

In [1]:



```
# This Python 3 environment comes with many helpful analytics libraries installed
# It is defined by the kaggle/python Docker image: https://github.com/kaggle/docker-python
# For example, here's several helpful packages to load

import numpy as np # linear algebra
import pandas as pd # data processing, CSV file I/O (e.g. pd.read_csv)

# Input data files are available in the read-only "../input/" directory
# For example, running this (by clicking run or pressing Shift+Enter) will list all files under the input directory

import os
for dirname, _, filenames in os.walk('/kaggle/input'):
    for filename in filenames:
        print(os.path.join(dirname, filename))

# You can write up to 20GB to the current directory (/kaggle/working/) that gets preserved as output when you create a version using "Save & Run All"
# You can also write temporary files to /kaggle/temp/, but they won't be saved outside of the current session
```

```
/kaggle/input/data-analysis/cdbrfss1999.csv
```

Data Analysis in Astronomy and Physics

Exercise Set 1

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1.Sampling

In [2]:

```
import os  
os.getcwd() # get current working dictory
```

Out[2]:

```
'/kaggle/working'
```

In [3]:

```
os.chdir('/kaggle/') #change dirctory to kaggle  
os.getcwd()
```

Out[3]:

```
'/kaggle'
```

In [4]:

```
os.listdir('/kaggle') # show the list of dictory 'kaggle'
```

Out[4]:

```
['src', 'lib', 'input', 'working']
```

In [5]:

```
os.listdir('/kaggle/input/data-analysis')
```

Out[5]:

```
['cdbrfss1999.csv']
```

Large data

- a. Take a sample of 30000 from this dataset and export it to an ASCII file. Make sure that your method allows to draw more than one sample from the population.
- b. Discuss your method to do so. Is your sampling a “good sample” in the sense that it is representative for the larger “population”?

solution:

- a.
 1. Get the total number of data in file by counting the number of rows and minus 1
 2. Set the desired sample size(30000)
 3. select (n-s) numbers randomly from range(1,n+1)
 4. skip the corresponding rows when reading the csv
- b. The method here is simple random sampling,each sample chosen by equal probability, so it's can be representative for larger "population"

In [6]:

```
import pandas as pd
import random

filename = 'input/data-analysis/cdbrfss1999.csv'
n = sum(1 for line in open(filename)) - 1 #number of rows in file (excludes header)
s = 30000 #desired sample size
skip = sorted(random.sample(range(1,n+1),n-s)) #select (n-s) numbers randomly from range(1,n+1). In next step, skip the corresponding row when reading the csv
cdbr = pd.read_csv(filename,skiprows = skip)
cdbr #display the 30000 samples by dataframe
```

/opt/conda/lib/python3.7/site-packages/IPython/core/interactiveshell.py:3457: DtypeWarning: Columns (8) have mixed types.Specify dtype option on import or set low_memory=False.

```
exec(code_obj, self.user_global_ns, self.user_ns)
```

Out[6]:

	STATE	GEOSTR	DENSTR	PSU	RECORD	IMONTH	IDAY	IYEAR	INTVID
0	1	1	1	10013	1	1	12	1999	33
1	1	1	1	10105	1	1	11	1999	40
2	1	1	1	10120	1	1	23	1999	33
3	1	1	1	10126	1	1	15	1999	6
4	1	1	1	10177	1	1	16	1999	0
...
29995	72	1	1	121031	1	12	19	1999	20
29996	72	1	1	121034	1	12	3	1999	6
29997	72	1	1	121048	1	12	5	1999	6
29998	72	1	1	121094	1	12	3	1999	6
29999	72	1	1	121095	1	12	1	1999	6

30000 rows × 282 columns

> Large columns

- a. Locate the columns corresponding to the variables genhlth, exerany, htf, hti, smoke100, weight, wt desire, age, and sex.
- b. Reduce your sample to include only these variables and export it to an ASCII file.

solution: export this Dataframe into an ASCII file that include only these variables

In [7]:

```
Large_Columns_b = cdbf[['GENHLTH', 'EXERANY', 'HTF', 'HTI', 'SMOKE100', 'WEIGHT', 'WTD
ESIRE', 'AGE', 'SEX']]
filename_destination= 'working/Large_Columns_b.csv'
Large_Columns_b.to_csv(filename_destination)
Large_Columns_b # display the Dataframe that question "Large Column b" required
```

Out[7]:

	GENHLTH	EXERANY	HTF	HTI	SMOKE100	WEIGHT	WTDESIRE	AGE	SEX
0	3	NaN	5	8	2	160	NaN	59	2
1	4	NaN	5	10	1	180	NaN	60	1
2	1	NaN	5	3	2	147	NaN	34	2
3	2	NaN	5	11	1	180	NaN	9	1
4	2	NaN	5	4	2	165	NaN	59	2
...
29995	4	NaN	5	8	1	190	NaN	58	1
29996	1	NaN	5	5	2	140	NaN	52	2
29997	2	NaN	5	2	2	145	NaN	42	2
29998	4	NaN	5	1	2	142	NaN	64	2
29999	3	NaN	5	0	2	127	NaN	67	2

30000 rows × 9 columns

c. How many cases and how many variables are there in your sample?

(c) 30,000 cases; 9 variables

d. What type of variable is genhlth?

(d) categorical, ordinal

e. What type of variable is weight?

(b) numerical, discrete

f. What type of variable is smoke100?

(c) categorical (not ordinal)

> One Bar chart

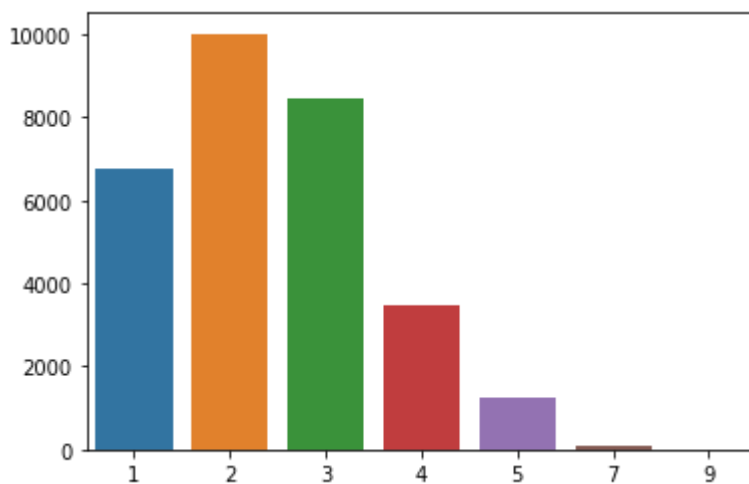
- Take all genhlth entries from your sample and draw a bar chart to visualize how the cases are distributed across the possible categories.

In [8]:

```
# x axis represents General Health. 1,2,3,4,5,7,9 means Excellent, Very good, Good, Fair, Poor, Don't know, Refused respectively
# y axis represents frequency
import numpy as np
import seaborn as sns
genhlth = Large_Columns_b['GENHLTH']
genhlth_data_count = np.unique(genhlth,return_counts = True)
sns.barplot(x = genhlth_data_count[0],y = genhlth_data_count[1])
```

Out[8]:

<AxesSubplot:>



Two Bar Charts

- Combine the smoke100 with the genhlth entries from your sample and draw two bar charts, one showing the health of the smokers and a second one showing the health of the non-smokers.

In [9]:

```

genhlth_count_ifsmoke = Large_Columns_b.groupby(['SMOKE100', 'GENHLTH']).SMOKE100
.agg('count')
genhlth_count_ifsmoke.describe()
#genhlth_count_ifsmoke['SMOKE100'].describe()
#sns.barplot(x = 'SMOKE100', y = 'count', hue = 'GENHLTH', data = genhlth_count_ifs
moke)

```

Out[9]:

```

count      25.000000
mean      1200.000000
std       1835.563288
min         1.000000
25%         4.000000
50%        22.000000
75%       1815.000000
max       5481.000000
Name: SMOKE100, dtype: float64

```

In [10]:

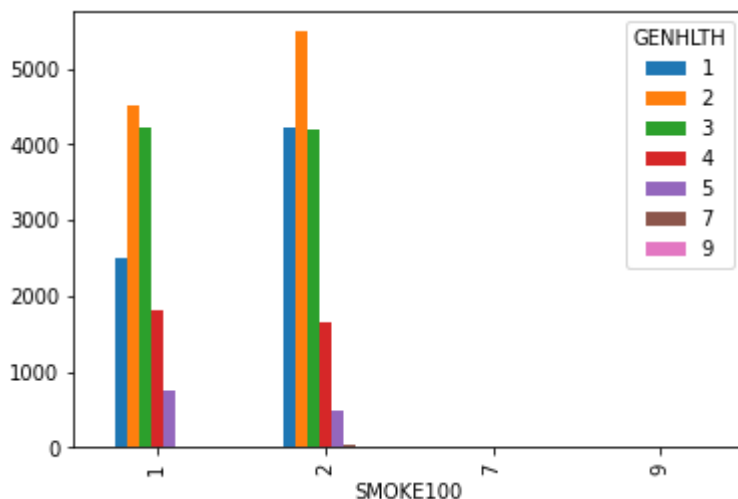
```

# x axis represents if smoke100, if so ,x=1; otherwise x=2. x=7,9 means 'not sur
e'/'refused' respectively. y axis means frequency
genhlth_count_ifsmoke.unstack().plot.bar()

```

Out[10]:

<AxesSubplot:xlabel='SMOKE100'>



> BMI

- Next let's consider a new variable bmi that doesn't show up directly in this data set: Body Mass Index (BMI).
- Compute the bmi for each case in your sample and add it to the sample (e.g. as additional column).
- Visualize the distribution of the BMI in your sample.

In [11]:

```

Large_Columns_b['BMI'] = Large_Columns_b['WEIGHT']/((Large_Columns_b['HTF']*12+L
arge_Columns_b['HTI'])*(Large_Columns_b['HTF']*12+Large_Columns_b['HTI']))*703
Large_Columns_b.drop(Large_Columns_b[Large_Columns_b.HTI > 11].index, inplace =
True) #drop the data whose height(iches) is 77 and 99(means Don't know and refused
respectively)
Large_Columns_b.drop(Large_Columns_b[Large_Columns_b.WEIGHT > 776].index, inplac
e = True)#drop the data whose weight is 777 and 999(means Don't know and refused r
espectively)
Large_Columns_b.drop(Large_Columns_b[Large_Columns_b.HTF > 7].index, inplace = T
rue)#drop the data whose height (feets) is 9(means refused)
BMI_data = Large_Columns_b['BMI']
BMI_data.drop(BMI_data[BMI_data == np.inf].index, inplace=True)# drop the data w
hose height is 0
BMI_data

```

```
/opt/conda/lib/python3.7/site-packages/ipykernel_launcher.py:1: Set
tingWithCopyWarning:
A value is trying to be set on a copy of a slice from a DataFrame.
Try using .loc[row_indexer,col_indexer] = value instead
```

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

```
"""Entry point for launching an IPython kernel.
/opt/conda/lib/python3.7/site-packages/pandas/core/frame.py:4913: S
ettingWithCopyWarning:
A value is trying to be set on a copy of a slice from a DataFrame
```

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

```
errors=errors,
/opt/conda/lib/python3.7/site-packages/pandas/core/generic.py:4153:
SettingWithCopyWarning:
A value is trying to be set on a copy of a slice from a DataFrame
```

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

```
self._update_inplace(obj)
```

Out[11]:

```
0      24.325260
1      25.824490
2      26.037037
3      25.102162
4      28.319092
...
29995   28.886246
29996   23.294675
29997   26.517950
29998   26.827734
29999   24.800278
Name: BMI, Length: 28830, dtype: float64
```

In [12]:

```
# It's a histogram, x axis represents BMI, y axis represents Frequency  
BMI_data.plot.hist('BMI', bins =60 , range=(0,60))
```

Out[12]:

<AxesSubplot:ylabel='Frequency'>

