## Heart Disease

## September 10, 2024

```
[41]: import pandas as pd
import numpy as np
import matplotlib.pyplot as plt
import seaborn as sns
pd.options.display.float_format='{:,.2f}'.format # to round all number to 0.2
sns.set()
```

About Dataset

Column Name

Description

sex

This column represents the gender of the individuals (female-male).

age

This column represents the age of the individuals in the dataset. Age is a crucial factor in assessing the risk of coronary heart disease.

education

This column represents the level of education of the individuals. It could be coded using categorical values indicating different levels of education attainment.

smokingStatus

This column likely represents the smoking status of the individuals, indicating whether they are smokers (yes) or non-smokers (no).

cigsPerDay

If an individual is a smoker, this column represents the number of cigarettes smoked per day.

**BPMeds** 

This column indicates whether the individual is taking blood pressure medications (binary: 0 for not taking, 1 for taking).

prevalentStroke

This column indicates whether an individual has had a stroke prior to the study (binary: 0 for no, 1 for yes).

prevalentHyp

This column indicates whether an individual has hypertension (binary: 0 for no, 1 for yes).

diabetes

This column indicates whether an individual has diabetes (binary: 0 for no, 1 for yes).

totChol

This column represents the total cholesterol level of the individuals.

sysBP

This column represents the systolic blood pressure of the individuals.

diaBP

This column represents the diastolic blood pressure of the individuals.

BMI

This column represents the Body Mass Index (BMI) of the individuals, which is a measure of body fat based on height and weight.

heartRate

This column represents the resting heart rate of the individuals.

glucose

This column represents the fasting blood glucose level of the individuals.

#### **CHDRisk**

This column likely represents the Ten-Year Coronary Heart Disease (CHD) Risk for each individual, which is the target variable that you may want to predict or analyze.

```
[42]: data=pd.read_csv('Heart_Disease.csv')
data.head()
```

F407						~ .		_	DD14 1		a.		
[42]:		sex	age	education	smoking	Status	cigsPe	rDay	BPMeds	prev	alentStro	ke	\
	0	${\tt male}$	39	4		no		0	0			0	
	1	female	46	2		no		0	0			0	
	2	male	48	1		yes		20	0			0	
	3	female	61	3		yes		30	0			0	
	4	female	46	3		yes		23	0			0	
		prevale	ntHyp	diabetes	totChol	sysBP	diaBP	BMI	heart	Rate	glucose	\	
	0		0	no	195	106.00	70.00	26.97	,	80	77		
	1		0	no	250	121.00	81.00	28.73	3	95	76		
	2		0	no	245	127.50	80.00	25.34	<u> </u>	75	70		
	3		1	no	225	150.00	95.00	28.58	3	65	103		
	4		0	no	285	130.00	84.00	23.10	)	85	85		

CHDRisk

0 no

```
1 no
2 no
3 yes
4 no

[43]: df=data.copy()
```

## 1 Data\_Size

```
[44]: print(f'shape of data is {df.shape[0]}, {df.shape[1]}')
```

shape of data is 3674 , 16

## 2 Data\_Types

```
[45]: df.dtypes
```

```
[45]: sex
                            object
      age
                             int64
      education
                             int64
      smokingStatus
                            object
      cigsPerDay
                             int64
      BPMeds
                             int64
      prevalentStroke
                             int64
      prevalentHyp
                             int64
      diabetes
                            object
      totChol
                             int64
      sysBP
                           float64
      diaBP
                           float64
      BMI
                           float64
      {\tt heartRate}
                             int64
      glucose
                             int64
      CHDRisk
                            object
```

dtype: object

[46]: df.info()

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 3674 entries, 0 to 3673
Data columns (total 16 columns):

#	Column	Non-Null Count	Dtype
0	sex	3663 non-null	object
1	age	3674 non-null	int64
2	education	3674 non-null	int64
3	${\tt smokingStatus}$	3661 non-null	object
4	cigsPerDay	3674 non-null	int64

```
5
    BPMeds
                     3674 non-null
                                    int64
    prevalentStroke 3674 non-null
 6
                                    int64
 7
    prevalentHyp
                     3674 non-null
                                  int64
 8
    diabetes
                     3674 non-null
                                    object
 9
    totChol
                     3674 non-null int64
 10 sysBP
                     3674 non-null
                                    float64
 11 diaBP
                     3674 non-null float64
 12 BMT
                     3674 non-null
                                    float64
 13 heartRate
                    3674 non-null int64
                     3674 non-null
14 glucose
                                    int64
 15 CHDRisk
                     3674 non-null
                                    object
dtypes: float64(3), int64(9), object(4)
memory usage: 459.4+ KB
```

## 3 Unique Values

```
[47]: for col in df.columns:
       print(f' {col}: \n number of unique value for each column {df[col].
     →nunique()} , \n unique values is {df[col].unique()}')
       print('='*100)
    sex:
    number of unique value for each column 2 ,
    unique values is ['male' 'female' nan]
   ______
   number of unique value for each column 39,
    unique values is [39 46 48 61 43 63 45 52 50 41 38 42 44 47 35 60 36 59 54 37
   53 49 65 51
    62 40 56 67 57 66 64 55 58 68 34 33 32 70 69]
    _____
    education:
    number of unique value for each column 4 ,
    unique values is [4 2 1 3]
   ______
   ================
    smokingStatus:
    number of unique value for each column 2 ,
    unique values is ['no' 'yes' nan]
   ______
   _____
    cigsPerDay:
    number of unique value for each column 33 ,
    unique values is [ 0 20 30 23 15 10 5 35 43 1 40 3 9 2 12 4 18 60 25 45
   8 13 11 7
```

```
______
BPMeds:
number of unique value for each column 2 ,
unique values is [0 1]
______
______
prevalentStroke:
number of unique value for each column 2 ,
unique values is [0 1]
______
prevalentHyp:
number of unique value for each column 2 ,
unique values is [0 1]
______
diabetes:
number of unique value for each column 2,
unique values is ['no' 'yes']
______
==============
totChol:
number of unique value for each column 241,
unique values is [195 250 245 225 285 228 205 313 260 254 247 294 332 221 232
291 190 234
215 270 272 295 226 209 214 178 233 180 243 237 311 208 252 261 179 267
216 240 266 255 185 220 235 212 223 300 302 175 189 258 202 183 274 170
210 197 326 188 256 244 193 239 296 269 275 268 265 173 273 290 278 264
282 257 241 288 200 213 303 246 150 187 286 154 279 293 259 219 230 320
312 165 159 174 242 301 167 308 325 229 236 224 253 464 248 171 186 227
249 176 196 310 164 135 238 207 342 287 182 352 284 203 262 155 323 206
283 319 194 340 328 222 368 218 276 339 231 198 201 277 304 177 199 292
305 152 161 168 181 251 271 217 370 439 145 263 330 157 398 162 314 166
160 281 289 355 307 156 329 143 211 298 334 192 184 204 280 191 163 318
353 360 335 158 346 169 140 324 600 315 392 322 306 309 149 137 172 317
358 345 391 410 297 338 148 372 366 333 327 344 144 390 321 405 359 350
336 380 299 124 371 113 354 382 364 341 133 367 153 432 351 337 363 331
316 361 453 347 373 385 119]
______
===============
number of unique value for each column 231,
unique values is [106. 121. 127.5 150. 130. 180. 138. 100. 141.5 162.
133. 131.
142. 124. 140. 112. 122. 139. 108. 148. 132. 137.5 102.
115. 147. 124.5 160. 153. 111. 116.5 206. 96. 179.5 119.
```

```
114. 143.5 158. 157. 123.5 126.5 136. 154. 190.
156.5 145.
112.5 110. 138.5 155. 151. 152. 179. 113. 200. 132.5 126.
134. 141.
          135. 187. 127. 160.5 105. 109. 128. 118.
                                                       117.5 149.
180.5 136.5 212. 191. 121.5 173.
                                144. 129.5 117. 125.
                                                       144.5 170.
           166. 177.5 129. 159. 130.5 107.5 189. 168.
137.
      94.
                                                       197.5 146.
174.
      98.
           131.5 101. 158.5 97.
                                151.5 97.5 120.
                                                  204.
                                                       157.5 140.5
171. 215.
           95. 156. 122.5 178. 146.5 113.5 197.
                                                   90.
                                                       109.5 165.
95.5 209. 162.5 295. 103. 134.5 115.5 174.5 163. 118.5 185. 220.
164. 120.5 98.5 161. 139.5 168.5 176. 163.5 128.5 167. 205.5 119.5
167.5 152.5 186. 183. 153.5 147.5 175. 142.5 192.
                                                   96.5 159.5 177.
102.5 244. 104. 213. 199. 184. 198. 114.5 125.5 111.5 105.5 143.
161.5 164.5 171.5 108.5 201. 148.5 172. 243. 145.5 187.5 99.
133.5 100.5 135.5 172.5 103.5 149.5 182.5 186.5 217. 196. 193.
155.5 92. 169. 166.5 202. 150.5 195. 232.
                                             85.5 184.5 188.
169.5 210. 181.5 188.5 176.5 92.5 202.5 154.5 83.5 106.5 170.5 93.
175.5 207.5 199.5 101.5 248. 99.5 85. 230. 214. 192.5 104.5 194.
93.5 207. 185.5]
```

\_\_\_\_\_\_

#### diaBP:

number of unique value for each column 142, unique values is [ 70. 81. 80. 95. 84. 110. 71. 89. 107. 76. 88. 94. 90. 78. 84.5 70.5 82. 68. 91. 121. 85.5 85. 92.5 74. 98. 101. 73. 83.5 92. 63. 114. 77.5 69. 66. 82.5 102. 79. 67.5 72.5 106. 75. 87. 99. 60. 86.5 104. 86. 61.5

71.5 76.5 64. 77. 88.5 105. 96. 97. 100. 106.5 93. 80.5 124.5 61. 83. 67. 74.5 66.5 65. 72. 99.5 122.5 57. 78.5 104.5 89.5 112. 55. 120. 118. 59. 133. 95.5 96.5 111. 135. 64.5 68.5 98.5 62. 117. 59.5 103. 75.5 73.5 69.5 87.5

108. 93.5 90.5 114.5 62.5 94.5 140. 124. 91.5 115. 109. 65.5 105.5 103.5 63.5 79.5 107.5 142.5 109.5 58. 97.5 116.5 100.5

116. 119. 81.5 54. 132. 101.5 136. 51. 128. 125. 130. 110.5

113. 53. 108.5 112.5 52. 48. 56. 60.5 115.5 127.5]

\_\_\_\_\_

number of unique value for each column 1297, unique values is [26.97 28.73 25.34 ... 26.7 43.67 19.71]

\_\_\_\_\_\_

#### \_\_\_\_\_

### heartRate:

number of unique value for each column 72 , unique values is [ 80 95 75 65 85 77 60 79 76 93 72 98 64 70 71 62 73 90

96 68 63 88 78 83 100 84 57 50 74 86 55 92 66 87 110 81 82 54 69 67 52 61 140 130 58 104 94 105 91 53 108 106 59 107 48 112 125 103 44 47 45 97 122 102 120 99 115 143 101 46]

```
---------------
     glucose:
     number of unique value for each column 138,
     unique values is [ 77 76 70 103 85 99 78 79 88 61 64 84 72 89
    113 75 83
      66 74
             63
                87 225
                      90 80 100 215
                                     98 95
                                           94 55
                                                  82 93 73
      68 97 104
                96 126 120 105
                             71 56
                                     60 117
                                            62 102 58
                                                      92 109 86 107
               57 91 132 150 59 81 115 140 112 118 114 160 110 123 108
                                     53 216 163 144 116 121 172 124 111
     145 122 137 106 127 205 130 101
                                 47
      40 186 223 325 44 156 268 50 274 292 255 136 206 131 148 43 173 386
     155 147 170 52 320 254 394 270 244 183 142 119 167 135 207 129 177 250
     294 125 332 368 348 370 193 191 256 235 210 260]
    _____
    ______
     CHDRisk:
     number of unique value for each column 2 ,
     unique values is ['no' 'yes']
    _____
     ______
    Categorizing Age into Age Groups
[48]: df['age_group'] = pd.cut(df['age'], bins=[0, 30, 40, 60, 80], labels=['0-30', __
      df.drop(columns='age',inplace=True)
    Converting Binary Medical Columns to Categorical Values for easier interpretation and analysis
[49]: df['BPMeds']=df['BPMeds'].astype('object')
     df['prevalentStroke'] = df['prevalentStroke'].astype('object')
     df['prevalentHyp']=df['prevalentHyp'].astype('object')
     df['BPMeds']=df['BPMeds'].replace([0,1],['no','yes'])
     df['prevalentStroke']=df['prevalentStroke'].replace([0,1],['no','yes'])
     df['prevalentHyp']=df['prevalentHyp'].replace([0,1],['no','yes'])
```

## 4 Show Data

```
[50]: df.head()
[50]:
             sex
                  education smokingStatus
                                             cigsPerDay BPMeds prevalentStroke
           male
                           4
                                                       0
                                                              no
                                         no
      1
         female
                           2
                                                       0
                                         no
                                                              no
                                                                               no
           male
                           1
                                                      20
                                        yes
                                                              no
                                                                               no
      3
        female
                           3
                                                      30
                                        yes
                                                              no
                                                                               no
      4 female
                           3
                                                      23
                                        yes
                                                              no
                                                                               no
```

```
prevalentHyp diabetes
                                totChol sysBP
                                                  diaBP
                                                          BMI
                                                                heartRate glucose \
                                     195 106.00
                                                  70.00 26.97
                                                                       80
                                                                                 77
      0
                   no
                            no
      1
                   no
                            no
                                     250 121.00
                                                  81.00 28.73
                                                                       95
                                                                                 76
      2
                                     245 127.50
                                                  80.00 25.34
                                                                       75
                                                                                 70
                   no
                            no
      3
                                     225 150.00
                                                  95.00 28.58
                                                                       65
                                                                                103
                  yes
                            no
                                     285 130.00
                                                  84.00 23.10
                                                                       85
                                                                                 85
                   no
                            no
        CHDRisk age_group
                     30-40
      0
             no
      1
                     40-60
             no
      2
                     40-60
             no
      3
                     60-80
            yes
      4
             no
                     40-60
[51]: df.tail()
                   education smokingStatus cigsPerDay BPMeds prevalentStroke
[51]:
              sex
      3669 male
                           3
                                                      25
                                                              no
                                        yes
                                                                               no
      3670 male
                           3
                                                      25
                                        yes
                                                              no
                                                                               no
      3671 male
                           3
                                                      25
                                        yes
                                                              no
                                                                               no
      3672 male
                           3
                                                      25
                                        yes
                                                              no
                                                                               no
      3673 male
                           2
                                                      25
                                        yes
                                                              no
                                                                               no
           prevalentHyp diabetes
                                    totChol sysBP
                                                     diaBP
                                                              {\tt BMI}
                                                                   heartRate
                                                                               glucose
      3669
                                        208 137.50
                                                     82.50 25.58
                      no
                                no
                                                                           75
                                                                                    63
      3670
                                        208 137.50
                                                     82.50 25.58
                                                                           75
                                                                                    63
                      no
                                no
      3671
                      no
                                no
                                        208 137.50
                                                     82.50 25.58
                                                                           75
                                                                                    63
      3672
                                        208 137.50
                                                     82.50 25.58
                                                                           75
                                                                                    63
                      no
                                no
      3673
                                        208 137.50 82.50 25.97
                                                                           69
                                                                                    68
                      no
                                no
           CHDRisk age_group
      3669
                yes
                        40-60
      3670
                        40-60
                yes
      3671
                        40-60
                yes
      3672
                        40-60
                yes
      3673
                        40-60
                yes
[52]: df.sample(2)
[52]:
                     education smokingStatus cigsPerDay BPMeds prevalentStroke
                sex
                              3
      1529
            female
                                          yes
                                                        20
                                                                no
                                                                                 no
      1801 female
                             1
                                          yes
                                                         9
                                                                                 no
                                                                no
           prevalentHyp diabetes totChol sysBP
                                                                   heartRate
                                                                               glucose \
                                                     diaBP
                                                              BMI
      1529
                                                                           94
                                        249 109.00
                                                     66.50 24.79
                                                                                    85
                      no
                                no
      1801
                      no
                                no
                                        309 130.00
                                                     86.00 22.37
                                                                           82
                                                                                    80
```

```
CHDRisk age_group
1529 no 40-60
1801 no 40-60
```

## 5 Missing\_Values

```
[53]: df.isna().sum()
[53]: sex
                          11
      education
                           0
                          13
      smokingStatus
      cigsPerDay
                           0
      BPMeds
                           0
      prevalentStroke
                           0
      prevalentHyp
                           0
                           0
      diabetes
      totChol
                           0
                           0
      sysBP
      diaBP
                           0
      BMI
                           0
      heartRate
                           0
      glucose
                           0
      CHDRisk
                           0
                           0
      age_group
      dtype: int64
[54]: df=df.dropna()
[55]: df.info()
     <class 'pandas.core.frame.DataFrame'>
     Index: 3652 entries, 0 to 3673
     Data columns (total 16 columns):
      #
          Column
                            Non-Null Count
                                             Dtype
                            _____
          ----
      0
                            3652 non-null
                                             object
          sex
      1
          education
                            3652 non-null
                                             int64
      2
          smokingStatus
                            3652 non-null
                                             object
      3
          cigsPerDay
                            3652 non-null
                                             int64
      4
          BPMeds
                            3652 non-null
                                             object
      5
          prevalentStroke
                            3652 non-null
                                             object
          prevalentHyp
                            3652 non-null
                                             object
          diabetes
                            3652 non-null
                                             object
      8
          totChol
                            3652 non-null
                                             int64
      9
          sysBP
                            3652 non-null
                                             float64
      10
          diaBP
                            3652 non-null
                                             float64
      11
          BMI
                            3652 non-null
                                             float64
```

```
12 heartRate
                     3652 non-null
                                     int64
 13 glucose
                     3652 non-null int64
 14 CHDRisk
                     3652 non-null
                                     object
 15 age_group
                     3652 non-null
                                     category
dtypes: category(1), float64(3), int64(5), object(7)
```

memory usage: 460.3+ KB

# 6 Duplicated\_Values

[56]:	uplicated_data=df[df.duplicated(keep=False)]									
	duplicated_data									
[56] •	gay adjugation emokingStatus cigsParDay RDMade provalentStroke \									

[56].		gov	oducation	amolri	ngC+2+110	ai maDa	~Dorr	DDModa	nnovolon+C+n	ماده	`	
[56]:	3118	sex male	education		-	Cigsre	25	no	prevalentStr		\	
	3658	male		3	yes		25 25	no		no no		
	3659	male		3	yes		25					
	3660	male		3	yes		25	no no		no no		
	3661	male		3	yes yes		25	no		no		
	3662	male		3	yes		25	no		no		
	3663	male		3	yes		25	no		no		
	3664	male		3	yes		25	no		no		
	3665	male		3	yes		25	no		no		
	3666	male		3	yes		25	no		no		
	3667	male		3	yes		25	no		no		
	3668	male		3	yes		25	no		no		
	3669	male		3	yes		25	no		no		
	3670	male		3	yes		25	no		no		
	3671	male		3	yes		25	no		no		
	3672	male	3	3	yes		25	no		no		
					v							
		preval	entHyp dia	abetes	totChol	sysBP	diaB	P BMI	I heartRate	glud	cose	\
	3118		no	no		137.50		0 25.58			63	
	3658		no	no	208	137.50		0 25.58			63	
	3659		no	no		137.50		0 25.58			63	
	3660		no	no		137.50		0 25.58			63	
	3661		no	no		137.50		0 25.58			63	
	3662		no	no		137.50		0 25.58			63	
	3663		no	no		137.50		0 25.58			63	
	3664		no	no		137.50		0 25.58			63	
	3665		no	no		137.50		0 25.58			63	
	3666		no	no		137.50		0 25.58			63	
	3667		no	no		137.50		0 25.58			63	
	3668		no	no		137.50		0 25.58			63	
	3669		no	no		137.50		0 25.58			63	
	3670		no	no		137.50		0 25.58			63	
	3671		no	no		137.50		0 25.58			63	
	3672		no	no	208	137.50	82.5	0 25.58	3 75		63	

```
CHDRisk age_group
      3118
                        40-60
               yes
      3658
                        40-60
               yes
      3659
               yes
                        40-60
      3660
               yes
                        40-60
      3661
                        40-60
               yes
      3662
                        40-60
               yes
      3663
                        40-60
               yes
      3664
               yes
                        40-60
      3665
                        40-60
               yes
      3666
               yes
                        40-60
      3667
                        40-60
               yes
      3668
                        40-60
               yes
      3669
               yes
                        40-60
      3670
                        40-60
               yes
      3671
                        40-60
               yes
      3672
               yes
                        40-60
[57]: df.duplicated().sum()
[57]: 15
[58]: df.drop_duplicates(inplace=True)
[59]: df.duplicated().any().sum()
[59]: 0
[60]: df.isna().sum()
[60]: sex
                          0
                          0
      education
      smokingStatus
                          0
      cigsPerDay
                          0
      BPMeds
                          0
      prevalentStroke
                          0
      prevalentHyp
                          0
      diabetes
                          0
      totChol
                          0
      sysBP
                          0
      diaBP
                          0
      BMI
                          0
      heartRate
                          0
      glucose
                          0
      CHDRisk
                          0
                          0
      age_group
```

dtype: int64

## 7 Statistical OverView

```
[61]: df.describe().T
[61]:
                                                 25%
                                                        50%
                                                               75%
                    count
                            mean
                                   std
                                          min
                                                                      max
                            1.98 1.02
      education 3,637.00
                                         1.00
                                                1.00
                                                       2.00
                                                              3.00
                                                                     4.00
      cigsPerDay 3,637.00
                            9.03 11.91
                                         0.00
                                                0.00
                                                       0.00
                                                             20.00
                                                                   70.00
      totChol
                 3,637.00 236.88 44.13 113.00 206.00 234.00 263.00 600.00
      sysBP
                 3,637.00 132.36 22.08 83.50 117.00 128.00 144.00 295.00
                                                             90.00 142.50
      diaBP
                 3,637.00 82.90 11.96 48.00 75.00
                                                      82.00
     BMI
                 3,637.00
                           25.79 4.06 15.54
                                               23.08
                                                      25.38
                                                             28.04 56.80
     heartRate 3,637.00
                           75.75 11.99
                                        44.00
                                               68.00 75.00
                                                             82.00 143.00
      glucose
                 3,637.00 81.81 23.77
                                        40.00 71.00 78.00 87.00 394.00
         Analysis
[62]: df.columns
[62]: Index(['sex', 'education', 'smokingStatus', 'cigsPerDay', 'BPMeds',
             'prevalentStroke', 'prevalentHyp', 'diabetes', 'totChol', 'sysBP',
             'diaBP', 'BMI', 'heartRate', 'glucose', 'CHDRisk', 'age_group'],
            dtype='object')
     1- univariate
     Count Plots for Categorical Variables
[63]: plt.figure(figsize=(20,20))
      for i , col in enumerate(df.select_dtypes('object')):
          plt.subplot(4,2,i+1)
          ax=sns.countplot(x=df[col],order=df[col].value_counts().index)
          ax.bar_label(ax.containers[0])# this line to show count for each col
          plt.title(f'CountPlot of {col}')
          plt.tight layout()
          print(f'Counts for {col}:\n', df[col].value_counts(), '\n')
      plt.show()
     Counts for sex:
      sex
     female
               2026
     male
               1611
     Name: count, dtype: int64
     Counts for smokingStatus:
```

smokingStatus

no 1857 yes 1780

Name: count, dtype: int64

Counts for BPMeds:

BPMeds

no 3527 yes 110

Name: count, dtype: int64

 ${\tt Counts\ for\ prevalentStroke:}$ 

prevalentStroke

no 3616 yes 21

Name: count, dtype: int64

Counts for prevalentHyp:

prevalentHyp

no 2502

yes 1135

Name: count, dtype: int64

Counts for diabetes:

diabetes

no 3540 yes 97

Name: count, dtype: int64

Counts for CHDRisk:

CHDRisk

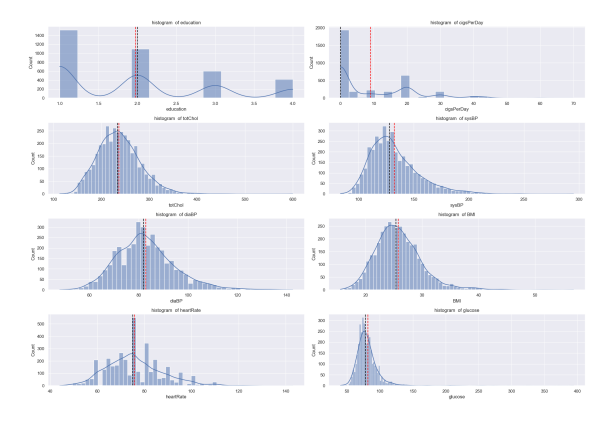
no 3084 yes 553

Name: count, dtype: int64



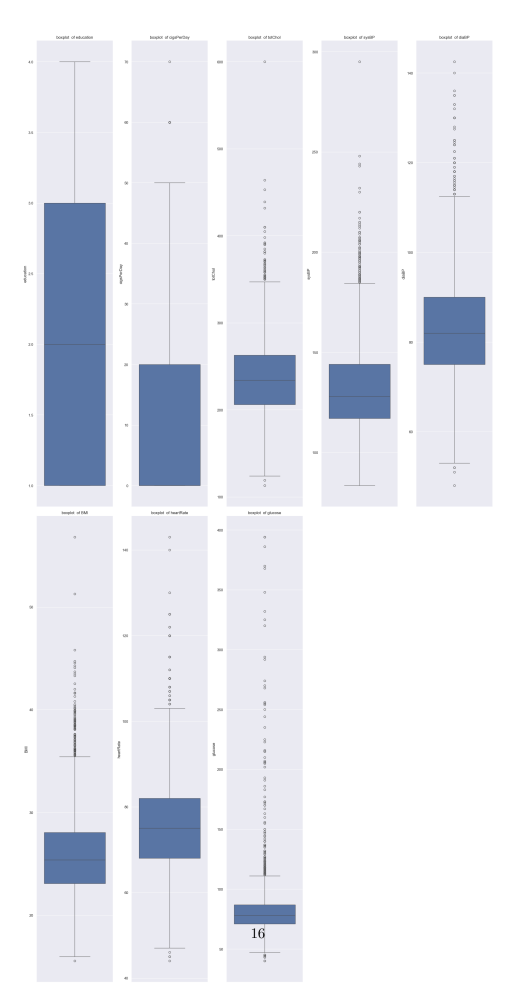
### Box Plots for numerical columns

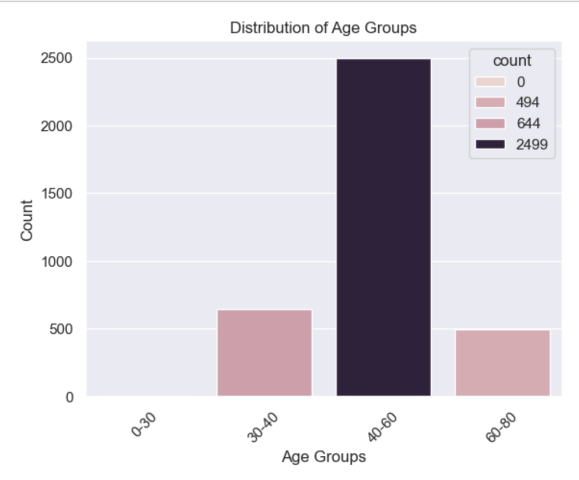
```
[65]: plt.figure(figsize=(20,20))
for i , col in enumerate(df.select_dtypes('number')):
    plt.subplot(6,2,i+1)
    ax=sns.histplot(df[col],kde=True)
    ax.axvline(df[col].mean(),color='red',linestyle='--')
    ax.axvline(df[col].median(),color='black',linestyle='--')
    plt.title(f'histogram of {col}')
    plt.tight_layout()
plt.show()
```



data is approximately normally distributed bec mean is closer to Median in histogram and descriptive stat

```
[64]: plt.figure(figsize=(20,40))
for i , col in enumerate(df.select_dtypes('number')):
    plt.subplot(2,5,i+1)
    ax=sns.boxplot(df[col])
    plt.title(f'boxplot of {col}')
    plt.tight_layout()
plt.show()
```





### 2- Bivariate

a. Categorical vs Categorical:

```
[67]: crossTab_df=pd.crosstab(df['sex'],df['CHDRisk'])
print(crossTab_df)
```

```
ax=crossTab_df.plot(kind='bar',title='Sex vs CHD Risk')
ax.bar_label(ax.containers[0])
ax.bar_label(ax.containers[1])
plt.show()
```

CHDRisk no yes sex female 1778 248 male 1306 305



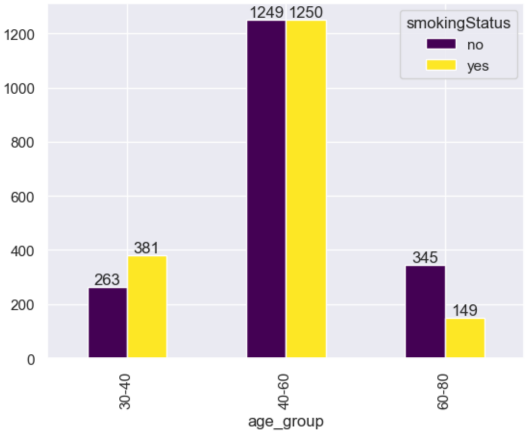
```
[]: crossTab_df=pd.crosstab(df['age_group'],df['CHDRisk'])
    print(crossTab_df)

ax=crossTab_df.plot(kind='bar',title='age_group vs CHD Risk',colormap='viridis')
ax.bar_label(ax.containers[0])
ax.bar_label(ax.containers[1])
```

```
plt.show()
```

smokingStatus	no	yes
age_group		
30-40	263	381
40-60	1249	1250
60-80	345	149

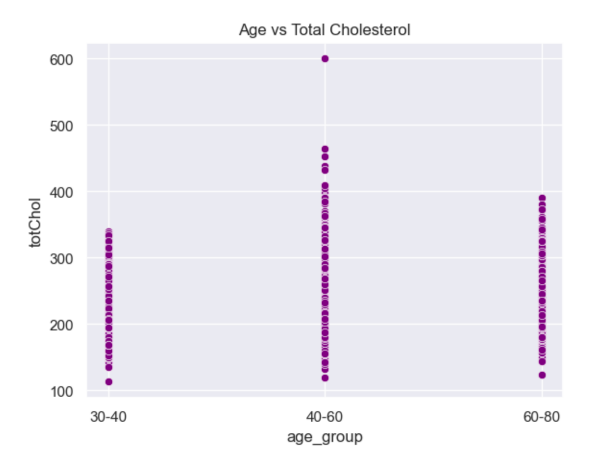




## b. Numerical vs Categorical

```
[69]: sns.scatterplot(data=df,x='age_group',y='totChol',color='purple')
plt.title('Age vs Total Cholesterol')
```

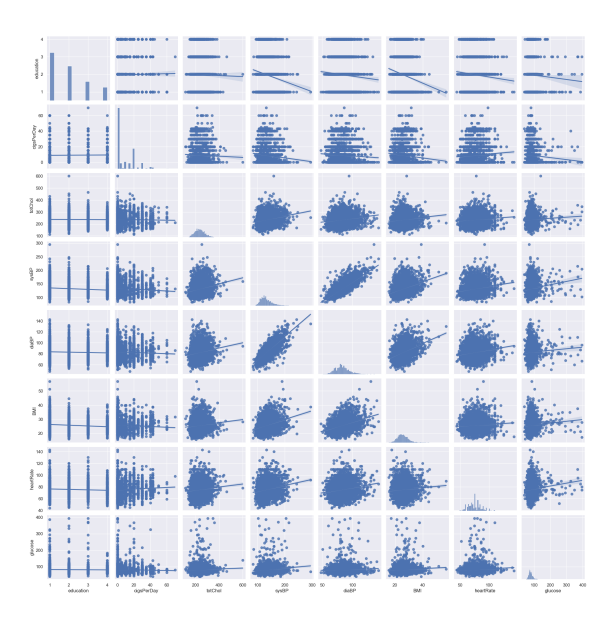
[69]: Text(0.5, 1.0, 'Age vs Total Cholesterol')

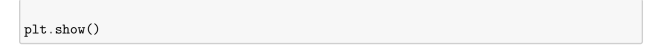


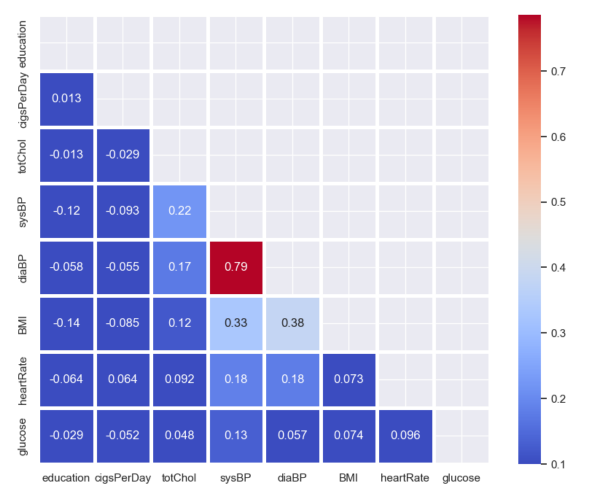
## 3- Multivariate Analysis

```
[71]: numerical_columns=df.select_dtypes('number')
sns.pairplot(data=numerical_columns,kind='reg')
```

[71]: <seaborn.axisgrid.PairGrid at 0x1f1c4b3cfb0>







there is a strong correlation between sysBP and diaBp

# 9 Health Metrics: Normal, Borderline, and High Ranges

Metric	Normal Range	Borderline/Elevate	${ m d High/Abnormal}$
Total	Less than $200 \text{ mg/dL}$	200-239  mg/dL	240 mg/dL and above
Cholesterol			
(totChol)			
Systolic Blood	90-120  mmHg	120-129  mmHg	130 mmHg and above
Pressure			
(sysBP)			

Metric	Normal Range	Borderline/Elevat	ed High/Abnormal
Diastolic Blood Pressure (diaBP)	60-80 mmHg	80-89 mmHg	90 mmHg and above
Body Mass Index (BMI)	18.5-24.9	25-29.9	30 and above (Obesity)
Heart Rate (heartRate)	60-100 bpm	-	Less than 60 bpm (Bradycardia), more than 100 bpm (Tachycardia)
Glucose	70-99  mg/dL (fasting)	100-125  mg/dL (Prediabetes)	126 mg/dL and above (Diabetes)

Number of Individuals Classified as Healthy Based on Key Health Metrics

```
[73]: healty_conditions=(
                     (df['sex'].isin(['male', 'female']))&
                     (df['smokingStatus'].isin(['no','yes']))&
                     (df['cigsPerDay'].between(0,20))&
                     (df['BPMeds']=='no')&
                     (df['prevalentStroke'] == 'no')&
                      (df['prevalentHyp']=='no')&
                      (df['diabetes']== 'no')&
                      (df['diaBP'].between(60,80))&
                      (df['sysBP'].between(90,120))&
                      (df['BMI'].between(18.5,24.9))&
                      (df['heartRate'].between(60,100))&
                      (df['glucose'].between(70,99))&
                      (df['totChol']<200)
      healthy_People=df[healty_conditions]
      length_healthy_people = len(healthy_People)
      print(f"Number of healthy people: {length_healthy_people}")
```

Number of healthy people: 100

# [74]: healthy\_People.sample(10)

[74]:		sex	education	smokingStatus	cigsPerDay	BPMeds	prevalentStroke	\
	1095	male	1	yes	13	no	no	
	1926	female	4	yes	15	no	no	
	3600	female	4	no	0	no	no	
	2702	female	2	yes	5	no	no	
	2267	female	2	yes	10	no	no	
	1518	female	3	yes	8	no	no	
	2504	female	2	no	0	no	no	
	2589	female	2	yes	5	no	no	
	3228	female	3	no	0	no	no	

3287	female		3	no		0 no			no	
	prevalent	:Нур с	diabetes	totChol	sysBP	diaBP	BMI	heartRate	glucose	\
1095		no	no	196	120.00	74.00	20.12	75	73	
1926		no	no	178	96.00	67.00	20.40	65	82	
3600		no	no	199	111.00	70.00	21.99	65	82	
2702		no	no	193	107.00	73.00	20.73	85	72	
2267		no	no	173	105.00	70.00	21.98	60	79	
1518		no	no	195	111.00	79.00	23.22	86	85	
2504		no	no	183	120.00	76.00	21.12	100	72	
2589		no	no	165	106.00	64.00	19.14	68	70	
3228		no	no	190	115.00	77.00	23.95	70	80	
3287		no	no	185	100.00	72.00	22.15	85	83	
	arran									
	CHDRisk a		-							
1095	no		0-60							
1926	no		0-40							
3600	no		0-60							
2702	no	30	0-40							
2267	no	40	0-60							
1518	no	40	0-60							
2504	no	40	0-60							
2589	no	30	0-40							
3228	no	30	0-40							
3287	no	30	0-40							

index of 1778 write yes in CHDRisk Although it is among the healthy people

#### 10 Distribution and Classification of Health Metrics and **CHDRisk**

• (barplot)

```
[75]: from matplotlib.ticker import FixedLocator
     # Create dictionaries for each category
     Smokers = {'NonSmokers': df[(df.cigsPerDay == 0)]['cigsPerDay'].count(),
                'Smokers': df[(df.cigsPerDay > 0) & (df.cigsPerDay <=_
      →20)]['cigsPerDay'].count(),
                'Dangerous Smokers': df[(df.cigsPerDay > 20) & (df.cigsPerDay <=__
       'High Smokers': df[(df.cigsPerDay > 50)]['cigsPerDay'].count()}
     Chol = {'LowChol': df[(df.totChol < 125)]['totChol'].count(),</pre>
             'NormalChol': df[(df.totChol > 125) & (df.totChol <= 200)]['totChol'].
       ⇔count(),
```

```
'DangerousChol': df[(df.totChol > 200) & (df.totChol <=__
 'HighChol': df[(df.totChol > 240)]['totChol'].count()}
SysBP = {'LowsysBP': df[(df.sysBP < 120)]['sysBP'].count(),
         'NormalsysBP': df[(df.sysBP >= 120) & (df.sysBP <= 129)]['sysBP'].
 ⇔count(),
         'DangeroussysBP': df[(df.sysBP >= 130) & (df.sysBP <= 179)]['sysBP'].
 ⇔count(),
         'HighsysBP': df[(df.sysBP >= 180)]['sysBP'].count()}
DiaBP = {'LowdiaBP': df[(df.diaBP < 80)]['diaBP'].count(),</pre>
         'NormaldiaBP': df[(df.diaBP >= 80) & (df.diaBP <= 84)]['diaBP'].
 ⇔count(),
         'DangerousdiaBP': df[(df.diaBP >= 85) & (df.diaBP <= 109)]['diaBP'].
 ⇔count(),
         'HighdiaBP': df[(df.diaBP >= 110)]['diaBP'].count()}
BMI = \{'LowBMI': df[(df.BMI < 18.5)]['BMI'].count(),
       'NormalBMI': df[(df.BMI >= 18.5) & (df.BMI <= 29.9)]['BMI'].count(),
       'DangerousBMI': df[(df.BMI >= 30) & (df.BMI <= 39.9)]['BMI'].count(),</pre>
       'HighBMI': df[(df.BMI >= 40)]['BMI'].count()}
HeartRate = {'LowHR': df[(df.heartRate < 60)]['heartRate'].count(),</pre>
             'NormalHR': df[(df.heartRate >= 60) & (df.heartRate <=__
 →100)]['heartRate'].count(),
             'DangerousHR': df[(df.heartRate > 100)]['heartRate'].count()}
Glo = {'LowGol': df[(df.glucose < 70)]['glucose'].count(),</pre>
       'NormalGol': df[(df.glucose >= 70) & (df.glucose <= 100)]['glucose'].
 ⇔count(),
       'DangerousGol': df[(df.glucose > 100)]['glucose'].count()}
BPMeds = {'Yes': df[(df.BPMeds == 'yes')]['BPMeds'].count(),
          'No': df[(df.BPMeds == 'no')]['BPMeds'].count()}
prevalentStroke = {'Yes': df[(df.prevalentStroke == 'yes')]['prevalentStroke'].
 ⇔count(),
                   'No': df[(df.prevalentStroke == 'no')]['prevalentStroke'].

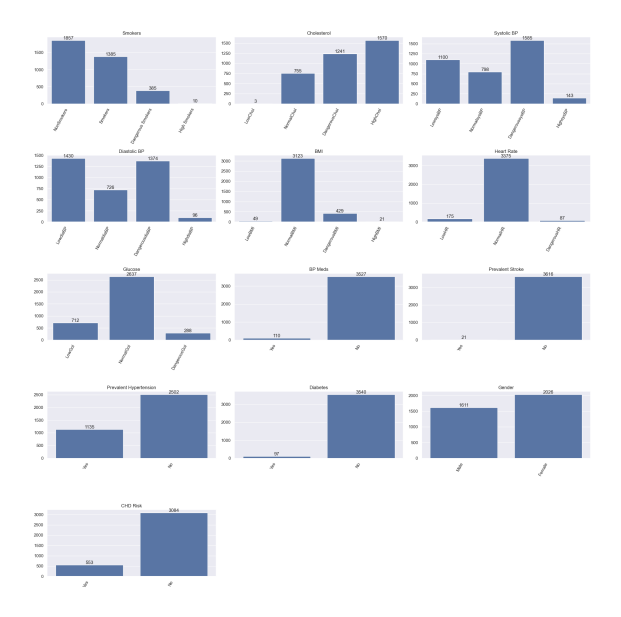
    count()}

prevalentHyp = {'Yes': df[(df.prevalentHyp == 'yes')]['prevalentHyp'].count(),
                'No': df[(df.prevalentHyp == 'no')]['prevalentHyp'].count()}
gender = {'Male': df[df['sex'] == 'male']['sex'].count(),
```

```
'Female': df[df['sex'] == 'female']['sex'].count()}
CHDRisk = {'Yes': df[(df.CHDRisk == 'yes')]['CHDRisk'].count(),
            'No': df[(df.CHDRisk == 'no')]['CHDRisk'].count()}
Diabetes = {'Yes': df[(df.diabetes == 'yes')]['diabetes'].count(),
             'No': df[(df.diabetes == 'no')]['diabetes'].count()}
# List of dictionaries and titles
dicts = [Smokers, Chol, SysBP, DiaBP, BMI, HeartRate, Glo, BPMeds, __
 ⇔prevalentStroke, prevalentHyp, Diabetes, gender, CHDRisk]
dicts_title = ['Smokers', 'Cholesterol', 'Systolic BP', 'Diastolic BP', 'BMI', |
 →'Heart Rate', 'Glucose', 'BP Meds', 'Prevalent Stroke', 'Prevalent<sub>□</sub>
 →Hypertension', 'Diabetes', 'Gender', 'CHD Risk']
# Plotting
plt.figure(figsize=(20, 20))
for e, dict_data in enumerate(dicts):
    d = pd.DataFrame(dict_data.items(), columns=['Classification', 'Value'])
    print(f" {dicts_title[e]}")
    print(d)
    print('=' * 50)
    plt.subplot(5, 3, e+1)
    ax = sns.barplot(data=d, x='Classification', y='Value')
    ax.bar_label(ax.containers[0])# to show count outside plot
    ax.set_title(dicts_title[e])
    ax.set_xlabel('')
    ax.set_ylabel('')
    ax.set_xticks(range(len(d['Classification'])))
    ax.set_xticklabels(d['Classification'], rotation=62, ha='right')
plt.tight_layout()
plt.show()
Smokers
      Classification Value
0
          NonSmokers
                      1857
             Smokers
                      1385
1
2 Dangerous Smokers
                        385
3
        High Smokers
                         10
```

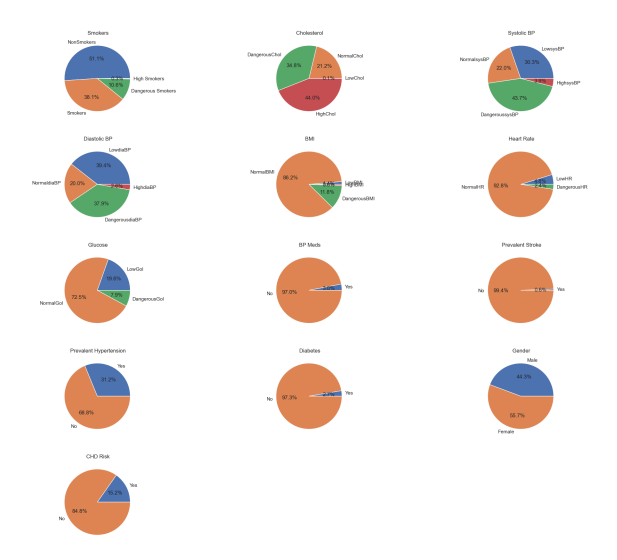
==	:========		-==
	Cholesterol	W= 1	
0	Classification LowChol	Value 3	
1	NormalChol	755	
2	DangerousChol	1241	
3	HighChol	1570	
==			-==
S	ystolic BP		
	Classification	Value	
0	LowsysBP	1100	
1	NormalsysBP	798	
2	DangeroussysBP	1585	
ა ==	HighsysBP	143 	) ===
_ <b>_</b>	iastolic BP		
ے	Classification	Value	3
0	LowdiaBP	1430	)
1	NormaldiaBP	726	3
2	DangerousdiaBP	1374	ŀ
3	HighdiaBP	96	3
==		======	-==
	MI Classification	Voler -	
0	Classification LowBMI	Value 49	
1	NormalBMI	3123	
2	DangerousBMI	429	
3	HighBMI	21	
==	:=====================================		-==
Н	leart Rate		
	Classification	Value	
0	LowHR	175	
1	NormalHR	3375	
2	DangerousHR	87	
==			===
	lucose Classification	Value	
0	LowGol	712	
1	NormalGol	2637	
2	DangerousGol	288	
==	- :===========		-==
В	P Meds		
		Value	
0	Yes	110	
1	No	3527 	
== n	revalent Stroke		-==
	Classification	Value	
	OTASSILICATION	varue	

0	Yes	21	
1	No	3616	
==		=====	
P	revalent Hypert	ension	
	Classification	Value	
0	Yes	1135	
1	No	2502	
==		======	
D	iabetes		
	Classification	Value	
0	Yes	97	
1	No	3540	
==	=========	======	
	ender		
		Value	
0	Male	1611	
1	Female	2026	
==		======	
	HD Risk		
		Value	
0	Yes	553	
1	No	3084	
==		======	



• (pie)

```
[76]: plt.figure(figsize=(20,20))
for i , dict_col in enumerate(dicts):
    d=pd.DataFrame(dict_col.items(),columns=['Classification','Value'])
    plt.subplot(6,3,i+1)
    plt.pie(d['Value'],labels=d['Classification'],autopct='%1.1f%%')
    plt.tight_layout()
    plt.title(f'\n \n {dicts_title[i]}')
plt.show()
```

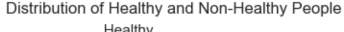


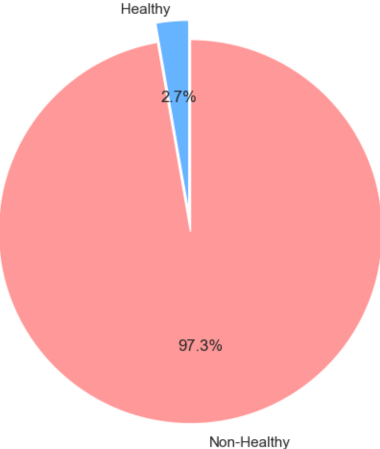
```
[77]: non_healthy_people=len(df)-length_healthy_people non_healthy_people
```

[77]: 3537

Distribution of Healthy vs. Non-Healthy Individuals

```
plt.axis('equal')
plt.show()
```





```
[79]: df.columns
```

percentage of males and females with various health conditions

```
[80]: disease_columns = ['smokingStatus', 'BPMeds', 'prevalentStroke', 'prevalentHyp', u diabetes', 'CHDRisk']

plt.figure(figsize=(15,10))
```

```
for i ,col in enumerate(disease_columns):
   gender_disease_count=df.groupby(['sex',col]).size().
 →reset_index(name='count')
     print(gender disease count)
   gender_total=df.groupby('sex').size().reset_index(name='total')
     print(gender total)
   merged_df=pd.merge(gender_disease_count,gender_total,on='sex')
    print(merged_df)
   merged_df['Percentage']=(merged_df['count'] / merged_df['total'])*100
     print(merged_df)
   print(f"\nPercentage of Males and Females with {col}:")
    #iterrows() function returns both the index and the row data.
   #If you are only interested in the row and do not need the index, you can
 →use _ to indicate that you are intentionally ignoring the index.
   for , row in merged df.iterrows():
       print(f"Gender: {row['sex']}, Disease Status: {row[col]}, Percentage:
 →{row['Percentage']:.2f}%")
   plt.subplot(3, 3, i+1)
   ax=sns.barplot(data=merged_df, x='sex', y='Percentage', hue=col)
   ax.bar label(ax.containers[0])# to show count outside plot
   ax.bar_label(ax.containers[1])
   ax.set_title(f'Percentage of Males and Females with {col}')
   ax.set ylabel('')
   ax.set_xlabel('')
plt.tight_layout()
plt.show()
```

```
Percentage of Males and Females with smokingStatus:
Gender: female, Disease Status: no, Percentage: 60.32%
Gender: female, Disease Status: yes, Percentage: 39.68%
Gender: male, Disease Status: no, Percentage: 39.42%
Gender: male, Disease Status: yes, Percentage: 60.58%

Percentage of Males and Females with BPMeds:
Gender: female, Disease Status: no, Percentage: 96.15%
Gender: female, Disease Status: yes, Percentage: 3.85%
Gender: male, Disease Status: no, Percentage: 98.01%
Gender: male, Disease Status: yes, Percentage: 1.99%
```

Percentage of Males and Females with prevalentStroke: Gender: female, Disease Status: no, Percentage: 99.41% Gender: female, Disease Status: yes, Percentage: 0.59% Gender: male, Disease Status: no, Percentage: 99.44% Gender: male, Disease Status: yes, Percentage: 0.56%

Percentage of Males and Females with prevalentHyp:

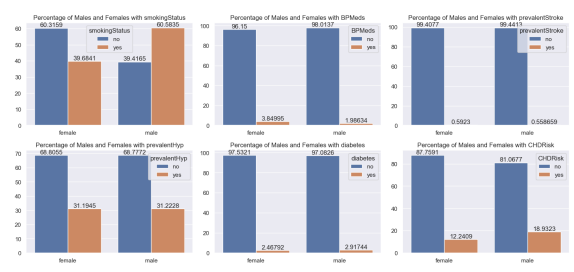
Gender: female, Disease Status: no, Percentage: 68.81% Gender: female, Disease Status: yes, Percentage: 31.19% Gender: male, Disease Status: no, Percentage: 68.78% Gender: male, Disease Status: yes, Percentage: 31.22%

Percentage of Males and Females with diabetes:

Gender: female, Disease Status: no, Percentage: 97.53% Gender: female, Disease Status: yes, Percentage: 2.47% Gender: male, Disease Status: no, Percentage: 97.08% Gender: male, Disease Status: yes, Percentage: 2.92%

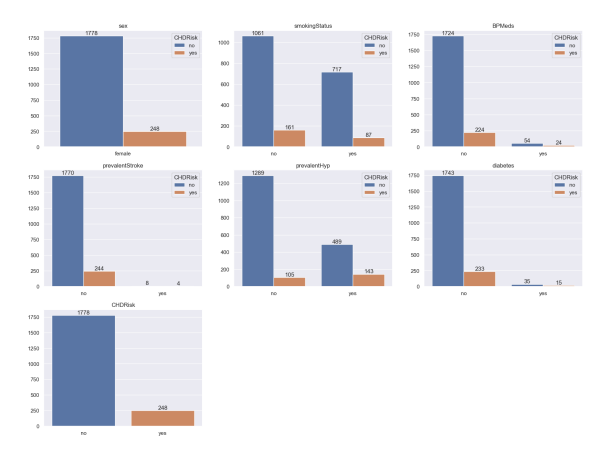
Percentage of Males and Females with CHDRisk:

Gender: female, Disease Status: no, Percentage: 87.76% Gender: female, Disease Status: yes, Percentage: 12.24% Gender: male, Disease Status: no, Percentage: 81.07% Gender: male, Disease Status: yes, Percentage: 18.93%



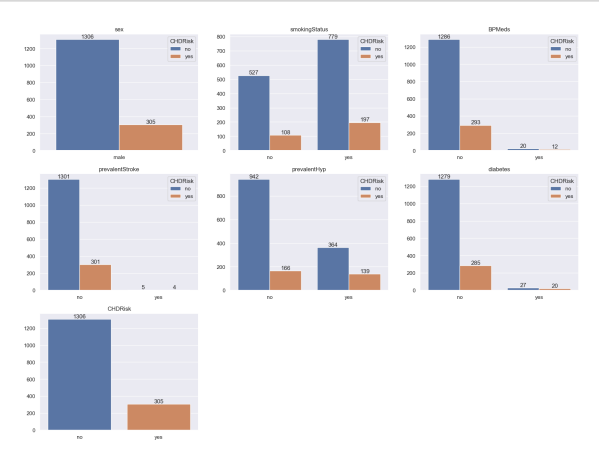
relationship between various health conditions and CHDRisk among females

```
plt.figure(figsize=(20,20))
for i ,col in enumerate(gender_female):
    plt.subplot(4,3,i+1)
    ax=sns.countplot(data=gender_female,x=col ,hue='CHDRisk')
    ax.bar_label(ax.containers[0])
    ax.bar_label(ax.containers[1])
    ax.set_title('\n\n' + col)
    ax.set_xlabel('')
    ax.set_ylabel('')
```



relationship between various health conditions and CHDRisk among males

```
plt.subplot(4,3,i+1)
ax=sns.countplot(data=gender_male,x=col ,hue='CHDRisk')
ax.bar_label(ax.containers[0])
ax.bar_label(ax.containers[1])
ax.set_title('\n\n' + col)
ax.set_xlabel('')
ax.set_ylabel('')
```



## 11 statistical tests

1-Chi-Square Test for Categorical Variables

• Chi-Square Test Use this to test for independence between two categorical variables.show if sex and CHDRisk are associated.

```
[83]: # Chi-square test
from scipy.stats import chi2_contingency
contingency = pd.crosstab(df['sex'], df['CHDRisk'])
chi2, p, dof, expected = chi2_contingency(contingency)
print(f"Chi-square: {chi2}, p-value: {p}")
```

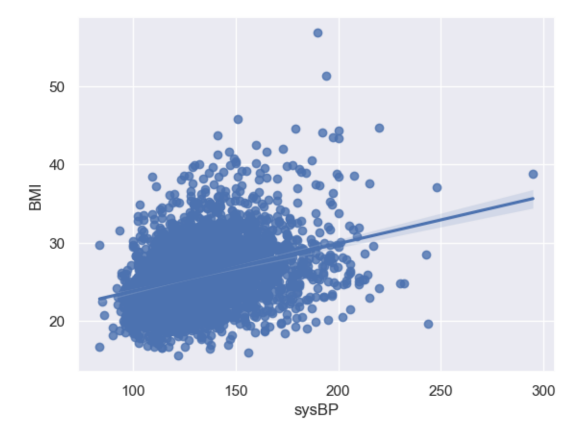
```
if p < 0.05:
    print("There is a significant relationship between age_group and CHDRisk")
else:
    print("No significant relationship between age_group and CHDRisk")</pre>
```

Chi-square: 30.64922901316927, p-value: 3.091494190770845e-08 There is a significant relationship between age\_group and CHDRisk

2-Pearson Correlation Coefficient: For testing the linear relationship between two continuous variables

```
[84]: from scipy.stats import pearsonr
    corr ,p_val =pearsonr(df['sysBP'],df['BMI'])
    print(f'Pearson Correlation={corr}, p={p}')
    if p < 0.05:
        print('There is a significant correlation between sysBP and BMI')
    else:
        print('No significant correlation between sysBP and BMI')
    sns.regplot(data=df,x='sysBP', y='BMI')
    plt.show()</pre>
```

Pearson Correlation=0.3292817576370576, p=3.091494190770845e-08 There is a significant correlation between sysBP and BMI



# 12 Logistic Regression Significance Test:

For testing whether predictor variables in a logistic regression are significant

```
[85]: from sklearn.preprocessing import LabelEncoder
      le=LabelEncoder()
      df['smokingStatus']=le.fit_transform(df['smokingStatus'])
      df['BPMeds']=le.fit_transform(df['BPMeds'])
      df['prevalentStroke'] = le.fit_transform(df['prevalentStroke'])
      df['prevalentHyp']=le.fit_transform(df['prevalentHyp'])
      df['diabetes']=le.fit_transform(df['diabetes'])
      df['age_group'] = le.fit_transform(df['age_group'])
[87]: df['sex']=df['sex'].replace('female',0)
      df['sex']=df['sex'].replace('male',1)
      df['CHDRisk']=df['CHDRisk'].replace('no',0)
      df['CHDRisk']=df['CHDRisk'].replace('yes',1)
      df.sample(5)
[87]:
                 education
                             smokingStatus
                                            cigsPerDay
                                                         BPMeds
                                                                prevalentStroke
            sex
      1229
              0
                          1
                                                              0
                                                      0
      1714
                                                      0
              0
                          1
                                         0
                                                              0
                                                                                0
      2579
                          2
                                         1
                                                     20
                                                              0
                                                                                0
      541
              1
                          3
                                         1
                                                      3
                                                              0
                                                                                0
      2314
              0
                          1
                                         0
                                                      0
                                                                               glucose \
            prevalentHyp diabetes
                                     totChol sysBP
                                                                   heartRate
                                                      diaBP
                                                              BMI
      1229
                                  0
                                         243 142.00 92.00 30.24
                                                                           70
                                                                                    85
                        1
      1714
                        0
                                  0
                                         193 134.00 88.00 25.77
                                                                           69
                                                                                    76
      2579
                        1
                                  0
                                         258 150.00 105.00 25.94
                                                                           83
                                                                                    60
      541
                                         200 105.00 68.00 23.30
                                  0
                                                                           65
                                                                                    68
      2314
                        0
                                         246 116.00 69.00 23.44
                                                                           65
                                                                                    78
            CHDRisk
                    age_group
      1229
                  0
                              1
                  0
      1714
                              1
      2579
                  0
                              1
      541
                  0
                              1
                              2
      2314
                  1
```

# [88]: import statsmodels.api as sm X=df.drop(columns=['CHDRisk']) X=sm.add\_constant(X) y=df['CHDRisk']

# [89]: logit\_model=sm.Logit(y,X)

# [90]: results=logit\_model.fit() print(results.summary())

Optimization terminated successfully.

Current function value: 0.381751

Iterations 7

### Logit Regression Results

Dep. Variable: Model: Method: Date: Time: converged: Covariance Type:	CHDRisk		Pseudo R-squ.: Log-Likelihood:		3637 3621 15 0.1044 -1388.4 -1550.3 6.669e-60
0.975]	coef	std err	z	P> z	[0.025
 const -4.757	-6.0047	0.636	-9.435	0.000	-7.252
sex 0.800	0.5872	0.108	5.416	0.000	0.375
education 0.017	-0.0796	0.050	-1.607	0.108	-0.177
smokingStatus	0.0172	0.157	0.109	0.913	-0.291
cigsPerDay 0.028	0.0161	0.006	2.584	0.010	0.004
BPMeds 0.613	0.1488	0.237	0.628	0.530	-0.316
prevalentStroke 1.794	0.8417	0.486	1.732	0.083	-0.111
prevalentHyp 0.522	0.2511	0.138	1.817	0.069	-0.020
diabetes 0.730	0.1092	0.317	0.345	0.730	-0.512
totChol	0.0030	0.001	2.720	0.007	0.001

0.005 sysBP	0.0201	0.004	5.346	0.000	0.013	
0.028 diaBP	-0.0092	0.006	-1.434	0.152	-0.022	
0.003						
BMI 0.033	0.0079	0.013	0.618	0.537	-0.017	
heartRate 0.003	-0.0048	0.004	-1.141	0.254	-0.013	
glucose 0.012	0.0073	0.002	3.243	0.001	0.003	
age_group 0.861	0.6689	0.098	6.823	0.000	0.477	

-----

===

if the p-value of a feature is less than 0.05, it means the feature is significant. . remove all features that p-value > 0.05

```
[91]: X.columns
```

```
[92]: X=df[['sex','cigsPerDay','totChol','glucose','sysBP','age_group']]
X=sm.add_constant(X)
y=df['CHDRisk']
logit_model=sm.Logit(y,X)
results=logit_model.fit()
print(results.summary())
```

Optimization terminated successfully.

Current function value: 0.383639

Iterations 7

Logit Regression Results

	======		======		======			
Dep. Variable	:		CHDI	Risk	No. Oh	servations:		3637
Model:			Lo	ogit	Df Res	siduals:		3630
Method:				MLE	Df Mod	del:		6
Date:		Tue,	10 Sep 2	2024	Pseudo	R-squ.:		0.09996
Time:			08:3	1:18	Log-Li	kelihood:		-1395.3
converged:				True	LL-Nu	11:		-1550.3
Covariance Ty	pe:		nonrol	oust	LLR p	-value:		6.124e-64
=========	=======	-===:		=====				
	coef		std err		Z	P> z	[0.025	0.975]
const	-7.1055	5	0.408	 -17	.414	0.000	-7.905	-6.306

```
0.5901
                             0.106
                                         5.567
                                                     0.000
                                                                  0.382
                                                                               0.798
sex
                0.0154
                             0.004
                                         3.756
                                                     0.000
                                                                  0.007
                                                                               0.023
cigsPerDay
totChol
                0.0030
                             0.001
                                         2.701
                                                     0.007
                                                                  0.001
                                                                               0.005
glucose
                0.0079
                             0.002
                                         4.602
                                                     0.000
                                                                  0.005
                                                                               0.011
sysBP
                                         9.669
                                                                               0.025
                0.0205
                             0.002
                                                     0.000
                                                                  0.016
                                                                               0.906
age_group
                0.7189
                             0.096
                                         7.510
                                                     0.000
                                                                  0.531
```

For each 1 unit increase in cigsPerDay, the odds of CHDRisk increase by 1.02 times, holding all else constant

For each 1 unit increase in totChol, the odds of CHDRisk increase by 1.0 times, holding all else constant

For each 1 unit increase in glucose, the odds of CHDRisk increase by 1.01 times, holding all else constant

For each 1 unit increase in sysBP, the odds of CHDRisk increase by 1.02 times, holding all else constant

For each 1 unit increase in age\_group, the odds of CHDRisk increase by 2.05 times, holding all else constant

#### 13 Prediction

```
[96]: X=df[['sex','cigsPerDay','totChol','glucose','sysBP','age_group']]
y=df['CHDRisk']

[97]: from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler
```

 $\label{eq:control_control_control} $$X_{\text{train},X_{\text{test},y_{\text{train},y_{\text{test}=\text{train}_{\text{test}_{\text{split}}}(X,y,\text{test}_{\text{size}=0}.)}$$$ $$\Rightarrow 2, \text{random}_{\text{state}=42,\text{stratify=y})$$$$ 

scaler=StandardScaler()

```
x_train_scaled=scaler.fit_transform(X_train)
x_test_scaled=scaler.transform(X_test)
```

```
[115]: from sklearn.svm import SVC
from sklearn.linear_model import LogisticRegression
from sklearn.ensemble import RandomForestClassifier
from sklearn.ensemble import GradientBoostingClassifier
from sklearn.neighbors import KNeighborsClassifier

models={
    'logistic_Model':LogisticRegression(),
    'random_model':RandomForestClassifier(),
    'gradient_model':GradientBoostingClassifier(),
    'knn':KNeighborsClassifier(),
    'svm_model':SVC()
}
```

# 14 GridSearch: to find the best hyperparameters

```
[99]: from sklearn.model_selection import GridSearchCV
      # Define the parameter grids for each model
      param_grids = {
          "logistic_Model": {
              'C': [0.01, 0.1, 1, 10, 100],
              'penalty': ['11', '12'],
              'solver': ['liblinear']
          },
          "random model": {
              'n_estimators': [50, 100, 200],
              'max_depth': [None, 10, 20, 30],
              'min_samples_split': [2, 5, 10]
          },
          "gradient_model": {
              'n_estimators': [50, 100, 200],
              'learning_rate': [0.01, 0.1, 0.2],
              'max_depth': [3, 5, 7]
          },
          "knn": {
              'n_neighbors': [3, 5, 7, 9],
              'weights': ['uniform', 'distance']
```

```
"svm_model": {
        'C': [0.1, 1, 10, 100],
        'kernel': ['linear', 'rbf'],
        'gamma': ['scale', 'auto']
    }
}
# Function to perform GridSearchCV
def perform_grid_search(model, param_grid):
    grid search = GridSearchCV(model, param grid, cv=5, scoring='accuracy', |
 \rightarrown jobs=-1)
    grid_search.fit(x_train_scaled, y_train)
    return grid_search
# GridSearchCV for each model
for name, model in models.items():
    print(f"Performing Grid Search for {name}...")
    grid_search = perform_grid_search(model, param_grids[name])
    print(f"Best parameters for {name}: {grid_search.best_params_}")
    print(f"Best accuracy for {name}: {grid search.best score }")
    print("-" * 50)
Performing Grid Search for logistic_Model...
Best parameters for logistic Model: {'C': 0.01, 'penalty': '12', 'solver':
'liblinear'}
Best accuracy for logistic_Model: 0.8514937511459681
Performing Grid Search for random model...
Best parameters for random_model: {'max_depth': 10, 'min_samples_split': 5,
'n estimators': 50}
Best accuracy for random_model: 0.8490864784617115
_____
Performing Grid Search for gradient_model...
Best parameters for gradient model: {'learning rate': 0.01, 'max_depth': 5,
'n_estimators': 50}
Best accuracy for gradient_model: 0.8484015591083036
_____
Performing Grid Search for knn...
Best parameters for knn: {'n_neighbors': 9, 'weights': 'uniform'}
Best accuracy for knn: 0.8470258057265883
Performing Grid Search for svm_model...
Best parameters for svm_model: {'C': 1, 'gamma': 'auto', 'kernel': 'rbf'}
Best accuracy for svm_model: 0.8490888443316713
```

#### 15 Data is Imbalanced:

weighted avg

0.82

0.85

```
[116]: from sklearn.metrics import confusion_matrix
       model=SVC()
       model.fit(x_train_scaled,y_train)
       y_pred=model.predict(x_test_scaled)
       print("Confusion Matrix:")
       print(confusion_matrix(y_test, y_pred))
      Confusion Matrix:
      [[617
               07
       Γ108
               3]]
      Total Negative Cases (TN + FP): 617 + 0 = 617 Total Positive Cases (FN + TP): 108 + 3 = 111.
      The imbalance is evident because the number of samples in the majority class (no CHD risk) is
      much larger than the number of samples in the minority class (CHD risk). Specifically, there are
      617 instances of the majority class compared to only 111 instances of the minority class.
      So ,we will use classification report not Confision Matrix
[117]: print(df['CHDRisk'].value_counts())
      CHDRisk
      0
            3084
      1
             553
      Name: count, dtype: int64
      it is also here number of 0 class > number of 1 class
[130]: from sklearn.metrics import accuracy_score, classification_report
       for name , model in models.items():
           model.fit(x_train_scaled,y_train)
           y_pred=model.predict(x_test_scaled)
           print(f"{name}:")
           print(f"Accuracy: {accuracy_score(y_test, y_pred)}")
           print(classification_report(y_test, y_pred))
           print("-" * 50)
      logistic_Model:
      Accuracy: 0.853021978021978
                     precision
                                    recall f1-score
                                                        support
                  0
                           0.86
                                      0.99
                                                 0.92
                                                             617
                  1
                           0.64
                                      0.08
                                                 0.14
                                                             111
                                                 0.85
                                                             728
           accuracy
                           0.75
                                      0.54
                                                 0.53
                                                             728
         macro avg
```

0.80

728

-----

	random	model	:
--	--------	-------	---

Accuracy: 0.8557692307692307

	precision	recall	f1-score	support
0	0.86	0.98	0.92	617
1	0.62	0.14	0.23	111
accuracy			0.86	728
macro avg	0.74	0.56	0.58	728
weighted avg	0.83	0.86	0.82	728

-----

gradient\_model:

Accuracy: 0.8502747252747253

	precision	recall	f1-score	support
0	0.86	0.99	0.92	617
1	0.56	0.08	0.14	111
accuracy			0.85	728
macro avg	0.71	0.53	0.53	728
weighted avg	0.81	0.85	0.80	728

\_\_\_\_\_

knn:

Accuracy: 0.8543956043956044

	precision	recall	f1-score	support
0	0.87	0.98	0.92	617
1	0.57	0.18	0.27	111
accuracy			0.85	728
macro avg	0.72	0.58	0.60	728
weighted avg	0.82	0.85	0.82	728

\_\_\_\_\_

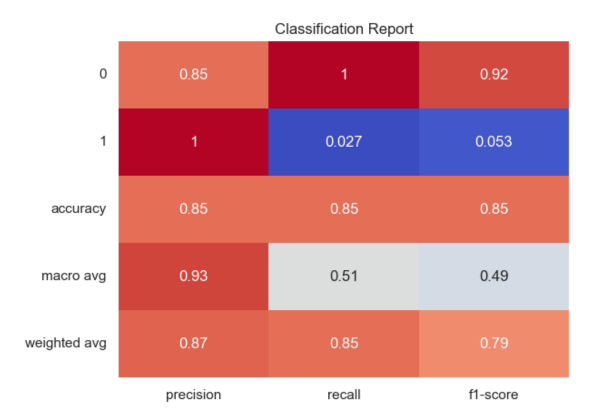
svm\_model:

Accuracy: 0.8516483516483516

	precision	recall	f1-score	support
0 1	0.85 1.00	1.00	0.92 0.05	617 111
accuracy macro avg	0.93	0.51	0.85 0.49	728 728
weighted avg	0.87	0.85	0.79	728

-----

[131]: Text(0.5, 1.0, 'Classification Report')



performance metrics for Logistic Regression model:

#### 15.0.1 Model Overview:

- Accuracy: 0.853 (85.3%)
  - Accuracy is the proportion of correctly predicted cases out of all the cases. In this case, the model correctly predicted **85.3%** of the instances in the dataset.

#### 15.0.2 Class 0 (No heart disease):

• Precision: 0.86 (86%)

Out of all the instances where the model predicted Class 0 (no heart disease), 86% of the predictions were correct.

Example: If the model predicted 100 people as not having heart disease (Class 0), 86 of them actually do not have heart disease.

#### • Recall: 0.99 (99%)

- Out of all the people who actually **do not have heart disease (Class 0)**, the model correctly identified **99**% of them.

Example: If there were 100 people who actually didn't have heart disease, the model correctly identified 99 of them.

#### • F1-score: 0.92 (92%)

- The F1-score is a balance between precision and recall. For Class 0, the F1-score is **92**%, indicating the model is performing very well in predicting **no heart disease**.

#### • Support: 617

This is the actual number of instances in the test set for Class 0. In this case, there are
 617 people who do not have heart disease.

#### 15.0.3 Class 1 (Heart disease):

#### • Precision: 0.64 (64%)

 Out of all the instances where the model predicted Class 1 (heart disease), 64% of the predictions were correct.

Example: If the model predicted 100 people as having heart disease, only 64 of them actually had heart disease.

#### • Recall: 0.08 (8%)

Out of all the people who actually have heart disease (Class 1), the model correctly identified only 8% of them.

Example: If there were 100 people who actually had heart disease, the model only correctly identified 8 of them.

#### • F1-score: 0.14 (14%)

- The F1-score is very low for **Class 1** (only **14%**). This indicates that while the model has some success in predicting people with heart disease (as seen in the 64% precision), it misses most of them (low recall of 8%).

#### • Support: 111

This is the actual number of instances in the test set for Class 1 (people with heart disease). There are 111 people in this category.

#### 15.0.4 Key Takeaways:

- The model performs **very well** in predicting **Class 0** (people who don't have heart disease), with high precision (86%) and excellent recall (99%).
- However, the model performs **poorly** in predicting **Class 1** (people who do have heart disease). The precision (64%) indicates that when it predicts heart disease, it's somewhat correct, but the recall (8%) shows that it **misses the majority** of people who actually have heart disease.

```
[120]: res = pd.DataFrame({'Actual': y_test, 'Predicted': y_pred})
print(res)
```

	Actual	Predicted
1519	0	0
1690	0	0
1703	0	0
2268	1	0
524	1	1
•••	•••	•••
118	0	0
1540	1	0
484	0	0
982	0	0
784	1	0

[728 rows x 2 columns]

# 16 Compare the ROC Curves of two Models.

```
print(f'AUC: {roc_auc_1}')

roc_auc_2 = roc_auc_score(y_test, y_prob_model_2)
print(f'AUC: {roc_auc_2}')

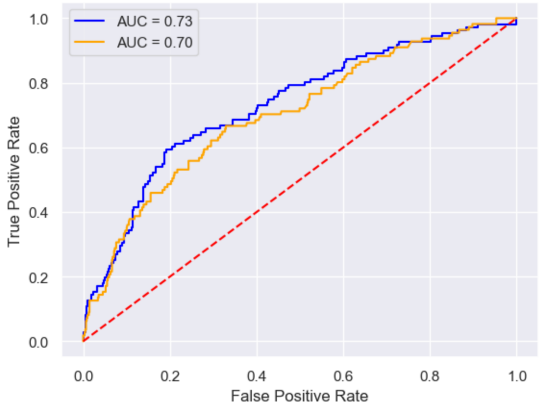
# Plot ROC curve
fpr1, tpr1, thresholds1 = roc_curve(y_test, y_prob_model_1)
fpr2, tpr2, thresholds2 = roc_curve(y_test, y_prob_model_2)

plt.plot(fpr1, tpr1, color='blue', label=f'AUC = {roc_auc_1:.2f}')
plt.plot(fpr2, tpr2, color='orange', label=f'AUC = {roc_auc_2:.2f}')

plt.plot([0, 1], [0, 1], color='red', linestyle='--')
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('Receiver Operating Characteristic (ROC) Curve')
plt.legend()
plt.show()
```

AUC: 0.7311752595383065 AUC: 0.7024106764787479





LogisticRegression is better than gradient

What is an ROC Curve?

The ROC (Receiver Operating Characteristic) Curve is a graphical representation of the performance of a classification model at different threshold values. It plots two metrics:

True Positive Rate (Recall) on the y-axis:

This measures how many actual positives (True Positives) are correctly identified.

<i>True Positive Rate (TPR) = True Positives / (True Positives + False Negatives)</i>

False Positive Rate on the x-axis:

This measures how many actual negatives are incorrectly classified as positives.

<i>False Positive Rate (FPR) = False Positives / (False Positives + True Negatives)</i>
Key Points:

The closer the ROC curve is to the top-left corner, the better the model's performance.

A random classifier would produce a diagonal line (FPR = TPR), representing an AUC of 0.5.

In summary, the ROC curve helps visualize the trade-off between the True Positive Rate and the False Positive Rate, while the AUC quantifies the overall performance of the model. A higher AUC means a better performing model.

Interpretation of AUC (Area Under the Curve):

AUC = 1.0: The model is perfect, distinguishing between classes with 100% accuracy.

- 0.9 AUC < 1.0: The model has excellent performance, meaning it is highly accurate in distinguishing between the classes.
- 0.8 AUC < 0.9: The model has good performance, but not perfect.
- 0.7 AUC < 0.8: The model has fair performance. It can distinguish between the classes better than random guessing, but there's room for improvement.
- 0.5 AUC < 0.7: The model has poor performance, only slightly better than random guessing.
- AUC = 0.5: The model has no discriminatory power and is equivalent to random guessing.

[]: