Einleitung

Der vorliegende Report dient zur Beschreibung des Projekts im Rahmen der Data Exploration Vorlesung. Die Abgabe umfasst ein GitHub Repositpory mit dem erstellten Code und dem Report in Form eines Jupyter Notebooks und einer PDF-Datei. Ziel dieses Projekts ist es anhand des, im Folgenden beschriebenen Datensatzes, eine exlorative Datenanalyse zu betreiben und ein Machine Learning Modell zu entwickeln, das zuverlässige Ergebnisse für eine binäre Klassifikation liefert.

Insalliere Requirements

```
In [ ]: ! pip install -r requirements.txt
```

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Requirement already satisfied: appnope==0.1.4 in ./venv/lib/python3.9/site
-packages (from -r requirements.txt (line 1)) (0.1.4)
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Requirement already satisfied: zipp==3.17.0 in ./venv/lib/python3.9/site-p
ackages (from -r requirements.txt (line 50)) (3.17.0)
```

In []: # importing all required libraries import pandas as pd import seaborn as sns import matplotlib.pyplot as plt import numpy as np # machine learning libraries from sklearn.ensemble import RandomForestClassifier from sklearn.linear_model import LogisticRegression from sklearn.metrics import accuracy_score, confusion_matrix, classificat from sklearn.model_selection import GridSearchCV from sklearn.metrics import roc_auc_score from sklearn.metrics import roc_curve from sklearn.svm import SVC from sklearn.model_selection import train_test_split # dummy classifier from sklearn.dummy import DummyClassifier

%matplotlib inline

Data Quality Check & Data Characterization

Die verwendeten Daten

Bei den verwendeten Daten handelt es sich um einen Kaggle Datensatz (https://www.kaggle.com/datasets/fedesoriano/heart-failure-prediction/data; letzter Abruf: 04.04.2024). Der Datensatz enthält Informationen von 918 Patienten und umfasst zwölf verschiedene Merkmale, darunter demografische Angaben wie Alter und Geschlecht, klinische Messungen wie Ruheblutdruck und maximale Herzfrequenz, sowie Informationen zu Symptomen wie Brustschmerzen und zuvor diagnostizierten Herzkrankheiten. Die Daten widerspiegeln auch medizinischen Tests wie Ruhe-Elektrokardiogrammen und Belastungsuntersuchungen.

Die Spalte 'HeartDisease' nimmt Werte von {0,1} an was für {normal, erkrankt} steht.

```
In []: # path in which the data is stored
data = "data/heart.csv"

In []: # reading the data
df = pd.read_csv(data)
df
```

		Age	Sex	ChestPainType	RestingBP	Cholesterol	FastingBS	RestingECG	Mi
	0	40	М	ATA	140	289	0	Normal	
	1	49	F	NAP	160	180	0	Normal	
	2	37	М	ATA	130	283	0	ST	
	3	48	F	ASY	138	214	0	Normal	
	4	54	М	NAP	150	195	0	Normal	
	•••		•••						
9	913	45	М	TA	110	264	0	Normal	
9	914	68	М	ASY	144	193	1	Normal	
•	915	57	М	ASY	130	131	0	Normal	
9	916	57	F	ATA	130	236	0	LVH	
,	917	38	М	NAP	138	175	0	Normal	

918 rows × 12 columns

Out[]:

Beschreibung der Attribute:

- Age: Alter des Patienten [Jahre]
- Sex: Geschlecht des Patienten [M: Männlich, F: Weiblich]
- ChestPainType: Brustschmerztyp [TA: Typische Angina, ATA: Atypische Angina, NAP: Nicht-Anginaler Schmerz, ASY: Asymptomatisch]
- RestingBP: Ruheblutdruck [mm Hg]
- Cholesterol: Serumcholesterin [mm/dl]
- FastingBS: Nüchternblutzucker [1: Wenn Nüchternblutzucker > 120 mg/dl, 0: Ansonsten]
- RestingECG: Ruheelektrokardiogrammergebnisse [Normal: Normal, ST: Mit ST-T-Wellen-Abnormalitäten (T-Wellen-Inversionen und/oder ST-Hebungen oder Senkungen von > 0,05 mV), LVH: Zeigt wahrscheinliche oder definitive linksventrikuläre Hypertrophie nach Estes-Kriterien]
- MaxHR: Maximale erreichte Herzfrequenz [Numerischer Wert zwischen 60 und 202]
- ExerciseAngina: Belastungsinduzierte Angina [J: Ja, N: Nein]
- Oldpeak: ST-Depression = ST [Numerischer Wert gemessen in Depression]
- ST_Slope: Die Steigung des Spitzen-Belastungs-ST-Segments [Up: Aufsteigend, Flat: Flach, Down: Absteigend]
- HeartDisease: Ausgabeklasse [1: Herzkrankheit, 0: Normal]

```
In []: # using the pandas method "describe()"" to get a describtion of the datas
# ".T" transposes the dataframe (rows and columns are switched)
df.describe().T
```

Out[]:		count	mean	std	min	25%	50%	75%	max
	Age	918.0	53.510893	9.432617	28.0	47.00	54.0	60.0	77.0
	RestingBP	918.0	132.396514	18.514154	0.0	120.00	130.0	140.0	200.0
	Cholesterol	918.0	198.799564	109.384145	0.0	173.25	223.0	267.0	603.0
	FastingBS	918.0	0.233115	0.423046	0.0	0.00	0.0	0.0	1.0
	MaxHR	918.0	136.809368	25.460334	60.0	120.00	138.0	156.0	202.0
	Oldpeak	918.0	0.887364	1.066570	-2.6	0.00	0.6	1.5	6.2
	HeartDisease	918.0	0.553377	0.497414	0.0	0.00	1.0	1.0	1.0

Bereits nachdem man sich die Beschreibung des Datensatzes anschaut, kann man feststellen, dass die SPalten "Cholesterol" und "RestingBP" unerwartete minimal Werte aufweisen (ruhe Puls und Cholisterinspiegel können keine Werte von 0 annehmen).

```
In []: # count null values
    null_values_count = (df['RestingBP'] == 0).sum()
    print("Anzahl der Nullwerte in der Spalte 'RestingBP':", null_values_coun
    Anzahl der Nullwerte in der Spalte 'RestingBP': 1
In []: # delete the only patient with the null value in RestingBP
    df = df[df['RestingBP'] != 0]
```

> Da es nur bei einem Patienten eine vermutliche Fehlmessung gab, wird dieser Patient aus dem Datensatz gelöscht.

```
In [ ]: # count null values
        null_values_count = (df['Cholesterol'] == 0).sum()
        print("Anzahl der Nullwerte in der Spalte 'Cholesterol':", null_values_co
```

Anzahl der Nullwerte in der Spalte 'Cholesterol': 171

Leider weisen dennoch 171 Patienten bei Cholesterol den Wert 0 auf. Dies war bei der initialen explorativen Datenanalyse nicht auf den ersten Blick ersichtlich. Da das löschen von 171 Einträgen problematisch ist, wird in den fehldenen Stellen der durschnittliche Cholesterol Wert des Datensatzes eingesetzt. Somit sollen erheblichere Verfälschungen im Machine Learning Model im nachhinein vermieden werden. Ein Modell, das mit Daten von Patienten mit einem Cholesterol Wert von 0 trainiert ist, ist in der Realität nicht nützlich.

```
In [ ]: | # we don't wont the 0 values, when calculating the mean value
        df_cleaned = df[df['Cholesterol'] != 0]
        # calculate mean value
        average_chol = round(df_cleaned['Cholesterol'].mean())
        print("Durchschnittlicher Cholesterinspiegel nach Entfernen von Nullwerte
```

Durchschnittlicher Cholesterinspiegel nach Entfernen von Nullwerten (ohne

```
Nachkommastellen): 245
In []: # replace 0 values with the mean value
        df.loc[df['Cholesterol'] == 0, 'Cholesterol'] = average_chol
In []: # check if the anomaly still exists
        df["Cholesterol"].min()
Out[]: 85
In [ ]: # checking for missung values in the dataframe
        missing_values = df.isnull().sum()
        missing_values
Out[]: Age
                           0
        Sex
                           0
        ChestPainType
        RestingBP
                           0
        Cholesterol
                           0
                           0
        FastingBS
        RestingECG
                           0
                           0
        MaxHR
        ExerciseAngina
                           0
                           0
        0ldpeak
        ST_Slope
                           0
        HeartDisease
                           0
        dtype: int64
In [ ]: # checking for duplicated rows in the dataframe
```

127.0.0.1:5500/analysis.html 6/51

duplicates = df.duplicated().sum()

df.max()

duplicates Out[]: 0 In []: # determining unique values of categorial columns in the dataframe categorical_columns = ['Sex', 'ChestPainType', 'RestingECG', 'ExerciseAng for col in categorical_columns: unique_values = df[col].unique() print(f"Eindeutige Werte für {col}:") print(unique values) Eindeutige Werte für Sex: ['M' 'F'] Eindeutige Werte für ChestPainType: ['ATA' 'NAP' 'ASY' 'TA'] Eindeutige Werte für RestingECG: ['Normal' 'ST' 'LVH'] Eindeutige Werte für ExerciseAngina: ['N' 'Y'] Eindeutige Werte für ST_Slope: ['Up' 'Flat' 'Down'] In []: # get dataframe info df.info() <class 'pandas.core.frame.DataFrame'> Index: 917 entries. 0 to 917 Data columns (total 12 columns): Non-Null Count Dtype # Column 917 non-null 0 Age int64 917 non-null 1 Sex object 2 ChestPainType 917 non-null object 3 RestingBP 917 non-null int64 4 Cholesterol 917 non-null int64 5 FastingBS 917 non-null int64 917 non-null 6 RestingECG object 7 917 non-null MaxHR int64 ExerciseAngina 917 non-null 8 object 917 non-null 9 Oldpeak float64 10 ST_Slope 917 non-null object 11 HeartDisease 917 non-null int64 dtypes: float64(1), int64(6), object(5) memory usage: 93.1+ KB In []: # getting the highest values of each column (categorial columns may be ig

```
77
Out[]: Age
         Sex
                             Μ
                            TΑ
         ChestPainType
         RestingBP
                            200
                            603
         Cholesterol
         FastingBS
                             1
         RestingECG
                            ST
         MaxHR
                            202
         ExerciseAngina
                             Υ
         0ldpeak
                            6.2
         ST_Slope
                            Up
         HeartDisease
                              1
         dtype: object
In [ ]: # same goes for this but for minimal values
        df.min()
Out[]:
                              28
        Age
                              F
         Sex
         ChestPainType
                             ASY
         RestingBP
                             80
         Cholesterol
                             85
         FastingBS
                              0
                             LVH
         RestingECG
         MaxHR
                              60
         ExerciseAngina
                              Ν
         0ldpeak
                            -2.6
         ST Slope
                           Down
                              0
         HeartDisease
         dtype: object
In [ ]: | # check how many unique elements the dataset contains in each column
        df.nunique()
                             50
Out[]: Age
         Sex
                              2
                              4
         ChestPainType
         RestingBP
                             66
         Cholesterol
                            221
                              2
         FastingBS
                              3
         RestingECG
                            119
         MaxHR
         ExerciseAngina
                             2
                             53
         0ldpeak
         ST_Slope
                              3
                              2
         HeartDisease
         dtype: int64
```

Die Analyse zur Datenqualität liefert auf den ersten Blick, bis auf die 2 Anomalien, kaum Mängel, da es keine fehlenden Einträge oder duplizierte Zeilen gibt. Auch die Spalten mit den kategorischen Werten liefern saubere und "aufgeräumte" Werte. Der Datensatz ist im Allgemeinen sehr gut gepflegt.

Exploratory Data Analysis

Im Folgenden werden die Daten analysiert und statistische Verteilungen und Merkmale, sowie Anhängigkeiten zwischen verschiedenen Attributen werden grafisch aufgezeigt.

```
In []: # visualize disease distribution in the dataset

colors_red_green = ["#9aff9a", "#ff3030"]

sns.countplot(x='HeartDisease', data=df, palette=colors_red_green)

plt.xlabel('Heart Disease')
plt.ylabel('Count')
plt.title('Distribution of Heart Disease')

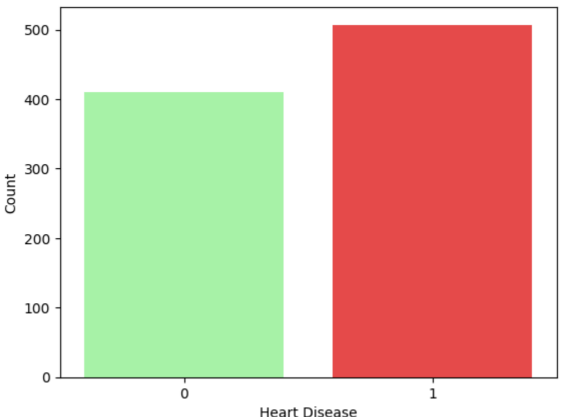
plt.show()
```

/var/folders/3l/_xvv3581559_krvl1r82px5w0000gn/T/ipykernel_18016/182032164 9.py:5: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be remove d in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.countplot(x='HeartDisease', data=df, palette=colors_red_green)





```
In [ ]: heart_disease_distribution = df['HeartDisease'].value_counts()
heart_disease_distribution
```

```
Out[]: HeartDisease

1 507

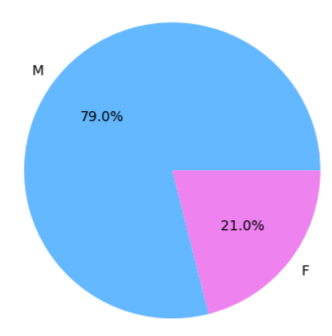
0 410

Name: count, dtype: int64
```

Die erste Visualisierung zeigt die Verteilung zwischen gesunden und kranken Patienten. Es ist eine leichte Inbalance der Werte vorhanden. Nach Absprache mit dem Dozenten kann diese Aufgrund ihrer leichten Ausprägung in diesem Fall ignoriert werden. Der Datensatz enthält 97 mehr betroffene als gesunde Patienten.xy

```
In []: # pie chart for sex distribution
    distribution = df["Sex"].value_counts()
    colors = ['#63b8ff', '#ee82ee']
    plt.title("Distribution of Sex")
    plt.pie(distribution, labels=distribution.index, colors=colors, autopct='
    plt.show()
```

Distribution of Sex



Dieses Kuchendiagramm zeigt die Verteilung der Geschlächter in den Daten. 79% der Patienten sind männlich und 21% sind weiblich.

```
In []: # create boxplots to display age distribution
    fig = plt.figure(figsize=(12, 6))
    gs = fig.add_gridspec(1, 3, width_ratios=[2, 1, 1])

# total age distribution
    ax1 = fig.add_subplot(gs[0])
    sns.boxplot(x=df["Age"], ax=ax1, color='#5c5c5c')
    ax1.set_title('total age distribution')

# female age distribution
```

```
ax2 = fig.add_subplot(gs[1])
sns.boxplot(x='Sex', y='Age', data=df[df['Sex'] == 'F'], ax=ax2, palette=
ax2.set_title('female age distribution')

# male age distribution
ax3 = fig.add_subplot(gs[2])
sns.boxplot(x='Sex', y='Age', data=df[df['Sex'] == 'M'], ax=ax3, palette=
ax3.set_title('male age distribution')

plt.tight_layout()
plt.show()
```

/var/folders/3l/_xvv3581559_krvl1r82px5w0000gn/T/ipykernel_18016/293609210 3.py:12: FutureWarning:

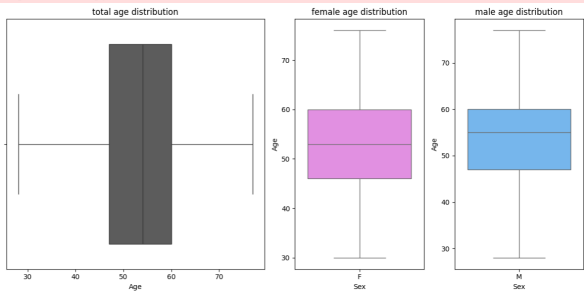
Passing `palette` without assigning `hue` is deprecated and will be remove d in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='Sex', y='Age', data=df[df['Sex'] == 'F'], ax=ax2, palette
=['#ee82ee'])

/var/folders/3l/_xvv3581559_krvl1r82px5w0000gn/T/ipykernel_18016/293609210 3.py:17: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be remove d in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='Sex', y='Age', data=df[df['Sex'] == 'M'], ax=ax3, palette
=['#63b8ff'])



Diese Boxplots zeigen die gesamte Altersverteilung sowie die Verteilung pro Geschlecht. Die Altersspanne liegt im Durschschnitt zwischen 48 und 60 Jahren.

```
In []: # print value_counts of age to identify outliers
    counts = df["Age"].value_counts()
    print("Counts for Age:")
    print(counts)
```

```
Counts for Age:
Age
54
      51
58
       42
55
      40
56
       38
57
       38
52
      36
51
      35
59
       35
62
      35
53
      33
      32
60
48
       31
61
      31
63
      30
50
      25
46
      24
41
      24
43
      24
64
      22
65
      21
49
      21
47
      19
44
       19
42
       18
45
      18
38
       16
67
       15
39
      15
66
      13
69
       13
40
       13
35
      11
37
       11
68
       10
34
        7
74
        7
70
        7
36
        6
        5
71
32
        5
72
        4
29
        3
        3
75
        2
33
        2
77
        2
76
        2
31
30
        1
28
        1
73
        1
```

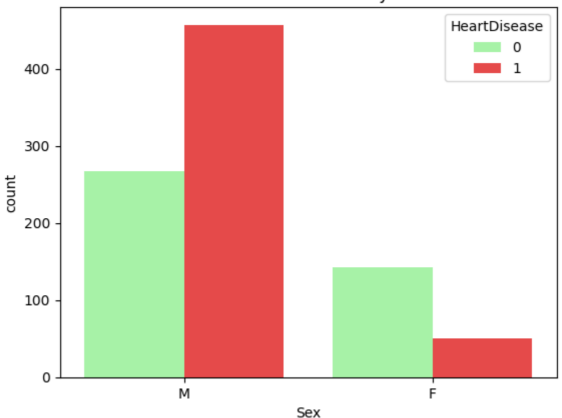
Name: count, dtype: int64

Es gibt auch Patienten die "sehr" jung oder alt sind. Der jüngste Patient ist 28 und der älteste ist 77. Es kommen allerdings wenige Personen in diesem Datensatz vor, die an diese Altersgrenzen stoßen.

Im folgenden werden die verschiedenen kategorischen Attribute je nach Haufigkeit der Erkrankungen dargestellt.

```
In []: # create countplot to display HeartDisease distribution by sex
sns.countplot(x='Sex', hue='HeartDisease', data=df, palette=['#9aff9a', '
    plt.xlabel("Sex")
    plt.title("Heart Disease Count by Sex")
    plt.show()
```





In diesem Datensatz gibt es innerhalt der männlichen Patientengruppe deutlich mehr Herzerkrankte, während es bei der weiblichen Gruppe weniger Betroffene gibt. Man beachte, dass der Datensatz mehr männliche Einträge enthält als weibliche.

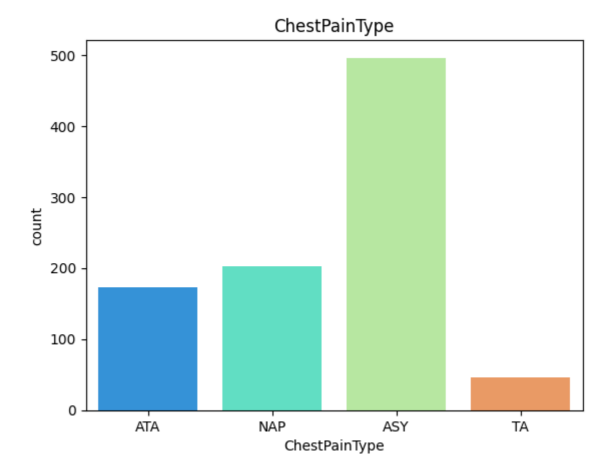
```
In []: # countplot to display ChestPainType distribution
    sns.countplot(x=df['ChestPainType'], palette="rainbow")
    plt.title('ChestPainType')

/var/folders/3l/_xvv3581559_krvl1r82px5w0000gn/T/ipykernel_18016/79895086
    1.py:2: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be remove d in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

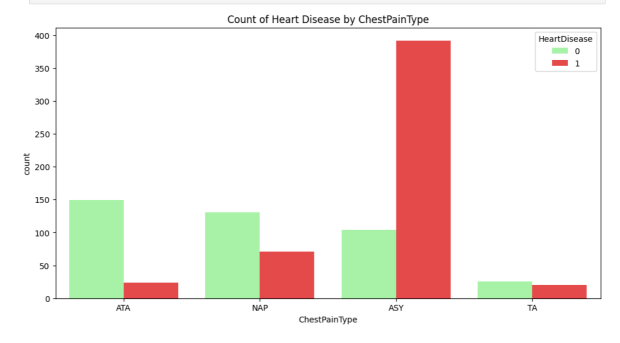
    sns.countplot(x=df['ChestPainType'], palette="rainbow")

Out[]: Text(0.5, 1.0, 'ChestPainType')
```



Die häufigste Ausprägung bei den Brustschmerzen sind die asymptomatischen Brustschmerzen. Die wenigsten Fälle beschreiben typical angina chest pain.

```
In []: # create a countplot showing the distribution of heart disease by ChestPa
    plt.figure(figsize=(12, 6))
    sns.countplot(x='ChestPainType', hue='HeartDisease', data=df, palette=col
    plt.title('Count of Heart Disease by ChestPainType')
    plt.show()
```



Interessanterweise zeigen Patienten mit asystomatischen Brustschmerzen am häufigsten eine Herzkrankheit auf. Bei der Gruppe TA gibt es in etwa gleich viele

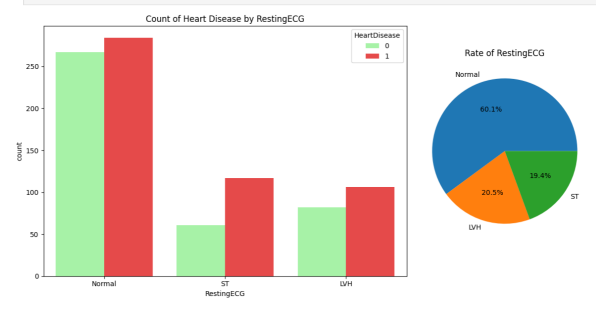
Gesunde wie Erkrankte. In den anderen beiden Gruppen überwiegt die Anzahl der gesunden Patienten.

```
In []: # create a figure with a grid of 1 row and 2 columns, with the second col
    fig = plt.figure(figsize=(12, 6))
    gs = fig.add_gridspec(1, 2, width_ratios=[2, 1])

# subplot 1: Countplot showing the distribution of heart disease by Resti
    ax1 = fig.add_subplot(gs[0])
    sns.countplot(x='RestingECG', hue='HeartDisease', data=df, palette=colors
    ax1.set_title('Count of Heart Disease by RestingECG')

# subplot 2: Pie chart illustrating the distribution of RestingECG values
    ax2 = fig.add_subplot(gs[1])
    types = df['RestingECG'].value_counts()
    ax2.pie(types, labels=types.index, autopct='%1.1f%*')
    ax2.set_title('Rate of RestingECG')

plt.tight_layout()
    plt.show()
```



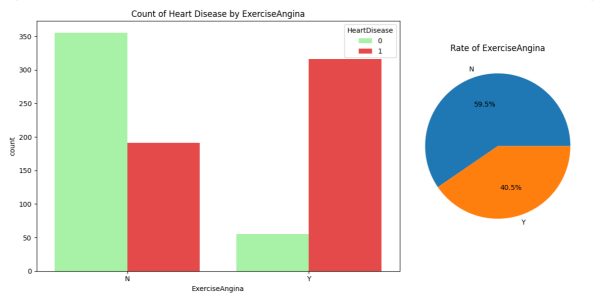
Die Ruheelektrokardiogrammergebnisse zeigen überwiegend normale Werte (60%). Die Gruppen LVH und ST (Beschreibung siehe oben) sind mit jeweils annähernd 20% seltener vertreten. Pro Gruppe gibt es allerdings stets mehr erkrankte als gesunde Patienten.

```
In []: # create a figure with a grid of 1 row and 2 columns, with the second col
    fig = plt.figure(figsize=(12, 6))
    gs = fig.add_gridspec(1, 2, width_ratios=[2, 1])

# subplot 1: Countplot showing the distribution of heart disease by Exerc
    ax1 = fig.add_subplot(gs[0])
    sns.countplot(x='ExerciseAngina', hue='HeartDisease', data=df, palette=co
    ax1.set_title('Count of Heart Disease by ExerciseAngina')

# subplot 2: Pie chart illustrating the distribution of ExerciseAngina in
    ax2 = fig.add_subplot(gs[1])
    types = df['ExerciseAngina'].value_counts()
    ax2.pie(types, labels=types.index, autopct='%1.1f%%')
    ax2.set_title('Rate of ExerciseAngina')
```

```
plt.tight_layout()
plt.show()
```



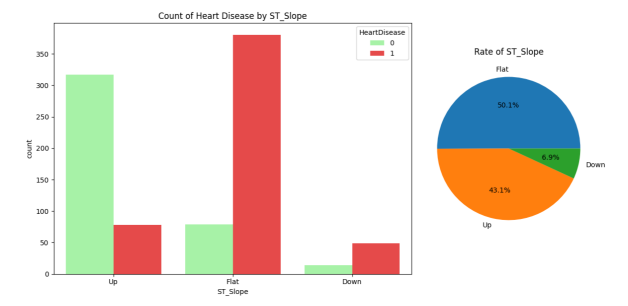
Die Mehrheit im Datensatz besitzt keine Belastungsinduzierte Brustschmerzen. Diejenigen Patienten die derartige Brustschmerzen aufweisen, haben jedoch signifikant öfter eine Herzerkranung als die andere Gruppe.

```
In []: # create a figure with a grid of 1 row and 2 columns, with the second col
    fig = plt.figure(figsize=(12, 6))
    gs = fig.add_gridspec(1, 2, width_ratios=[2, 1])

# subplot 1: Countplot showing the distribution of heart disease by ST_Sl
    ax1 = fig.add_subplot(gs[0])
    sns.countplot(x='ST_Slope', hue='HeartDisease', data=df, palette=colors_r
    ax1.set_title('Count of Heart Disease by ST_Slope')

# subplot 2: Pie chart illustrating the distribution of ST_Slope in the d
    ax2 = fig.add_subplot(gs[1])
    types = df['ST_Slope'].value_counts()
    ax2.pie(types, labels=types.index, autopct='%1.1f%*')
    ax2.set_title('Rate of ST_Slope')

plt.tight_layout()
    plt.show()
```



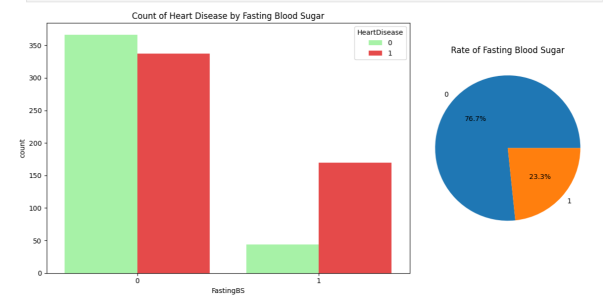
ST_Slope beschreibt die Steigung des peak exercise ST Segments. Wie man der Visualisierung entnehmen kann, gibt es überwiegend flache und und steigende ST Segmente. Bis auf den steigenden Segmenten gibt es in jeder Gruppe deutlich mehr Herzerkrankte. Wie es scheint, sind Patienten mit einer steigenden Kurve wahrscheinlicher gesund.

```
In []: fig = plt.figure(figsize=(12, 6))
    gs = fig.add_gridspec(1, 2, width_ratios=[2, 1])

# countplot for heart disease by fasting blood sugar
    ax1 = fig.add_subplot(gs[0])
    sns.countplot(x='FastingBS', hue='HeartDisease', data=df, palette=colors_ax1.set_title('Count of Heart Disease by Fasting Blood Sugar')

# pie chart for the distribution of fasting blood sugar
    ax2 = fig.add_subplot(gs[1])
    types = df['FastingBS'].value_counts()
    ax2.pie(types, labels=types.index, autopct='%1.1f%*')
    ax2.set_title('Rate of Fasting Blood Sugar')

plt.tight_layout()
    plt.show()
```

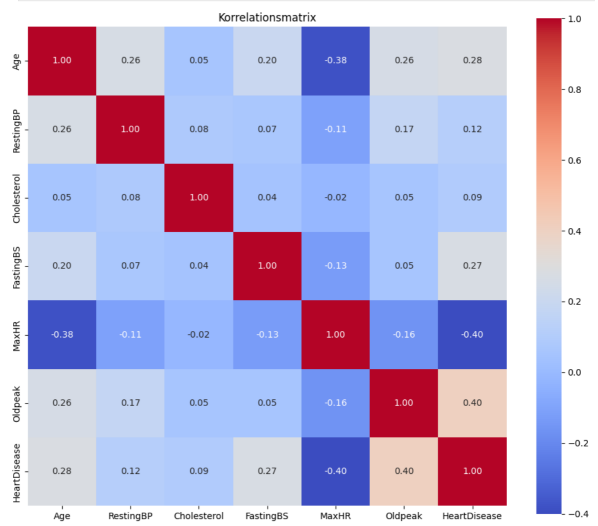


Das Attribut FastingBS beschreibt den nüchternen Blutzuckerspiegel eines Patienten, während Werte von 1 einen Blutzuckerspiegel von > 120 mg/dl kennzeichnen. Werte darunter sind mit 0 beschrieben. Der Großteil der Patienten fällt unter die Gruppe 0. In dieser Gruppe gibt es annähernd gleich viele Patienten mit, sowie ohne Krankheit. In der Gruppe mit dem höheren Blutzuckerspiegel haben weitaus mehr Patienten eine Herzkrankheit.

```
In []: # create corelation matrix

correlations = df.corr(numeric_only=True)

plt.figure(figsize=(12, 10))
sns.heatmap(correlations, annot=True, cmap='coolwarm', fmt=".2f", square=plt.title('Korrelationsmatrix')
plt.show()
```



Um die Korrelationen zwischen den einzelnen Attributen zu ermittlen, wird diese Korrelationsmatrix erstellt. Die stärkste Korrelation weisen die Attribute Oldpeak und Heartdisease auf. Das lässt darauf schließen, dass sich je nach Gruppe innerhalb des Attributs Oldpeak eine genauere Aussage über den Gesundheitszustands eines Patienten fallen lässt. Weitere, jedoch schwächere Korrelationen (>= 0.20) herrschen zwischen den Attributen Age und Heartdisease, Age und Oldpeak, RestingPB und Age, FastingBS und Age, MaxHR und Cholesterol. Auffällig ist die negative Korrelation zwsichen MaxHR und Heartdisease. Der Wert -0,40 besagt, dass ein Patient mit

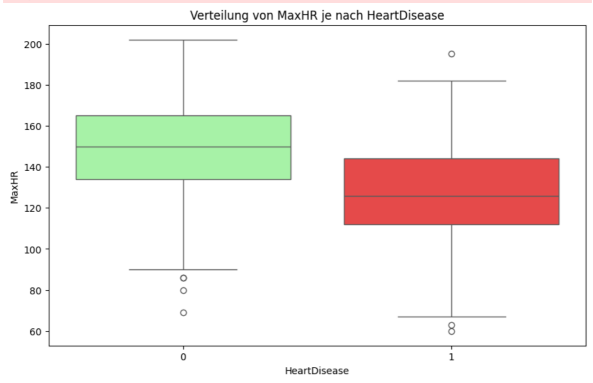
Herzerkrankung einen tendenziell niedrigeren Maximalen Puls hat. Dies erschien auf den ersten Blick merkwürdig, da die Annahme herrschte, Herzerkrankte menschen hätten einen höheren Puls.

```
In []: # create box plot
   plt.figure(figsize=(10, 6))
   sns.boxplot(x='HeartDisease', y='MaxHR', data=df, palette=colors_red_gree
   plt.title('Verteilung von MaxHR je nach HeartDisease')
   plt.xlabel('HeartDisease')
   plt.ylabel('MaxHR')
   plt.show()
```

/var/folders/31/_xvv3581559_krvl1r82px5w0000gn/T/ipykernel_18016/99511433 4.py:3: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be remove d in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='HeartDisease', y='MaxHR', data=df, palette=colors_red_gre
en)



Doch dieser Boxplot bestätigt den Wert in der Korrelationsmatrix.

Outlier Detection

In den nächsten Schritten werden die Outlier im Datensatz analysiert.

```
Detect outliers in the given DataFrame.
    Parameters:

    data (DataFrame): The DataFrame containing the data.

    Returns:
    - outliers (list): A list of indices corresponding to the outliers in
    outliers = []
    for col in data.columns:
        q1 = data[col].quantile(0.25)
        q3 = data[col].quantile(0.75)
        iqr = q3 - q1
        lower_bound = q1 - 1.5 * iqr
        upper_bound = q3 + 1.5 * iqr
        outlier_indices = data[(data[col] < lower_bound) | (data[col] > u
        outliers.extend(outlier_indices)
    return outliers
# find outliers
outliers_indices = detect_outliers(df[numeric_cols])
# remove duplicated indices
outliers_indices = list(set(outliers_indices))
print("Indices of outliers:", outliers_indices)
print("Outlier rows:")
print(df.iloc[outliers_indices])
```

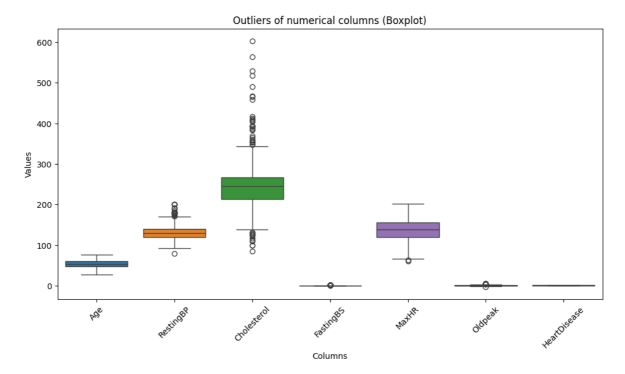
```
Indices of outliers: [515, 516, 518, 521, 522, 531, 532, 536, 537, 538, 2
8, 541, 30, 544, 546, 547, 36, 549, 550, 38, 553, 554, 556, 557, 559, 563,
52, 564, 58, 571, 826, 573, 574, 575, 577, 579, 580, 69, 582, 68, 584, 58
5, 76, 589, 78, 592, 593, 594, 595, 84, 86, 599, 604, 605, 606, 607, 97, 9
8, 610, 612, 613, 102, 103, 616, 105, 108, 109, 621, 624, 112, 117, 120, 6
32, 123, 639, 128, 132, 644, 650, 658, 659, 660, 149, 666, 667, 155, 160,
673, 672, 675, 165, 166, 679, 682, 686, 182, 185, 187, 189, 190, 702, 701,
718, 208, 210, 725, 728, 732, 734, 224, 738, 227, 744, 238, 752, 241, 242,
759, 247, 250, 256, 771, 774, 263, 775, 780, 782, 784, 785, 274, 275, 278,
790, 791, 793, 795, 284, 796, 799, 802, 803, 294, 295, 296, 297, 298, 299,
300, 809, 302, 303, 304, 305, 306, 308, 309, 820, 311, 312, 313, 314, 315,
316, 317, 318, 319, 320, 321, 322, 323, 324, 325, 838, 327, 328, 329, 330,
331, 842, 333, 334, 335, 843, 337, 338, 339, 340, 341, 342, 855, 343, 344,
850, 347, 349, 350, 869, 871, 872, 365, 880, 370, 372, 887, 888, 377, 378,
900, 389, 901, 390, 393, 394, 395, 396, 397, 398, 399, 400, 401, 402, 915,
403, 404, 405, 406, 407, 408, 409, 411, 410, 412, 413, 414, 415, 420, 422,
423, 424, 425, 430, 437, 441, 442, 443, 444, 448, 454, 457, 458, 460, 914,
469, 472, 473, 908, 475, 476, 477, 478, 480, 481, 482, 485, 486, 911, 491,
496, 498, 500, 503, 504, 505, 508, 511]
Outlier rows:
```

Age Sex ChestPainType RestingBP Cholesterol FastingBS RestingECG 516 68 NAP 195 1 Normal Μ 150 517 65 **ASY** 150 235 Normal Μ 0 519 63 Μ **ASY** 96 305 0 ST 0 522 50 Μ ASY 144 349 LVH 523 59 Μ ASY 124 160 0 Normal . 504 62 М **ASY** 210 1 Normal 158 505 55 Μ NAP 136 245 1 ST 506 75 **ASY** 136 225 0 Μ Normal 509 58 Μ ASY 110 198 0 Normal 35 0 ST 512 Μ NAP 123 161

	MaxHR	ExerciseAngina	0ldpeak	ST_Slope	HeartDisease
516	132	N	0.0	Flat	1
517	120	Υ	1.5	Flat	1
519	121	Υ	1.0	Up	1
522	120	Υ	1.0	Up	1
523	117	Υ	1.0	Flat	1
504	112	Υ	3.0	Down	1
505	131	Υ	1.2	Flat	1
506	112	Υ	3.0	Flat	1
509	110	N	0.0	Flat	1
512	153	N	-0.1	Up	0

[275 rows x 12 columns]

```
In []: # box plot to display outliers
   plt.figure(figsize=(12, 6))
   sns.boxplot(data=df[numeric_cols])
   plt.xticks(rotation=45)
   plt.title('Outliers of numerical columns (Boxplot)')
   plt.xlabel('Columns')
   plt.ylabel('Values')
   plt.show()
```



Die Outlier Detection liefert bis gute Ergebnisse. Dadurch, dass die Anomalien bei der Analyse der Datenqulität behoben worden, gibt es keine weiteren erheblichen Einschränkungen in den Daten. Während Serum Cholesterol Werte von > 600 äußert gefährlich erscheinen, sind diese in der Realität dennoch möglich.

Machine Learning

Der vorliegende Datensatz liefert ein binäres Klassifikationsproblem. Um einen ersten Ansatz für die Auswahl eines endgültigen Classifiers zu ermitteln, wurden im weitern Verlauf 3 verschiedene Classifier getestet.

Diese wären:

- Randomforest
- Logistic Regression
- Support Vector Machine

Die Auswahl dieser drei Classifie beruht auf ihrer Effektivität bei binären Klassifikationsproblemen. Jeder Algorithmus bietet spezifische Vorzüge und kann unterschiedliche Aspekte des Problems abdecken.

Metriken

Für die Bewertung des Models werden insbesondere die folgenden Metriken verwendet:

- Recall & Accuracy
- F1-Score
- ROC-AUC-Score

Der Recall bewertet die Fähigkeit des Modells, positive Instanzen korrekt zu identifizieren, was besonders wichtig ist, um sicherzustellen, dass keine relevanten Fälle übersehen werden.

Der F1-Score ist ein harmonisches Mittelmaß zwischen Precision und Recall und ermöglicht eine ausgewogene Bewertung von False Positives und False Negatives. Dieser ist in den meisten Fällen ein gutes Maß, da es häufig auf die Balance zwischen Recall und Precision ankommt.

Der ROC-AUC-Score bewertet die Fähigkeit des Modells, zwischen den Klassen zu unterscheiden, indem er die Fläche unter der ROC-Kurve misst, wobei ein höherer Wert auf eine bessere Leistung hinweist. Das Vorliegende Modell soll eine möglichst hohe Unterscheidungskraft haben.

Feature Engineering

Da die Kategorischen Attribute nicht vom Classifier erkannt werden, wird hier ein One-Hot-Encoding angewandt. One-Hot-Encoding ist eine Methode zur Umwandlung von kategorischen Variablen in ein binäres Format, das von Algorithmus besser verstanden werden kann.

```
In [ ]: # featrue engineering
df_encoded = pd.get_dummies(df, columns=["Sex", "ChestPainType", "Resting")
```

Abgesehen von dieser Änderung ist im Vorliegenden Datensatz kein weiteres Feature Engineering notwendig, da die Qualität der Daten mit Hilfe der Outlier Detection bereits erhöht wurde (ersetzen der 0-Werte bei Cholesterol durch den Durchschnittswert). Außerdem ist es schwierig, ohne tieferes, medizinisches Fachwissen mehr Wert für das Modell zu schaffen.

Im nächsten Schritt werden die target und feature Variablen festgelegt.

```
In []: # preparation for train/test split
    target = df_encoded["HeartDisease"]
    features = df_encoded.drop("HeartDisease", axis=1)
```

Train-/Testsplit

Um einen Bias im Machine Learning Model zu vermeiden, splittet man den Datensatz auf in Trainings- und Testdaten. Der Trainingsdatensatz wird verwendet, um das tatsächliche Modell zu erstellen, das der Algorithmus verwenden wird, wenn er neuen Daten ausgesetzt ist.

Das Testset ist der letzte Datensatz, der verwendet wird. Die Genauigkeit bei der Vorhersage des Testsets entspricht der Genauigkeit des ML-Algorithmus.

Für den train/test Split wird ein Verhältnis von 80/20 gewählt.

```
In []: # train/test split (80%/20%)
features_train, features_test, target_train, target_test = train_test_spl
```

Dummy Classifier

Um nachher die erzielten Scores zu beurteilen, wird zunächste ein Dummy Classifier gebaut. Dieser trifft zufällige Vorhersagen basierend auf der Verteilung im Datensatz.

```
In []: # build and train Dummy Classifier
dummy_clf = DummyClassifier(strategy="uniform")
dummy_clf.fit(features_train, target_train)

# get prediction
target_pred = dummy_clf.predict(features_test)

# get accuracy
accuracy = accuracy_score(target_test, target_pred)
print("Genauigkeit des Dummy Classifiers:", accuracy)
```

Genauigkeit des Dummy Classifiers: 0.4945652173913043

Classifier

Die folgende Methoden werden verwendet, um die drei gewählten Classifier zur fitten, die Scores anzuzeigen und jeweils die Confusion Matrix auszuegebn.

```
In [ ]: def model(classifier):
            Train the classifier on the training data and evaluate its performance
            Parameters:
            - classifier: The classifier model to be trained and evaluated.
            Returns:
            None
            classifier.fit(features_train, target_train)
            prediction = classifier.predict(features_test)
            print("Accuracy: {:.2%}".format(accuracy_score(target_test, prediction))
            print("ROC_AUC Score: {:.2%}".format(roc_auc_score(target_test, predi
        def model_evaluation(classifier):
            Evaluate the classifier using various performance metrics and visuali
            Parameters:
            - classifier: The trained classifier model.
            Returns:
            None
            # disply confusion Matrix
            cm = confusion_matrix(target_test, classifier.predict(features_test))
            names = ['True Neg', 'False Pos', 'False Neg', 'True Pos']
```

```
counts = [value for value in cm.flatten()]
percentages = ['{:.2%}'.format(value) for value in cm.flatten() / np.
labels = [f'{v1}\n{v2}\n{v3}' for v1, v2, v3 in zip(names, counts, pe
labels = np.asarray(labels).reshape(2, 2)
sns.heatmap(cm, annot=labels, cmap=colors, fmt='')

# show classification Report
print(classification_report(target_test, classifier.predict(features_
```

ML: Random Forrest

Der Random Forest Classifier ist ein Algorithmus für die Klassifizierung, der auf der Kombination mehrerer Entscheidungsbäume basiert. Er eignet sich gut für die Vorhersage von Herzkrankheiten aufgrund seiner Fähigkeit, mit verschiedenen Datentypen umzugehen und robuste Ergebnisse zu liefern. Er heißt "Random" Forest, da beim Algorithmus zwei zufällige Prozesse ablaufen. Zum einen das Bootstrapping zum anderen die Feature Auswahl beim erstellen der Entscheidungsbäume. Der Algorithmus baut also eine Vielzahl an Bäumen, die auf zufälligen Daten des Datensatzes basieren.

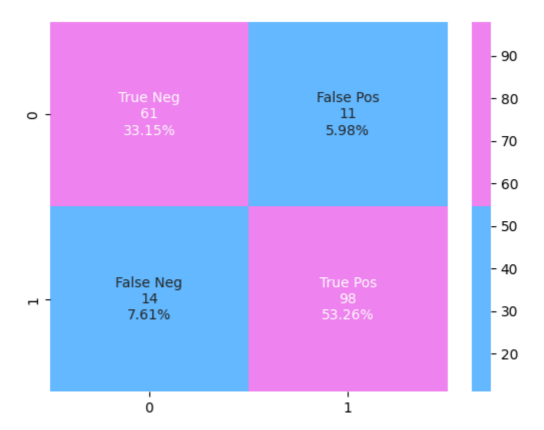
```
In []: # define RFC
    forest = RandomForestClassifier()

# get scores
    model(forest)

Accuracy: 86.41%
    ROC_AUC Score: 86.11%
```

In []: # get evalutation
model_evaluation(forest)

	precision	recall	f1–score	support
0	0.81	0.85	0.83	72
1	0.90	0.88	0.89	112
accuracy			0.86	184
macro avg	0.86	0.86	0.86	184
weighted avg	0.87	0.86	0.86	184



ML: Logistic Regression

Die logistische Regression ist ein Algorithmus zur Klassifizierung, der die Wahrscheinlichkeit für das Eintreten eines Ereignisses basierend auf einer oder mehreren unabhängigen Variablen schätzt. Dabei nutzt sie die logistische Funktion, um die Vorhersage zwischen 0 und 1 zu skalieren. Sie eignet sich gut für binäre Klassifizierungsaufgaben wie die Vorhersage von Herzkrankheiten.

```
In []: # defining LRC
    classifier_lr = LogisticRegression(max_iter=10000)
# get scores
model(classifier_lr)
```

Accuracy: 86.41% ROC_AUC Score: 86.86%

In []: # get scores
model_evaluation(classifier_lr)

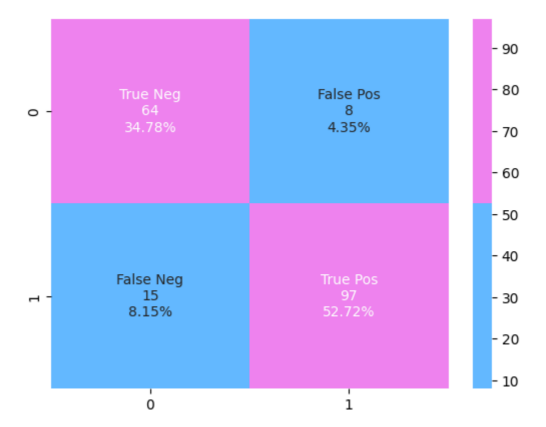
	precision	recall	f1-score	support
0 1	0.79 0.92	0.89 0.85	0.84 0.88	72 112
accuracy macro avg	0.86	0.87	0.86 0.86	184 184
weighted avg	0.87	0.86	0.87	184



ML: Support Vector Machine

Die Support Vector Machine ist ebenfalls ein Klassifizierungsalgorithmus, der darauf abzielt, eine Trennung zwischen den verschiedenen Klassen zu finden, indem er die beste Entscheidungsgrenze (Hyperplane) zwischen den Datenpunkten sucht. Sie funktioniert, indem sie den Abstand zwischen den Datenpunkten maximiert und gleichzeitig eine minimale Fehlerrate aufweist. SVM eignet sich gut für datengetriebene Anwendungen mit komplexen Entscheidungsgrenzen und kann auch mit nicht-linearen Daten umgehen, indem sie den sogenannten Kernel-Trick anwendet. In Bezug auf Herzkrankheiten eignet sich die SVM, wenn die Daten gut separierbar sind und klare Entscheidungsgrenzen zwischen den Klassen existieren.

	precision	recall	f1-score	support
0 1	0.81 0.92	0.89 0.87	0.85 0.89	72 112
accuracy	0 . 87	0 . 88	0.88 0.87	184 184
macro avg weighted avg	0.88	0.88	0.88	184



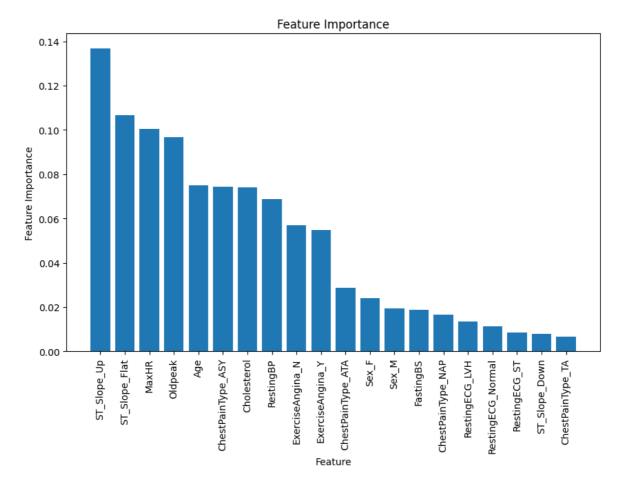
Die Scores aller Classifier liefern deutlich präzisere Ergebnisse als der Dummy-Classifier.

Feature Importance für den RFC

Um zu visualisieren, welche Features den größten Einfluss haben (am Beispiel des RFC) wird der folgende Plot erstellt.

```
In []: # plot for feature importances
    importances = forest.feature_importances_
    indices = np.argsort(importances)[::-1]
    feature_names = features_train.columns

plt.figure(figsize=(10, 6))
    plt.title("Feature Importance")
    plt.bar(range(features_train.shape[1]), importances[indices], align="cent plt.xticks(range(features_train.shape[1]), feature_names[indices], rotati plt.xlabel("Feature")
    plt.ylabel("Feature Importance")
    plt.show()
```



Da es sich um relativ wenige Features handelt, wurde sich beim Training nicht auf einzelne Features konzentriert.

Gridsearch Prameter Tuning

Das Parameter-Tuning wird nur für den Random Forest Classifier (RFC) durchgeführt. Der RFC ist bekannt für seine Vielseitigkeit und Robustheit, insbesondere bei binären Klassifikationsproblemen wie im vorliegenden Fall. Durch das Feintuning seiner Hyperparameter kann die Vorhersagegenauigkeit weiter optimiert und potenzielles Overfitting reduziert werden. Dies ermöglicht eine präzisere Identifizierung von Herzkrankheiten, was in medizinischen Anwendungen von entscheidender Bedeutung ist.

Das Parameter-Tuning wird mithilfe von Grid Search durchgeführt, einem Ansatz zur systematischen Suche nach den besten Hyperparameter-Kombinationen für ein Machine Learning Modell. Grid Search durchläuft vordefinierte Kombinationen von Hyperparametern und bewertet die Leistung des Modells anhand einer bestimmten Metrik für jede Kombination. In unserem Fall optimieren wir den Receiver Operating Characteristic Area Under Curve (ROC AUC) Score. Der ROC AUC Score ist eine Metrik, die die Fähigkeit eines Modells bewertet, zwischen den Klassen zu unterscheiden und die Trade-offs zwischen True Positive Rate und False Positive Rate darstellt. Für das binäre Klassifikationsproblem mit Herzkrankheiten ist es wichtig, dass das Modell eine hohe Unterscheidungskraft zwischen kranken und gesunden Patienten aufweist, weshalb der ROC AUC Score optimiert wird.

```
In []: # defining the grid search parameters
param_grid = {
         'n_estimators': [100, 200, 300],
         'max_depth': [None, 10, 20, 30],
         'min_samples_split': [2, 5, 10],
         'min_samples_leaf': [1, 2, 4]
}
```

Mittels Gridsearch wird die beste Kombination folgender Parameter ermittelt: Mittels Gridsearch wird die beste Kombination folgender Parameter ermittelt:

- n_estimators: Die Anzahl der Bäume im Random Forest.
- max_depth: Die maximale Tiefe jedes Entscheidungsbaums.
- min_samples_split: Minimale Anzahl von Beispielen, um einen Knoten zu teilen.
- min_samples_leaf: Minimale Anzahl von Beispielen in einem Blatt.

```
In [ ]: # search for the best combination
    grid_search.fit(features_train, target_train)
```

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Fitting 5 folds for each of 108 candidates, totalling 540 fits [CV 1/5] END max_depth=None, min_samples_leaf=1, min_samples_split=2, n_es timators=100;, score=0.951 total time= 0.1s [CV 2/5] END max_depth=None, min_samples_leaf=1, min_samples_split=2, n_es timators=100;, score=0.934 total time= 0.1s [CV 3/5] END max_depth=None, min_samples_leaf=1, min_samples_split=2, n_es timators=100;, score=0.903 total time= 0.1s [CV 4/5] END max_depth=None, min_samples_leaf=1, min_samples_split=2, n_es timators=100;, score=0.901 total time= 0.2s
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                                       0.3s
[CV 4/5] END max_depth=10, min_samples_leaf=1, min_samples_split=5, n_esti
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0.3s

mators=200;, score=0.902 total time=

```
[CV 1/5] END max_depth=10, min_samples_leaf=1, min_samples_split=5, n_esti
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[CV 5/5] END max_depth=10, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.893 total time=
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mators=200;, score=0.900 total time=
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[CV 5/5] END max_depth=10, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.896 total time=
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                                       0.4s
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mators=100;, score=0.901 total time=
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```
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[CV 1/5] END max_depth=10, min_samples_leaf=4, min_samples_split=10, n_est
imators=300;, score=0.959 total time=
                                        0.4s
[CV 2/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.943 total time=
                                       0.2s
```

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```
[CV 3/5] END max_depth=10, min_samples_leaf=4, min_samples_split=10, n_est
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                                       0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
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                                       0.3s
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[CV 5/5] END max_depth=10, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.4s
[CV 1/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.952 total time=
                                       0.4s
[CV 3/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.899 total time=
                                       0.2s
[CV 2/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.933 total time=
                                       0.2s
[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.896 total time=
                                       0.2s
[CV 4/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.900 total time=
                                       0.2s
[CV 3/5] END max depth=20, min samples leaf=1, min samples split=2, n esti
mators=300;, score=0.903 total time=
                                       0.5s
[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.891 total time=
                                       0.4s
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                                       0.1s
[CV 3/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
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                                       0.2s
[CV 1/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.955 total time=
                                       0.3s
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mators=100;, score=0.955 total time=
                                       0.2s
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                                       0.3s
[CV 2/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.938 total time=
                                       0.4s
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                                        0.1s
[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
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[CV 4/5] END max_depth=20, min_samples_leaf=1, min_samples_split=2, n_esti
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[CV 2/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
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                                       0.3s
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[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=10, n_est
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[CV 4/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
mators=300;, score=0.903 total time=
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imators=100;, score=0.903 total time=
                                        0.1s
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mators=300;, score=0.954 total time=
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                                        0.3s
```

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                                        0.4s
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mators=100;, score=0.940 total time=
                                       0.1s
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                                        0.2s
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mators=300;, score=0.902 total time=
                                       0.5s
[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=5, n_esti
mators=300;, score=0.897 total time=
                                       0.4s
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imators=200;, score=0.896 total time=
                                      0.3s
[CV 3/5] END max_depth=20, min_samples_leaf=1, min_samples_split=10, n_est
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                                        0.4s
[CV 5/5] END max_depth=20, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.899 total time=
                                        0.4s
[CV 3/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.913 total time=
                                       0.2s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.902 total time=
                                       0.1s
[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.959 total time=
                                       0.3s
[CV 5/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.895 total time=
                                       0.1s
[CV 5/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.896 total time=
                                       0.3s
[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.958 total time=
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[CV 3/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.905 total time=
                                       0.3s
[CV 2/5] END max_depth=20, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.937 total time=
                                        0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.900 total time=
                                        0.4s
[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.957 total time=
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mators=100;, score=0.903 total time=
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                                       0.1s
[CV 5/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.896 total time=
                                       0.2s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
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                                       0.3s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.899 total time=
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[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.959 total time=
                                       0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.905 total time=
                                       0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.902 total time=
                                       0.2s
```

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```
[CV 2/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
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[CV 3/5] END max_depth=20, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.909 total time=
                                       0.4s
[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.955 total time=
                                       0.3s
[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.957 total time=
                                       0.4s
[CV 3/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.902 total time=
                                       0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=10, n_est
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                                      0.1s
[CV 2/5] END max_depth=20, min_samples_leaf=2, min_samples_split=10, n_est
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                                      0.2s
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                                     0.2s
[CV 5/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.896 total time=
                                       0.3s
[CV 5/5] END max depth=20, min samples leaf=2, min samples split=2, n esti
mators=300;, score=0.897 total time=
                                       0.4s
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                                        0.1s
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[CV 3/5] END max_depth=20, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.906 total time=
                                        0.3s
[CV 5/5] END max_depth=20, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.895 total time=
                                        0.3s
[CV 2/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
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[CV 4/5] END max_depth=20, min_samples_leaf=2, min_samples_split=5, n_esti
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[CV 4/5] END max_depth=20, min_samples_leaf=4, min_samples_split=2, n_esti
mators=100;, score=0.902 total time=
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[CV 2/5] END max_depth=20, min_samples_leaf=4, min_samples_split=2, n_esti
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[CV 1/5] END max_depth=20, min_samples_leaf=2, min_samples_split=10, n_est
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```

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```
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                                       0.3s
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                                        0.4s
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                                        0.4s
[CV 1/5] END max_depth=20, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.960 total time=
                                       0.3s
[CV 5/5] END max_depth=20, min_samples_leaf=4, min_samples_split=2, n_esti
mators=200;, score=0.896 total time=
                                       0.3s
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                                       0.3s
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mators=100;, score=0.902 total time=
                                       0.1s
[CV 3/5] END max depth=20, min samples leaf=4, min samples split=5, n esti
mators=100;, score=0.906 total time=
                                       0.2s
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                                       0.1s
[CV 5/5] END max_depth=20, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.897 total time=
                                       0.4s
[CV 1/5] END max_depth=20, min_samples_leaf=4, min_samples_split=5, n_esti
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                                       0.2s
[CV 5/5] END max_depth=20, min_samples_leaf=4, min_samples_split=5, n_esti
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                                       0.4s
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                                       0.4s
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imators=100;, score=0.937 total time=
                                        0.2s
[CV 1/5] END max_depth=20, min_samples_leaf=4, min_samples_split=5, n_esti
mators=300;, score=0.960 total time=
                                       0.4s
[CV 2/5] END max_depth=20, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.2s
```

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```
[CV 3/5] END max_depth=20, min_samples_leaf=4, min_samples_split=5, n_esti
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imators=200;, score=0.908 total time=
                                      0.3s
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imators=200;, score=0.960 total time=
                                        0.3s
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mators=300;, score=0.896 total time=
                                       0.4s
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                                        0.4s
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                                        0.4s
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                                       0.2s
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                                       0.1s
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                                       0.2s
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                                       0.2s
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                                        0.4s
[CV 4/5] END max_depth=20, min_samples_leaf=4, min_samples_split=10, n_est
imators=300;, score=0.906 total time=
                                        0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.892 total time=
                                       0.3s
[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.931 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.938 total time=
                                       0.2s
[CV 3/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.900 total time=
                                       0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=200;, score=0.905 total time=
                                       0.3s
[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.936 total time=
                                       0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.884 total time=
                                       0.2s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=100;, score=0.904 total time=
                                       0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.897 total time=
                                       0.4s
[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.939 total time=
                                       0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.905 total time=
                                       0.3s
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[CV 1/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.956 total time=
                                      0.2s
[CV 1/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.950 total time=
                                       0.4s
[CV 1/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=300;, score=0.953 total time=
                                       0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.896 total time=
                                       0.4s
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mators=300;, score=0.901 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=1, min_samples_split=2, n_esti
mators=300;, score=0.901 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.904 total time=
                                       0.3s
[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
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[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=200;, score=0.896 total time=
                                       0.3s
[CV 3/5] END max depth=30, min samples leaf=1, min samples split=10, n est
imators=100;, score=0.902 total time=
                                        0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=100;, score=0.903 total time=
                                        0.2s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
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                                       0.4s
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                                        0.3s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=100;, score=0.898 total time=
                                        0.2s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=200;, score=0.898 total time=
                                        0.2s
[CV 3/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
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                                       0.3s
[CV 1/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
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                                        0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
mators=300;, score=0.903 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=5, n_esti
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[CV 2/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
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                                        0.4s
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mators=100;, score=0.904 total time=
                                       0.1s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.898 total time=
                                       0.1s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.955 total time=
                                       0.2s
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imators=200;, score=0.937 total time=
                                        0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=200;, score=0.902 total time=
                                        0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.903 total time=
                                        0.4s
[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=100;, score=0.900 total time=
                                       0.1s
[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
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[CV 1/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.956 total time=
                                        0.5s
[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.937 total time=
                                       0.2s
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[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.957 total time=
                                      0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.903 total time=
                                       0.3s
[CV 3/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.904 total time=
                                        0.4s
[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.934 total time=
                                       0.1s
[CV 5/5] END max_depth=30, min_samples_leaf=1, min_samples_split=10, n_est
imators=300;, score=0.898 total time=
                                        0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.908 total time=
                                       0.2s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.960 total time=
                                       0.5s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=200;, score=0.895 total time=
                                       0.3s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.907 total time=
                                       0.2s
[CV 4/5] END max depth=30, min samples leaf=2, min samples split=5, n esti
mators=100;, score=0.898 total time=
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[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
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                                       0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.895 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.905 total time=
                                       0.3s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=100;, score=0.893 total time=
                                       0.2s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
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[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.896 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.938 total time=
                                       0.4s
[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.938 total time=
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[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.893 total time=
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[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=2, n_esti
mators=300;, score=0.904 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=200;, score=0.937 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=100;, score=0.901 total time=
                                        0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.904 total time=
                                       0.4s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.957 total time=
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[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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                                        0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.901 total time=
                                        0.2s
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[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.938 total time=
                                       0.3s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
mators=300;, score=0.902 total time=
                                       0.4s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.960 total time=
                                        0.3s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=100;, score=0.935 total time=
                                       0.1s
[CV 1/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=300;, score=0.961 total time=
                                        0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=200;, score=0.898 total time=
                                        0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=100;, score=0.902 total time=
                                       0.1s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=5, n_esti
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                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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[CV 3/5] END max depth=30, min samples leaf=4, min samples split=2, n esti
mators=100;, score=0.909 total time=
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[CV 5/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
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                                       0.1s
[CV 3/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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                                        0.5s
[CV 5/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
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                                        0.5s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
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                                       0.3s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=100;, score=0.961 total time=
                                       0.1s
[CV 5/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=200;, score=0.897 total time=
                                       0.2s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=200;, score=0.907 total time=
                                       0.3s
[CV 2/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=300;, score=0.937 total time=
                                        0.4s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=100;, score=0.960 total time=
                                       0.1s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=200;, score=0.930 total time=
                                       0.2s
[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.903 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=100;, score=0.910 total time=
                                       0.2s
[CV 4/5] END max_depth=30, min_samples_leaf=2, min_samples_split=10, n_est
imators=300;, score=0.903 total time=
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[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.936 total time=
                                       0.5s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=100;, score=0.933 total time=
                                       0.1s
[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=200;, score=0.904 total time=
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[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.960 total time=
                                       0.4s
[CV 5/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=100;, score=0.895 total time=
                                       0.2s
[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
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                                       0.2s
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mators=200;, score=0.936 total time=
                                       0.2s
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[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=200;, score=0.903 total time=
                                      0.3s
[CV 5/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.897 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=2, n_esti
mators=300;, score=0.908 total time=
                                       0.4s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=200;, score=0.963 total time=
                                       0.3s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=300;, score=0.961 total time=
                                       0.4s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=100;, score=0.938 total time=
                                      0.1s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=200;, score=0.912 total time=
                                       0.3s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=5, n_esti
mators=300;, score=0.910 total time=
                                       0.4s
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                                       0.3s
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mators=300;, score=0.898 total time=
                                       0.4s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=100;, score=0.904 total time=
                                        0.1s
[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=100;, score=0.960 total time=
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[CV 1/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.2s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.2s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
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[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.4s
[CV 2/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
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                                        0.3s
[CV 5/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=300;, score=0.900 total time=
                                        0.3s
[CV 4/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=300;, score=0.903 total time=
                                        0.3s
[CV 3/5] END max_depth=30, min_samples_leaf=4, min_samples_split=10, n_est
imators=300;, score=0.907 total time=
                                        0.4s
             GridSearchCV
  ▶ estimator: RandomForestClassifier
```

Out[]:

RandomForestClassifier

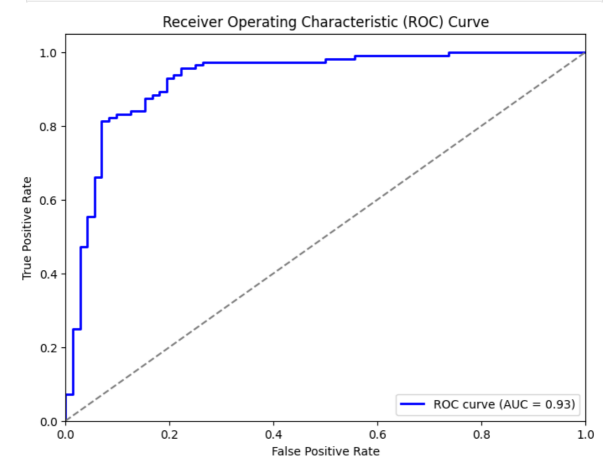
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Die beste Parameter Kombination:

```
In [ ]: # get best combination and result
        print("Beste Hyperparameter-Kombinationen: ", grid_search.best_params_)
       Beste Hyperparameter-Kombinationen: {'max_depth': 20, 'min_samples_leaf':
       4, 'min_samples_split': 10, 'n_estimators': 100}
In [ ]: # best params
        best_params = grid_search.best_params_
        # forest with best params
        best_forest = RandomForestClassifier(**best_params)
In [ ]: # fit the best model
        best_forest.fit(features_train, target_train)
Out[]:
                              RandomForestClassifier
        RandomForestClassifier(max_depth=20, min_samples_leaf=4, min_samp
        les split=10)
In [ ]: # show scores
        train_score = best_forest.score(features_train, target_train)
        test score = best forest.score(features test, target test)
        print("Trainingsgenauigkeit:", train_score)
        print("Testgenauigkeit:", test_score)
       Trainingsgenauigkeit: 0.903137789904502
       Testgenauigkeit: 0.8478260869565217
In [ ]: # predictions for train and test data
        train_predictions = best_forest.predict_proba(features_train)[:, 1] # pr
        test predictions = best forest.predict proba(features test)[:, 1] # prob
        # calculate roc_auc-score
        train_roc_auc = roc_auc_score(target_train, train_predictions)
        test_roc_auc = roc_auc_score(target_test, test_predictions)
        # result
        print("Trainings-ROC-AUC-Score:", train_roc_auc)
        print("Test-ROC-AUC-Score:", test_roc_auc)
       Trainings-ROC-AUC-Score: 0.9740918283274661
       Test-ROC-AUC-Score: 0.9268353174603174
In [ ]: | target_pred_proba = best_forest.predict_proba(features_test)[:, 1]
        # compute ROC curve and ROC-AUC score
        fpr, tpr, thresholds = roc_curve(target_test, target_pred_proba)
        roc_auc = roc_auc_score(target_test, target_pred_proba)
        # plot ROC curve
        plt.figure(figsize=(8, 6))
        plt.plot(fpr, tpr, color='blue', lw=2, label='ROC curve (AUC = %0.2f)' %
        plt.plot([0, 1], [0, 1], color='gray', linestyle='--')
        plt.xlim([0.0, 1.0])
        plt.ylim([0.0, 1.05])
        plt.xlabel('False Positive Rate')
```

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```
plt.ylabel('True Positive Rate')
plt.title('Receiver Operating Characteristic (ROC) Curve')
plt.legend(loc="lower right")
plt.show()
```



Evaluation und Ergebnisdarstellung

Basierend auf der Evaluation der drei Klassifikationsmodelle – Random Forest Classifier, Logistische Regression und Support Vector Machine – erzielte der RFC vor dem Parameter-Tuning die beste Leistung mit einer Genauigkeit von 88.59% und einem ROC AUC Score von 88.27% (Diese Werte können bei erneutem ausführen des Notebooks abweichen, da der Train-Test split einen randomstate von 42 besitzt). Außerdem kamen beim RFC die wenigsten False Negatives zum Vorschein, was bei der Erkennung von Krankheiten besonders wichtig ist. Der precision, recall und f1-score für beide Klassen (Herzkrankheit und Normal) zeigen eine ausgeglichene Leistung des Modells. Nach dem Parameter-Tuning wurden die Hyperparameter des RFC optimiert, wodurch eine verbesserte Leistung mit einer ROC AUC Score von 92.78% erzielt wurde (möglicherweise ebenfalls abweichende Werte). Dies unterstreicht die Wirksamkeit des gewählten Ansatzes und die Fähigkeit des Modells, zwischen Herzkrankheit und Normalzustand zu unterscheiden.

Vorhersage-Demo

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korrekt als Herzerkrankung an.

```
In [ ]: selected data point = features train.iloc[0:1, :]
        selected_target = target_train.iloc[0]
        # make the prediction for the chosen data point
        prediction = forest.predict(selected_data_point)
        # display the chosen data point, the true class and the predicted class
        print("Ausgewählter Datenpunkt:")
        print(selected_data_point)
        print("\nWahre Klasse des ausgewählten Datenpunkts:", selected_target)
        print("\nVorhersage für den ausgewählten Datenpunkt:", prediction)
       Ausgewählter Datenpunkt:
            Age RestingBP Cholesterol FastingBS MaxHR Oldpeak Sex_F Sex_M
       795
             42
                       120
                                    240
                                                 1
                                                      194
                                                               0.8 False
                                                                            True
            ChestPainType_ASY ChestPainType_ATA ChestPainType_NAP \
       795
                                           False
                        False
                                                               True
            ChestPainType_TA RestingECG_LVH RestingECG_Normal RestingECG_ST \
       795
                                       False
                                                           True
                       False
                                                                         False
            ExerciseAngina_N ExerciseAngina_Y ST_Slope_Down ST_Slope_Flat \
       795
                        True
                                         False
                                                         True
                                                                       False
            ST_Slope_Up
       795
                  False
       Wahre Klasse des ausgewählten Datenpunkts: 0
       Vorhersage für den ausgewählten Datenpunkt: [0]
        Wie man anhand des Beispiels sieht, erkennt das Model den gegeben Datenpunkt
```

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