

BioMLStudio

Machine Learning Analysis Report

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1. Dataset Summary

Property	Value
Dataset Name	seq_001_affected.fasta
Dataset Type	dna
Total Samples	40
Features	71
File Size	0.01 MB

2. Preprocessing Steps

Step 1: Load and Clean Data

- Loaded 40 sequences from FASTA
- Removed invalid characters
- Extracted labels from headers
- Label distribution: {'affected': 20, 'normal': 20}

Step 2: Handle Missing Values

- No missing values found

Step 3: Feature Engineering (Biological)

- Added 6 engineered features

Step 4: Sequence Encoding (kmer)

- Applied k-mer encoding (k=3)

Step 5: Feature Normalization (standard)

- Normalized 71 features using standard scaling

Step 6: Data Splitting

- Encoded target variable (2 classes)
- Split data: 28 train, 4 val, 8 test

3. Model Selection & Training

Model	Training Time	Score
Logistic Regression	2.63s	1.0000
Random Forest	0.18s	1.0000
Gradient Boosting	0.07s	1.0000

Best Model: Logistic Regression

4. Performance Metrics

Training Metrics:

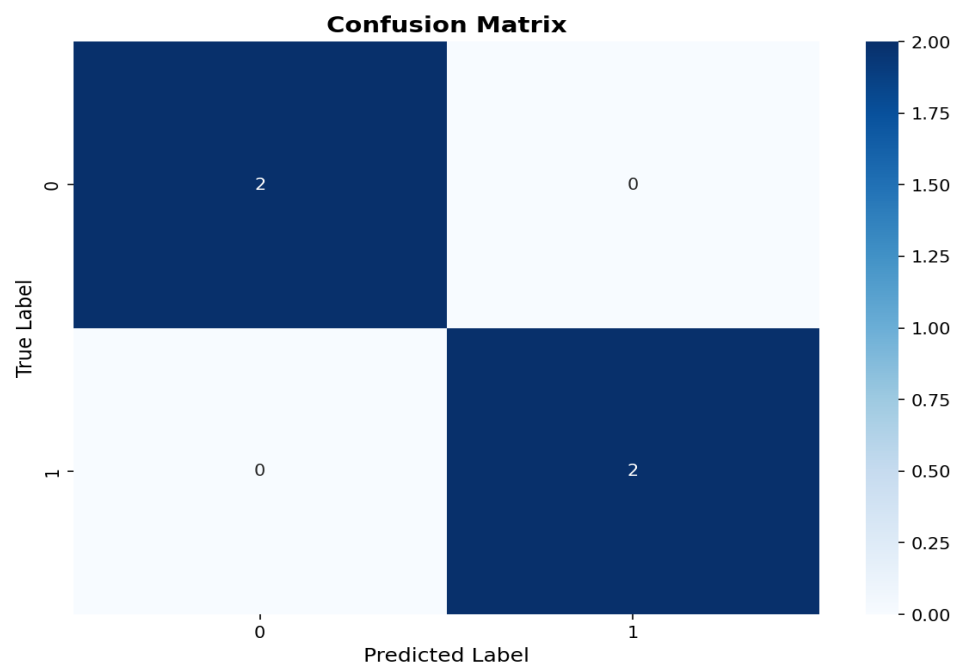
Metric	Value
Accuracy	1.0000
Precision	1.0000
Recall	1.0000
F1 Score	1.0000
Roc Auc	1.0000

Validation Metrics:

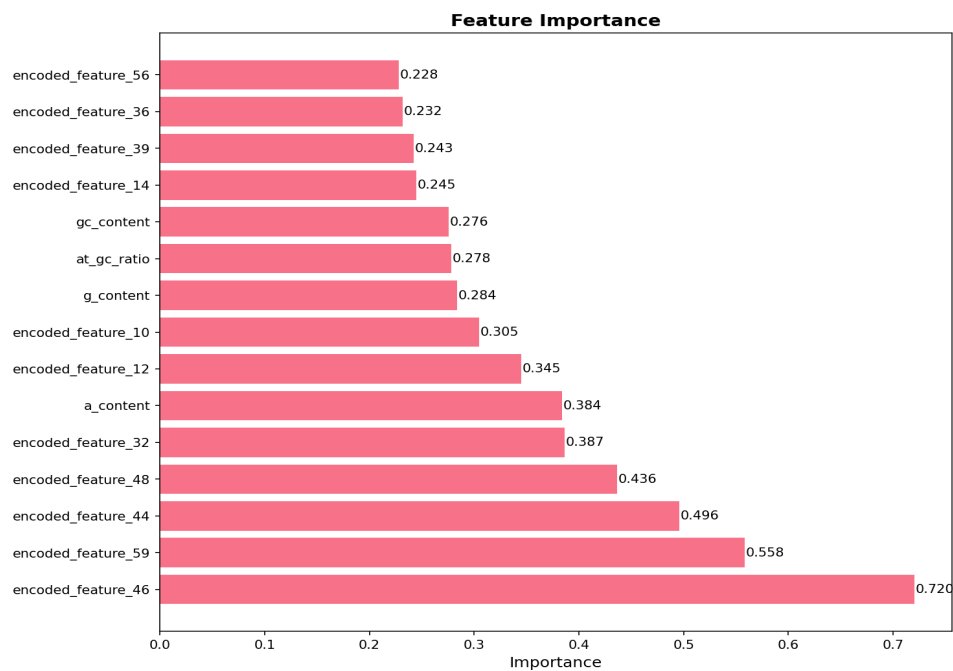
Metric	Value
Accuracy	1.0000
Precision	1.0000
Recall	1.0000
F1 Score	1.0000
Roc Auc	1.0000

5. Visualizations

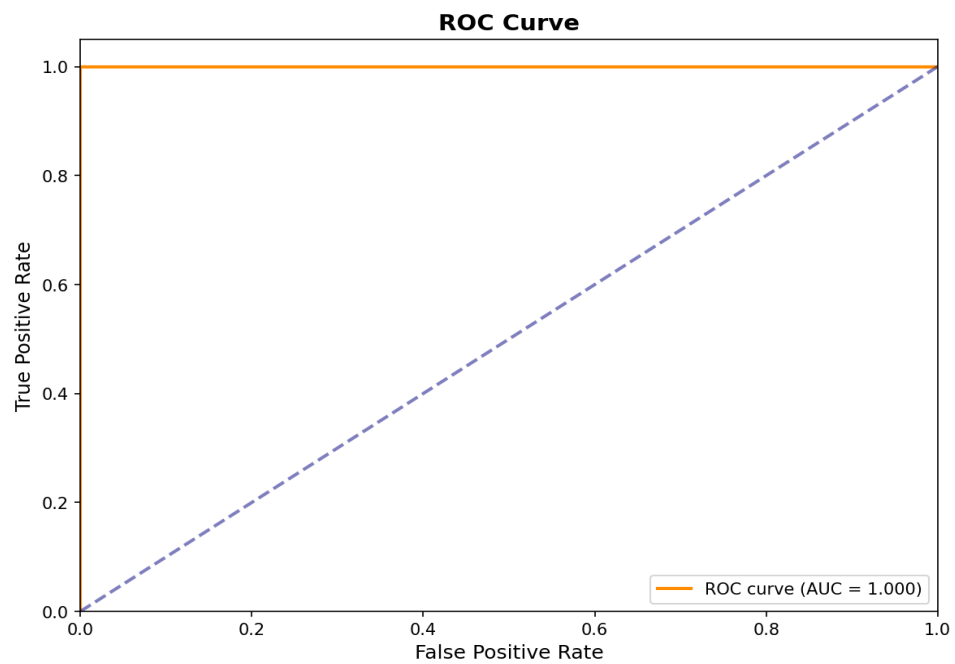
Confusion Matrix



Feature Importance



Roc Curve



6. Training Summary

Total training time: 4.43 seconds

Key Events:

[SUCCESS] Logistic Regression - Score: 1.0000

[SUCCESS] Random Forest - Score: 1.0000

[SUCCESS] Gradient Boosting - Score: 1.0000

[SUCCESS] Best model: Logistic Regression (Score: 1.0000)

Report generated by BioMLStudio - AI-Based No-Code Platform for Bioinformatics

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