

# **GALLSTONE DISEASE PREDICTION WITH EXPLAINABLE ARTIFICIAL INTELLIGENCE**

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**CMSC 177**

# OVERVIEW

- Introduction
- Literary Review
- Methodology
- Result
- Conclusion
- Recommendation

# BACKGROUND

- Gallstones are quite common and influenced by age, comorbidities, and body composition.
- The growth of clinical data creates opportunities for machine learning-based early prediction.
- Early and accurate diagnosis can greatly improve quality of life.

# TRADITIONAL METHODS

- Diagnosis mostly through imaging after symptoms appear.
- Risk assessments are often subjective and limited.
- Blood tests alone aren't predictive.
- These approaches miss early or asymptomatic cases.

# MACHINE LEARNING

- Offers early, data-driven prediction.
- Models used: Random Forest, SVM, Gradient Boosting, Logistic Regression
- Use of SHAP and LIME for model transparency and clinical trust.

# EXPECTED CONTRIBUTION

- Goal: Predict gallstones using explainable ML
- Benefits: early detection, support decision-making.
- Steps: feature selection → model training → evaluation → interpretation

# LITERATURE REVIEW

## Machine Learning in Medical Diagnostics

Machine learning is being used more and more to accurately identify a variety of medical conditions, enabling quicker, data-driven diagnoses that enhance early intervention and lower errors.

## Machine Learning for Gallstone Prediction

Gallstone risk can be reliably predicted by models that integrate metabolic data, inflammation markers, and body composition. They provide cheap, non-invasive substitutes for conventional imaging.

# LITERATURE REVIEW

## Explainable AI (XAI)

By clarifying the decision-making process of models, XAI tools such as SHAP and LIME develop clinician trust and promote the moral and open application of AI in healthcare.

## XAI in Practice

Research on gallstones is still in early stages, but XAI has advanced knowledge of other illnesses. By using it in gallstone models, important risk factors are highlighted and individualized care is supported.

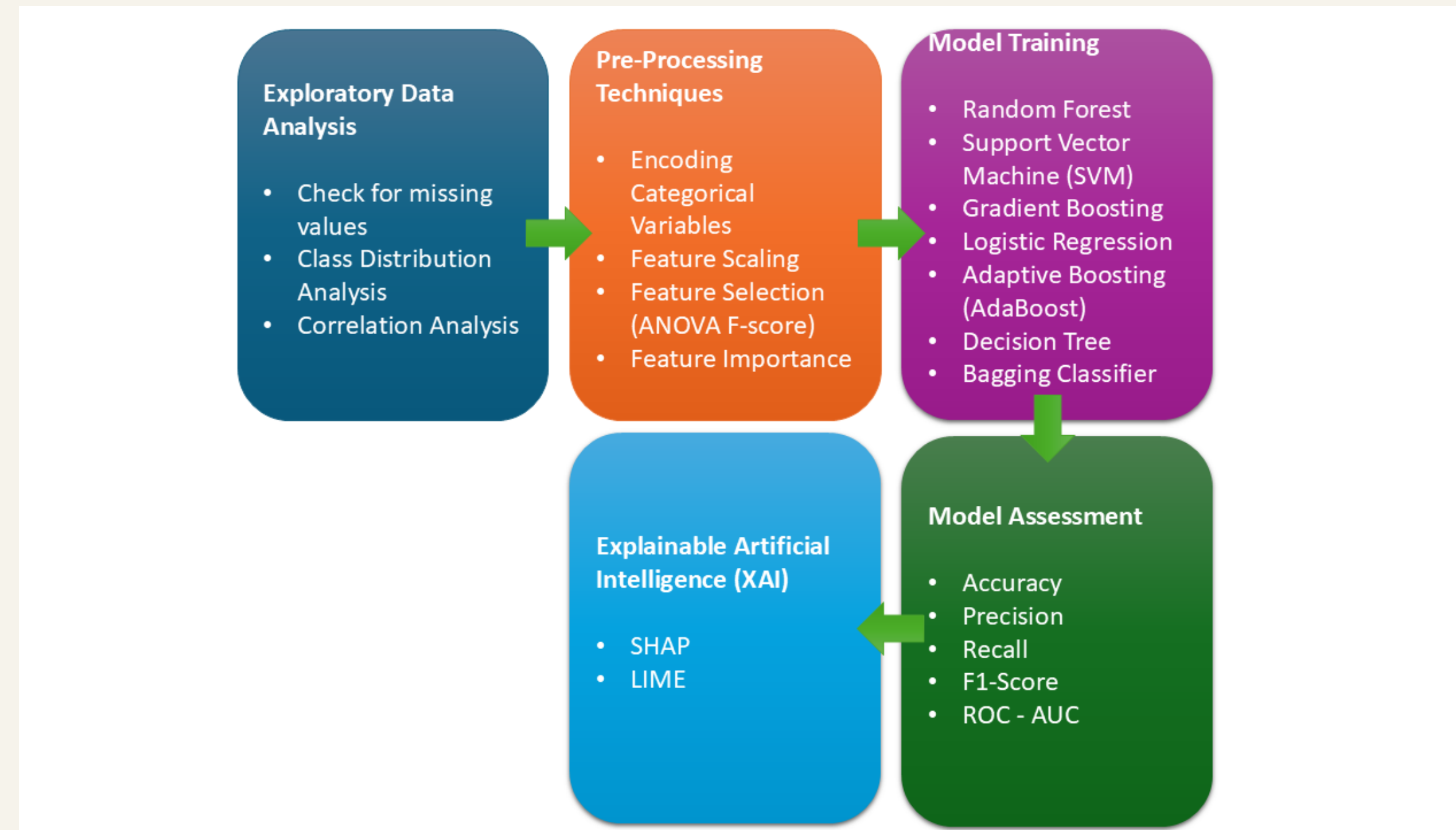
# LITERATURE REVIEW

## Effective Models and Features

Complex, unbalanced data can be effectively handled by algorithms such as Random Forest and Gradient Boosting. For improved performance and clarity, feature selection guarantees that models concentrate on the most crucial clinical variables.

# METHODOLOGY

## Machine Learning Pipeline



# I. DATASET DESCRIPTION

The dataset used in this study was sourced from a publicly available repository for gallstone disease classification. It consists of 319 patient records, each containing real-valued clinical and laboratory features, with the target label **Gallstone Status** indicating the presence or absence of gallstones.

The dataset was confirmed to have no missing values and was balanced across target classes. Each record contains features derived from medical parameters such as lean mass, extracellular water, and vitamin levels, which have shown predictive value in prior studies. This structure enabled robust machine learning development without the need for oversampling techniques.

# 2. DATA PRE-PROCESSING

## Initial Checks & Findings

### Checks Conducted:

- Missing values
- Duplicates
- Outlier presence
- Class distribution

### Findings:

- No missing or duplicate values
- Dataset was balanced
- Clinical outliers were present but retained for relevance

## Data Cleaning & Transformation

### Cleaning Steps:

- Verified dataset integrity (319 complete records)
- No imputation or oversampling needed

### Transformation:

- Applied standard scaling (zero mean, unit variance)
- Ensured uniformity across numeric clinical features

## Notes on Outliers and Balance

- Outliers retained after boxplot analysis
- Dataset balance → no need for SMOTE
- Preprocessing supported model convergence and interpretability

# 2. DATA PRE-PROCESSING

## Exploratory Data Analysis (EDA)

### Descriptive Statistics

- Computed mean, median, standard deviation, min, and max
- Helped analyze key features like Lean Mass, ECF/TBW, and CRP
- Supported data normalization (standard scaling)

### Correlation Analysis & Feature Selection

- Generated correlation heatmaps to explore inter-feature relationships
- Informed feature selection process using ANOVA F-value (SelectKBest)
- Threshold optimized via cross-validation for best model performance

### Visualization & Outlier Detection

- Used histograms and boxplots to explore distributions
- Assessed clinical outliers and retained them for relevance
- Helped validate class balance and variable spread

### Split Strategy

- Dataset divided into 70% training and 30% testing using stratified split
- Ensured balanced target distribution across training and test sets

## Descriptive Statistics

```

print("\nDescriptive statistics for numerical features:")
# display.display(df[num_features].describe())
display(df[num_features].describe())
✓ 0.0s

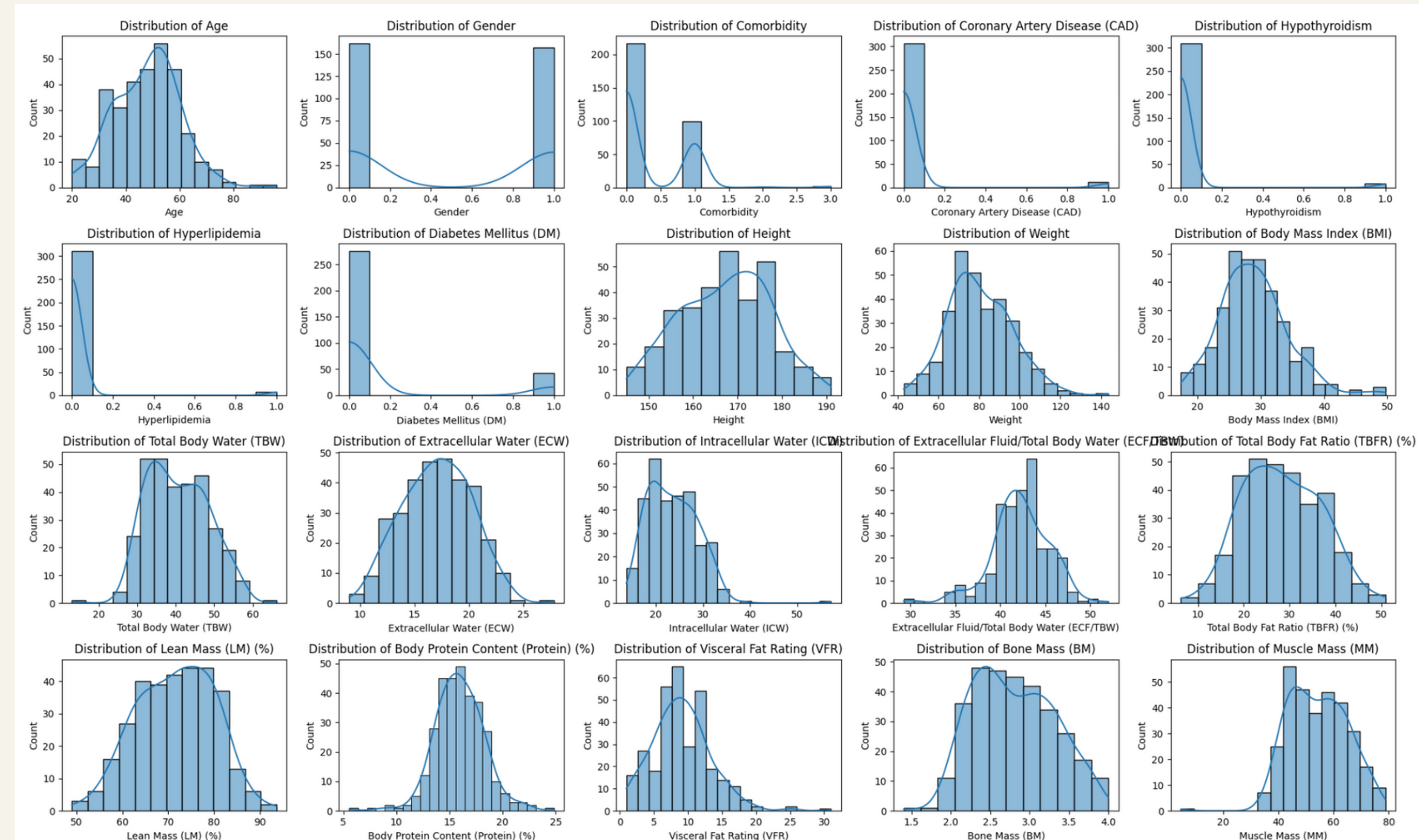
```

Descriptive statistics for numerical features:

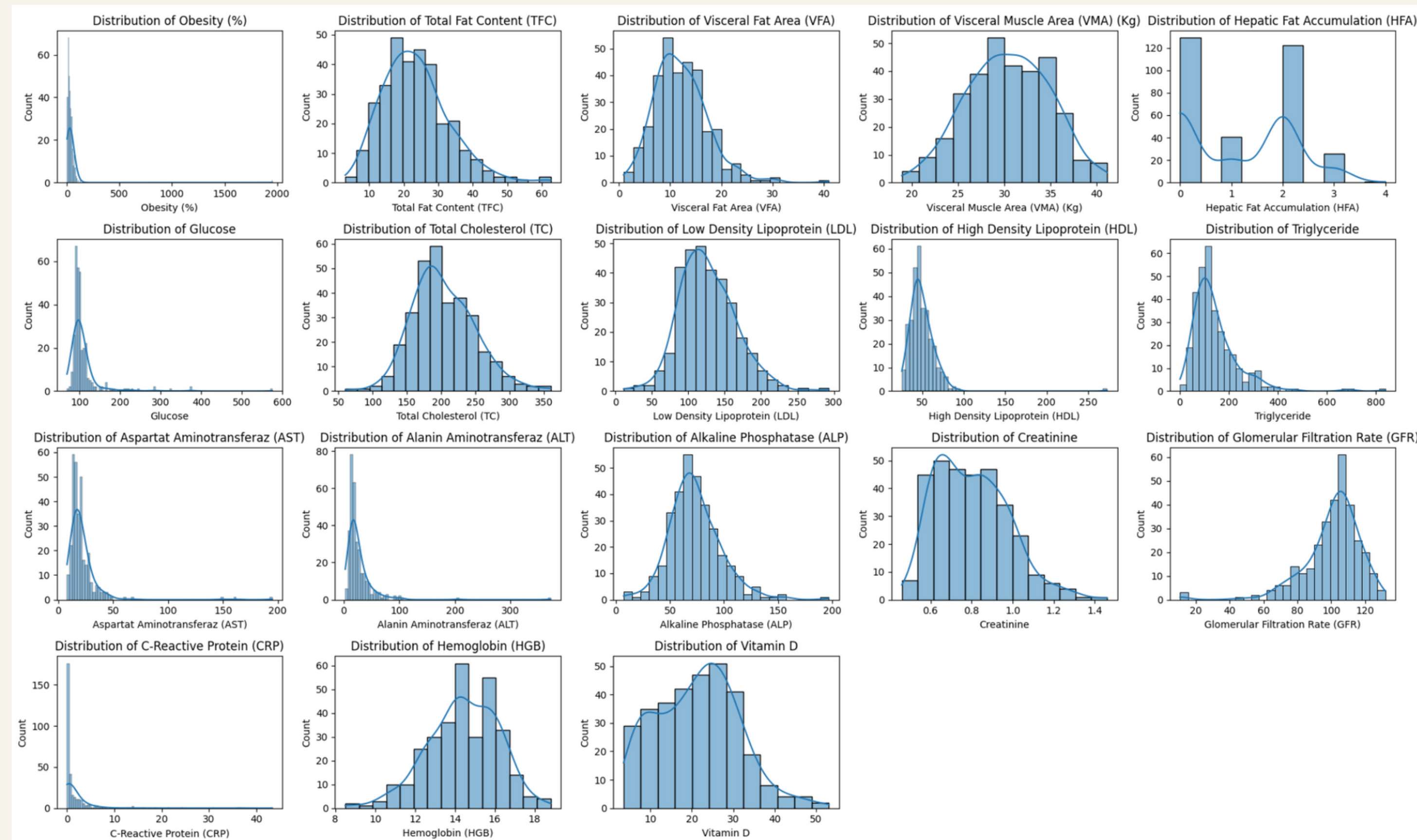
	Age	Gender	Comorbidity	Coronary Artery Disease (CAD)	Hypothyroidism	Hyperlipidemia	Diabetes Mellitus (DM)	Height	Weight	Body Mass Index (BMI)	...
count	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	...
mean	48.068966	0.492163	0.335423	0.037618	0.028213	0.025078	0.134796	167.15674	80.564890	28.877116	...
std	12.114558	0.500724	0.517340	0.190568	0.165841	0.156609	0.342042	10.05303	15.709069	5.313707	...
min	20.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	145.00000	42.900000	17.400000	...
25%	38.500000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	159.50000	69.600000	25.250000	...
50%	49.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	168.00000	78.800000	28.300000	...
75%	56.000000	1.000000	1.000000	0.000000	0.000000	0.000000	0.000000	175.00000	91.250000	31.850000	...
max	96.000000	1.000000	3.000000	1.000000	1.000000	1.000000	1.000000	191.00000	143.500000	49.700000	...

High Density Lipoprotein (HDL)	Triglyceride	Aspartat Aminotransferaz (AST)	Alanin Aminotransferaz (ALT)	Alkaline Phosphatase (ALP)	Creatinine	Glomerular Filtration Rate (GFR)	C-Reactive Protein (CRP)	Hemoglobin (HGB)	Vitamin D
319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000	319.000000
49.475549	144.502163	21.684953	26.855799	73.112539	0.800611	100.818903	1.853856	14.418182	21.401411
17.718701	97.904493	16.697605	27.884413	24.181069	0.176433	16.971396	4.989591	1.775815	9.981659
25.000000	1.390000	8.000000	3.000000	7.000000	0.460000	10.600000	0.000000	8.500000	3.500000
40.000000	83.000000	15.000000	14.250000	58.000000	0.650000	94.170000	0.000000	13.300000	13.250000
46.500000	119.000000	18.000000	19.000000	71.000000	0.790000	104.000000	0.215000	14.400000	22.000000
56.000000	172.000000	23.000000	30.000000	86.000000	0.920000	110.745000	1.615000	15.700000	28.060000
273.000000	838.000000	195.000000	372.000000	197.000000	1.460000	132.000000	43.400000	18.800000	53.100000

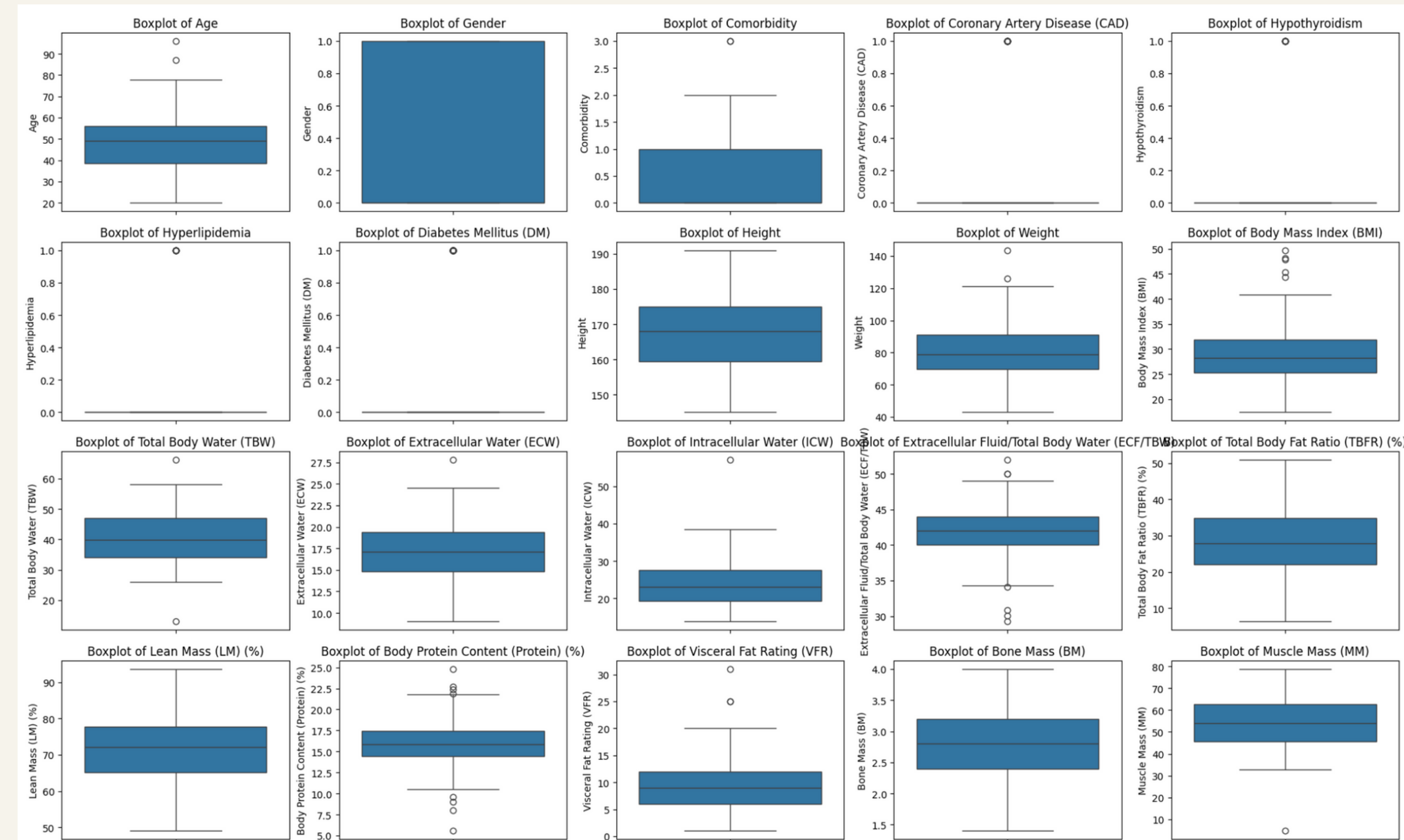
# Histograms of Numerical Features



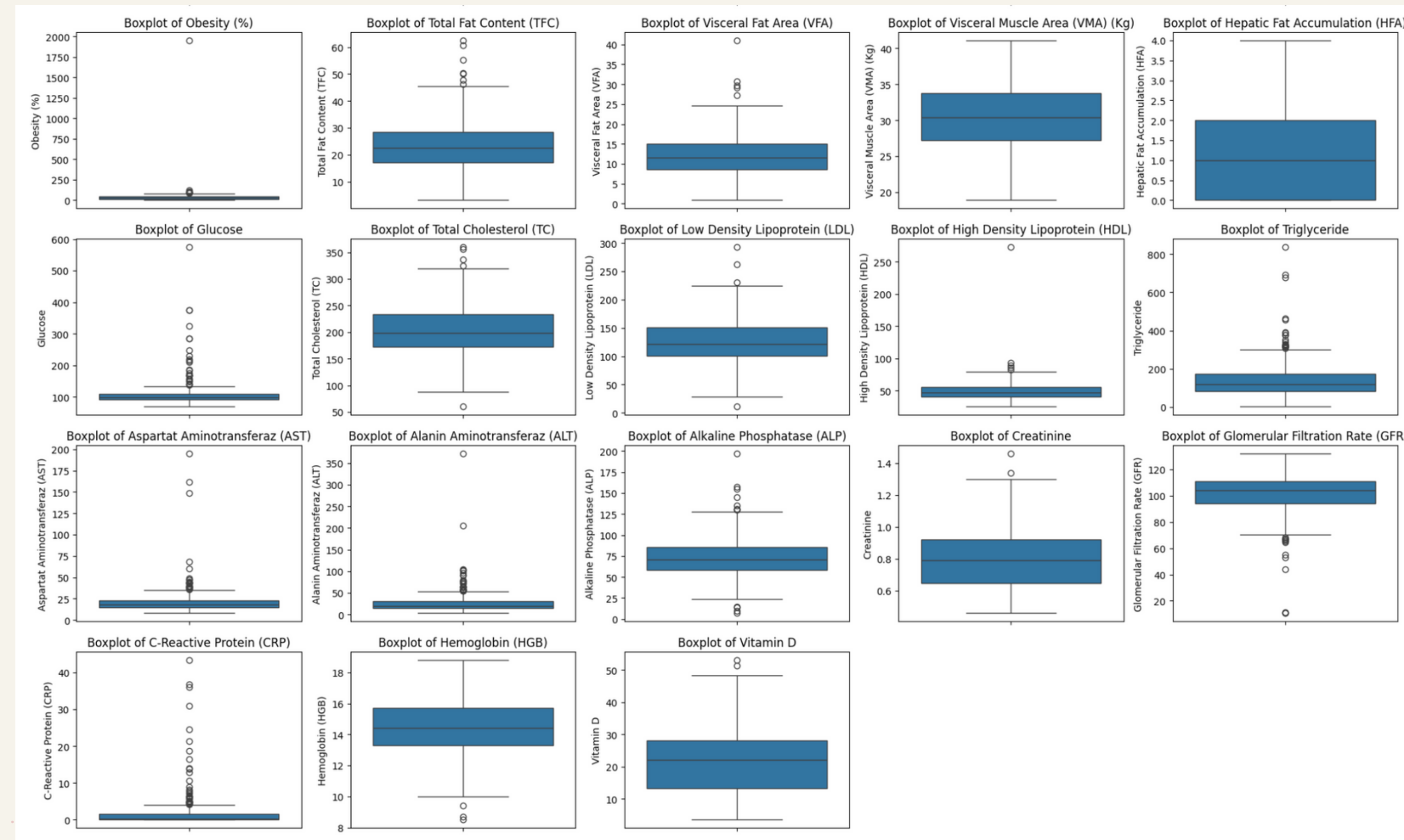
# Histograms of Numerical Features

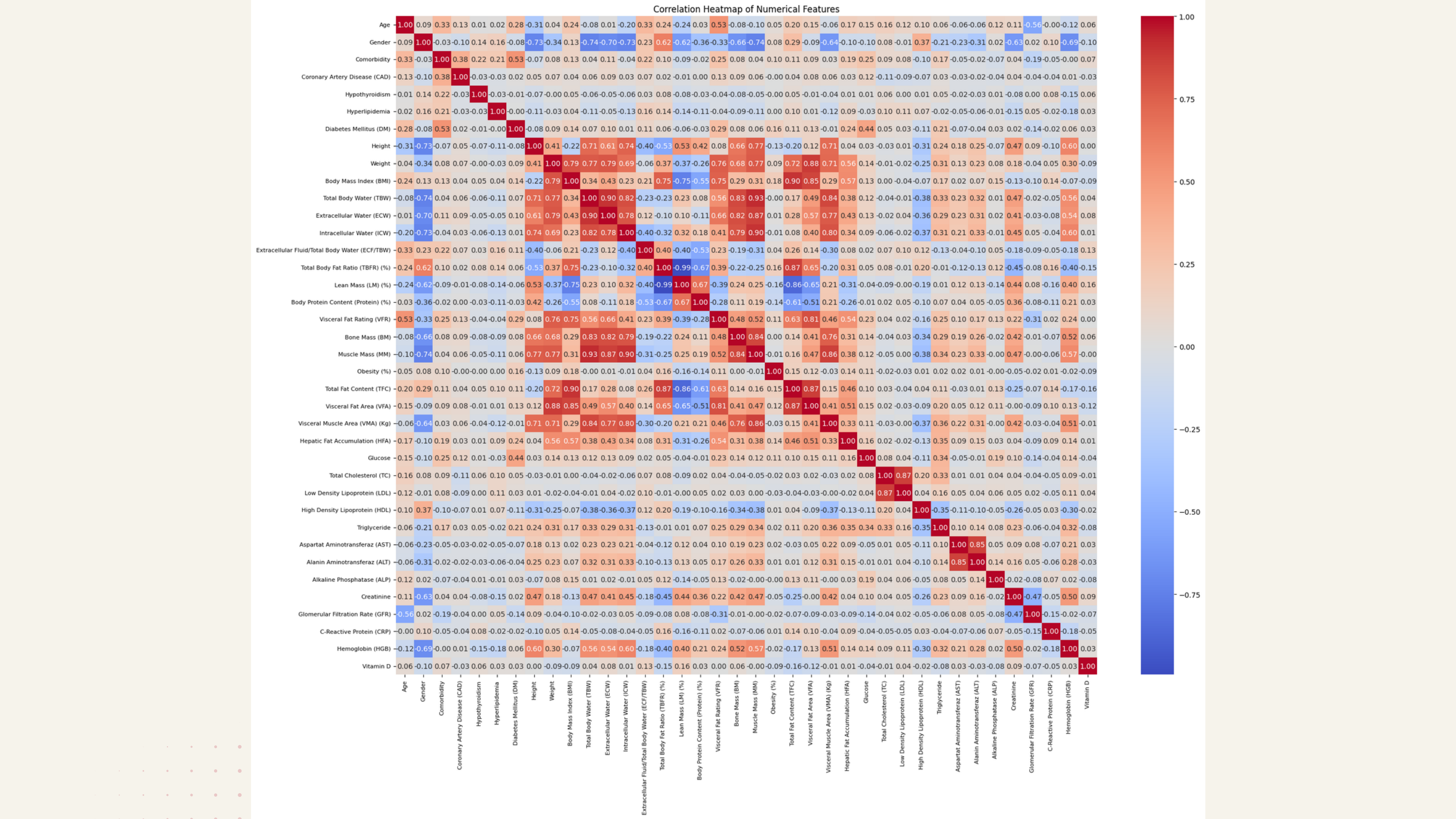


# Boxplots of Numerical Features

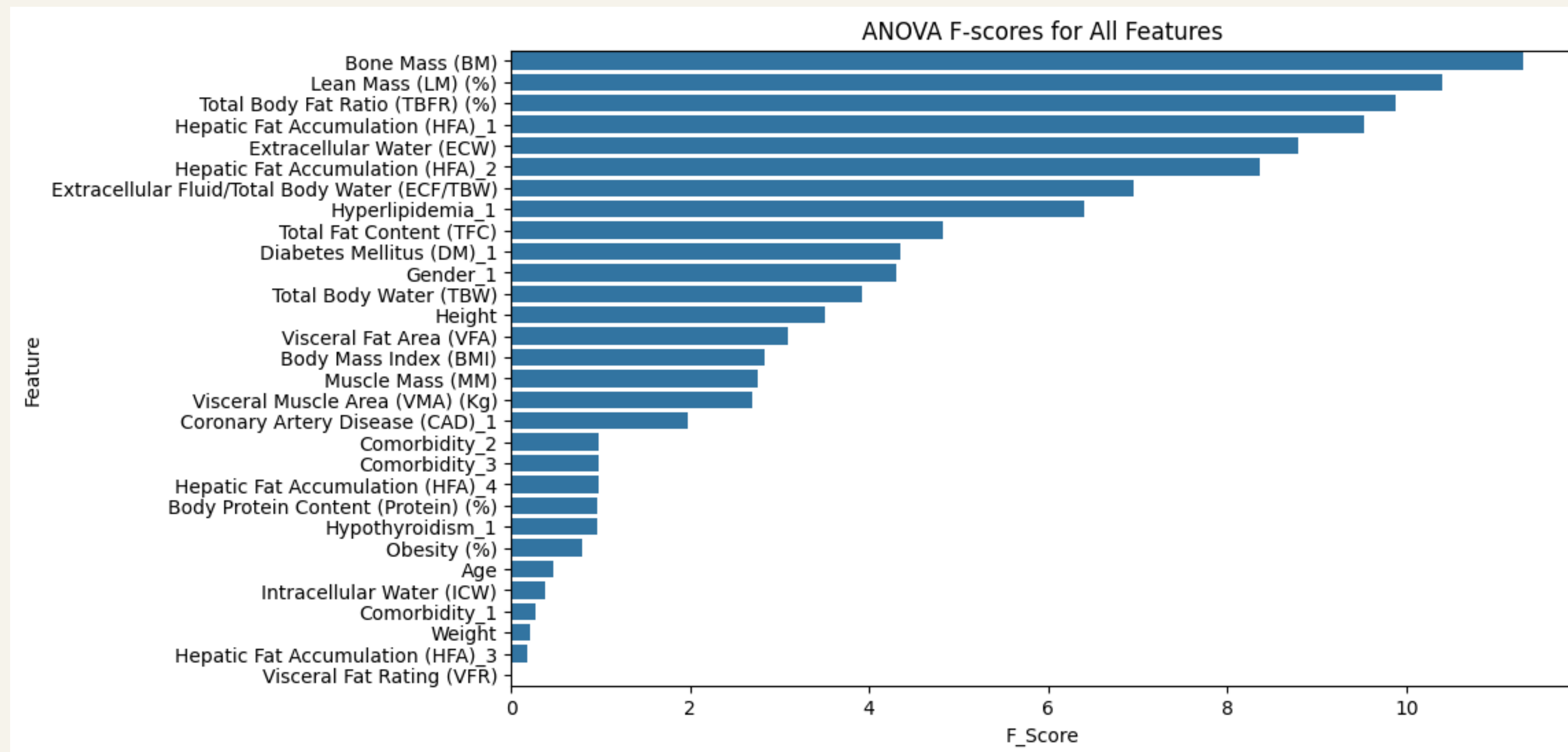


# Boxplots of Numerical Features





## ANOVA F-Scores



### ANOVA F-Score Threshold

- A threshold at the 75<sup>th</sup> percentile was set to retain the most relevant features.
- After application, the top 8 features were selected

# MACHINE LEARNING MODELS

## Train-Test Split

A Train-Test split of 70-30 was selected.

## ML Algorithms

- Random Forest
- Support Vector Machine
- Gradient Boosting
- Logistic Regression
- Adaptive Boosting
- Decision Tree
- Bagging Classifier

## Performance Metrics

- Accuracy
- Precision
- Recall
- F1-Score
- AUC

## Validation

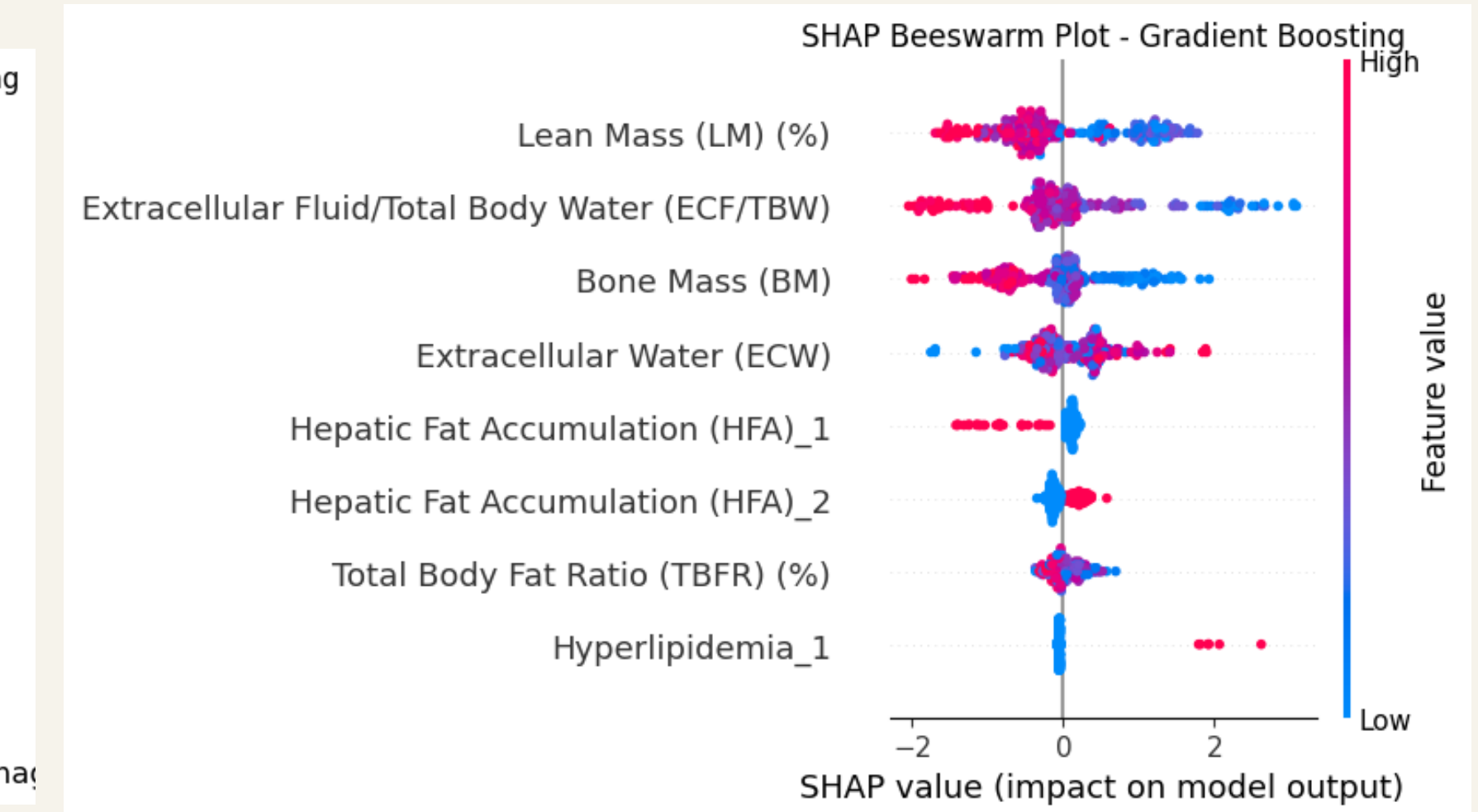
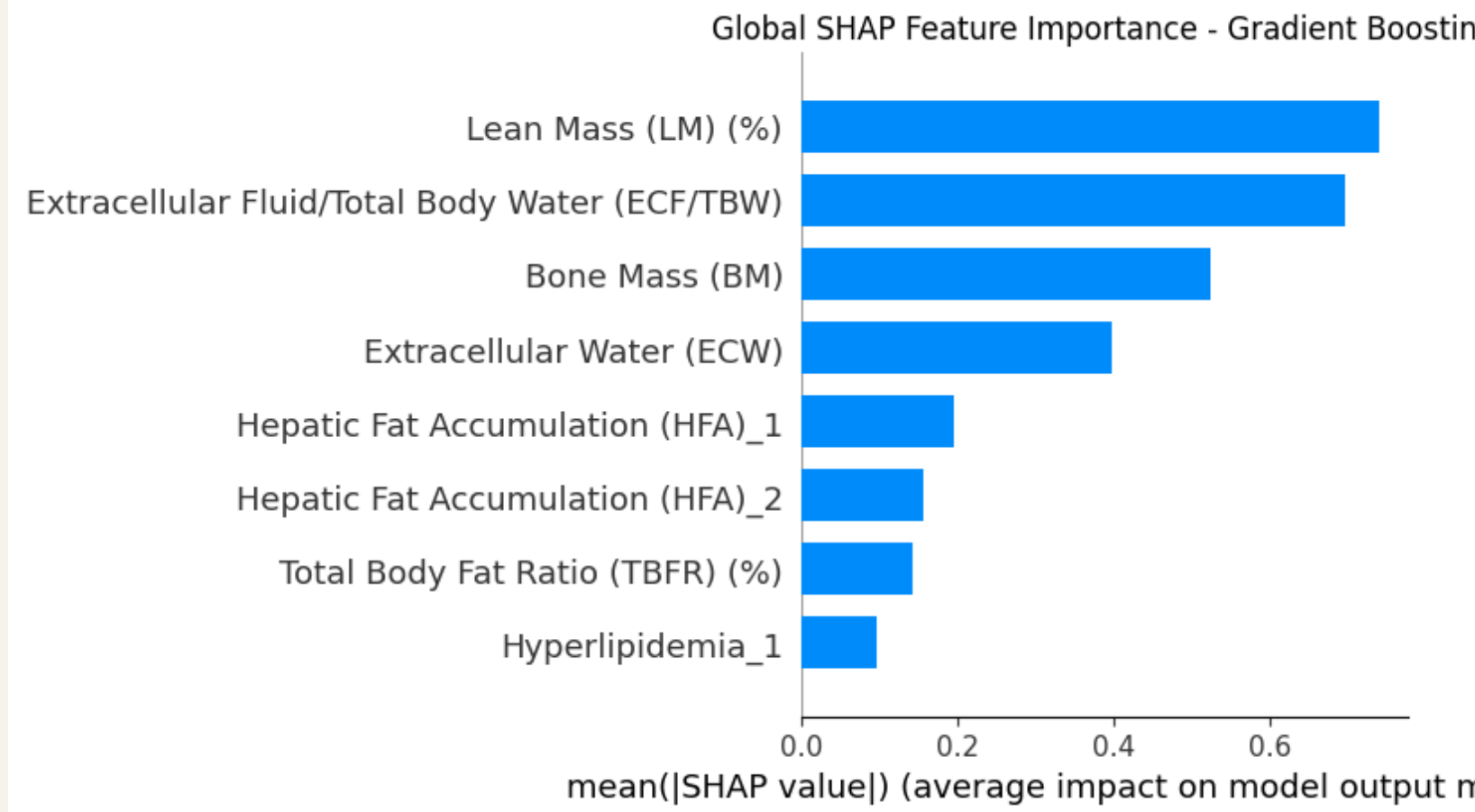
5-fold cross-validation was implemented to reduce overfitting

# RESULT

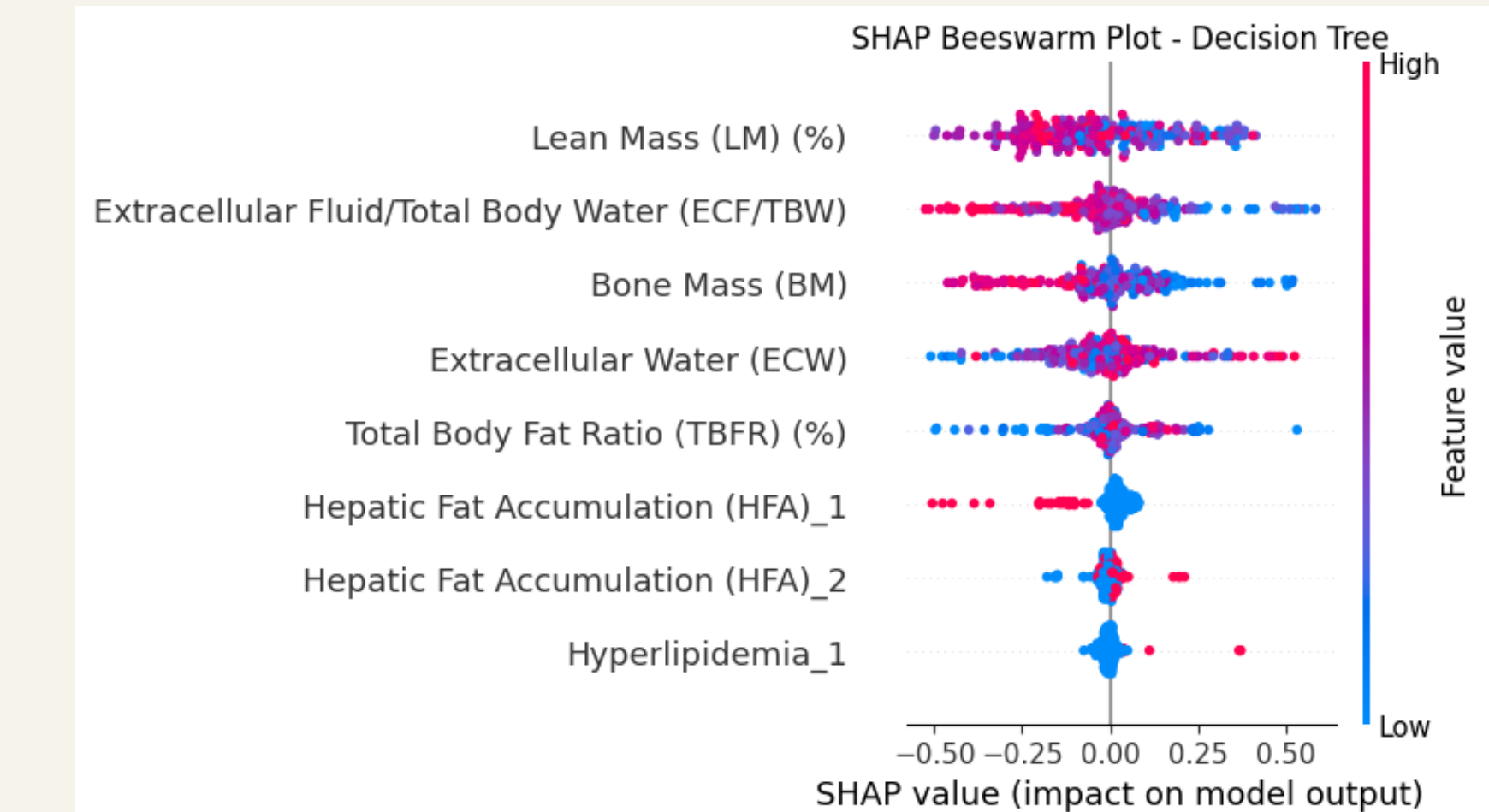
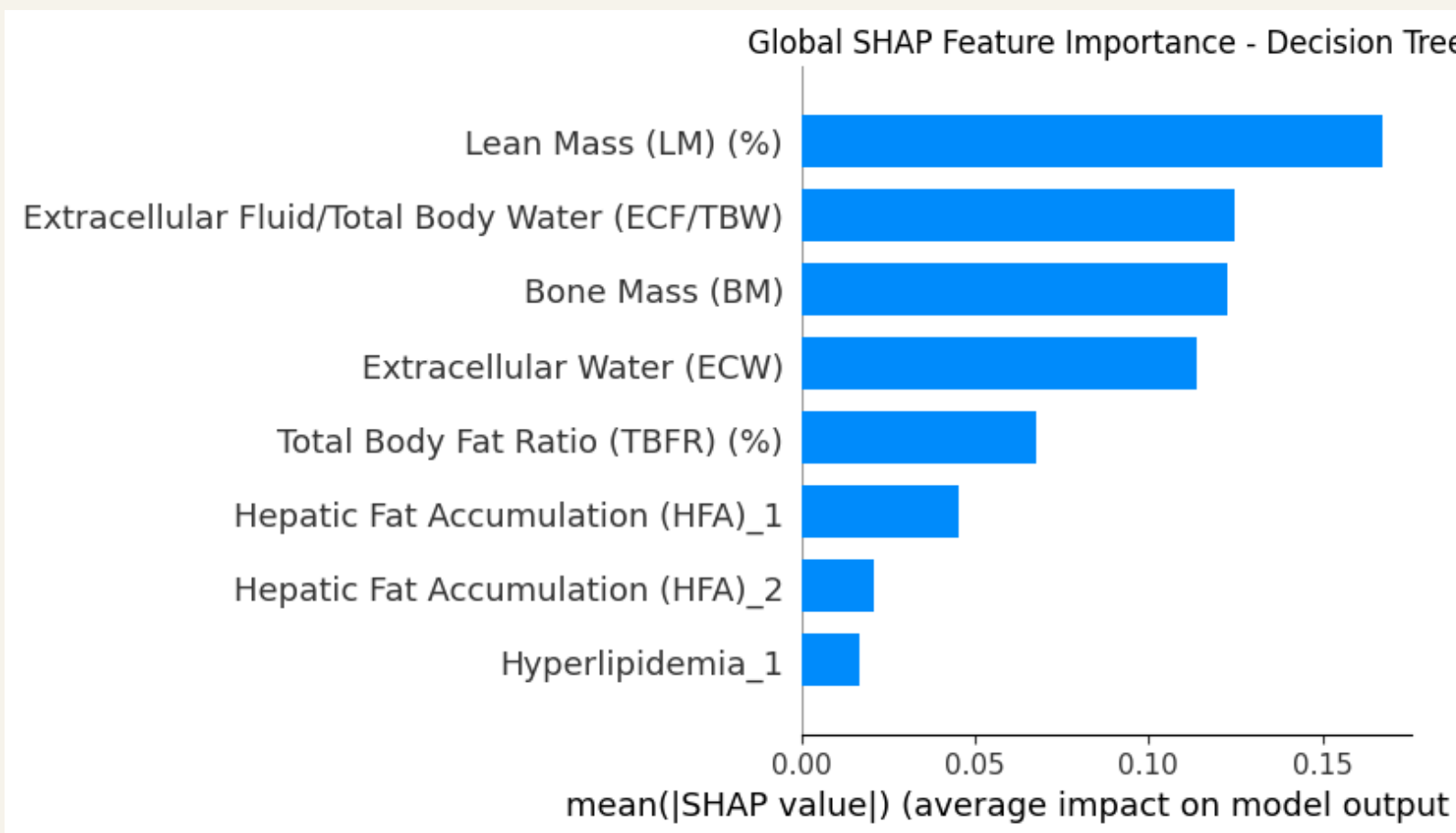
ML Model	Accuracy	Precision	Recall	F1 Score	AUC
RF	0.75	0.79	0.69	0.73	0.79
SVM	0.75	0.94	0.53	0.68	0.82
GB	0.72	0.71	0.75	0.73	0.81
LR	0.77	0.81	0.69	0.75	0.86
AB	0.69	0.70	0.66	0.68	0.79
DT	0.73	0.74	0.72	0.73	0.73
BC	0.73	0.76	0.69	0.72	0.75

Best model in terms of relevant metrics (recall): Gradient Boosting

# XAI EXPLAINERS: SHAP

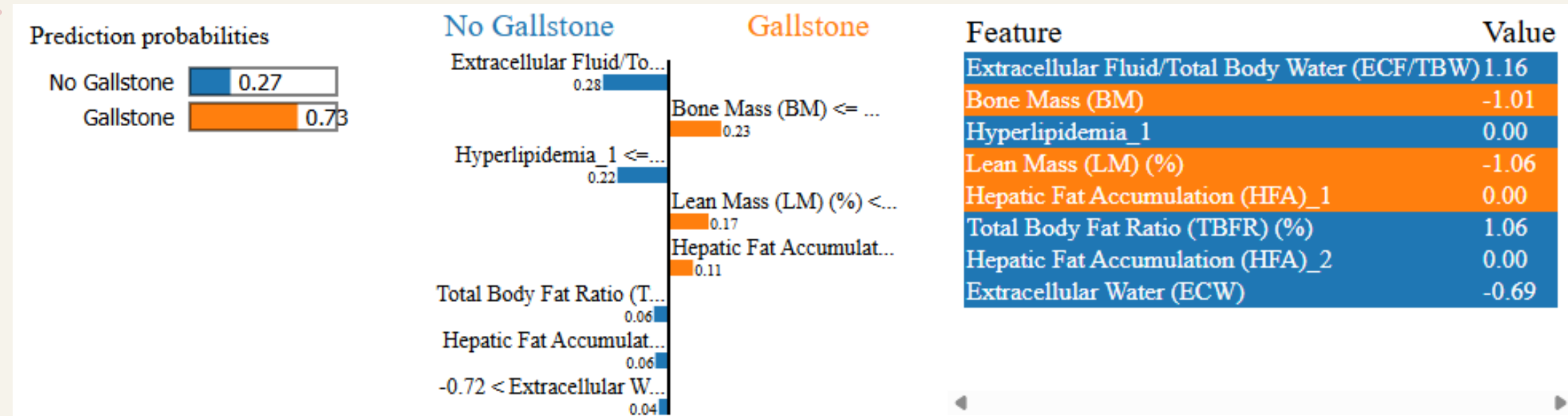


# XAI EXPLAINERS: SHAP

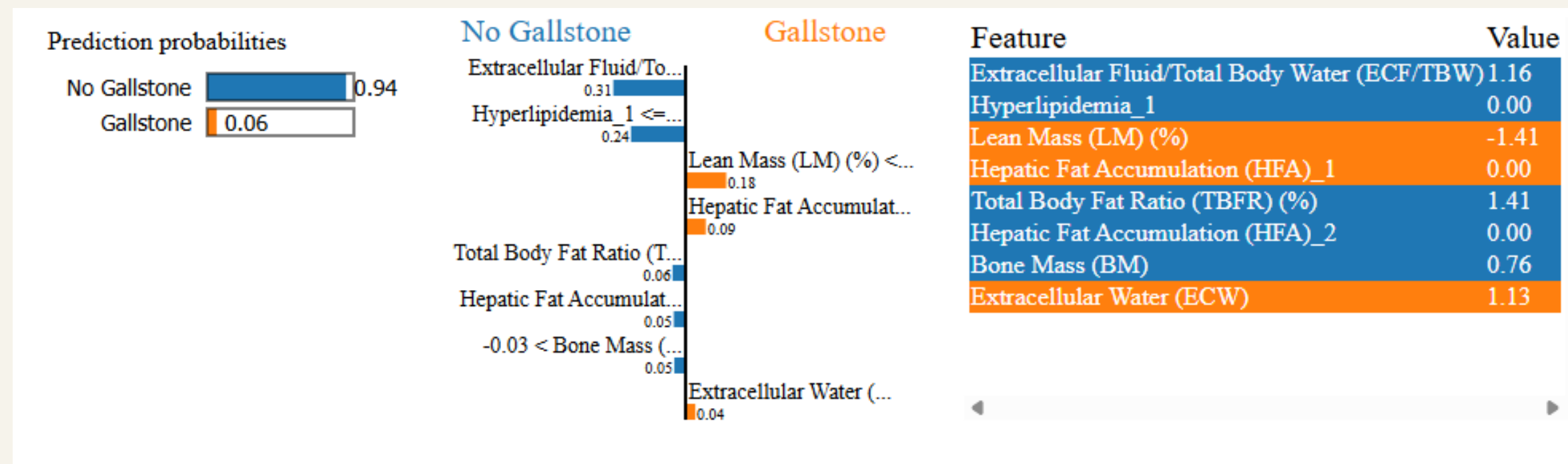


# XAI EXPLAINERS: LIME

Positive  
Sample



Negative  
Sample



# CONCLUSION

This study used clinical and physiological data to evaluate the predictive power of several machine learning models for gallstone disease. With a 75% recall rate, gradient boosting performed the best. By adding SHAP and LIME, the model became more transparent and the results were easier to interpret clinically. All things accounted for, the models hold potential for early diagnosis and individualized risk assessment in patient care.

# RECOMMENDATION

## Expand Dataset Diversity

To increase the generalizability and dependability of the model, incorporate data from more sources and larger populations.

## Incorporate Advanced Features

To find more profound predictive patterns, use imaging, genetic information, and longitudinal records.

## Conduct External Validation

To confirm the models' applicability and practical impact, test them on independent datasets or in actual clinical settings.

## Improve Preprocessing Techniques

Improve input quality and model performance by implementing more thorough data-cleaning and transformation techniques.

## Explore Additional Modeling Approaches

To possibly boost predictive power, try out new approaches and ensemble techniques like stacking.

# RECOMMENDATION

## ● Conduct External Validation

To confirm the models' applicability and practical impact, test them on independent datasets or in actual clinical settings.

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