# cardiovascular-risk-prediction

April 19, 2024

#	
Cardiovascular Risk Prediction	

## About Project:

The dataset is from an ongoing cardiovascular study on residents of the town of Framingham, Massachusetts. The classification goal is to predict whether the patient has a 10-year risk of future coronary heart disease (CHD). The dataset provides the patients' information. It includes over 4,000 records and 15 attributes.

#### Variables:

• Each attribute is a potential risk factor. There are both demographic, behavioral, and medical risk factors.

#### Objective:

Predict the overall risk of heart disease using Classification regression

# 1 Data Description

## Demographic

- 1. **Sex:** male or female("M" or "F")
- 2. **Age:** Age of the patient; (Continuous Although the recorded ages have been truncated to whole numbers, the concept of age is continuous)

#### **Behavioral**

- 3. is\_smoking: whether or not the patient is a current smoker ("YES" or "NO")
- 4. Cigs Per Day: the number of cigarettes that the person smoked on average in one day.(can be considered continuous as one can have any number of cigarettes, even half a cigarette.)

#### Medical(history)

- 5. **BP Meds:** whether or not the patient was on blood pressure medication (Nominal)
- 6. **Prevalent Stroke:** whether or not the patient had previously had a stroke (Nominal)
- 7. **Prevalent Hyp:** whether or not the patient was hypertensive (Nominal)
- 8. **Diabetes:** whether or not the patient had diabetes (Nominal)

## Medical(current)

- 9. **Tot Chol:** total cholesterol level (Continuous)
- 10. Sys BP: systolic blood pressure (Continuous)
- 11. **Dia BP:** diastolic blood pressure (Continuous)
- 12. **BMI:** Body Mass Index (Continuous)
- 13. **Heart Rate:** heart rate (Continuous In medical research, variables such as heart rate though in fact discrete, yet are considered continuous because of large number of possible values.)
- 14. **Glucose:** glucose level (Continuous)

#### Predict variable (desired target)

15. **TenYearCHD:** 10-year risk of coronary heart disease CHD(binary: 1 means "Yes", 0 means "No") - DV

## 2 What is Cardiovascular Disease?

Cardiovascular disease can refer to a number of conditions:

#### Heart disease

Heart and blood vessel disease (also called heart disease) includes numerous problems, many of which are related to a process called atherosclerosis.

Atherosclerosis is a condition that develops when a substance called plaque builds up in the walls of the arteries. This buildup narrows the arteries, making it harder for blood to flow through. If a blood clot forms, it can block the blood flow. This can cause a heart attack or stroke.

#### Heart attack

A heart attack occurs when the blood flow to a part of the heart is blocked by a blood clot. If this clot cuts off the blood flow completely, the part of the heart muscle supplied by that artery begins to die.

Most people survive their first heart attack and return to their normal lives, enjoying many more years of productive activity. But experiencing a heart attack does mean that you need to make some changes.

The medications and lifestyle changes that your doctor recommends may vary according to how badly your heart was damaged, and to what degree of heart disease caused the heart attack.

Learn more about heart attack.

#### Stroke

An ischemic stroke (the most common type of stroke) occurs when a blood vessel that feeds the brain gets blocked, usually from a blood clot.

When the blood supply to a part of the brain is cut off, some brain cells will begin to die. This can result in the loss of functions controlled by that part of the brain, such as walking or talking.

A hemorrhagic stroke occurs when a blood vessel within the brain bursts. This is most often caused by uncontrolled hypertension (high blood pressure).

Some effects of stroke are permanent if too many brain cells die after being starved of oxygen. These cells are never replaced.

The good news is that sometimes brain cells don't die during stroke — instead, the damage is temporary. Over time, as injured cells repair themselves, previously impaired function improves. (In other cases, undamaged brain cells nearby may take over for the areas of the brain that were injured.)

Either way, strength may return, speech may get better and memory may improve. This recovery process is what stroke rehabilitation is all about.

When it comes to spotting stroke and getting help, the faster, the better. That's because prompt treatment may make the difference between life and death — or the difference between a full recovery and long-term disability. Use the letters in F.A.S.T to spot a stroke. F is for face drooping. A is for arm weakness. S is for speech difficulty. T is for time to call 911.

Learn more about stroke.

#### Heart failure

Heart failure, sometimes called congestive heart failure, means the heart isn't pumping blood as well as it should. Heart failure does not mean that the heart stops beating — that's a common misperception. Instead, the heart keeps working, but the body's need for blood and oxygen isn't being met.

Heart failure can get worse if left untreated. If your loved one has heart failure, it's very important to follow the doctor's orders.

Learn more about heart failure.

#### Arrhythmia

Arrhythmia refers to an abnormal heart rhythm. There are various types of arrhythmias. The heart can beat too slow, too fast or irregularly.

Bradycardia, or a heart rate that's too slow, is when the heart rate is less than 60 beats per minute. Tachycardia, or a heart rate that's too fast, refers to a heart rate of more than 100 beats per minute.

An arrhythmia can affect how well your heart works. With an irregular heartbeat, your heart may not be able to pump enough blood to meet your body's needs.

Learn more about arrhythmia.

#### Heart valve problems

When heart valves don't open enough to allow the blood to flow through as it should, a condition called stenosis results. When the heart valves don't close properly and thus allow blood to leak through, it's called regurgitation. If the valve leaflets bulge or prolapse back into the upper chamber, it's a condition called prolapse. Discover more about the roles your heart valves play in healthy circulation.

Learn more about heart valve disease.

# 3 Data preparation phase

## 3.1 Importing required libraries

We imported several libraries for the project: 1. numpy: To work with arrays 2. pandas: To work with csv files and dataframes 3. matplotlib: To create charts using pyplot, define parameters using rcParams and color them with cm.rainbow 4. warnings: To ignore all warnings which might be showing up in the notebook due to past/future depreciation of a feature 5. train\_test\_split: To split the dataset into training and testing data 6. StandardScaler: To scale all the features, so that the Machine Learning model better adapts to the dataset

Next, we imported all the necessary Machine Learning algorithms.

```
import warnings
warnings.filterwarnings('ignore')

import pandas as pd
import numpy as np

import matplotlib.pyplot as plt
import seaborn as sns
from pylab import rcParams
rcParams['figure.figsize'] = (10, 6)
%matplotlib inline

from sklearn.model_selection import train_test_split
from sklearn.preprocessing import OneHotEncoder
from sklearn.model_selection import GridSearchCV
from sklearn.model_selection import RandomizedSearchCV
from sklearn.model_selection import train_test_split
from sklearn.model_selection import train_test_split
from sklearn import ensemble
```

#### 3.2 Loading data from csv file to dataframe

```
[2]: # from google.colab import drive
# drive.mount('/content/drive')

[3]: path = './data_cardiovascular_risk.csv'

[4]: # using pandas library and 'read_csv' function to read csv file
dataset = pd.read_csv(path)

# Make a copy of data
df = dataset.copy()
```

# 3.3 Data Exploration

```
[5]: #number of rows and columns
     df.shape
[5]: (3390, 17)
       • This Dataset has 3390 observations in it with 17 columns(features)
[6]: df.columns.tolist()
[6]: ['id',
      'age',
      'education',
      'sex',
      'is_smoking',
      'cigsPerDay',
      'BPMeds',
      'prevalentStroke',
      'prevalentHyp',
      'diabetes',
      'totChol',
      'sysBP',
      'diaBP',
      'BMI',
      'heartRate',
      'glucose',
      'TenYearCHD']
[7]: # Here showing in Dataset the First Five rows by using head() method
     df.head()
[7]:
                  education sex is_smoking
                                             cigsPerDay
                                                          BPMeds
                                                                  prevalentStroke
        id
            age
             64
                        2.0
                              F
                                        YES
                                                     3.0
                                                             0.0
     1
         1
             36
                        4.0
                              М
                                         NO
                                                     0.0
                                                             0.0
                                                                                  0
     2
         2
             46
                        1.0
                              F
                                        YES
                                                    10.0
                                                             0.0
                                                                                  0
     3
                                                    20.0
                                                             0.0
                                                                                  0
         3
             50
                        1.0
                              Μ
                                        YES
         4
             64
                        1.0
                              F
                                        YES
                                                    30.0
                                                             0.0
                                                                                  0
                                                                 heartRate glucose
        prevalentHyp
                       diabetes
                                 totChol
                                          sysBP
                                                   diaBP
                                                            BMI
     0
                                    221.0
                                           148.0
                                                    85.0
                                                                       90.0
                                                                                80.0
                    0
                              0
                                                            NaN
     1
                    1
                              0
                                    212.0 168.0
                                                    98.0 29.77
                                                                       72.0
                                                                                75.0
     2
                    0
                              0
                                    250.0 116.0
                                                    71.0
                                                          20.35
                                                                       88.0
                                                                                94.0
     3
                                    233.0 158.0
                                                                       68.0
                                                                                94.0
                    1
                              0
                                                   88.0
                                                          28.26
                    0
                              0
                                    241.0 136.5
                                                   85.0 26.42
                                                                       70.0
                                                                                77.0
        TenYearCHD
     0
```

```
1 0
2 0
3 1
4 0
```

```
[8]: # Here showing in Dataset the Last Five rows by using tail() method df.tail()
```

```
[8]:
                                                   cigsPerDay
                                                                BPMeds
                       education sex is_smoking
              id
                  age
     3385
           3385
                   60
                              1.0
                                    F
                                               NO
                                                           0.0
                                                                    0.0
                                                           0.0
     3386
                              1.0
                                    F
                                                                    0.0
           3386
                   46
                                               NO
                              3.0
                                                           3.0
     3387
           3387
                   44
                                    Μ
                                              YES
                                                                    0.0
                              1.0
                                                           0.0
     3388
           3388
                   60
                                    Μ
                                               NO
                                                                    NaN
     3389
           3389
                   54
                              3.0
                                    F
                                               NO
                                                           0.0
                                                                    0.0
                             prevalentHyp
                                                       totChol
                                                                 sysBP
           prevalentStroke
                                             diabetes
                                                                         diaBP
                                                                                   BMI
                                                                 123.5
                                                                          79.0
     3385
                                          0
                                                    0
                                                          261.0
                                                                                29.28
                          0
                                                                 102.0
     3386
                          0
                                          0
                                                    0
                                                          199.0
                                                                          56.0
                                                                                 21.96
                                                          352.0 164.0
     3387
                          0
                                          1
                                                    0
                                                                         119.0
                                                                                28.92
     3388
                           0
                                                          191.0
                                                                 167.0
                                                                                 23.01
                                          1
                                                    0
                                                                         105.0
                                                          288.0 124.0
     3389
                           0
                                          0
                                                                          77.0
                                                                                29.88
           heartRate
                       glucose
                                 TenYearCHD
     3385
                 70.0
                         103.0
                 80.0
                                           0
     3386
                          84.0
                 73.0
                          72.0
                                           1
     3387
                                           0
     3388
                 80.0
                          85.0
     3389
                 79.0
                           92.0
                                           0
```

Before any analysis, we just wanted to take a look at the data. So, we used the info() method.

# [9]: #information of the dataset df.info()

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 3390 entries, 0 to 3389
Data columns (total 17 columns):

#	Column	Non-Null Count	Dtype
0	id	3390 non-null	int64
1	age	3390 non-null	int64
2	education	3303 non-null	float64
3	sex	3390 non-null	object
4	is_smoking	3390 non-null	object
5	cigsPerDay	3368 non-null	float64
6	BPMeds	3346 non-null	float64
7	prevalentStroke	3390 non-null	int64

```
3390 non-null
8
    prevalentHyp
                                     int64
9
    diabetes
                     3390 non-null
                                     int64
10
   totChol
                     3352 non-null
                                     float64
11
    sysBP
                     3390 non-null
                                     float64
    diaBP
                     3390 non-null
12
                                     float64
13
   BMI
                     3376 non-null
                                     float64
                     3389 non-null
14 heartRate
                                     float64
   glucose
                     3086 non-null
15
                                     float64
16 TenYearCHD
                     3390 non-null
                                     int64
```

dtypes: float64(9), int64(6), object(2)

memory usage: 450.4+ KB

As you can see from the output above, there are a total of 16 features and 1 target variable. Also, there are some missing values so we need to take care of null values. Next, we used describe() method.

```
[10]: #description of the data in the dataset df.describe(include='all')
```

[10]:		id	age	education	Sex	is_smoking	cigsPerDay	\
[10].	count	3390.000000	3390.000000	3303.000000	3390	3390	3368.000000	`
	unique	NaN	NaN	NaN	2	2	NaN	
	top	NaN	NaN	NaN	F	NO	NaN	
	freq	NaN	NaN	NaN	1923	1703	NaN	
	mean	1694.500000	49.542183	1.970936	NaN	NaN	9.069477	
	std	978.753033	8.592878	1.019081	NaN	NaN	11.879078	
	min	0.000000	32.000000	1.000000	NaN	NaN	0.000000	
	25%	847.250000	42.000000	1.000000	NaN	NaN	0.000000	
	50%	1694.500000	49.000000	2.000000	NaN	NaN	0.000000	
	75%	2541.750000	56.000000	3.000000	NaN	NaN	20.000000	
	max	3389.000000	70.000000	4.000000	NaN	NaN	70.000000	
		BPMeds	prevalentStro	oke prevaler	ntHyp	diabetes	totChol	\
	count	3346.000000	3390.0000	3390.00	0000	3390.000000	3352.000000	
	unique	NaN	N	VaN	NaN	NaN	NaN	
	top	NaN	N	NaN	NaN	NaN	NaN	
	freq	NaN	N	NaN	NaN	NaN	NaN	
	mean	0.029886	0.0064	190 0.31	L5339	0.025664	237.074284	
	std	0.170299	0.0803	309 0.46	64719	0.158153	45.247430	
	min	0.000000	0.0000	0.00	0000	0.000000	107.000000	
	25%	0.000000	0.0000	0.00	00000	0.000000	206.000000	
	50%	0.000000	0.0000	0.00	0000	0.000000	234.000000	
	75%	0.000000	0.0000	1.00	00000	0.000000	264.000000	
	max	1.000000	1.0000	1.00	00000	1.000000	696.000000	
		sysBP	diaBP	BMI	hear		glucose \	
	count	3390.00000	3390.000000 3	3376.000000	3389.0	000000 3086	.000000	
	unique	NaN	NaN	NaN		NaN	NaN	

top	NaN	NaN	NaN	NaN	NaN
_					
freq	NaN	NaN	NaN	NaN	NaN
mean	132.60118	82.883038	25.794964	75.977279	82.086520
std	22.29203	12.023581	4.115449	11.971868	24.244753
min	83.50000	48.000000	15.960000	45.000000	40.000000
25%	117.00000	74.500000	23.020000	68.000000	71.000000
50%	128.50000	82.000000	25.380000	75.000000	78.000000
75%	144.00000	90.000000	28.040000	83.000000	87.000000
max	295.00000	142.500000	56.800000	143.000000	394.000000
	${\tt TenYearCHD}$				
count	3390.000000				
unique	NaN				
top	NaN				
freq	NaN				
mean	0.150737				
std	0.357846				
min	0.000000				

The method revealed that the range of each variable is different. The maximum value of age is 70 but for chol it is 696. Thus, feature scaling must be performed on the dataset.

# 4 Data Preprocessing and Basic EDA

0.00000

0.00000

0.000000

25%

50%

75%

max

We will drop the education and id columns because it has no correlation with heart disease.

```
[11]: | # Removing columns such as 'id' & 'education'
      df.drop(['id', 'education'],axis=1,inplace=True)
[12]: df.head()
[12]:
          age sex is_smoking
                                cigsPerDay
                                             BPMeds
                                                      prevalentStroke
                                                                        prevalentHyp
           64
                F
                          YES
                                       3.0
                                                0.0
      0
      1
                                       0.0
                                                0.0
           36
                Μ
                                                                     0
                                                                                     1
                           NO
      2
           46
                F
                          YES
                                      10.0
                                                0.0
                                                                     0
                                                                                     0
      3
           50
                М
                          YES
                                      20.0
                                                0.0
                                                                     0
                                                                                     1
           64
                F
                          YES
                                      30.0
                                                0.0
                                                                     0
                                                                                     0
                                                                  glucose
         diabetes
                    totChol
                               sysBP
                                      diaBP
                                                BMI
                                                      heartRate
                                                                            TenYearCHD
      0
                 0
                       221.0
                              148.0
                                       85.0
                                                NaN
                                                            90.0
                                                                     80.0
                                                                                      1
                 0
                       212.0
                              168.0
                                                            72.0
                                                                     75.0
                                                                                      0
      1
                                       98.0
                                              29.77
      2
                 0
                       250.0
                              116.0
                                       71.0
                                              20.35
                                                            88.0
                                                                     94.0
                                                                                      0
```

3	0	233.0	158.0	88.0	28.26	68.0	94.0	1
4	0	241.0	136.5	85.0	26.42	70.0	77.0	0

Before we go ahead, an important step to do is to convert our string feature into an integer.

We will name

- In sex feature M will be converted to 1 and F will be converted to 0.
- In is smoking feature YES will be converted to 1 and NO will be converted to 0.

```
[13]: # Applying function to convert string data to an integer
      df['sex'] = df['sex'].apply(lambda x : 1 if x == "M" else 0)
      df['is_smoking'] = df['is_smoking'].apply(lambda x : 1 if x == "YES" else 0)
[14]: # checking the distribution
      df['sex'].value_counts()
[14]: sex
      0
           1923
      1
           1467
      Name: count, dtype: int64
[15]: # checking the distribution
      df['is_smoking'].value_counts()
[15]: is_smoking
      0
           1703
      1
           1687
      Name: count, dtype: int64
```

[16]: df.info()

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 3390 entries, 0 to 3389
Data columns (total 15 columns):

#	Column	Non-Null Count	Dtype
0	age	3390 non-null	int64
1	sex	3390 non-null	int64
2	is_smoking	3390 non-null	int64
3	cigsPerDay	3368 non-null	float64
4	BPMeds	3346 non-null	float64
5	prevalentStroke	3390 non-null	int64
6	${\tt prevalentHyp}$	3390 non-null	int64
7	diabetes	3390 non-null	int64
8	totChol	3352 non-null	float64
9	sysBP	3390 non-null	float64
10	diaBP	3390 non-null	float64
11	BMI	3376 non-null	float64

```
      12 heartRate
      3389 non-null float64

      13 glucose
      3086 non-null float64

      14 TenYearCHD
      3390 non-null int64
```

dtypes: float64(8), int64(7)

memory usage: 397.4 KB

# 4.1 Missing Value Analysis

Handling missing data is important as many machine learning algorithms do not support data with missing values.

```
[17]: # checking for missing values
df.isnull().sum()
```

[17]:	age	0
	sex	0
	is_smoking	0
	cigsPerDay	22
	BPMeds	44
	prevalentStroke	0
	prevalentHyp	0
	diabetes	0
	totChol	38
	sysBP	0
	diaBP	0
	BMI	14
	heartRate	1
	glucose	304
	TenYearCHD	0
	dtype: int64	

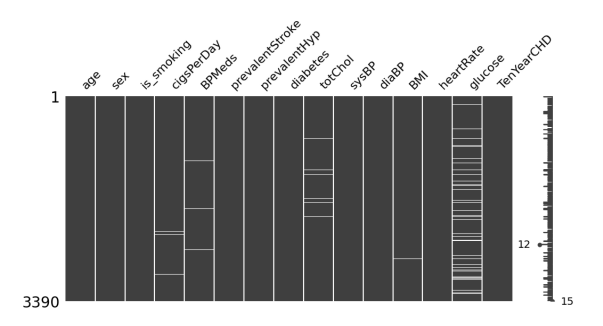
Visualize missing values (NaN) values using  $\bf Missingno\ Library > The\ next\ single-line\ code\ will$  visualize the location of missing values.

```
[18]: # Program to visualize missing values in dataset

# Importing the Missingno libraries
import missingno as msno

# Visualize missing values as a matrix
msno.matrix(df,figsize=(12, 5))
```

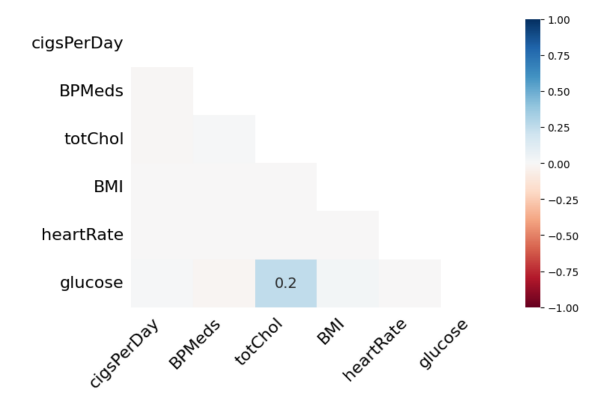
[18]: <Axes: >



These trends give an idea about how the features are correlated with one another. But to get a better idea about correlations we need to use heatmaps.

```
[19]: # Visualize the correlation between the number of # missing values in different columns as a heatmap msno.heatmap(df,figsize=(8, 5))
```

[19]: <Axes: >



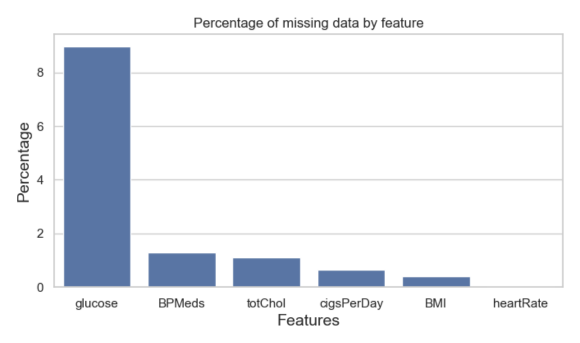
• There is no strong Correlation between any features.

Let's check percentage of missing data of each features

```
Total Percentage
glucose
              304
                     8.967552
BPMeds
               44
                     1.297935
totChol
               38
                     1.120944
cigsPerDay
                     0.648968
               22
BMI
               14
                     0.412979
heartRate
                1
                     0.029499
```

```
[21]: # Visualizing the percentage of missing data plt.figure(figsize=(8,4))
```

```
sns.set(style="whitegrid")
sns.barplot(x=missingdf.index, y=missingdf['Percentage'], data = missingdf)
plt.title('Percentage of missing data by feature')
plt.xlabel('Features', fontsize=14)
plt.ylabel('Percentage', fontsize=14)
plt.show()
```



Total number of rows with missing values is 386 since it is only 11 percent of the entire dataset the rows with missing values are excluded.

At 8.97%, the blood glucose entry has the highest percentage of missing data. The other features have very few missing entries.

Since the missing entries account for only 11% of the total data so, we can drop these entries without losing alot of data.

```
[24]: # Now checking for missing values
      df.isnull().any()
[24]: age
                          False
      sex
                          False
      is_smoking
                          False
      cigsPerDay
                          False
      BPMeds
                          False
      prevalentStroke
                          False
      prevalentHyp
                          False
      diabetes
                          False
      totChol
                          False
      sysBP
                          False
      diaBP
                          False
      BMI
                          False
                          False
      heartRate
                          False
      glucose
      TenYearCHD
                          False
      dtype: bool
[25]: # Checking for any duplicate values
      len(df[df.duplicated()])
[25]: 0
[26]: # statistical measures about the data
      df.describe()
[26]:
                                   sex
                                          is_smoking
                                                        cigsPerDay
                                                                         BPMeds \
                      age
                                        3004.000000
      count
             3004.000000
                           3004.000000
                                                      3004.000000
                                                                    3004.000000
      mean
               49.521305
                              0.448402
                                            0.492676
                                                         9.078562
                                                                       0.030293
      std
                8.595076
                              0.497413
                                            0.500030
                                                        11.890855
                                                                       0.171421
                              0.000000
                                            0.000000
                                                         0.000000
                                                                       0.000000
      min
               32.000000
      25%
               42.000000
                              0.000000
                                            0.000000
                                                         0.000000
                                                                       0.000000
      50%
               49.000000
                              0.000000
                                            0.000000
                                                         0.000000
                                                                       0.000000
      75%
               56.000000
                              1.000000
                                            1.000000
                                                        20.000000
                                                                       0.000000
               70.000000
                              1.000000
                                            1.000000
                                                        70.000000
                                                                       1.000000
      max
                                                                totChol
             prevalentStroke
                               prevalentHyp
                                                 diabetes
                                                                                sysBP
      count
                 3004.000000
                                3004.000000
                                              3004.000000
                                                           3004.000000
                                                                         3004.000000
      mean
                     0.005992
                                   0.314913
                                                 0.027297
                                                             237.222370
                                                                          132.591711
      std
                     0.077189
                                   0.464559
                                                 0.162974
                                                                           22.270882
                                                              45.267516
      min
                     0.000000
                                   0.000000
                                                 0.000000
                                                             113.000000
                                                                           83.500000
      25%
                     0.000000
                                   0.000000
                                                 0.000000
                                                             206.000000
                                                                          117.000000
```

[23]: # Excluding the missing values

df.dropna(axis=0,inplace=True)

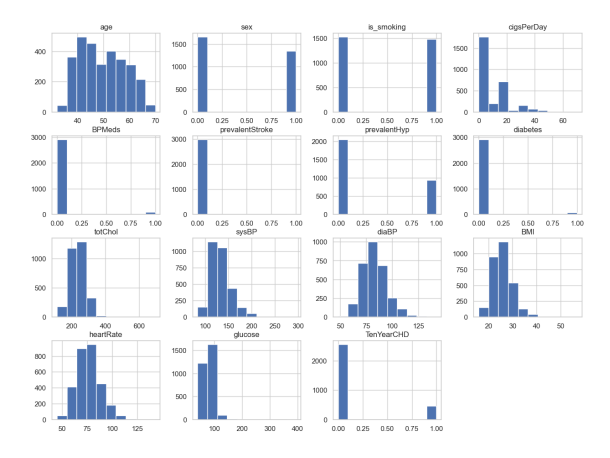
50% 75% max	0.000 0.000 1.000	000 1.00	0000 0.00	234.00 0000 265.00 0000 696.00	0000 144.000000
	diaBP	BMI	heartRate	glucose	TenYearCHD
count	3004.000000	3004.000000	3004.000000	3004.000000	3004.000000
mean	82.923103	25.824041	75.816911	81.993009	0.152130
std	12.038348	4.121130	11.937110	24.125814	0.359207
min	48.000000	15.960000	45.000000	40.000000	0.000000
25%	74.500000	23.057500	68.000000	71.000000	0.000000
50%	82.000000	25.410000	75.000000	78.000000	0.000000
75%	90.000000	28.072500	83.000000	87.000000	0.000000
max	142.500000	56.800000	143.000000	394.000000	1.000000

The features described in the above data set are:

- Count: tells us the number of NoN-empty rows in a feature.
- Mean: tells us the mean value of that feature.
- Std: tells us the Standard Deviation Value of that feature.
- Min: tells us the minimum value of that feature.
- 25%, 50%, and 75%: are the percentile/quartile of each features.
- Max: tells us the maximum value of that feature.

Now, Let's visualise Data Distribution

```
[27]: # plot histogram to see the distribution of the data
fig = plt.figure(figsize = (15,11))
ax = fig.gca()
df.hist(ax = ax)
plt.show()
```



• From above distribution plot we can say that the data on the **prevalent stroke**, **diabetes**, and **blood pressure meds(BPMeds)** are poorly balanced.

# 4.2 Analysing features

Target Variable Analysis:

[28]: # checking the distribution of Target Variable df.TenYearCHD.value\_counts()

[28]: TenYearCHD 0 2547

1 457

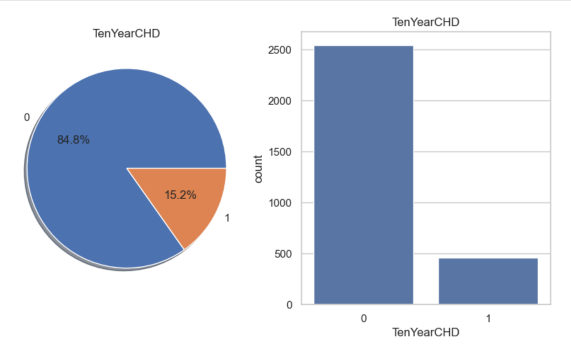
Name: count, dtype: int64

- 0 -> Person do not have risk of coronary heart disease
- 1 -> Person have risk of coronary heart disease

There are 2547 patients without heart disease and 457 patients with the disease.

[29]: #Plotting pie chart of target variable f,ax=plt.subplots(1,2,figsize=(10,5))

```
df['TenYearCHD'].value_counts().plot.pie(autopct='%1.1f%%',ax=ax[0],shadow=True)
ax[0].set_title('TenYearCHD')
ax[0].set_ylabel('')
sns.countplot(x='TenYearCHD',data=df,ax=ax[1])
ax[1].set_title('TenYearCHD')
plt.show()
```



• We can see above that we have the imbalanced data set as the number of people without the disease greatly exceeds the number of people with the disease.

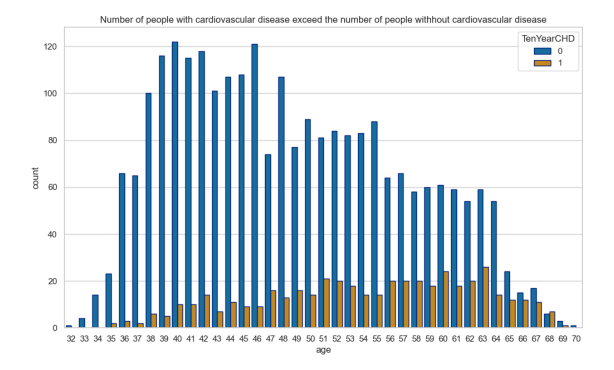
Let's look at the number of people with cardiovascular disease exceed the number of people withhout cardiovascular disease respect to age.

```
[30]: # Visualising the target and age variable
plt.figure(figsize=[12,7])
plt.title("Number of people with cardiovascular disease exceed the number of

→people withhout cardiovascular disease")
sns.countplot(x='age', hue='TenYearCHD',data=df, palette='colorblind',

→edgecolor=sns.color_palette('dark', n_colors=1)[0])
```

[30]: <Axes: title={'center': 'Number of people with cardiovascular disease exceed the number of people withhout cardiovascular disease'}, xlabel='age', ylabel='count'>



- As we can see in above plot The people with **the highest risk** of developing heart disease are between the **ages of 51 and 63**.
- Because the number of sick people generally increases with age.

# 4.3 Categorical variable comparisons with Target Variable (TenYearCHD):

We will use [Stacked Bar chart](https://businessq-software.com/2017/02/21/stacked-bar-chart-definition-and-examples-businessq/#:~:text=A%20stacked%20bar%20graph%20(or,or%20categories%20of%20tha for Comparison between Categorical variable and Target Variable.

**Stacked Bar Chart:** A stacked bar graph (or stacked bar chart) is a chart that uses bars to show comparisons between categories of data, but with ability to break down and compare parts of a whole. Each bar in the chart represents a whole, and segments in the bar represent different parts or categories of that whole.

```
[31]: from operator import add
def stacked_barchart(data, title = None, ylabel = None, xlabel = None):
    # Function to plot stacked bar chart
    default_colors = ['#006400', '#FF0000', '#228B22']
    # From raw value to percentage
    totals = data.sum(axis=1)
    bars = ((data.T / totals) * 100).T
    r = list(range(data.index.size))

# Plot
```

```
barWidth = 0.95
      names = data.index.tolist()
      bottom = [0] * bars.shape[0]
       # Create bars
      color_index = 0
      plots = []
      for bar in bars.columns:
                 plots.append(plt.bar(r, bars[bar], bottom=bottom,__
Good of a color o
                 bottom = list(map(add, bottom, bars[bar]))
                 color_index = 0 if color_index >= len(default_colors) else color_index

_ | |
→+ 1
       # Custom x axis
      plt.title(title)
      plt.xticks(r, names)
      plt.xlabel(data.index.name if xlabel is None else xlabel)
      plt.ylabel(data.columns.name if ylabel is None else ylabel)
      ax = plt.gca()
      y_labels = ax.get_yticks()
      ax.set_yticklabels([str(y) + '%' for y in y_labels])
      flat_list = [item for sublist in data.T.values for item in sublist]
      for i, d in zip(ax.patches, flat_list):
                 data label = str(d) + " (" + str(round(i.get height(), 2)) + "%)"
                 ax.text(i.get_x() + 0.45, i.get_y() + 5, data_label_{,u}
⊸horizontalalignment='center', verticalalignment='center', fontdict =□

dict(color = 'white', size = 20))
      for item in ([ax.title]):
                 item.set_fontsize(27)
      for item in ([ax.xaxis.label, ax.yaxis.label] + ax.get_xticklabels() + ax.

    get_yticklabels()):
                 item.set_fontsize(24)
      legend = ax.legend(plots, bars.columns.tolist(), fancybox=True)
      plt.setp(legend.get_texts(), fontsize='20')
```

Now, Let's Visualzing each category with respect to target variable

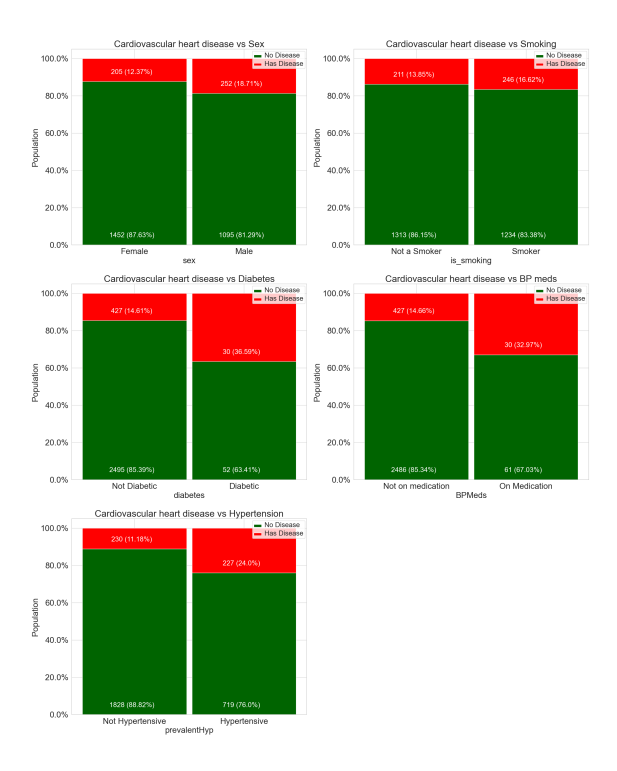
```
[32]: # Visualzing each category with respect to target variable
fig = plt.gcf()
fig.set_size_inches(27, 35)
grid_rows = 3
```

```
grid_cols = 2
#Plot sex vs disease outcome
plt.subplot(grid_rows, grid_cols, 1)
temp = df[['sex', 'TenYearCHD']].groupby(['sex', 'TenYearCHD']).size().

¬unstack('TenYearCHD')
temp.rename(index={0:'Female', 1:'Male'}, columns={0:'No Disease', 1:'Hasu
 ⇔Disease'}, inplace = True)
stacked_barchart(temp, title = 'Cardiovascular heart disease vs Sex', ylabel = L
 #Plot smoking satus vs disease outcome
plt.subplot(grid_rows, grid_cols, 2)
temp = df[['is_smoking','TenYearCHD']].groupby(['is_smoking','TenYearCHD']).
 ⇔size().unstack('TenYearCHD')
temp.rename(index={0:'Not a Smoker', 1:'Smoker'}, columns={0:'No Disease', 1:
stacked_barchart(temp, title = 'Cardiovascular heart disease vs Smoking', __
 #Plot diabetes vs disease outcome
plt.subplot(grid rows, grid cols, 3)
temp = df[['diabetes','TenYearCHD']].groupby(['diabetes','TenYearCHD']).size().

unstack('TenYearCHD')

temp.rename(index={0:'Not Diabetic', 1:'Diabetic'}, columns={0:'No Disease', 1:
 stacked barchart(temp, title = 'Cardiovascular heart disease vs Diabetes', |
 ⇔ylabel = 'Population')
#Plot BP meds vs disease outcome
plt.subplot(grid_rows, grid_cols, 4)
temp = df[['BPMeds','TenYearCHD']].groupby(['BPMeds','TenYearCHD']).size().
 temp.rename(index={0:'Not on medication', 1:'On Medication'}, columns={0:'Nou
 ⇔Disease', 1:'Has Disease'}, inplace = True)
stacked_barchart(temp, title = 'Cardiovascular heart disease vs BP meds', __
 ⇔ylabel = 'Population')
#Plot Hypertension vs disease outcome
plt.subplot(grid_rows, grid_cols, 5)
temp = df[['prevalentHyp','TenYearCHD']].groupby(['prevalentHyp','TenYearCHD']).
 ⇔size().unstack('TenYearCHD')
temp.rename(index={0:'Not Hypertensive', 1:'Hypertensive'}, columns={0:'Nou
→Disease', 1: 'Has Disease'}, inplace = True)
stacked_barchart(temp, title = 'Cardiovascular heart disease vs Hypertension', u
 ⇔ylabel = 'Population')
```



From the above categorical variables comparison plot we can conclude that,

- Slightly more males are suffering from Cardiovascular heart disease than females.
- The people who have Cardiovascular heart disease is almost equal between smokers and non smokers.
- The percentage of people who have Cardiovascular heart disease is higher among the diabetic

- patients and also those patients with prevalent hypertension have more risk of Cardiovascular heart disease compare to those who don't have hypertensive problem.
- The percentage of people who are on medication of blood pressure have more risk of Cardiovascular heart disease compare to those who are not on medication.

Now, Let's see the Correlation between the all features

```
[33]: # let's find all feature's correlation with Heatmap
    # correlation
    plt.figure(figsize=(20,8))
    correlation = df.corr()
    sns.heatmap(abs(correlation), annot = True, cmap='YlGnBu')
    plt.title('Correlation between the all features')
```

[33]: Text(0.5, 1.0, 'Correlation between the all features')



#### From the above correlation plot we can conclude that,

- There are no features with more than 0.2 correlation with the Ten year risk of developing CHD and this shows that the features a poor predictors. However the features with the highest correlations are age, prevalent hypertension(prevalentHyp) and systolic blood pressure(sysBP).
- Also there are a couple of features that are highly correlated with each other and it makes no sense to use both of them in building a machine learning model.

#### These includes:

- Blood glucose and diabetes;
- systolic and diastolic blood pressures;
- cigarette smoking and the number of cigarretes smoked per day.

Therefore we need to carry out feature selection to pick the best features.

# 5 Feature Engineering/Selection

#### Tree-based: SelectFromModel

**SelectFromModel** is an Embedded method. Embedded methods use algorithms that have built-in feature selection methods.

Here,

We have used RandomForest() to select features based on feature importance. We calculate feature importance using node impurities in each decision tree.

In Random forest, the final feature importance is the average of all decision tree feature importance.

```
[34]: #define the features

X = df.iloc[:,:-1]

y = df.iloc[:,-1]
```

7 selected features

```
[36]: # Important or top Features
embeded_rf_feature
```

```
[36]: ['age', 'totChol', 'sysBP', 'diaBP', 'BMI', 'heartRate', 'glucose']
```

The top features are:

- 1. Age
- 2. Total cholesterol
- 3. Systolic blood pressure
- 4. Diastolic blood pressure
- 5. BMI
- 6. Heart rate
- 7. Blood glucose

Statistics on the top feature:

```
[37]: # Importing statsmodels
import statsmodels.api as sm
```

**Statsmodels** is a Python module that provides classes and functions for the estimation of many different statistical models, as well as for conducting statistical tests, and statistical data exploration.

```
[38]: # Splitting the dependent and independent variables
top_features = df[embeded_rf_feature]
y = df['TenYearCHD']
```

```
[39]: # Fit the data

result = sm.Logit(y,top_features).fit()

# The summary table below, gives us a descriptive summary about the regression

→results

print(result.summary())
```

Optimization terminated successfully.

Current function value: 0.415336

Iterations 6

Logit Regression Results

========		=======	=======	========	========	
Dep. Variab	ole:	TenYear	CHD No.	Observations	:	3004
Model:		Log	git Df R	esiduals:		2997
Method:		1	MLE Df M	odel:		6
Date:	Fr	i, 19 Apr 20	024 Pseu	do R-squ.:		0.02592
Time:		15:46	:54 Log-	Likelihood:		-1247.7
converged:		T	rue LL-N	ull:		-1280.9
Covariance	Type:	nonrob	ust LLR j	p-value:		2.236e-12
========	========	========	=======	========	========	
	coef	std err	z	P> z	[0.025	0.975]
age	0.0226	0.006	3.584	0.000	0.010	0.035
totChol	-0.0018	0.001	-1.554	0.120	-0.004	0.000
sysBP	0.0245	0.004	6.722	0.000	0.017	0.032
diaBP	-0.0297	0.006	-4.601	0.000	-0.042	-0.017
BMI	-0.0544	0.013	-4.082	0.000	-0.081	-0.028
heartRate	-0.0301	0.004	-7.336	0.000	-0.038	-0.022
glucose	0.0055	0.002	3.060	0.002	0.002	0.009

In the output, **Iterations** refer to the number of times the model iterates over the data, trying to optimise the model.

Explanation of some of the terms in the summary table:

- **coef** : the coefficients of the independent variables in the regression equation.
- Log-Likelihood: the natural logarithm of the Maximum Likelihood Estimation (MLE) function. MLE is the optimisation process of finding the set of parameters which result in best

fit.

- **LL-Null**: the value of log-likelihood of the model when no independent variable is included(only an intercept is included).
- **Pseudo R-squ.**: a substitute for the R-squared value in Least Squares linear regression. It is the ratio of the log-likelihood of the null model to that of the full model.

```
[40]: # Checking the odds ratio of top features
params = result.params
conf = result.conf_int()
conf['Odds Ratio'] = params
conf.columns = ['5%', '95%', 'Odds Ratio']
print(np.exp(conf))
```

```
95% Odds Ratio
                 5%
           1.010284
                    1.035552
                                 1.022840
age
           0.995854 1.000481
totChol
                                 0.998165
           1.017473 1.032083
sysBP
                                 1.024752
diaBP
           0.958523 0.983091
                                 0.970729
BMI
           0.922652 0.972119
                                 0.947062
heartRate
          0.962550 0.978167
                                 0.970327
           1.001974 1.009041
                                 1.005501
glucose
```

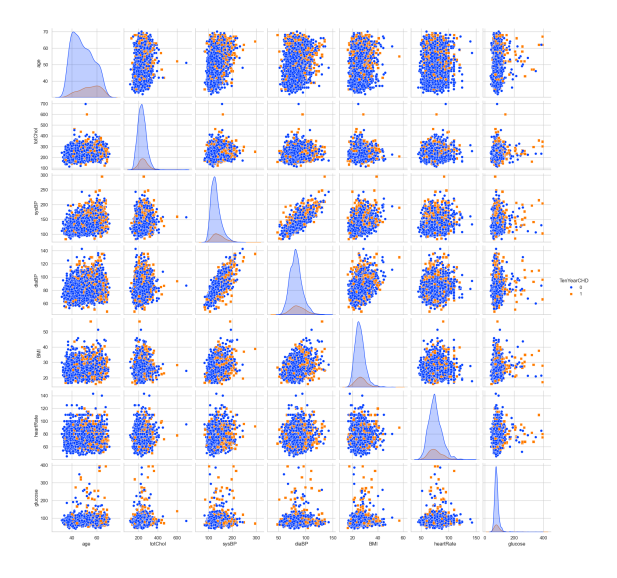
From above table we can conclude that the odds of getting cardiovascular heart disease risk increases with about 2% for every increase in age and systolic blood pressure(sysBP).

The other factors show no significant positive odds.

#### Pair plots

```
[41]: # Plotting pairplot of top features vs target variable sns.pairplot(df, hue = 'TenYearCHD', markers=["o", "s"], vars = □ → embeded_rf_feature, palette='bright')
```

[41]: <seaborn.axisgrid.PairGrid at 0x1e0f933cb60>



# 6 Modelling and predicting with Machine Learning

Since our dataset is imbalanced i.e for every positive case there are about 5-6 negative cases. We may end up with a classifier that is biased to the negative cases. The classifier may have a high accuracy but poor a precision and recall.

To handle this problem we will balance the dataset using the **Synthetic Minority Oversampling Technique (SMOTE)**.

# SMOTE :Synthetic Minority Oversampling Technique

SMOTE is an oversampling technique where the synthetic samples are generated for the minority class. This algorithm helps to overcome the overfitting problem posed by random oversampling. It focuses on the feature space to generate new instances with the help of interpolation between the positive instances that lie together.

Working Procedure: > At first the total no. of oversampling observations, N is set up. Generally,

it is selected such that the binary class distribution is 1:1. But that could be tuned down based on need. Then the iteration starts by first selecting a positive class instance at random. Next, the KNN's (by default 5) for that instance is obtained. At last, N of these K instances is chosen to interpolate new synthetic instances. To do that, using any distance metric the difference in distance between the feature vector and its neighbors is calculated. Now, this difference is multiplied by any random value in (0,1] and is added to the previous feature vector. This is pictorially represented below:

# **SMOTE** algorithm works in 4 simple steps:

- Choose a minority class as the input vector
- Find its k nearest neighbors (k\_neighbors is specified as an argument in the SMOTE() function)
- Choose one of these neighbors and place a synthetic point anywhere on the line joining the point under consideration and its chosen neighbor
- Repeat the steps until data is balanced

```
[42]: from imblearn.over_sampling import SMOTE

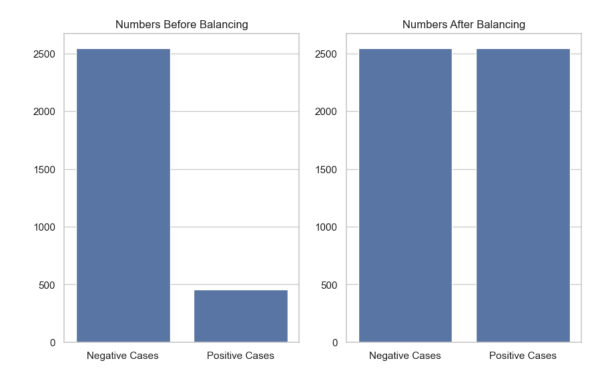
smote = SMOTE()

X = df[embeded_rf_feature]
y = df.iloc[:,-1]
# fit predictor and target variable
x_smote, y_smote = smote.fit_resample(X, y)

print('Original dataset shape', len(df))
print('Resampled dataset shape', len(y_smote))
```

Original dataset shape 3004 Resampled dataset shape 5094

```
[43]: from collections import Counter
  labels = ["Negative Cases", "Positive Cases"]
  plt.figure(figsize=(10,6))
  plt.subplot(1,2,1)
  sns.barplot(x=labels, y=list(dict(Counter(y)).values()))
  plt.title("Numbers Before Balancing")
  plt.subplot(1,2,2)
  sns.barplot(x=labels,y= list(dict(Counter(y_smote)).values()))
  plt.title("Numbers After Balancing")
  plt.show()
```



As seen after applying SMOTE, the new dataset is much more balanced.

## Splitting the data to Training and Testing set

```
[44]: # First let's create our new dataset
     df_new = pd.concat([pd.DataFrame(x_smote), pd.DataFrame(y_smote)], axis=1)
     df_new.columns = ['age', 'totChol', 'sysBP', 'diaBP', 'BMI', 'heartRate',
      df_new.head()
[44]:
            totChol sysBP diaBP
                                      BMI
                                          heartRate glucose
                                                              TenYearCHD
        age
               212.0 168.0
                              98.0 29.77
                                                72.0
                                                        75.0
     0
         36
                                                                       0
               250.0 116.0
                              71.0 20.35
                                                88.0
                                                        94.0
                                                                       0
     1
         46
     2
         50
               233.0 158.0
                              88.0 28.26
                                                68.0
                                                        94.0
                                                                       1
     3
         64
               241.0 136.5
                              85.0 26.42
                                                70.0
                                                        77.0
                                                                       0
               272.0 182.0 121.0 32.80
         61
                                                85.0
                                                        65.0
                                                                       1
[45]: X_new = df_new[embeded_rf_feature]
     y_new = df_new["TenYearCHD"]
     X_train, X_test, Y_train, Y_test = train_test_split(X_new, y_new, test_size=0.
       →2,random_state=42)
     print("Training features have {0} records and Testing features have {1} records.
           format(X_train.shape[0], X_test.shape[0]))
```

Training features have 4075 records and Testing features have 1019 records.

## 7 Models:

The four algorithms that we will be using are:

- 1. Logistic Regression
- 2. Random Forrest
- 3. XGBoost
- 4. Support Vector Machine

Here, we will be using **GridsearchCV** search algorithm for above algorithms

1. Logistic Regression

Logistic regression aims to measure the relationship between a categorical dependent variable and one or more independent variables (usually continuous) by plotting the dependent variables' probability scores.

```
[47]: # search for optimun parameters using gridsearch

params = {'penalty':['11','12'],

'C': [1e-6,1e-5,1e-4,1e-3,1e-2,1e-1,1,10,100,1e-3,1e+4,1e+5,1e+6],

'class_weight':['balanced',None]}

logistic_clf = GridSearchCV(LogisticRegression(),param_grid=params,cv=10,

⇒scoring='roc_auc')
```

```
[48]: #training the classifier logistic_clf.fit(X_train,Y_train) logistic_clf.best_params_
```

```
[48]: {'C': 0.0001, 'class_weight': 'balanced', 'penalty': '12'}
```

```
[49]: #making predictions
logistic_predict = logistic_clf.predict(X_test)
```

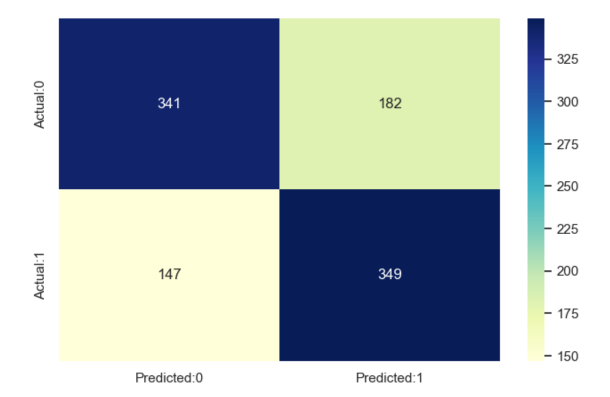
```
[50]: logistic_accuracy = accuracy_score(Y_test,logistic_predict)
print(f"Using logistic regression we get an accuracy of

Ground(logistic_accuracy*100,2)}%")
```

Using logistic regression we get an accuracy of 67.71%

Train ROC-AUC score : 0.6633128834355828 Test ROC-AUC score : 0.677134445534838

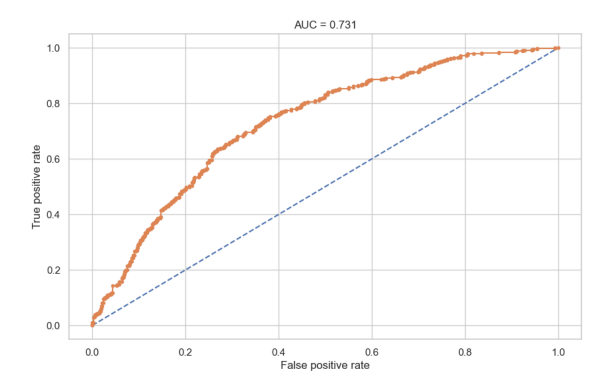
[52]: <Axes: >



# [53]: print(classification\_report(Y\_test,logistic\_predict))

```
precision
                            recall f1-score
                                                support
           0
                    0.70
                              0.65
                                         0.67
                                                    523
           1
                    0.66
                              0.70
                                         0.68
                                                     496
                                         0.68
                                                    1019
    accuracy
                              0.68
                                         0.68
                                                    1019
   macro avg
                    0.68
weighted avg
                    0.68
                              0.68
                                         0.68
                                                    1019
```

```
[54]: # ROC curve and AUC
      probs = logistic_clf.predict_proba(X_test)
      # keep probabilities for the positive outcome only
      probs = probs[:, 1]
      # calculate AUC
      log_auc = roc_auc_score(Y_test, probs)
      # calculate roc curve
      fpr, tpr, thresholds = roc_curve(Y_test, probs)
      # plot curve
      sns.set_style('whitegrid')
      plt.figure(figsize=(10,6))
      plt.plot([0, 1], [0, 1], linestyle='--')
      plt.plot(fpr, tpr, marker='.')
      plt.ylabel('True positive rate')
      plt.xlabel('False positive rate')
      plt.title(f"AUC = {round(log_auc,3)}")
      plt.show()
```



#### 2. Random Forest

Random forests are a way of averaging multiple deep decision trees, trained on different parts of the same training set, with the goal of reducing the variance. This comes at the expense of a small increase in the bias and some loss of interpretability, but generally greatly boosts the performance in the final model.

```
[55]: # search for optimun parameters using gridsearch
params_rf = {
    'max_depth': [4, 6, 8],
    'min_samples_leaf': [40, 50],
    'min_samples_split': [50, 100, 150],
    'n_estimators': [50, 80, 100]
}

random_clf = GridSearchCV(RandomForestClassifier(),param_grid=params_rf,cv=10, using elements)
scoring='roc_auc')
```

```
[56]: #training the classifier
random_clf.fit(X_train,Y_train)
random_clf.best_params_
```

```
'min_samples_split': 50,
'n_estimators': 100}
```

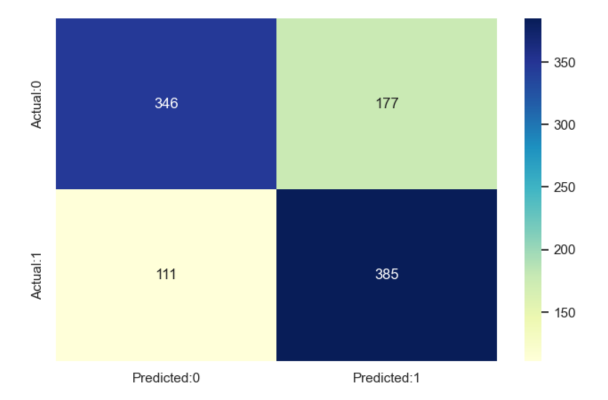
```
[57]: #making predictions
random_predict = random_clf.predict(X_test)
```

```
[58]: random_accuracy = accuracy_score(Y_test,random_predict)
print(f"Using Random Forest we get an accuracy of

→{round(random_accuracy*100,2)}%")
```

Using Random Forest we get an accuracy of 71.74%

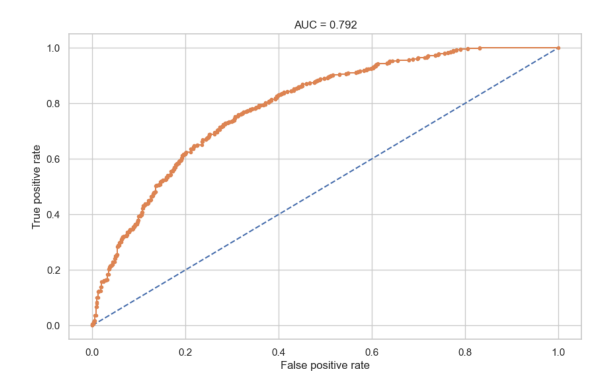
## [59]: <Axes: >



```
[60]: print(classification_report(Y_test,random_predict))
```

```
precision
                         recall f1-score
                                              support
                   0.76
           0
                             0.66
                                       0.71
                                                  523
           1
                   0.69
                             0.78
                                       0.73
                                                  496
   accuracy
                                       0.72
                                                 1019
                                       0.72
                                                 1019
  macro avg
                   0.72
                             0.72
weighted avg
                   0.72
                             0.72
                                       0.72
                                                 1019
```

```
[61]: # ROC curve and AUC
      probs1 = random_clf.predict_proba(X_test)
      # keep probabilities for the positive outcome only
      probs1 = probs1[:, 1]
      # calculate AUC
      ran_auc = roc_auc_score(Y_test, probs1)
      # calculate roc curve
      fpr, tpr, thresholds = roc_curve(Y_test, probs1)
      # plot curve
      sns.set_style('whitegrid')
      plt.figure(figsize=(10,6))
      plt.plot([0, 1], [0, 1], linestyle='--')
      plt.plot(fpr, tpr, marker='.')
      plt.ylabel('True positive rate')
      plt.xlabel('False positive rate')
      plt.title(f"AUC = {round(ran_auc,3)}")
      plt.show()
```



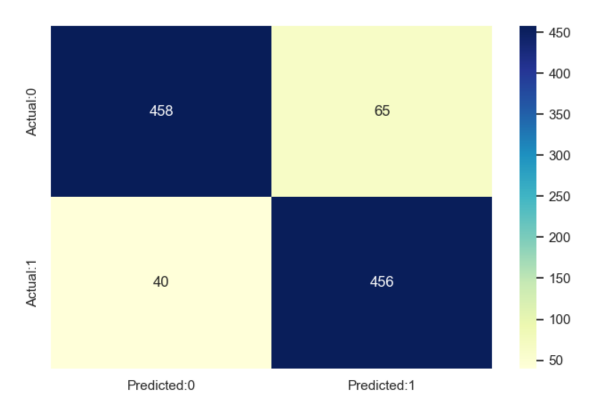
## 3. XGBoost

**XGBoost** stands for eXtreme Gradient Boosting. The name xgboost, though, actually refers to the engineering goal to push the limit of computations resources for boosted tree algorithms

```
[65]: xgb_accuracy = accuracy_score(Y_test,xgb_predict)
print(f"Using XG boost we get an accuracy of {round(xgb_accuracy*100,2)}%")
```

Using XG boost we get an accuracy of 89.7%

#### [66]: <Axes: >

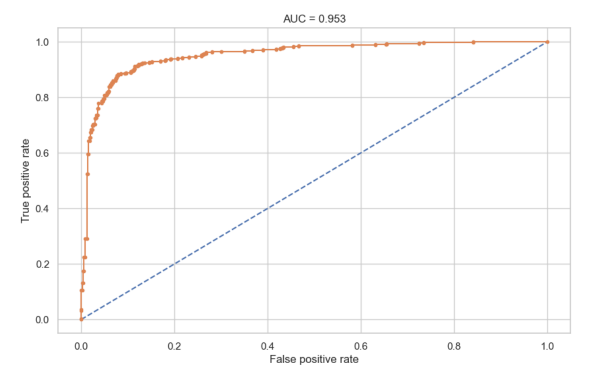


# [67]: print(classification\_report(Y\_test, xgb\_predict))

	precision	recall	f1-score	support
0	0.92	0.88	0.90	523
1	0.88	0.92	0.90	496
accuracy			0.90	1019
macro avg	0.90	0.90	0.90	1019

weighted avg 0.90 0.90 0.90 1019

```
[68]: # ROC curve and AUC
      probs2 = xgb_clf.predict_proba(X_test)
      # keep probabilities for the positive outcome only
      probs2 = probs2[:, 1]
      # calculate AUC
      xgb_auc = roc_auc_score(Y_test, probs2)
      # calculate roc curve
      fpr, tpr, thresholds = roc_curve(Y_test, probs2)
      # plot curve
      sns.set_style('whitegrid')
      plt.figure(figsize=(10,6))
      plt.plot([0, 1], [0, 1], linestyle='--')
      plt.plot(fpr, tpr, marker='.')
      plt.ylabel('True positive rate')
      plt.xlabel('False positive rate')
      plt.title(f"AUC = {round(xgb_auc,3)}")
      plt.show()
```



#### 4. Support Vector Machine

Support vector machines (SVMs) are powerful yet flexible supervised machine learning al-

gorithms which are used both for classification and regression. But generally, they are used in classification problems.

An SVM model is basically a representation of different classes in a hyperplane in multidimensional space. The hyperplane will be generated in an iterative manner by SVM so that the error can be minimized. The goal of SVM is to divide the datasets into classes to find a maximum marginal hyperplane (MMH).

```
[69]: # Grid search for optimum parameters
    Cs = [0.001, 0.01, 0.1, 1, 10]
    gammas = [0.001, 0.01, 0.1, 1]
    param_grid = {'C': Cs, 'gamma' : gammas}
    svm_clf = GridSearchCV(SVC(kernel='rbf', probability=True), param_grid, cv=10)

[70]: #training the classifier
    svm_clf.fit(X_train,Y_train)
    svm_clf.best_params_

[70]: {'C': 10, 'gamma': 0.01}

[71]: #making predictions
    svm_predict = svm_clf.predict(X_test)
[72]: svm_accuracy = accuracy score(Y test,svm_predict)
```

Using Support Vector Machine we get an accuracy of 92.74%

→{round(svm\_accuracy\*100,2)}%")

print(f"Using Support Vector Machine we get an accuracy of  $\Box$ 

```
[73]: # confusion matrix of SVM

cm=confusion_matrix(Y_test,svm_predict)

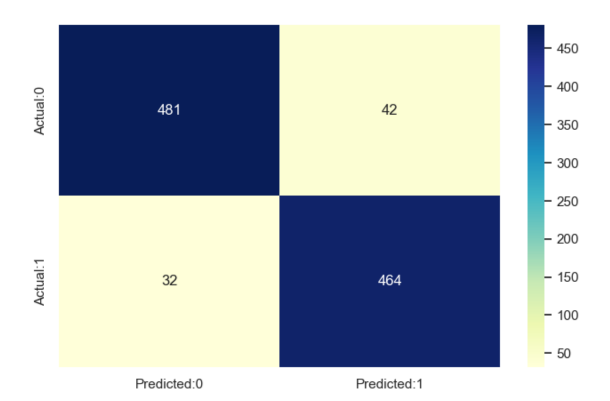
conf_matrix=pd.DataFrame(data=cm,columns=['Predicted:0','Predicted:

$\infty$1'],index=['Actual:0','Actual:1'])

plt.figure(figsize = (8,5))

sns.heatmap(conf_matrix, annot=True,fmt='d',cmap="YlGnBu")
```

[73]: <Axes: >



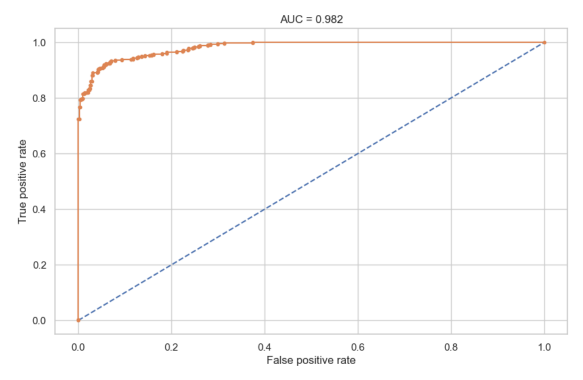
# [74]: print(classification\_report(Y\_test, svm\_predict))

support	f1-score	recall	precision	
523	0.93	0.92	0.94	0
496	0.93	0.94	0.92	1
1019	0.93			2.coura.cu
1019	0.93	0.93	0.93	accuracy macro avg
1019	0.93	0.93	0.93	weighted avg

```
[75]: # ROC curve and AUC
probs3 = svm_clf.predict_proba(X_test)
# keep probabilities for the positive outcome only
probs3 = probs3[:, 1]
# calculate AUC
svc_auc = roc_auc_score(Y_test, probs3)

# calculate roc curve
fpr, tpr, thresholds = roc_curve(Y_test, probs3)
# plot curve
sns.set_style('whitegrid')
```

```
plt.figure(figsize=(10,6))
plt.plot([0, 1], [0, 1], linestyle='--')
plt.plot(fpr, tpr, marker='.')
plt.ylabel('True positive rate')
plt.xlabel('False positive rate')
plt.title(f"AUC = {round(svc_auc,3)}")
plt.show()
```



#### Lets collect all our best models

```
"Support vector machine":{'Test Accuracy':round(svm_accuracy,__

42),'Precision': round(precision_score(Y_test, svm_predict), 2),'Recall':_

4round(recall_score(Y_test, svm_predict), 2),'F1 Score':_

4round(f1_score(Y_test, svm_predict), 2), 'AUC':round(svc_auc, 2)}

}).T
```

#### [77]: Performance\_df

[77]:	Te	est Accuracy	Precision	Recall	F1 Score	AUC
Logistic r	egression	0.68	0.66	0.70	0.68	0.73
Random For	est	0.72	0.69	0.78	0.73	0.79
XG Boost		0.90	0.88	0.92	0.90	0.95
Support ve	ctor machine	0.93	0.92	0.94	0.93	0.98

Observation from above table: \* XG Boost, Support vector machine gives highest Accuracy, Recall, Precision and AUC score. \* Highest recall is given by Support vector machine \* Highest AUC is given by Support vector machine

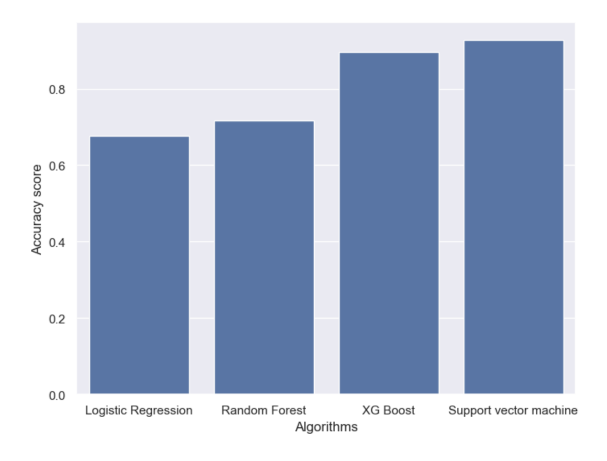
Overall we can say that **Support vector machine** is the best model that can be used for the risk prediction of Cardiovasular heart disease.

#### Let's Plot the Accuracy and AUC score graph of each algorithm

Accuracy Score plot

```
[78]: # Storing accuracies of each algorithm in a list
scores = [logistic_accuracy,random_accuracy,xgb_accuracy,svm_accuracy]
# Naming the algorithms and storing in a list
algorithms = ["Logistic Regression","Random Forest","XG Boost","Support vector
→machine"]
# Visualize the algorithms
sns.set(rc={'figure.figsize':(8,6)})
plt.xlabel("Algorithms")
plt.ylabel("Accuracy score")
sns.barplot(x=algorithms,y=scores)
```

[78]: <Axes: xlabel='Algorithms', ylabel='Accuracy score'>



# AUC Score plot

```
[79]: # Storing AUC score of each algorithm in a list

auc_scores = [log_auc,ran_auc,xgb_auc,svc_auc]

# Naming the algorithms and storing in a list

algorithms = ["Logistic Regression","Random Forest","XG Boost","Support vector

machine"]

# Visualize the algorithms

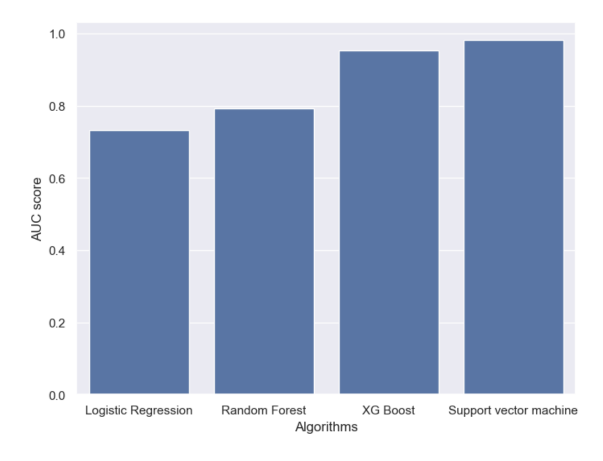
sns.set(rc={'figure.figsize':(8,6)})

plt.xlabel("Algorithms")

plt.ylabel("AUC score")

sns.barplot(x=algorithms,y=auc_scores)
```

[79]: <Axes: xlabel='Algorithms', ylabel='AUC score'>



From both the graphs we can say that the best performing model is **Support Vector Machine** algorithm.

# 8 Conclusion:

- The people who have Cardiovascular heart disease is almost equal between smokers and non smokers.
- The top features in predicting the ten year risk of developing Cardiovasular Heart Disease are 'age', 'totChol', 'sysBP', 'diaBP', 'BMI', 'heartRate', 'glucose'.
- The Support vector machine with the radial kernel is the best performing model in terms of accuracy and the F1 score and Its high AUC-score shows that it has a high true positive rate.
- Balancing the dataset by using the SMOTE technique helped in improving the models' sensitivity.
- With more data(especially that of the minority class) better models can be built.

```
[]:
```

[]:[