# PROJECT REPORT HEART FAILURE PREDICTION



Submitted By: -

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#### **UNDERTAKING**

I declare that the work presented in this project titled "Heart Failure Prediction", submitted to the All India Council of Robotics and Automation, for the award of the Internship in Data Science, is my original work. I have not plagiarized or submitted the same work for the award of any other Internship. In case this undertaking is found incorrect, I accept that my Project may be unconditionally withdrawn.

Saima Samreen 3GN18CS080

# **CERTIFICATE**

This is to certify that the work contained in the project titled "Heart Failure Prediction", by Saima Samreen, has been carried out under my supervision and that this work has not been submitted elsewhere for internship.

All India Council of Robotics and Automation

**Data Science** 

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#### **PREFACE**

Heart failure — sometimes known as congestive heart failure — occurs when the heart muscle doesn't pump blood as well as it should. When this happens, blood often backs up and fluid can build up in the lungs, causing shortness of breath.

Certain heart conditions, such as narrowed arteries in the heart (coronary artery disease) or high blood pressure, gradually leave the heart too weak or stiff to fill and pump blood properly.

Proper treatment can improve the signs and symptoms of heart failure and may help some people live longer. Lifestyle changes — such as losing weight, exercising, reducing salt (sodium) in your diet and managing stress — can improve your quality of life. However, heart failure can be life-threatening. People with heart failure may have severe symptoms, and some may need a heart transplant or a ventricular assist device (VAD).

One way to prevent heart failure is to prevent and control conditions that can cause it, such as coronary artery disease, high blood pressure, diabetes and obesity.

So, the main goal here is to predict whether the individual has any heart problems, further leading towards heart failure. People with cardiovascular disease or who are at high cardiovascular risk (due to the presence of one or more risk factors such as hypertension, diabetes, hyperlipidaemia or already established disease) need early detection and management wherein a machine learning model can be of great help.

## <u>ACKNOWLEDGEMENT</u>

I take upon this opportunity to acknowledge the many people whose prayers and support meant a lot to me.

I am deeply indebted to my mentor Mr. Sumit Chatterjee who motivated me along the way.

I would like to thank all my teachers who help me in this project.

I further thank my friends.

My heartfelt thanks to parents who supported me a lot. I owe my sincere gratitude towards the almighty God.

Finally, I would like to wind up by paying my heartfelt thanks to AICRA institute who provided me with this great opportunity.

Saima Samreen

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## DATASET DESCRIPTION

This dataset contains the medical records of 299 patients who had heart failure, collected during their follow-up period, where each patient profile has 13 clinical features.

These 13 clinical attributes are pre-processed and trained to predict that there is the presence or absence of the heart disease. The attributes are in the form of numeric type which represents the age of the patient in the numeric values, Age is the most important risk factor in growing the heart diseases; it doubles the risk in "ADOLESCENCE".

Dataset Attributes: -

Thirteen (13) clinical features: -

1)age: age of the patient (years)

2)anaemia: decrease of red blood cells or haemoglobin (boolean) 3)high blood pressure:

if the patient has hypertension (boolean)

4)creatinine phosphokinase (CPK): level of the CPK enzyme in the blood

(mcg/L)

5) diabetes: if the patient has diabetes (boolean)

6) ejection fraction: percentage of blood leaving the heart at each contraction

(percentage)

7) platelets: platelets in the blood (kiloplatelets/mL)

8)sex: woman or man (binary)

9)serum creatinine: level of serum creatinine in the blood (mg/dL)

10)serum sodium: level of serum sodium in the blood (mEq/L)

11) smoking: if the patient smokes or not (boolean)

12)time: follow-up period (days)

13)[target] death event: if the patient deceased during the follow-up period (boolean)

## **METHODOLOGY**

#### **Logistic Regression: -**

Logistic regression is the supervised learning algorithm, which is used to predict the categorical variables or discrete values. It can be used for the classification problems in machine learning, and the output of the logistic regression algorithm can be either Yes or NO, 0 or 1, Red or Blue, etc.

#### K-Nearest Neighbour (KNN): -

K-Nearest Neighbour is a supervised learning algorithm that can be used for both classification and regression problems. This algorithm works by assuming

the similarities between the new data point and available data points. Based on these similarities, the new data points are put in the most similar categories.

#### **Decision Tree Algorithm: -**

A decision tree is a supervised learning algorithm that is mainly used to solve the classification problems but can also be used for solving the regression problems. It can work with both categorical variables and continuous variables. It shows a tree-like structure that includes nodes and branches and starts with the root node that expand on further branches till the leaf node.

#### Random Forest Algorithm: -

Random forest is the supervised learning algorithm that can be used for both classification and regression problems in machine learning. It is an ensemble learning technique that provides the predictions by combining the multiple classifiers and improve the performance of the model.

#### **Gradient Boosting Algorithm: -**

Gradient Boosting is a popular boosting algorithm. In gradient boosting, each predictor corrects its predecessor's error. In contrast to Adaboost, the weights of the training instances are not tweaked, instead, each predictor is trained using the residual errors of predecessor as labels.

# **SYSTEM ARCHITECTURE**

- 1. The initial procedure for this project is the collection of data. We have collected the data set from Kaggle, which is available; it is an open-source.
- 2. The next step after data collection is data pre-processing. In this step, the data is cleansed by removing unnecessary values. It also removes the missing/ null/ corrupted values.
- 3. After Data is cleansed, the next step is dividing the data, we divide the data into two sets, Training data and testing data. The values must deal before we start to construct the training model. By using training data, we build a model for prediction.
- 4. Now, we must calculate the accuracy of the model.

5. The final step is predicting the disease.

## **IMPLEMENTATION**

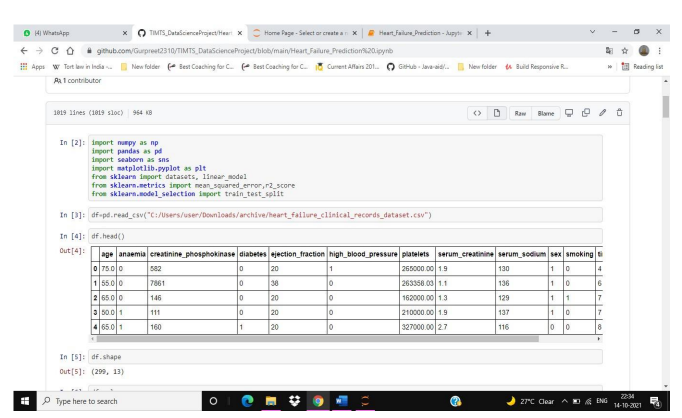
This work used Python programming for this project, as it is a high-level programming language and it has vast libraries and Python automates tasks and makes it efficient. Firstly, we need to install Python then we need to import some libraries, they are:

- 1) Numpy: Numpy is used for multi-dimensional arrays, it does element to element operations and it also has different methods for processing arrays.
- 2) Panda: Pandas is one of the highly used python libraries, it provides high performance. It manipulates data and it makes data analysis fast and easy.
- 3) Sklearn: It is most useful library, this library contains lot of efficient tools, it is used to build models like statistical modelling including classification, regression, clustering. After loading required packages, we divide dataset as training and testing as follows, here 80 % of dataset is taken as training and remaining 20 % as to perform test.

#### **IMPORT LIBRARIES**

First, let's import all the modules, functions and objects we are going to use

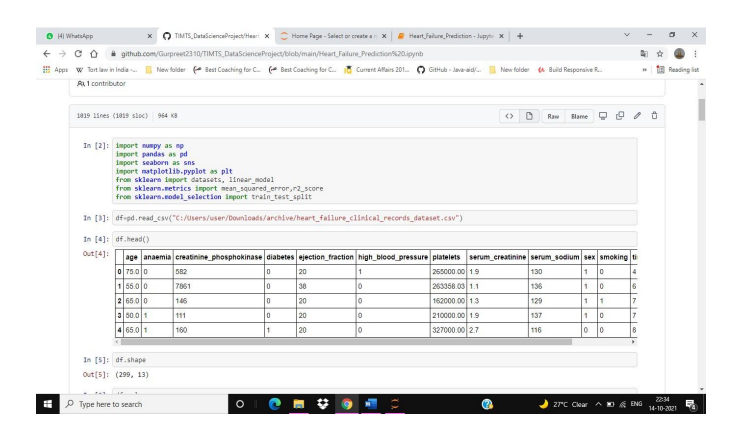
```
import numpy as np import
pandas as pd import seaborn as
sns import matplotlib.pyplot as
plt
from sklearn import datasets, linear_model
from sklearn.metrics import mean_squared_error,r2_score from
sklearn.model_selection import train_test_split
```



#### **Load Dataset**

We can load the data directly from the UCI Machine Learning repository. We are using pandas to load the data. We will also use pandas next to explore the data both with descriptive statistics and data visualization.

df=pd.read\_csv("C:/Users/user/Downloads/archive/heart\_failure\_clinical\_records\_da
taset.csv")

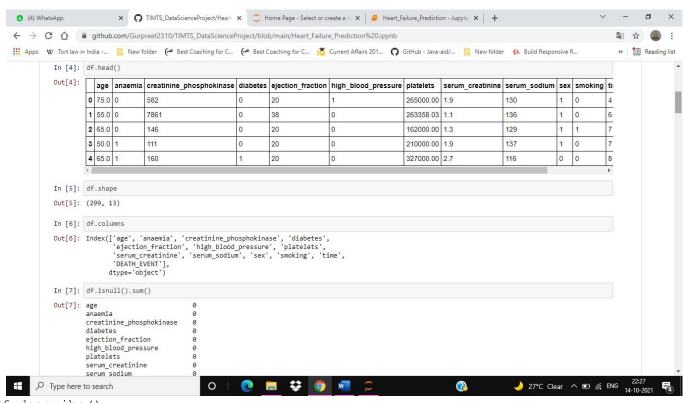


## **SUMMARIZE THE DATASET**

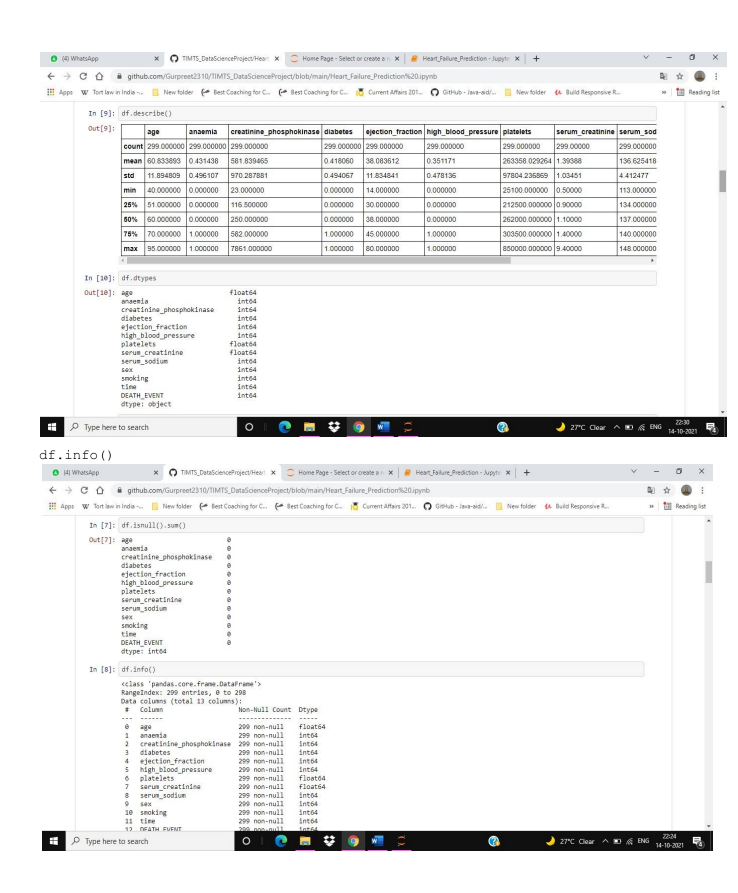
Now it is time to look at the data.

In this step we are going to look at the data a few different ways.

print(df.shape) print(df.head(10))



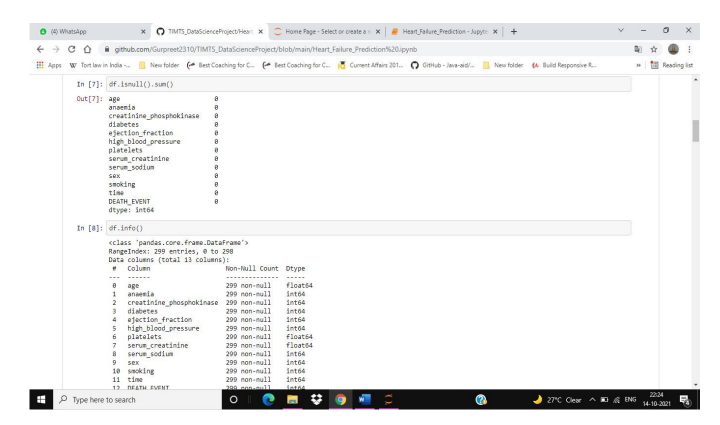
df.describe()



# **DATA PRE-PROCESSING**

In data pre-processing, it is pivotal to identify and correctly handle the missing values, failing to do this, you might draw inaccurate and faulty conclusions and inferences from the data.

heart data.isnull().sum()

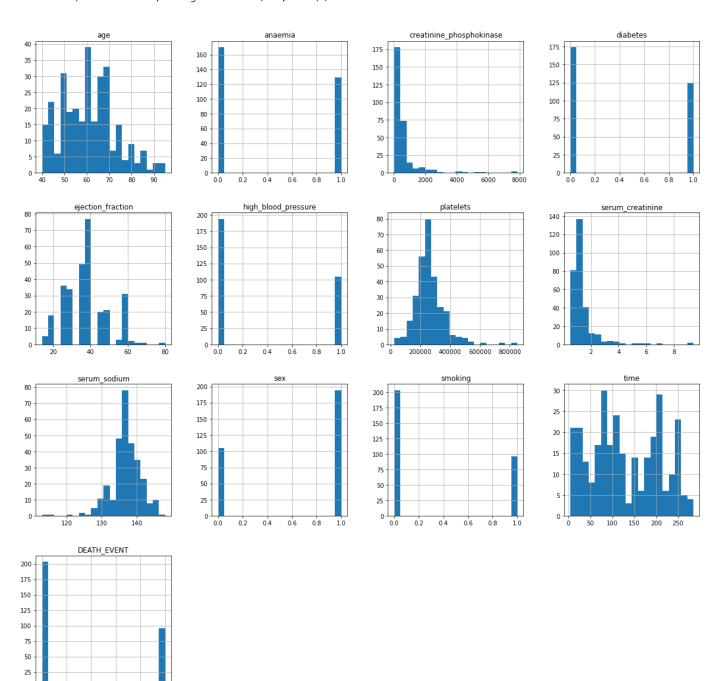


As we can see from above image, there are no missing values (null values) in the dataset so, the data pre-processing techniques are not required here.

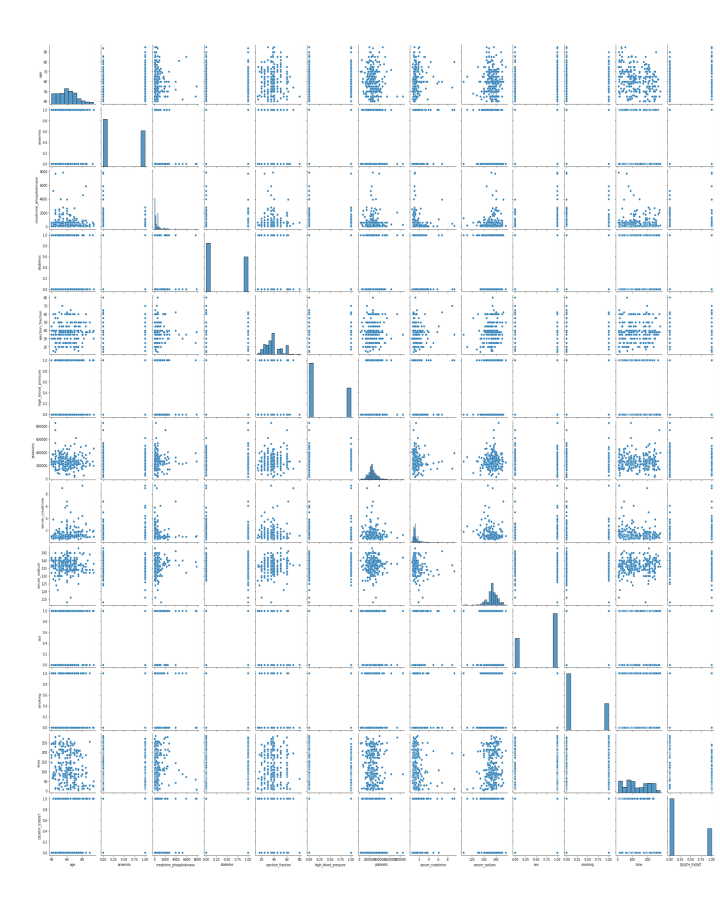
#### DATA VISUALIZATION

We now have a basic idea about the data. We need to extend that with some visualizations.

df.hist(bins = 20, figsize = (20, 20))



sns.pairplot(df)



corr\_matrix = df.corr()
plt.figure(figsize=(12,12,))
sns.heatmap(corr\_matrix, annot=True) plt.show()

age -	- 1	0.088	-0.082	-0.1	0.06	0.093	-0.052	0.16	-0.046	0.065	0.019	-0.22	0.25
anaemia -	0.088	1	-0.19	-0.013	0.032	0.038	-0.044	0.052	0.042	-0.095	-0.11	-0.14	0.066
creatinine_phosphokinase -	-0.082	-0.19	1	-0.0096	-0.044	-0.071	0.024	-0.016	0.06	0.08	0.0024	-0.0093	0.063
diabetes -	-0.1	-0.013	-0.0096	1	-0.0049	-0.013	0.092	-0.047	-0.09	-0.16	-0.15	0.034	-0.0019
ejection_fraction -	0.06	0.032	-0.044	-0.0049	1	0.024	0.072	-0.011	0.18	-0.15	-0.067	0.042	-0.27
high_blood_pressure -	0.093	0.038	-0.071	-0.013	0.024	1	0.05	-0.0049	0.037	-0.1	-0.056	-0.2	0.079
platelets -	-0.052	-0.044	0.024	0.092	0.072	0.05	1	-0.041	0.062	-0.13	0.028	0.011	-0.049
serum_creatinine -	0.16	0.052	-0.016	-0.047	-0.011	-0.0049	-0.041	1	-0.19	0.007	-0.027	-0.15	0.29
serum_sodium <sup>-</sup>	-0.046	0.042	0.06	-0.09	0.18	0.037	0.062	-0.19	1	-0.028	0.0048	0.088	-0.2
sex -	0.065	-0.095	0.08	-0.16	-0.15	-0.1	-0.13	0.007	-0.028	1	0.45	-0.016	-0.0043
smoking -	0.019	-0.11	0.0024	-0.15	-0.067	-0.056	0.028	-0.027	0.0048	0.45	1	-0.023	-0.013
time -	-0.22	-0.14	-0.0093	0.034	0.042	-0.2	0.011	-0.15	0.088	-0.016	-0.023	1	-0.53
DEATH_EVENT -	0.25	0.066	0.063	-0.0019	-0.27	0.079	-0.049	0.29	-0.2	-0.0043	-0.013	-0.53	1
	- age	anaemia -	creatinine_phosphokinase -	diabetes -	ejection_fraction -	high_blood_pressure -	platelets -	serum_creatinine -	serum_sodium -	- xas	smoking -	time -	DEATH_EVENT -

-1.0

- 0.8

- 0.6

- 0.4

- 0.2

- 0.0

#### CREATE A VALIDATION DATASET

We need to know that the model we created is good.

Later, we will use statistical methods to estimate the accuracy of the models that we create on unseen data. We also want a more concrete estimate of the accuracy of the best model on unseen data by evaluating it on actual unseen data.

That is, we are going to hold back some data that the algorithms will not get to see, and we will use this data to get a second and independent idea of how accurate the best model might be.

We will split the loaded dataset into two, 85% of which we will use to train, evaluate and select among our models, and 15% that we will hold back as a validation dataset.

```
x= df.drop(['DEATH_EVENT'],axis=1) y=df.DEATH_EVENT

x_train,x_test,y_train,y_test=train_test_split(x,y,test_size=0.15,random_state=25)
print(x_train.shape)
print(y_train.shape)
print(x_test.shape)
print(y_test.shape)
```

## **BUILD MODELS**

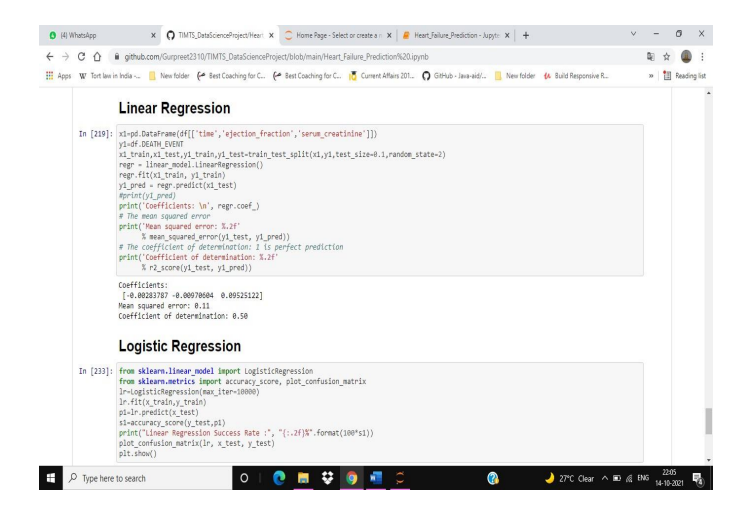
We don't know which algorithms would be good on this problem or what configurations to use.

We get an idea from the plots that some of the classes are partially linearly separable in some dimensions, so we are expecting generally good results.

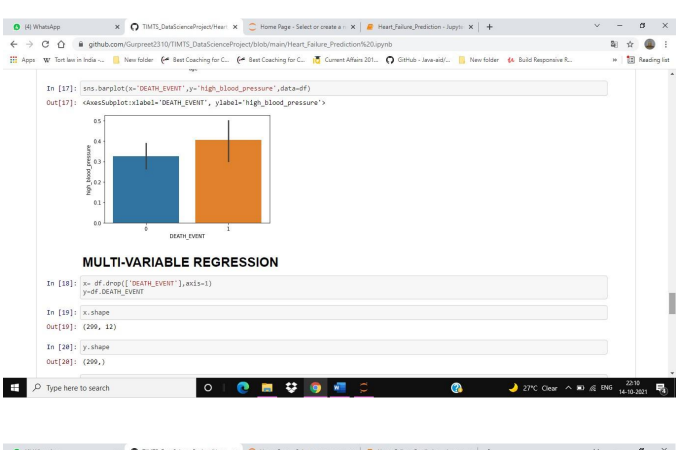
#### Let's test 3 different models:

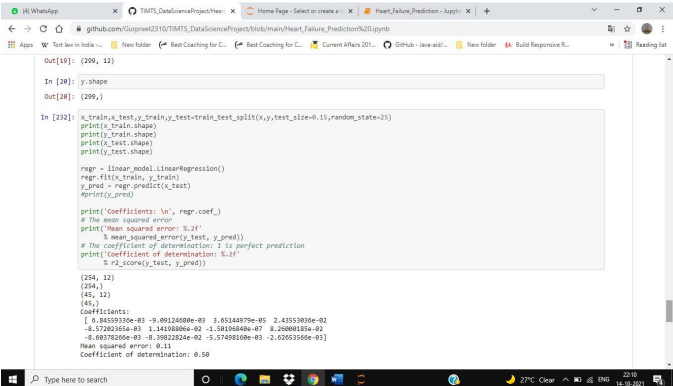
- Linear Regression Model
- Multi-Variable Regression Model
- Logistic Regression Model

# **Linear Regression**



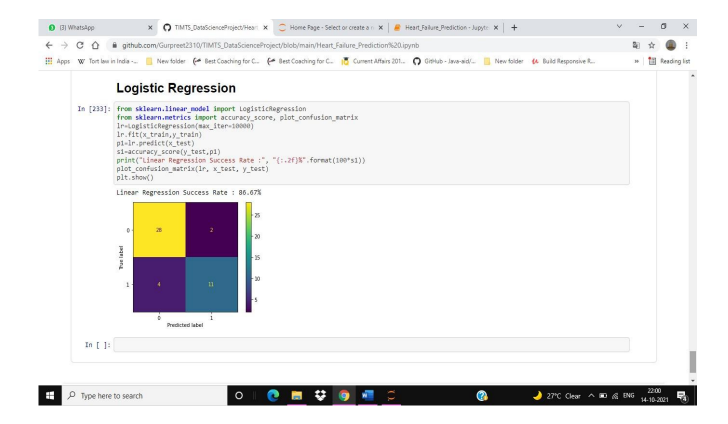
# Multi-Variable Regression





# **Logistic Regression**

from sklearn.linear\_model import LogisticRegression from
sklearn.metrics import accuracy\_score, plot\_confusion\_matrix
lr=LogisticRegression(max\_iter=10000) lr.fit(x\_train,y\_train)
p1=lr.predict(x\_test) s1=accuracy\_score(y\_test,p1) print("Linear
Regression Success Rate :", "{:.2f}%".format(100\*s1))
plot\_confusion\_matrix(lr, x\_test, y\_test) plt.show()



RESULT=86.67%

CONCLUSION

Heart is the most essential organ of the human body and day by day the loss of Human Life is increasing exponentially due to heart failure. It was experimentally found that the Global pandemic Corona Virus causes heart injury among a lot of patients. Hence there is an urgent need for research to focus into the causes for heart failure and to design a robust prediction system to detect at early stage so that loss of life can be avoided.

Even though there were many heart diseases prediction systems available at present but each one has its own limitations. The main objective of this research work is to overcome the difficulty faced by other researchers and to build a robust system which works efficiently and will able to predict accurately the possibility of heart attack at very early stage.

This research work could able to design a very robust and accurate model to predict the possibility of heart failure in the current scenario. By using the Logistic Regression this model could able to predict with an accuracy of about 86.6% which is highest as compared to other algorithms.

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