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Course :- DLBDSME01 Model Engineering Module



Acknowledgments

UCI ML Repository - Breast Cancer Wisconsin Dataset

IU Internationale Hochschule – DLBDSME01 Model Engineering Module

# Breast Cancer Prediction – Interpretable Machine Learning

This project builds an **interpretable machine learning model** to predict whether a breast tumor is **benign or** malignant, using the Wisconsin Breast Cancer Dataset and IU csv file.

It follows the CRISP-DM methodology and includes a Streamlit web app for medical professionals to interact with the model and understand its predictions.

The dataset consists of 569 records representing samples from breast tissue biopsies, each identified by a unique ID. The primary goal is to classify whether a tumor is malignant (cancerous) or benign (non-cancerous) based on a set of diagnostic features derived from medical imaging.

Each record includes a diagnosis column, which contains either an M (malignant) or B (benign) label. There are 357 benign and 212 malignant cases, indicating a moderately imbalanced dataset favoring benign outcomes.

The dataset contains 30 numeric features grouped into three sets:

Mean Values (e.g., radius\_mean, texture\_mean) - representing the average measurement for each characteristic.

Standard Error values (with \_se suffix) – indicating the variation or uncertainty of those measurements.

Worst-case values (with \_worst suffix) – the most extreme observed values across the tumor sample.

These features are based on ten core physical properties of cell nuclei, including:

Radius (distance from center to edge)

Texture (variation in grey-level intensity)

Perimeter

Area

Smoothness (local variations in radius)

Compactness, Concavity, Concave Points, Symmetry, and Fractal Dimension.

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Statistical analysis shows that benign tumors generally have lower values for features like radius, area, and concavity compared to malignant ones. For instance, the radius\_mean ranges from approximately 7 to 28, and the average is around 14, with higher values more often associated with malignant cases.

Lastly, the dataset includes a column labeled Unnamed: 32, which contains no values and can be safely removed during preprocessing.

In summary, the data is clean (no missing values), well-structured, and rich in diagnostic information, making it highly suitable for building predictive models. The balance of interpretability and numerical depth makes it a strong candidate for applying explainable machine learning techniques in a medical context.

## 

- Develop a classification model with **F1 score > 0.95**
- Focus on interpretability using SHAP
- Provide a prototype **GUI** for prediction and explanation

Do not run template.py because it will create new files and folder again and all code will be lost use template.py if you want to create files in another project

libraries (	(you d	can s	see in	requi	rement	t.txt)

pandas		
numpy		
scikit-learn		
joblib		
streamlit		
shap		
matplotlib		
seaborn		
pyyaml		

# **Important Notes**

src - It is consist of all main files to run

run.py - It runs evaluate\_model.py , explain\_model.py , train\_model.py it will run all three functions

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app - It consists of two apps which are based on Logistic regression and RandomForestClassifier to run both apps you need to type streamlit run appname(logistic\_reg.py in command prompt or terminal

notebooks - This notebook consist of eda.ipynb and model.ipynb

data - csv file for training model

Logger - I have created logger file to keep track of code which is in logs folder

#### How to run

You need to install necessary libraries which i have written in requirement.txt to run the project or you can create a new **enviroment** To see result in a form of browser you need to go in data/reports/breast\_cancer\_profiling\_report.html and from there you need to run live server

#### github project link :-

https://github.com/mandylegend/mandylegend-IU-Breast-Cancer-Model\_Project/tree/main you can fetch project from here and updated files

### Project Structure

```
breast-cancer-prediction/
- README.md
                             # This file
— LICENSE
                             # Project license (optional)
                             # Ignored files (e.g., data/, .pkl)
- .gitignore
— data/
   - raw/
                             # Original dataset (CSV)
   - reports/
                           # HTML file
notebooks/
  ├─ 01_eda.ipynb
                            # Exploratory Data Analysis
 - src/
   preprocessing/
      └─ clean_data.py # Cleaning functions
    — models/
      train_model.py # Train and save model
        - evaluate_model.py # Evaluate metrics and F1
      explain_model.py # SHAP-based explanations
    — utils/
      └─ helpers.py
   Logger
```

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