_dis_ds = dataset.loc[dataset["indicator_category"] == "Chronic Disease"]
4.And then we remove empty columns and unnecessary columns
In [7]: chronic_dis_ds.drop(['indicator_category','bchc_req_meth','source','methods','notes','source = 1, inplace = True)
5. Now we remove all the rows which has NaN or NA values
In [8]: chronic_dis_ds.dropna(axis=0, how='any',inplace= True)
```

In [10]: chronic\_dis\_ds.head(3)

In [9]: chronic\_dis\_ds.to\_csv("data/chronic\_disease.csv")

Out[10]:		indicator year	sex \		
	4136	Asthma Emergency Department Visit Rate (Age-Ad 2010	) Both		
	4137	Asthma Emergency Department Visit Rate (Age-Ad 2010	) Both		
	4139	Asthma Emergency Department Visit Rate (Age-Ad 2010	) Both		
		race_ethnicity value place 95p	place 95pc_con_lvl-low		
	4136	All 99.4 Chicago, Il	98.1		
	4407	A33 400 0 T W (G3 1 G + ) NW	407 5		

4137	All	109.0	Las Vegas	(Clark County),	NV 107.5
4139	Asian/PI	12.9		Chicago,	Il 10.8
	95pc con lvl-h	igh			

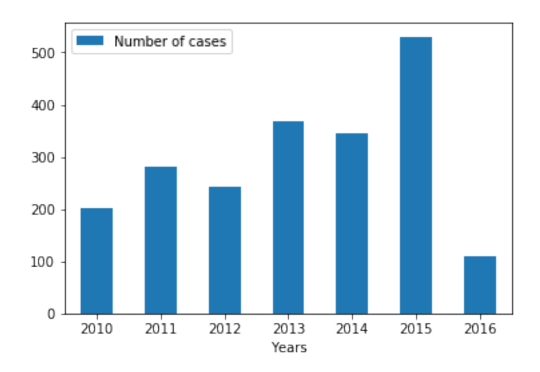
	appc_con_tvt-uign
4136	100.6
4137	110.5
4139	15.1

**Analysis** 

First we'll see how many patients have been reported for cancer in respective years from 2010 to 2016.

Following is the process to do the same

```
In [13]: c_year_2011=chronic_dis_ds[chronic_dis_ds['year']==2011]
                              c_year_2011_count=c_year_2011['year'].count()
                              c_year_2012=chronic_dis_ds[chronic_dis_ds['year']==2012]
                              c_year_2012_count=c_year_2012['year'].count()
                              c_year_2013=chronic_dis_ds[chronic_dis_ds['year']==2013]
                              c_year_2013_count=c_year_2013['year'].count()
                              c_year_2014=chronic_dis_ds[chronic_dis_ds['year']==2014]
                              c_year_2014_count=c_year_2014['year'].count()
                              c_year_2015=chronic_dis_ds[chronic_dis_ds['year']==2015]
                              c_year_2015_count=c_year_2015['year'].count()
                              c_year_2016=chronic_dis_ds[chronic_dis_ds['year']==2016]
                              c_year_2016_count=c_year_2016['year'].count()
In [14]: fig1 = pd.DataFrame({'Years':['2010', '2011', '2012','2013','2014','2015','2016'],
                                                                                                    'Number of cases': [c_year_2010_count, c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2010_count,c_year_2011_count,c_year_2010_count,c_year_2010_count,c_year_2011_count,c_year_2010_count,c_year_2011_count,c_year_2010_count,c_year_2011_count,c_year_2010_count,c_year_2011_count,c_year_2010_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_2011_count,c_year_20
                                                                                                                                                  , c_year_2013_count,c_year_2014_count,c_year_2015_
                             ax = fig1.plot.bar(x='Years', rot=0)
```

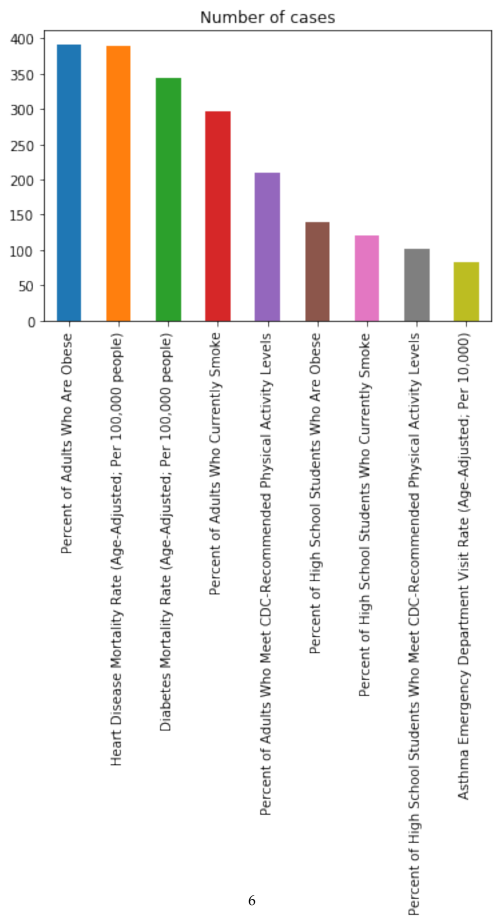


Now we calculate the number of cases for each type of cancer. In order to that we will group according to the indicator and take the count.

```
In [15]: sorted_cd = chronic_dis_ds['indicator'].value_counts()
        sorted_cd
Out[15]: Percent of Adults Who Are Obese
        Heart Disease Mortality Rate (Age-Adjusted; Per 100,000 people)
        Diabetes Mortality Rate (Age-Adjusted; Per 100,000 people)
        Percent of Adults Who Currently Smoke
        Percent of Adults Who Meet CDC-Recommended Physical Activity Levels
        Percent of High School Students Who Are Obese
        Percent of High School Students Who Currently Smoke
        Percent of High School Students Who Meet CDC-Recommended Physical Activity Levels
         Asthma Emergency Department Visit Rate (Age-Adjusted; Per 10,000)
        Name: indicator, dtype: int64
```

And we plot a histogram to see.

```
In [16]: labels=list(chronic_dis_ds.columns)
         sorted_cd = chronic_dis_ds['indicator'].value_counts().plot(title='Number of cases', :
         plt.show()
         #label=list(group.columns)
```



Now we find out the distribution of cancer patients with respect to the race and ethnicity.

```
In [17]: all=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="All"]
    all_count=all.race_ethnicity.count()

    asian=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="Asian/PI"]
    asian_count=asian.race_ethnicity.count()

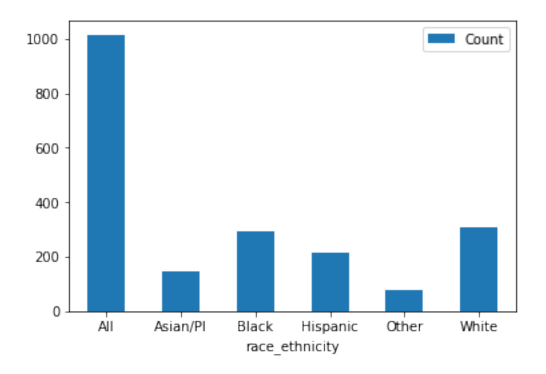
    black=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="Black"]
    black_count=black.race_ethnicity.count()

    hispanic=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="Hispanic"]
    hispanic_count=hispanic.race_ethnicity.count()

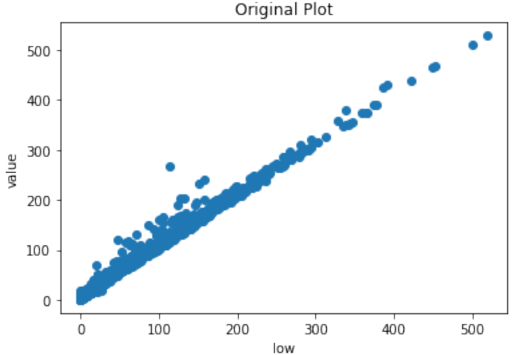
    other=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="Other"]
    other_count=other.race_ethnicity.count()

    white=chronic_dis_ds[chronic_dis_ds['race_ethnicity']=="White"]
    white_count=white.race_ethnicity.count()

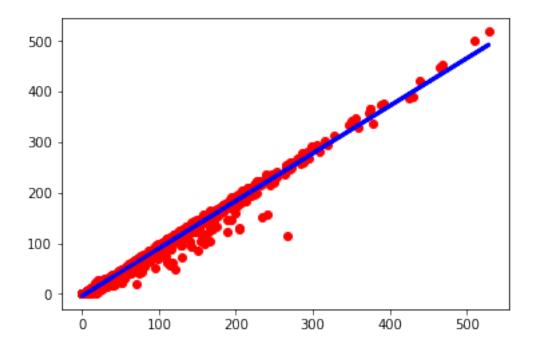
In [18]: fig2 = pd.DataFrame({'race_ethnicity':['All', 'Asian/PI', 'Black','Hispanic','Other', ax = fig2.plot.bar(x='race_ethnicity', rot=0)
```



```
In [19]: chronic_dis_ds=chronic_dis_ds.rename(columns={'95pc_con_lvl-low':'low','95pc_con_lvl-
In [20]: x='95pc_con_lvl-low'
         y='95pc_con_lvl-high'
         ds=chronic_dis_ds.drop(['indicator','year','sex','race_ethnicity'
                          ,'place','high'],
                              axis = 1)
In [21]: ds.head()
Out [21]:
               value
                        low
         4136
                99.4
                       98.1
         4137
               109.0
                      107.5
         4139
                12.9
                       10.8
         4141
                47.4
                       44.0
         4143
               226.9
                      223.6
In [22]: #very simple plotting
         fig = plt.figure(1)
         ax1 = fig.add_subplot(111)
         ax1.set_xlabel('low')
         ax1.set_ylabel('value')
         ax1.set_title('Original Plot')
         ax1.scatter('low', 'value', data = ds);
```



```
In [23]: x_y = np.array(ds)
        x, y = x_y[:,0], x_y[:,1]
         # Reshaping
         x, y = x.reshape(-1,1), y.reshape(-1, 1)
         # Linear Regression Object
         lin_regression = LinearRegression()
         # Fitting linear model to the data
         lin_regression.fit(x,y)
         # Get slope of fitted line
         m = lin_regression.coef_
         # Get y-Intercept of the Line
         b = lin_regression.intercept_
         # Get Predictions for original x values
         # you can also get predictions for new data
         predictions = lin_regression.predict(x)
         # following slope intercept form
         print ("formula: y = \{0\}x + \{1\}".format(m, b))
formula: y = [[0.93916792]]x + [-4.02478764]
In [24]: plt.scatter(x, y, color='red')
        plt.plot(x, predictions, color='blue',linewidth=3)
        plt.show()
```



In []: