# **Extensive Analysis + Visualization with Python**

Hello friends.

**Heart disease** or **Cardiovascular disease (CVD)** is a class of diseases that involve the heart or blood vessels. Cardiovascular diseases are the leading cause of death globally. This is true in all areas of the world except Africa. Together CVD resulted in 17.9 million deaths (32.1%) in 2015. Deaths, at a given age, from CVD are more common and have been increasing in much of the developing world, while rates have declined in most of the developed world since the 1970s.

So, in this kernel, I have conducted **Exploratory Data Analysis** or **EDA** of the heart disease dataset. **Exploratory Data Analysis** or **EDA** is a critical first step in analyzing a new dataset. The primary objective of EDA is to analyze the data for distribution, outliers and anomalies in the dataset. It enable us to direct specific testing of the hypothesis. It includes analysing the data to find the distribution of data, its main characteristics, identifying patterns and visualizations. It also provides tools for hypothesis generation by visualizing and understanding the data through graphical representation.

I hope you learn and enjoy this kernel.

So, your upvote would be highly appreciated.

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# 1. Introduction to EDA

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Several questions come to mind when we come across a new dataset. The below list shed light on some of these questions:-

- What is the distribution of the dataset?
- Are there any missing numerical values, outliers or anomalies in the dataset?
- What are the underlying assumptions in the dataset?
- Whether there exists relationships between variables in the dataset?
- How to be sure that our dataset is ready for input in a machine learning algorithm?
- How to select the most suitable algorithm for a given dataset?

So, how do we get answer to the above questions?

The answer is **Exploratory Data Analysis**. It enable us to answer all of the above questions.

# 2. Objectives of EDA

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The objectives of the EDA are as follows:-

i. To get an overview of the distribution of the dataset.

- ii. Check for missing numerical values, outliers or other anomalies in the dataset.
- iii.Discover patterns and relationships between variables in the dataset.
- iv. Check the underlying assumptions in the dataset.

# 3. Types of EDA

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EDA is generally cross-classified in two ways. First, each method is either non-graphical or graphical. Second, each method is either univariate or multivariate (usually bivariate). The non-graphical methods provide insight into the characteristics and the distribution of the variable(s) of interest. So, non-graphical methods involve calculation of summary statistics while graphical methods include summarizing the data diagrammatically.

There are four types of exploratory data analysis (EDA) based on the above crossclassification methods. Each of these types of EDA are described below:-

### i. Univariate non-graphical EDA

The objective of the univariate non-graphical EDA is to understand the sample distribution and also to make some initial conclusions about population distributions. Outlier detection is also a part of this analysis.

### ii. Multivariate non-graphical EDA

Multivariate non-graphical EDA techniques show the relationship between two or more variables in the form of either cross-tabulation or statistics.

### iii. Univariate graphical EDA

In addition to finding the various sample statistics of univariate distribution (discussed above), we also look graphically at the distribution of the sample. The non-graphical methods are quantitative and objective. They do not give full picture of the data. Hence, we need graphical methods, which are more qualitative in nature and presents an overview of the data.

### iv. Multivariate graphical EDA

There are several useful multivariate graphical EDA techniques, which are used to look at the distribution of multivariate data. These are as follows:-

- Side-by-Side Boxplots
- Scatterplots
- Heat Maps and 3-D Surface Plots

Enough of theory, now let the journey begin.

The first step in the EDA journey is to import the libraries.

# 4. Import libraries

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```
In [5]: # This Python 3 environment comes with many helpful analytics libraries installed
# It is defined by the kaggle/python docker image: https://github.com/kaggle/docke
# For example, here's several helpful packages to load i

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

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

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

# Any results you write to the current directory are as saved as output.
```

We can see that the input folder contains one input file named heart.csv.

```
In [351]: import seaborn as sns
   import matplotlib.pyplot as plt
   import scipy.stats as st
   %matplotlib inline
   sns.set(style="whitegrid")

In [8]: # ignore warnings
   import warnings
   warnings.filterwarnings('ignore')
```

I have imported the libraries. The next step is to import the datasets.

# 5. Import dataset

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I will import the dataset with the usual pandas read\_csv() function which is used to import CSV (Comma Separated Value) files.

```
In [9]: df = pd.read_csv(r'C:\Users\gadel\OneDrive\Desktop\Nareshit DataScience by Prakask
In [13]: df # to check wheather we imported the dataset or not we need to print the filenam
```

Out[13]:		age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	tl
	0	63	1	3	145	233	1	0	150	0	2.3	0	0	
	1	37	1	2	130	250	0	1	187	0	3.5	0	0	
	2	41	0	1	130	204	0	0	172	0	1.4	2	0	
	3	56	1	1	120	236	0	1	178	0	0.8	2	0	
	4	57	0	0	120	354	0	1	163	1	0.6	2	0	
	•••													
	298	57	0	0	140	241	0	1	123	1	0.2	1	0	
	299	45	1	3	110	264	0	1	132	0	1.2	1	0	
	300	68	1	0	144	193	1	1	141	0	3.4	1	2	
	301	57	1	0	130	131	0	1	115	1	1.2	1	1	
	302	57	0	1	130	236	0	0	174	0	0.0	1	1	

303 rows × 14 columns



# 6. Exploratory Data Analysis

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Evertything has been set up. Now let the actual Practicle Begin.

# Check shape of the dataset

• It is a good idea to first check the shape of the dataset.

```
In [15]: # print the shape
print('The shape of the datase : ', df.shape)
```

The shape of the datase: (303, 14)

Now, we can see that the dataset contains 303 instances and 14 variables.

### Preview the dataset

```
In [17]: # preview datase
df.head()
```

Out[17]:		age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	ca	thal
	0	63	1	3	145	233	1	0	150	0	2.3	0	0	1
	1	37	1	2	130	250	0	1	187	0	3.5	0	0	2
	2	41	0	1	130	204	0	0	172	0	1.4	2	0	2
	3	56	1	1	120	236	0	1	178	0	0.8	2	0	2
	4	57	0	0	120	354	0	1	163	1	0.6	2	0	2
	4													•

# Summary of dataset

```
In [19]: # summary of dataset
     df.info()
```

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 303 entries, 0 to 302
Data columns (total 14 columns):

#	Column	Non-Null Count	Dtype
0	age	303 non-null	int64
1	sex	303 non-null	int64
2	ср	303 non-null	int64
3	trestbps	303 non-null	int64
4	chol	303 non-null	int64
5	fbs	303 non-null	int64
6	restecg	303 non-null	int64
7	thalach	303 non-null	int64
8	exang	303 non-null	int64
9	oldpeak	303 non-null	float64
10	slope	303 non-null	int64
11	ca	303 non-null	int64
12	thal	303 non-null	int64
13	target	303 non-null	int64

dtypes: float64(1), int64(13)
memory usage: 33.3 KB

# Dataset description

- The dataset contains several columns which are as follows
  - age : age in years
  - sex: (1 = male; 0 = female)
  - cp : chest pain type
  - trestbps: resting blood pressure (in mm Hg on admission to the hospital)
  - chol: serum cholestoral in mg/dl
  - fbs: (fasting blood sugar > 120 mg/dl) (1 = true; 0 = false)
  - restecg : resting electrocardiographic results
  - thalach : maximum heart rate achieved
  - exang : exercise induced angina (1 = yes; 0 = no)
  - oldpeak : ST depression induced by exercise relative to rest
  - slope : the slope of the peak exercise ST segment

- ca: number of major vessels (0-3) colored by flourosopy
- thal: 3 = normal; 6 = fixed defect; 7 = reversable defect
- target: 1 or 0

### Check the data types of columns

- The above df.info() command gives us the number of filled values along with the data types of columns.
- If we simply want to check the data type of a particular column, we can use the following command.

```
In [21]:
         df.dtypes
Out[21]:
                        int64
         age
                        int64
         sex
         ср
                        int64
         trestbps
                        int64
         chol
                        int64
         fbs
                        int64
         restecg
                        int64
         thalach
                        int64
         exang
                        int64
                     float64
         oldpeak
         slope
                        int64
                        int64
         ca
         thal
                        int64
                        int64
         target
         dtype: object
```

### Important points about dataset

- sex is a character variable. Its data type should be object. But it is encoded as (1 = male; 0 = female). So, its data type is given as int64.
- Same is the case with several other variables fbs , exang and target .
- fbs (fasting blood sugar) should be a character variable as it contains only 0 and 1 as values (1 = true; 0 = false). As it contains only 0 and 1 as values, so its data type is given as int64.
- exang (exercise induced angina) should also be a character variable as it contains only 0 and 1 as values (1 = yes; 0 = no). It also contains only 0 and 1 as values, so its data type is given as int64.
- target should also be a character variable. But, it also contains 0 and 1 as values. So, its data type is given as int64.

### Statistical properties of dataset

```
In [25]: df.describe()
```

Out[25]:

	age	sex	ср	trestbps	chol	fbs	reste
count	303.000000	303.000000	303.000000	303.000000	303.000000	303.000000	303.0000
mean	54.366337	0.683168	0.966997	131.623762	246.264026	0.148515	0.5280
std	9.082101	0.466011	1.032052	17.538143	51.830751	0.356198	0.5258
min	29.000000	0.000000	0.000000	94.000000	126.000000	0.000000	0.0000
25%	47.500000	0.000000	0.000000	120.000000	211.000000	0.000000	0.0000
50%	55.000000	1.000000	1.000000	130.000000	240.000000	0.000000	1.0000
75%	61.000000	1.000000	2.000000	140.000000	274.500000	0.000000	1.0000
max	77.000000	1.000000	3.000000	200.000000	564.000000	1.000000	2.0000
4							•

### Important points to note

- The above command df.describe() helps us to view the statistical properties of numerical variables. It excludes character variables.
- If we want to view the statistical properties of character variables, we should run the following command -

```
df.describe(include=['object'])
```

• If we want to view the statistical properties of all the variables, we should run the following command -

```
df.describe(include='all')
```

#### View column names

# 7. Univariate analysis

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# Analysis of target feature variable

- Our feature variable of interest is target .
- It refers to the presence of heart disease in the patient.

- It is integer valued as it contains two integers 0 and 1 (0 stands for absence of heart disease and 1 for presence of heart disease).
- So, in this section, I will analyze the target variable.

### Check the number of unique values in target variable

```
In [43]: df['target'].nunique()
Out[43]: 2
```

We can see that there are 2 unique values in the target variable.

### View the unique values in target variable

```
In [49]: df['target'].unique()
Out[49]: array([1, 0], dtype=int64)
```

#### Comment

So, the unique values are 1 and 0. (1 stands for presence of heart disease and 0 for absence of hear disease).

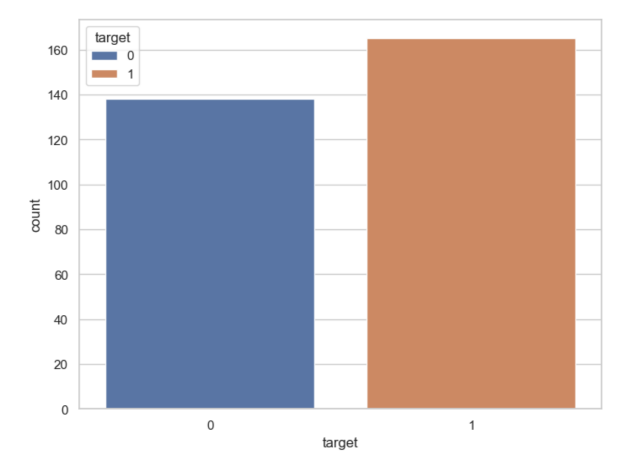
# Frequency distribution of target variable

#### Comment

- 1 stands for presence of heart disease. So, there are 165 patients suffering from heart disease.
- Similarly, 0 stands for absence of heart disease. So, there are 138 patients who do not have any heart disease.
- We can visualize this information below.

# Visualize frequency distribution of target variable

```
In [79]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="target", data=df,hue="target")
    plt.show()
```



- The above plot confirms the findings that -
  - There are 165 patients suffering from heart disease, and
  - There are 138 patients who do not have any heart disease.

# Frequency distribution of target variable wrt sex

```
In [75]: df.groupby('sex')['target'].value_counts()

Out[75]: sex target
    0     1     72
          0     24
    1    0     114
          1     93
    Name: count, dtype: int64
```

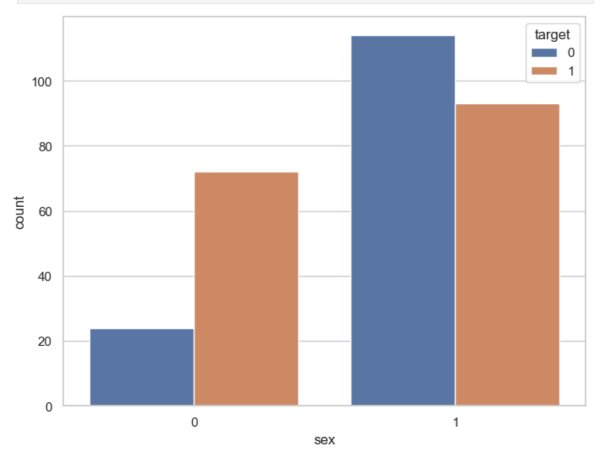
#### Comment

- sex variable contains two integer values 1 and 0 : (1 = male; 0 = female).
- target variable also contains two integer values 1 and 0 : (1 = Presence of heart disease; 0 = Absence of heart disease)
- So, out of 96 females 72 have heart disease and 24 do not have heart disease.

- Similarly, out of 207 males 93 have heart disease and 114 do not have heart disease.
- We can visualize this information below.

We can visualize the value counts of the sex variable wrt target as follows -

```
In [81]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="sex", hue="target", data=df)
    plt.show()
```



### Interpretation

- We can see that the values of target variable are plotted wrt sex : (1 = male; 0 = female).
- target variable also contains two integer values 1 and 0 : (1 = Presence of heart disease; 0 = Absence of heart disease)
- The above plot confirms our findings that -
  - Out of 96 females 72 have heart disease and 24 do not have heart disease.
  - Similarly, out of 207 males 93 have heart disease and 114 do not have heart disease.

Alternatively, we can visualize the same information as follows:

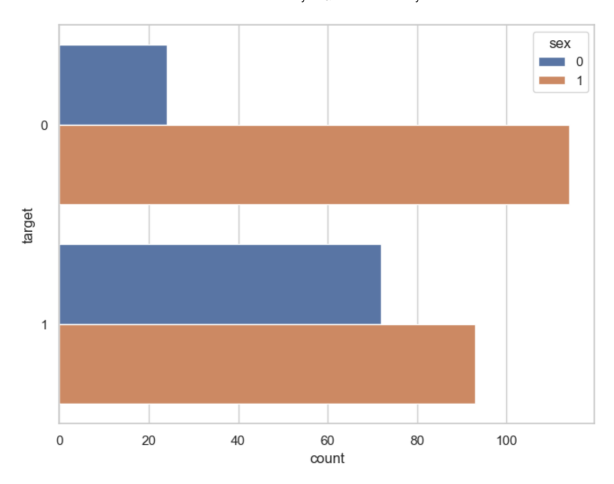


### Comment

- The above plot segregate the values of target variable and plot on two different columns labelled as (sex = 0, sex = 1).
- I think it is more convinient way of interpret the plots.

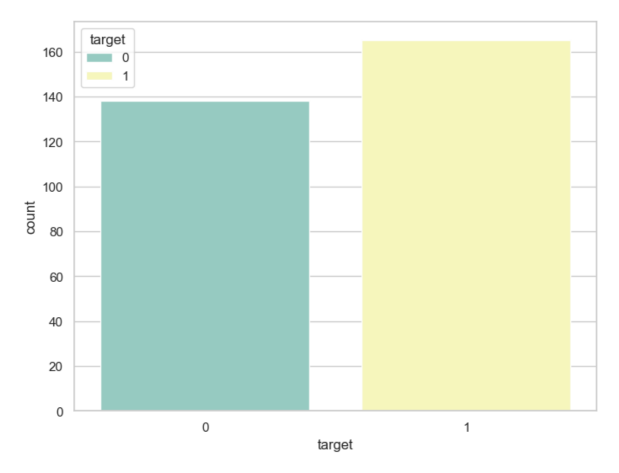
We can plot the bars horizontally as follows:

```
In [93]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(y="target", hue="sex", data=df)
    plt.show()
```

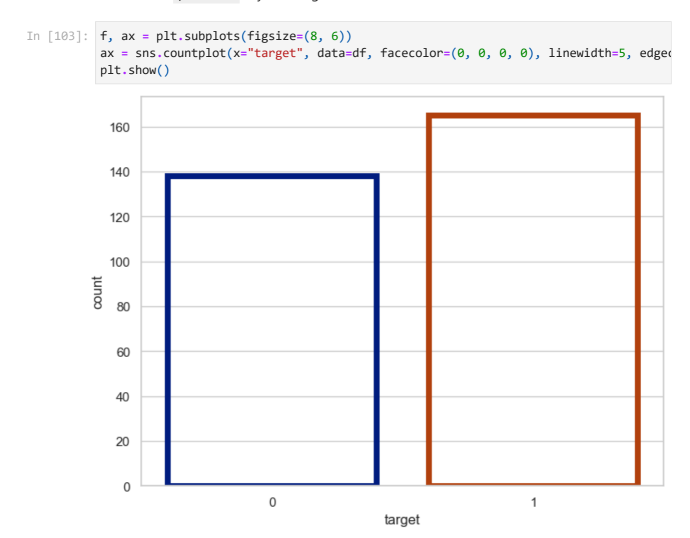


We can use a different color palette as follows :

```
In [100]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="target", hue="target", data=df, palette="Set3")
    plt.show()
```



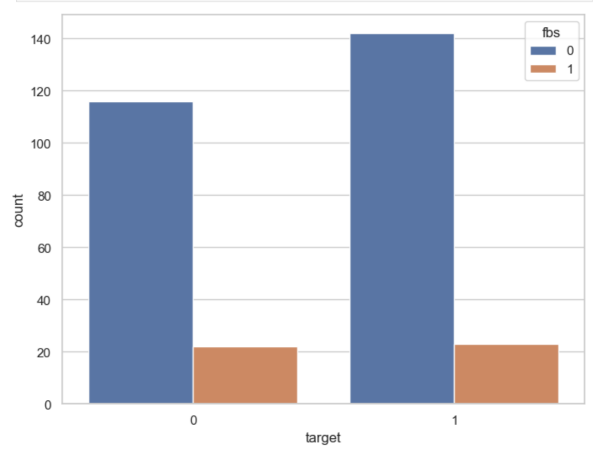
We can use plt.bar keyword arguments for a different look :



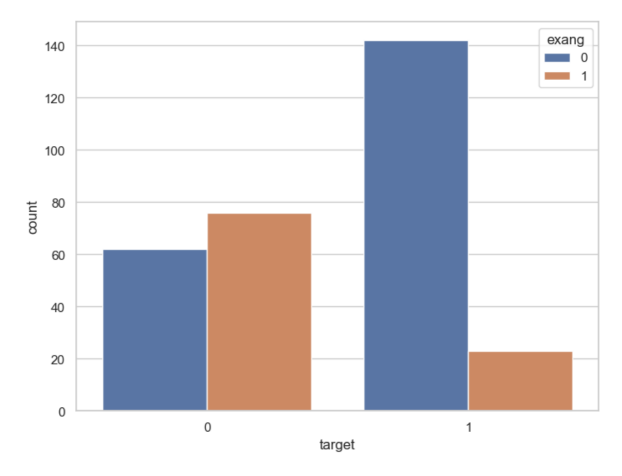
### Comment

- I have visualize the target values distribution wrt sex .
- We can follow the same principles and visualize the target values distribution wrt fbs (fasting blood sugar) and exang (exercise induced angina).

```
In [106]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="target", hue="fbs", data=df)
    plt.show()
```



```
In [112]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="target", hue="exang", data=df)
    plt.show()
```



# **Findings of Univariate Analysis**

Findings of univariate analysis are as follows:-

- Our feature variable of interest is target .
- It refers to the presence of heart disease in the patient.
- It is integer valued as it contains two integers 0 and 1 (0 stands for absence of heart disease and 1 for presence of heart disease).
- 1 stands for presence of heart disease. So, there are 165 patients suffering from heart disease.
- Similarly, 0 stands for absence of heart disease. So, there are 138 patients who do not have any heart disease.
- There are 165 patients suffering from heart disease, and
- There are 138 patients who do not have any heart disease.
- Out of 96 females 72 have heart disease and 24 do not have heart disease.
- Similarly, out of 207 males 93 have heart disease and 114 do not have heart disease.

# 8. Bivariate Analysis

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### **Estimate correlation coefficients**

Our dataset is very small. So, I will compute the standard correlation coefficient (also called Pearson's r) between every pair of attributes. I will compute it using the df.corr() method as follows:-

```
In [117]: correlation = df.corr()
```

The target variable is target . So, we should check how each attribute correlates with the target variable. We can do it as follows:-

```
In [120]:
          correlation['target'].sort_values(ascending=False)
Out[120]: target
                      1.000000
          ср
                      0.433798
                      0.421741
          thalach
                      0.345877
          slope
                      0.137230
          restecg
          fbs
                     -0.028046
          chol
                     -0.085239
          trestbps -0.144931
                     -0.225439
          age
                     -0.280937
          sex
          thal
                     -0.344029
          ca
                     -0.391724
          oldpeak
                     -0.430696
                     -0.436757
          exang
          Name: target, dtype: float64
```

#### Interpretation of correlation coefficient

- The correlation coefficient ranges from -1 to +1.
- When it is close to +1, this signifies that there is a strong positive correlation. So, we can see that there is no variable which has strong positive correlation with target variable.
- When it is clsoe to -1, it means that there is a strong negative correlation. So, we can see that there is no variable which has strong negative correlation with target variable.
- When it is close to 0, it means that there is no correlation. So, there is no correlation between target and fbs.
- We can see that the cp and thalach variables are mildly positively correlated with target variable. So, I will analyze the interaction between these features and target variable.

# Analysis of target and cp variable

# Explore cp variable

- cp stands for chest pain type.
- First, I will check number of unique values in cp variable.

```
In [125]: df['cp'].nunique()
```

Out[125]: 4

So, there are 4 unique values in cp variable. Hence, it is a categorical variable.

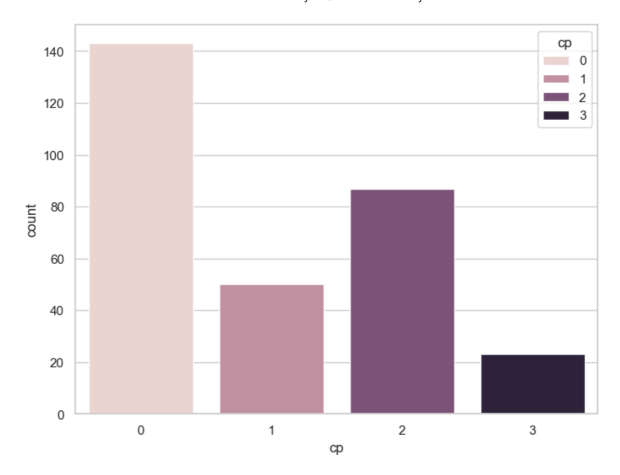
Now, I will view its frequency distribution as follows:

#### Comment

• It can be seen that cp is a categorical variable and it contains 4 types of values - 0, 1, 2 and 3.

# Visualize the frequency distribution of cp variable

```
In [168]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.countplot(x="cp", hue="cp",data=df)
    plt.show()
```



# Frequency distribution of target variable wrt cp

In [173]:	<pre>df.groupby('cp')['target'].value_counts()</pre>					
Out[173]:	ср	target				
		0	104			
		1	39			
	1	1	41			
		0	9			
	2	1	69			
		0	18			
	3	1	16			
		0	7			
	Nam	e: count,	dtype:	int64		

### Comment

- cp variable contains four integer values 0, 1, 2 and 3.
- target variable contains two integer values 1 and 0 : (1 = Presence of heart disease; 0 = Absence of heart disease)
- So, the above analysis gives target variable values categorized into presence and absence of heart disease and groupby cp variable values.
- We can visualize this information below.

We can visualize the value counts of the cp variable wrt target as follows -



0

0

- We can see that the values of target variable are plotted wrt cp.
- target variable contains two integer values 1 and 0 : (1 = Presence of heart disease; 0 = Absence of heart disease)

ср

• The above plot confirms our above findings,

Alternatively, we can visualize the same information as follows:



Analysis of target and thalach variable

3

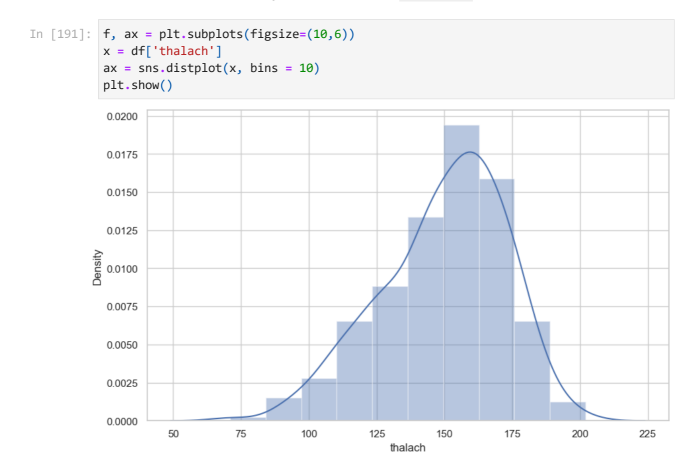
### Explore thalach variable

- thalach stands for maximum heart rate achieved.
- I will check number of unique values in thalach variable as follows:

```
In [187]: df['thalach'].nunique()
Out[187]: 91
```

- So, number of unique values in thalach variable is 91. Hence, it is numerical variable.
- I will visualize its frequency distribution of values as follows:

### Visualize the frequency distribution of thalach variable

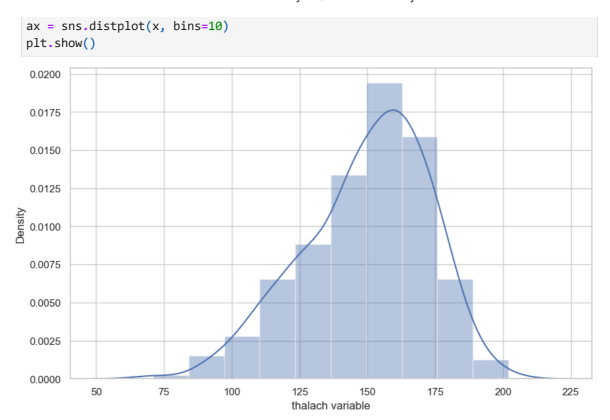


#### Comment

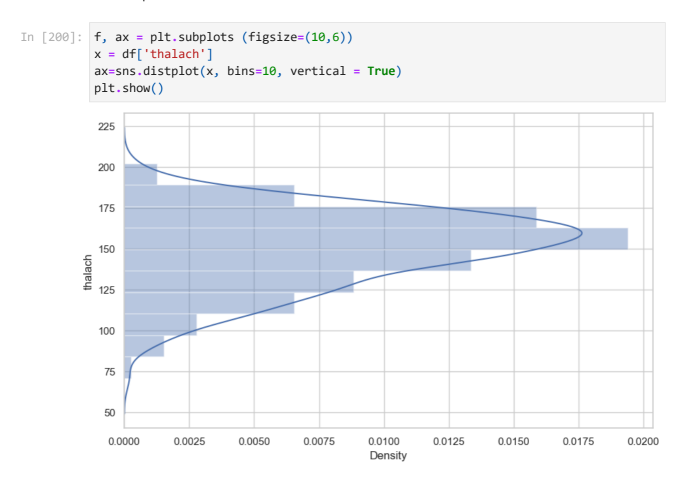
• We can see that the thalach variable is slightly negatively skewed.

We can use Pandas series object to get an informative axis label as follows:

```
In [195]:
    f, ax = plt.subplots(figsize = (10,6))
    x = df['thalach']
    x = pd.Series(x, name='thalach variable')
```



We can plot the distribution on the vertical axis as follows:-



# Seaborn Kernel Density Estimation (KDE) Plot

• The kernel density estimate (KDE) plot is a useful tool for plotting the shape of a distribution.

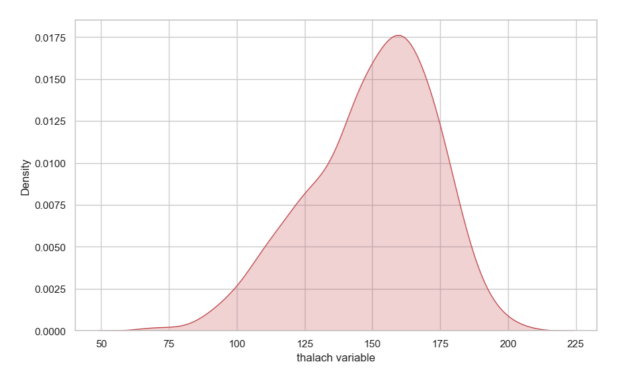
- The KDE plot plots the density of observations on one axis with height along the other axis.
- We can plot a KDE plot as follows:

```
In [203]: f, ax = plt.subplots(figsize=(10,6))
            x = df['thalach']
            x = pd.Series(x, name="thalach variable")
            ax = sns.kdeplot(x)
            plt.show()
            0.0175
            0.0150
            0.0125
          Density
0.0100
            0.0075
            0.0050
            0.0025
            0.0000
                                  75
                       50
                                             100
                                                        125
                                                                   150
                                                                                          200
                                                                                                     225
                                                                              175
```

We can shade under the density curve and use a different color as follows:

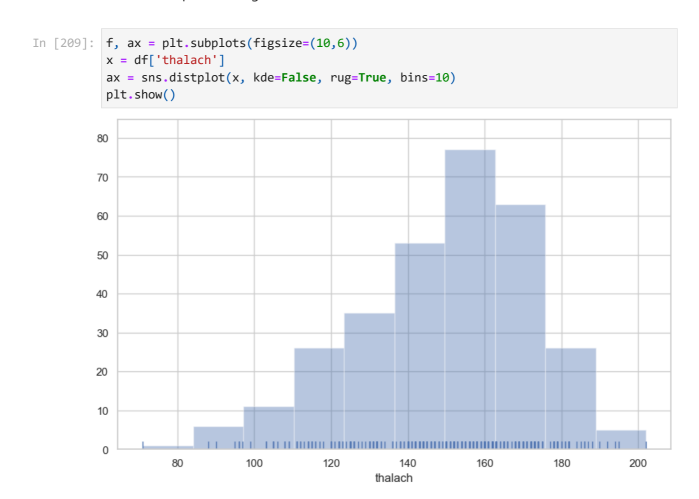
```
In [206]:
    f, ax = plt.subplots(figsize=(10,6))
    x = df['thalach']
    x = pd.Series(x, name="thalach variable")
    ax = sns.kdeplot(x, shade=True, color='r')
    plt.show()
```

thalach variable

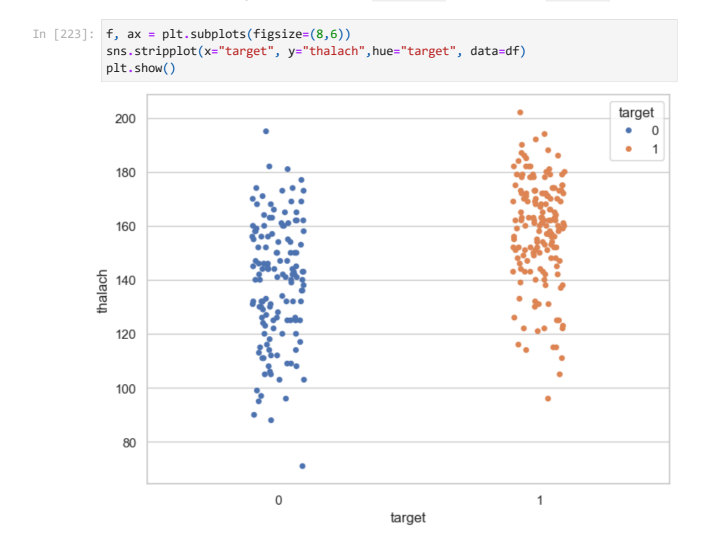


# Histogram

- A histogram represents the distribution of data by forming bins along the range of the data and then drawing bars to show the number of observations that fall in each bin.
- We can plot a histogram as follows:



# Visualize frequency distribution of thalach variable wrt target

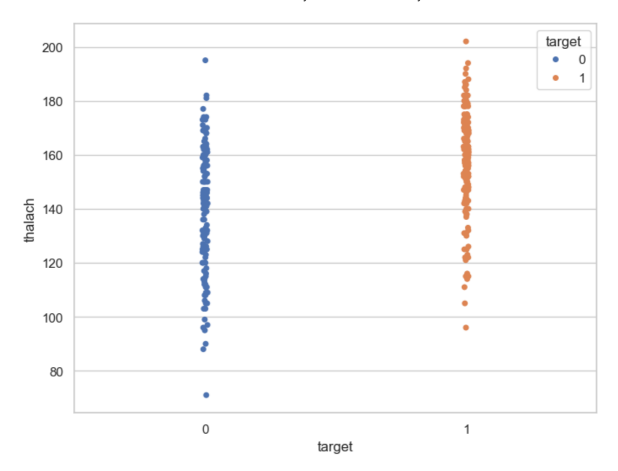


### Interpretation

• We can see that those people suffering from heart disease (target = 1) have relatively higher heart rate (thalach) as compared to people who are not suffering from heart disease (target = 0).

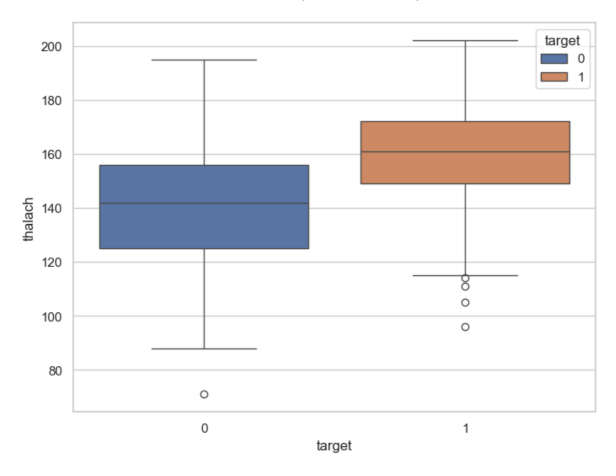
We can add jitter to bring out the distribution of values as follows:

```
In [228]: f, ax = plt.subplots(figsize=(8, 6))
sns.stripplot(x="target", y="thalach",hue="target", data=df, jitter = 0.01)
plt.show()
```



# Visualize distribution of thalach variable wrt target with boxplot

```
In [231]: f, ax = plt.subplots(figsize=(8, 6))
    sns.boxplot(x="target", y = "thalach", hue="target", data=df)
    plt.show()
```



The above boxplot confirms our finding that people suffering from heart disease (target = 1) have relatively higher heart rate (thalach) as compared to people who are not suffering from heart disease (target = 0).

# **Findings of Bivariate Analysis**

Findings of Bivariate Analysis are as follows -

- There is no variable which has strong positive correlation with target variable.
- There is no variable which has strong negative correlation with target variable.
- There is no correlation between target and fbs .
- The cp and thalach variables are mildly positively correlated with target variable.
- We can see that the thalach variable is slightly negatively skewed.
- The people suffering from heart disease (target = 1) have relatively higher heart rate (thalach) as compared to people who are not suffering from heart disease (target = 0).
- The people suffering from heart disease (target = 1) have relatively higher heart rate (thalach) as compared to people who are not suffering from heart disease (target = 0).

# 9. Multivariate analysis

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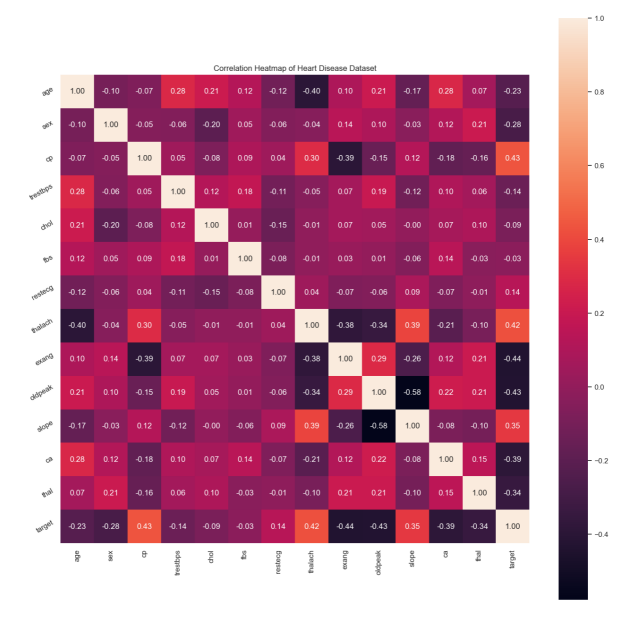
• The objective of the multivariate analysis is to discover patterns and relationships in the dataset.

# Discover patterns and relationships

- An important step in EDA is to discover patterns and relationships between variables in the dataset.
- I will use heat map and pair plot to discover the patterns and relationships in the dataset.
- First of all, I will draw a heat map.

# **Heat Map**

```
In [358]: plt.figure(figsize=(16,16))
   plt.title('Correlation Heatmap of Heart Disease Dataset')
   a = sns.heatmap(correlation, square=True, annot=True, fmt='.2f', linecolor='white'
   a.set_xticklabels(a.get_xticklabels(), rotation=90)
   a.set_yticklabels(a.get_yticklabels(), rotation=30)
   plt.show()
```

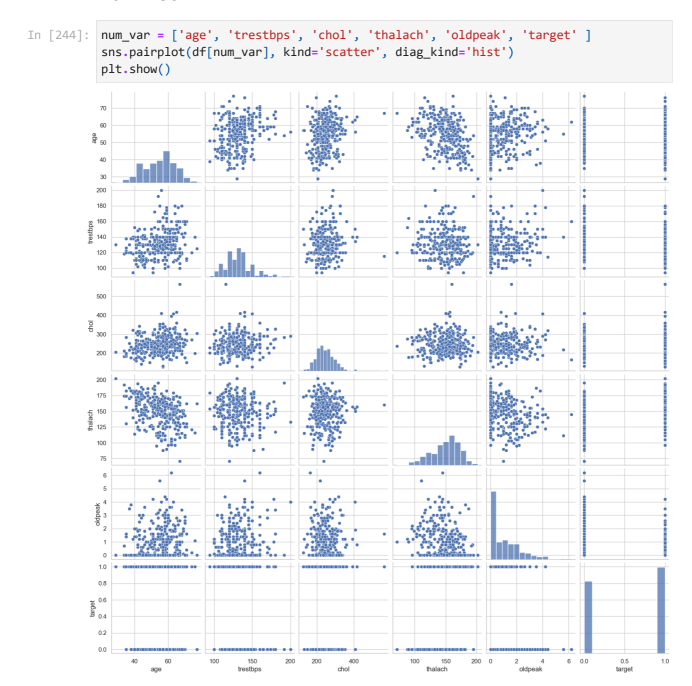


From the above correlation heat map, we can conclude that :-

- target and cp variable are mildly positively correlated (correlation coefficient = 0.43).
- target and thalach variable are also mildly positively correlated (correlation coefficient = 0.42).
- target and slope variable are weakly positively correlated (correlation coefficient = 0.35).
- target and exang variable are mildly negatively correlated (correlation coefficient = -0.44).
- target and oldpeak variable are also mildly negatively correlated (correlation coefficient = -0.43).
- target and ca variable are weakly negatively correlated (correlation coefficient = -0.39).

• target and thal variable are also waekly negatively correlated (correlation coefficient = -0.34).

# **Pair Plot**



#### Comment

- I have defined a variable num\_var . Here age , trestbps , chol`, `thalach` and `oldpeak are numerical variables and target is the categorical variable.
- So, I wll check relationships between these variables.

# Analysis of age and other variables

# Check the number of unique values in age variable

```
In [249]: df['age'].nunique()
Out[249]: 41
```

### View statistical summary of age variable

```
In [254]:
          df['age'].describe()
Out[254]: count
                   303.000000
                   54.366337
          mean
          std
                    9.082101
          min
                    29.000000
          25%
                   47.500000
          50%
                   55.000000
          75%
                    61.000000
                    77.000000
          Name: age, dtype: float64
```

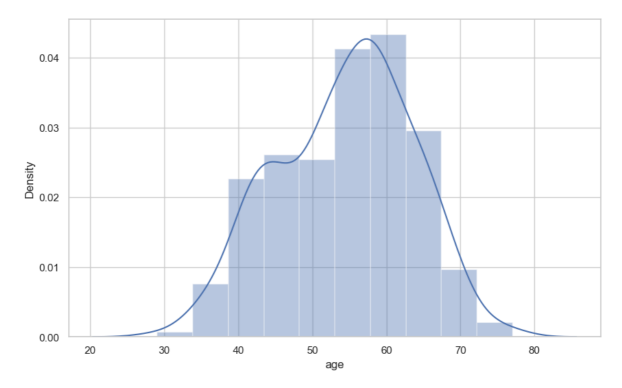
### Interpretation

- The mean value of the age variable is 54.37 years.
- The minimum and maximum values of age are 29 and 77 years.

### Plot the distribution of age variable

Now, I will plot the distribution of age variable to view the statistical properties.

```
In [258]: f, ax = plt.subplots(figsize=(10,6))
x = df['age']
ax = sns.distplot(x, bins=10)
plt.show()
```

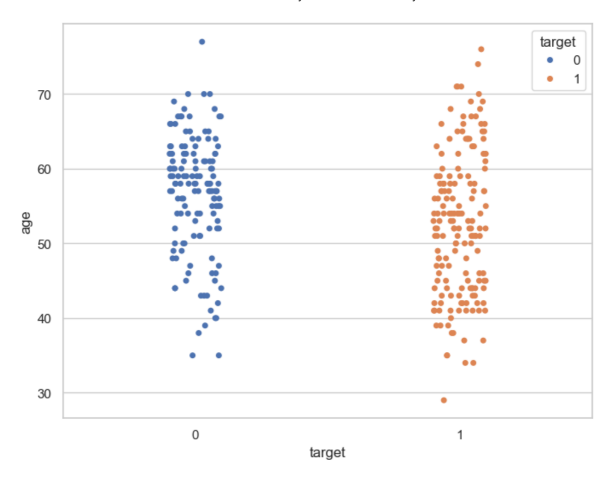


• The age variable distribution is approximately normal.

# Analyze age and target variable

Visualize frequency distribution of age variable wrt target

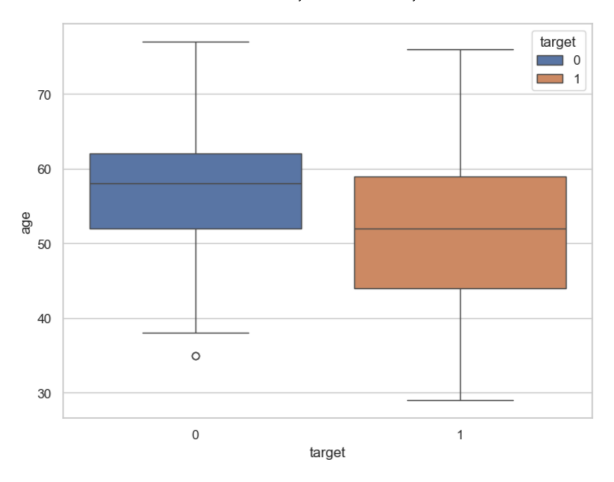
```
In [265]: f, ax = plt.subplots(figsize=(8, 6))
    sns.stripplot(x="target", y="age",hue="target", data=df)
    plt.show()
```



• We can see that the people suffering from heart disease (target = 1) and people who are not suffering from heart disease (target = 0) have comparable ages.

# Visualize distribution of age variable wrt target with boxplot

```
In [269]: f, ax = plt.subplots(figsize=(8, 6))
sns.boxplot(x="target", y="age", hue="target", data=df)
plt.show()
```

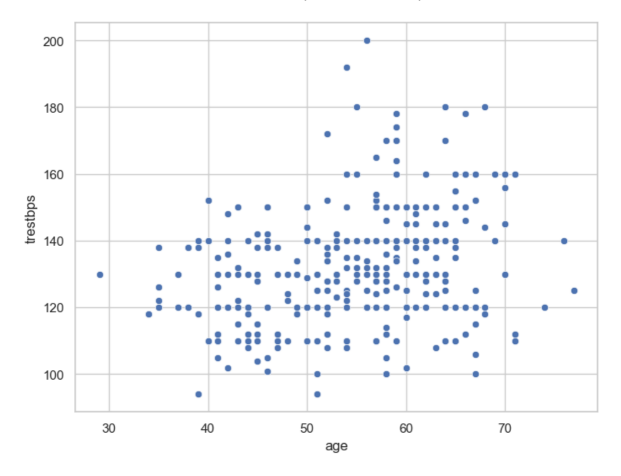


- The above boxplot tells two different things:
  - The mean age of the people who have heart disease is less than the mean age of the people who do not have heart disease.
  - The dispersion or spread of age of the people who have heart disease is greater than the dispersion or spread of age of the people who do not have heart disease.

# Analyze age and trestbps variable

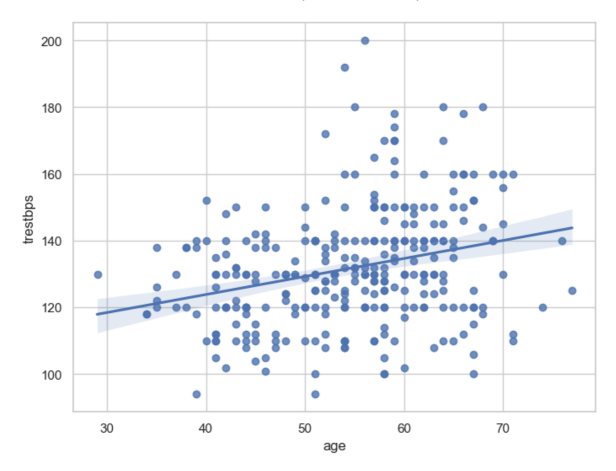
I will plot a scatterplot to visualize the relationship between age and trestbps variable.

```
In [278]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.scatterplot(x="age", y = "trestbps", data=df)
    plt.show()
```



• The above scatter plot shows that there is no correlation between age and trestbps variable.

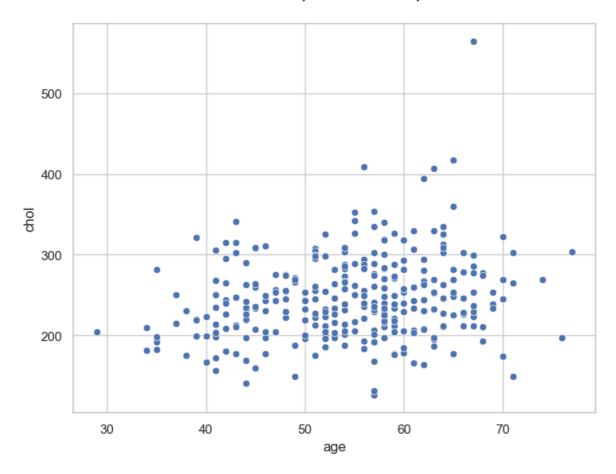
```
In [281]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.regplot(x="age", y="trestbps", data=df)
    plt.show()
```



• The above line shows that linear regression model is not good fit to the data.

# Analyze age and chol variable

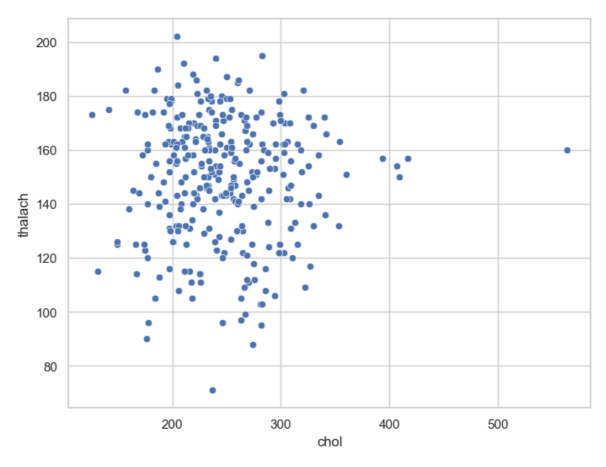
```
In [287]: f, ax = plt.subplots(figsize=(8, 6))
    ax = sns.scatterplot(x="age", y = "chol", data=df)
    plt.show()
```

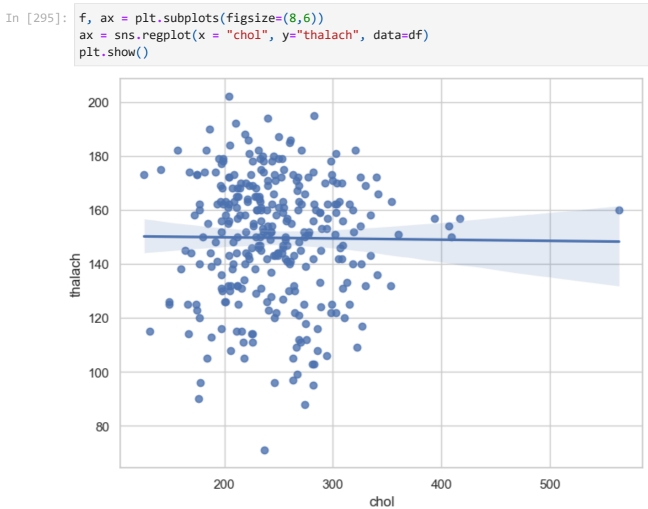


• The above plot confirms that there is a slighly positive correlation between age and chol variables.

# Analyze chol and thalach variable

```
In [293]: f, ax = plt.subplots(figsize=(8,6))
    ax = sns.scatterplot(x="chol",y="thalach",data=df)
    plt.show()
```





• The above plot shows that there is no correlation between chol and thalach variable.

# 10. Dealing with missing values

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- In Pandas missing data is represented by two values:
  - **None**: None is a Python singleton object that is often used for missing data in Python code.
  - NaN: NaN (an acronym for Not a Number), is a special floating-point value recognized by all systems that use the standard IEEE floating-point representation.
- There are different methods in place on how to detect missing values.

# Pandas isnull() and notnull() functions

- Pandas offers two functions to test for missing data isnull() and notnull(). These are simple functions that return a boolean value indicating whether the passed in argument value is in fact missing data.
- Below, I will list some useful commands to deal with missing values.

# Useful commands to detect missing values

#### df.isnull()

The above command checks whether each cell in a dataframe contains missing values or not. If the cell contains missing value, it returns True otherwise it returns False.

#### • df.isnull().sum()

The above command returns total number of missing values in each column in the dataframe.

#### df.isnull().sum().sum()

It returns total number of missing values in the dataframe.

#### • df.isnull().mean()

It returns percentage of missing values in each column in the dataframe.

#### • df.isnull().any()

It checks which column has null values and which has not. The columns which has null values returns TRUE and FALSE otherwise.

#### df.isnull().any().any()

It returns a boolean value indicating whether the dataframe has missing values or not. If dataframe contains missing values it returns TRUE and FALSE otherwise.

#### • df.isnull().values.any()

It checks whether a particular column has missing values or not. If the column contains missing values, then it returns TRUE otherwise FALSE.

#### • df.isnull().values.sum()

It returns the total number of missing values in the dataframe.

```
In [299]:
          # check for missing values
          df.isnull().sum()
Out[299]: age
          sex
          ср
                      0
          trestbps 0
                      0
          chol
          fbs
          restecg
                      0
          thalach
                      0
          exang
          oldpeak
                      0
          slope
          ca
          thal
          target
          dtype: int64
```

### Interpretation

We can see that there are no missing values in the dataset.

# 11. Check with ASSERT statement

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- We must confirm that our dataset has no missing values.
- We can write an **assert statement** to verify this.
- We can use an assert statement to programmatically check that no missing, unexpected 0 or negative values are present.
- This gives us confidence that our code is running properly.
- Assert statement will return nothing if the value being tested is true and will throw an AssertionError if the value is false.
- Asserts

- assert 1 == 1 (return Nothing if the value is True)
- assert 1 == 2 (return AssertionError if the value is False)

```
In [303]: # assert that there are no missing values in the dataframe

assert pd.notnull(df).all().all()

In [305]: # assert all values are greater than or equal to 0

assert pd.notnull(df).all().all()
```

- The above two commands do not throw any error. Hence, it is confirmed that there are no missing or negative values in the dataset.
- All the values are greater than or equal to zero.

# 12. Outlier detection

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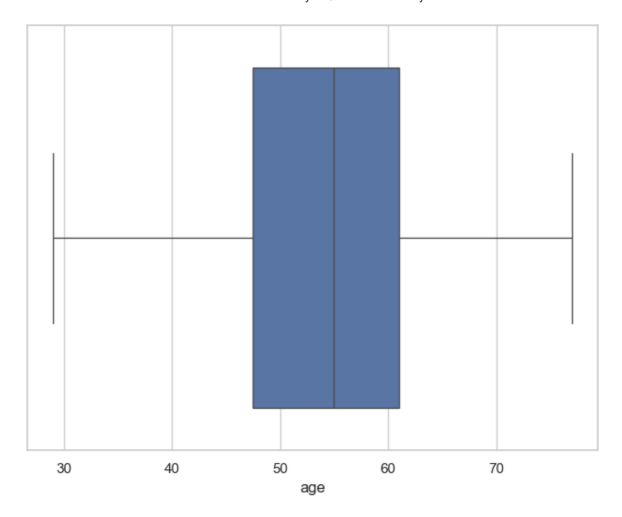
I will make boxplots to visualise outliers in the continuous numerical variables : -

```
age, trestbps, chol, thalach and oldpeak variables.
```

# age variable

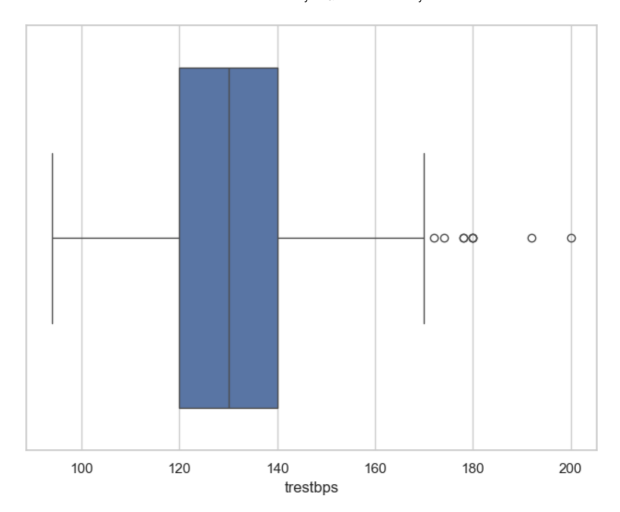
```
df['age'].describe()
In [311]:
Out[311]: count
                   303.000000
                    54.366337
          mean
          std
                     9.082101
          min
                    29.000000
          25%
                    47.500000
          50%
                    55.000000
          75%
                    61.000000
                    77.000000
          max
          Name: age, dtype: float64
          Box-plot of age variable
```

```
In [314]: f, ax = plt.subplots(figsize=(8,6))
    ax = sns.boxplot(x=df['age'])
    plt.show()
```



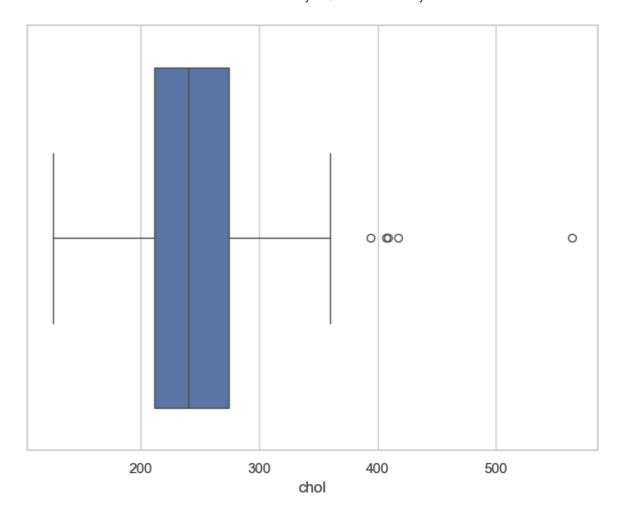
# trestbps variable

```
In [317]: df['trestbps'].describe()
Out[317]: count
                    303.000000
                   131.623762
           mean
           std
                    17.538143
                    94.000000
           min
           25%
                    120.000000
           50%
                    130.000000
           75%
                    140.000000
                    200.000000
           Name: trestbps, dtype: float64
          Box-plot of trestbps variable
In [322]: f,ax = plt.subplots(figsize=(8,6))
          ax = sns.boxplot(x=df["trestbps"])
          plt.show()
```



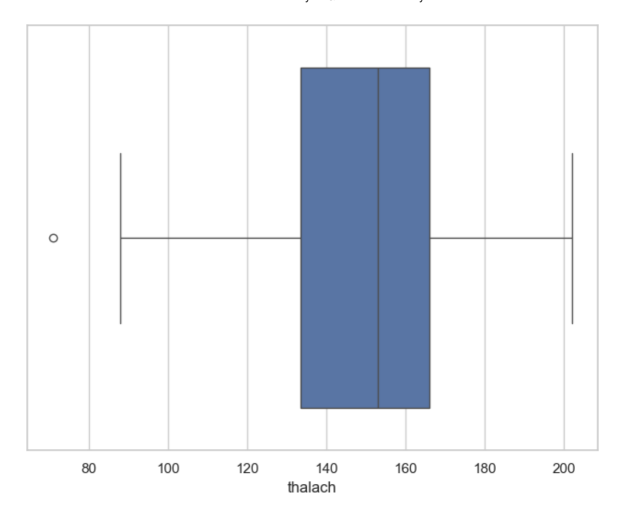
# chol variable

```
In [325]: df['chol'].describe()
Out[325]: count
                    303.000000
                    246.264026
           mean
           std
                    51.830751
                    126.000000
           min
           25%
                    211.000000
           50%
                    240.000000
           75%
                    274.500000
                    564.000000
           Name: chol, dtype: float64
          Box-plot of chol variable
In [328]: f,ax = plt.subplots(figsize=(8,6))
          ax = sns.boxplot(x=df['chol'])
          plt.show()
```



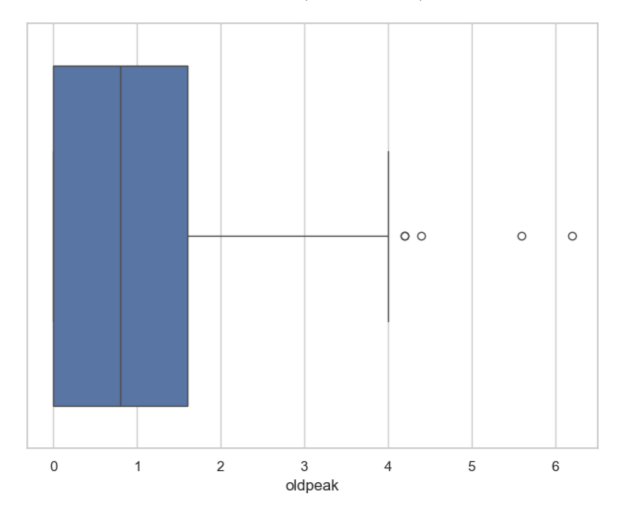
# thalach variable

```
In [331]: df['thalach'].describe()
Out[331]: count
                    303.000000
                    149.646865
           mean
           std
                     22.905161
                     71.000000
           min
           25%
                    133.500000
           50%
                    153.000000
           75%
                    166.000000
                    202.000000
           {\sf max}
           Name: thalach, dtype: float64
          Box-plot of thalach variable
In [336]: f,ax = plt.subplots(figsize=(8,6))
          ax = sns.boxplot(x=df['thalach'])
          plt.show()
```



# oldpeak variable

```
In [339]: df['oldpeak'].describe()
Out[339]: count
                    303.000000
                     1.039604
           mean
           std
                     1.161075
                     0.000000
           min
           25%
                     0.000000
           50%
                     0.800000
           75%
                     1.600000
                     6.200000
           Name: oldpeak, dtype: float64
          Box-plot of oldpeak variable
In [342]: f, ax = plt.subplots(figsize=(8, 6))
          sns.boxplot(x=df["oldpeak"])
          plt.show()
```



### **Findings**

- The age variable does not contain any outlier.
- trestbps variable contains outliers to the right side.
- chol variable also contains outliers to the right side.
- thalach variable contains a single outlier to the left side.
- oldpeak variable contains outliers to the right side.
- Those variables containing outliers needs further investigation.

# 13. Conclusion

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So, our EDA journey has come to an end.

In this kernel, we have explored the heart disease dataset. In this kernel, we have implemented many of the strategies presented in the book **Think Stats - Exploratory Data Analysis in Python by Allen B Downey**. The feature variable of interest is target variable. We have analyzed it alone and check its interaction with other variables. We have also discussed how to detect missing data and outliers.

I hope you like this kernel on EDA journey.

Thanks

# 14. References

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The following references are used to create this kernel

- Think Stats Exploratory Data Analysis in Python by Allen B Downey
- Seaborn API reference
- Other kernel

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