Wine Quality Prediction using the CRISP-DM Methodology

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Abstract

This paper presents a detailed approach to predicting wine quality using the CRISP-DM (Cross-Industry Standard Process for Data Mining) methodology. We explore multiple machine learning models, from simple regression to ensemble methods, to develop a model that predicts wine quality based on its chemical properties. Through the CRISP-DM framework, we systematically address each phase, from data understanding to deployment, while discussing challenges such as preprocessing errors, model optimization issues, and prediction calibration. Our final model—a calibrated weighted ensemble—improved performance, demonstrating the utility of CRISP-DM in practical machine learning applications.

1 Introduction

Predicting wine quality based on chemical properties is a widely studied problem in data science. This study applies the CRISP-DM methodology, which provides a structured approach for developing machine learning models. The CRISP-DM process ensures that every critical aspect of a data science project is covered, including business understanding, data preparation, model building, and evaluation.

This paper details each step of the CRISP-DM process and highlights the challenges encountered, such as errors in data preprocessing, the handling of optimized models, and model calibration for better prediction accuracy.

2 Business Understanding

The objective of this study is to predict the quality of red wine based on its chemical attributes, such as acidity, sugar content, and alcohol levels. Wine producers and distributors can benefit from such a predictive model by optimizing production processes and improving quality control.

The prediction task was framed as a regression problem, where the goal was to predict a continuous quality score between 0 and 10 using the wine's chemical features.

3 Data Understanding

The dataset comes from Kaggle's Wine Quality Dataset, containing chemical properties of red wine along with an expert-rated quality score. This dataset includes 11 independent variables (features) and one dependent variable (wine quality).

3.1 Initial Exploration

We began by exploring the dataset to understand its structure and identify potential issues:

- **Descriptive Statistics**: We examined the distribution of each feature. For instance, we observed outliers in features such as *citric acid* and *residual sugar*, which required further handling.
- Correlation Analysis: A correlation matrix was generated to assess the relationships between features. Notably, *alcohol* was positively correlated with wine quality, while *volatile acidity* showed a negative correlation.

3.2 Key Insights

The correlation heatmap revealed several important relationships:

- Alcohol had a strong positive correlation with wine quality.
- Fixed acidity showed little to no correlation with wine quality.
- Volatile acidity exhibited a negative correlation, suggesting that higher acidity negatively impacts quality.

4 Data Preparation

Data preparation was crucial to ensure the models received high-quality inputs. This phase involved outlier handling, feature scaling, and feature selection.

4.1 Outlier Detection and Capping

Outliers in variables such as *residual sugar* and *citric acid* could skew the model's performance. We applied capping at the 1st and 99th percentiles to mitigate this issue.

Challenge: Initially, outlier capping was applied *after* scaling, which distorted the results. Upon realizing the mistake, we reordered the steps, first capping outliers and then applying scaling.

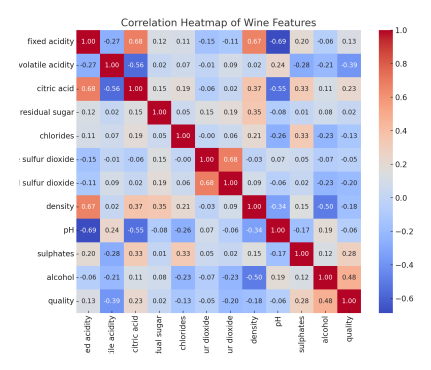


Figure 1: Correlation Heatmap of Wine Features

4.2 Feature Scaling

After addressing outliers, we used *StandardScaler* to standardize the features, ensuring all variables were on a similar scale. This is especially important for models sensitive to feature magnitudes, such as Support Vector Machines (SVM).

4.3 Feature Selection with Recursive Feature Elimination (RFE)

To reduce the dimensionality of the dataset, we applied *Recursive Feature Elimination* (RFE) using Random Forest as the base estimator. This process helped identify the most important features:

- Alcohol
- Volatile Acidity
- Sulphates
- Total Sulfur Dioxide

• Chlorides

5 Modeling

We experimented with multiple machine learning models, comparing their performance to predict wine quality.

5.1 Linear Regression

We began with *Linear Regression* as a baseline model. This model provided a simple benchmark, but the R² score was relatively low, indicating that linearity alone could not fully capture the relationships in the dataset.

5.2 Random Forest

Next, we used *Random Forest*, an ensemble model known for handling non-linear relationships and capturing feature interactions. This model performed significantly better than Linear Regression, with an R² score of approximately 0.493.

5.3 XGBoost

We then applied XGBoost, a gradient boosting model that builds trees sequentially and optimizes for speed and accuracy. XGBoost outperformed Random Forest, achieving an \mathbb{R}^2 score of 0.514.

5.4 Support Vector Machines (SVM)

While SVM is typically powerful for classification tasks, its performance in this regression problem was suboptimal compared to Random Forest and XGBoost. The model's complexity made it less suitable for this dataset.

5.5 K-Nearest Neighbors (KNN)

The K-Nearest Neighbors model also performed poorly. Since KNN is a distance-based model, it struggled with high-dimensional data, leading to lower accuracy and longer computation times.

5.6 Hyperparameter Tuning

After identifying Random Forest and XGBoost as the best-performing models, we used RandomizedSearchCV to optimize their hyperparameters. Initially, we encountered issues loading the optimized models due to version mismatches. After adjusting the local environment to match the versions used by ChatGPT, the models were successfully optimized and performed better than their default versions.

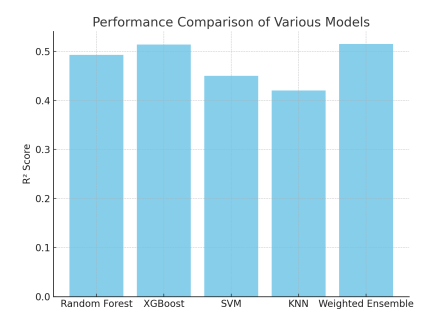


Figure 2: Performance Comparison of Various Models

6 Ensemble Modeling

To further improve performance, ChatGPT suggested creating a weighted averaging ensemble that combined the predictions from both Random Forest and XGBoost. The weights were based on the R² scores of the individual models.

$$y_{ensemble} = w_{rf} \cdot y_{rf} + w_{xgb} \cdot y_{xgb} \tag{1}$$

The ensemble model achieved an $\mathbf{R^2}$ score of 0.515, outperforming each model individually.

7 Model Calibration

Although the ensemble model performed well, it had a tendency to:

- Overestimate lower-quality wines
- \bullet Underestimate higher-quality wines

To address this, we applied *Isotonic Regression* for model calibration. After calibration, the model's \mathbb{R}^2 score improved to 0.524, and the predictions became more accurate, particularly for extreme cases.

8 Evaluation

The evaluation phase involved cross-validation and residual analysis to ensure model robustness and to identify areas for improvement.

8.1 Cross-Validation

We performed 5-fold cross-validation to evaluate the generalizability of the model. The mean ${\bf R^2}$ score across all folds was 0.436, indicating that the model generalized well and was not overfitting.

8.2 Residual Analysis

Residual analysis confirmed that the model did not exhibit significant bias. However, the errors were more pronounced at the extremes of the quality scale.

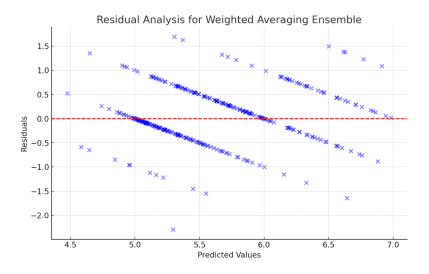


Figure 3: Residual Analysis for Ensemble Model

9 Challenges and Solutions

Several challenges emerged during the project:

• **Preprocessing Issues**: We initially applied scaling before capping outliers, which led to distorted results. Reordering these steps resolved the issue.

- Model Optimization Errors: When uploading the optimized Random Forest and XGBoost models, version mismatches caused errors. Aligning my local environment with ChatGPT's environment allowed the optimized models to be loaded successfully.
- **Prediction Bias**: The ensemble model's tendency to overestimate low-quality wines and underestimate high-quality wines

10 Deployment

The Deployment phase of the CRISP-DM methodology is where the final model is integrated into a real-world environment, allowing end-users to utilize it for predictions. In this project, the deployment of the trained and calibrated weighted ensemble model is achieved through a RESTful API, which can be accessed via HTTP requests to predict wine quality based on chemical properties.

10.1 Model Deployment Strategy

The goal of the deployment phase is to make the predictive model accessible and scalable for real-time use. The following steps outline how the deployment was implemented:

- Model Saving: The best-trained models (Random Forest, XGBoost, and Isotonic Calibration) were saved as serialized files using the joblib library. These models are then loaded within a web service for use during inference.
- API Setup: A web-based REST API was created using the Flask framework. This API allows users to send requests with chemical properties of wine, and it responds with the predicted wine quality.
- Weighted Averaging Ensemble: The deployment script incorporates the final ensemble model, where predictions from both Random Forest and XGBoost are weighted based on their individual performance. The weighted prediction is then calibrated using the Isotonic Regression model to ensure more accurate predictions for extreme wine quality values.
- Real-Time Prediction: The API accepts POST requests containing the wine's chemical features and returns the predicted wine quality score, providing real-time predictions to end-users.

10.2 Example Code for Deployment

Below is an outline of the Flask API code used to deploy the model:

```
from flask import Flask, request, jsonify
import joblib
import numpy as np
app = Flask(__name__)
# Load the pre-trained models
rf_model = joblib.load('best_random_forest.pkl')
xgb_model = joblib.load('best_xgboost.pkl')
calibration_model = joblib.load('calibration_isotonic.pkl')
# Define the feature list
FEATURES = ['alcohol', 'volatile_acidity', 'sulphates', 'total_sulfur_dioxide', 'chlorides']
# Define the ensemble weights
rf_weight = 0.49
xgb\_weight = 0.51
@app.route('/predict', methods=['POST'])
def predict():
    data = request.json
    features = [data[feature] for feature in FEATURES]
    features_array = np.array(features).reshape(1, -1)
    # Predict using both models
    rf_pred = rf_model.predict(features_array)
    xgb_pred = xgb_model.predict(features_array)
    # Weighted average ensemble prediction
    ensemble_pred = (rf_weight * rf_pred) + (xgb_weight * xgb_pred)
   # Calibrate the prediction using isotonic regression
    calibrated_pred = calibration_model.predict(ensemble_pred)
    # Return the prediction as JSON
    response = {'predicted_quality': float(calibrated_pred[0])}
    return jsonify(response)
if __name__ == '__main__':
    app.run(debug=True)
```

10.3 API Workflow

The API accepts a JSON payload containing the chemical features of a new wine sample and returns the predicted quality. The prediction workflow is as follows:

- 1. The user sends a POST request with the chemical properties of wine, including features like alcohol, volatile acidity, sulphates, total sulfur dioxide, and chlorides.
- 2. The API retrieves the saved Random Forest and XGBoost models, computes their predictions, and combines them using a weighted average.
- 3. The ensemble prediction is calibrated using the Isotonic Regression model to account for over- or underestimation in extreme quality predictions.
- 4. The final calibrated prediction is returned to the user as a JSON response.

10.4 Deployment Options

The Flask API can be deployed using several scalable cloud options, such as:

- AWS EC2 or Lambda: The Flask API can be containerized using Docker and deployed on an AWS EC2 instance or Lambda function for serverless deployment.
- **Heroku:** Another deployment option is Heroku, where the Flask app can be easily hosted as a web service.
- **Docker:** The entire deployment pipeline can be Dockerized, ensuring consistency across different environments and simplifying the deployment process.

10.5 Future Deployment Considerations

For future iterations, the deployment could incorporate additional scalability features, such as:

- Integrating a message queue system like RabbitMQ or Kafka to handle large-scale prediction requests.
- Implementing a load balancer to manage API traffic efficiently.
- Monitoring and logging system performance to ensure smooth operation and to track prediction trends over time.

This deployment provides a user-friendly and scalable solution for predicting wine quality, making the model accessible for real-time use in production environments.

11 Conclusion

By following the CRISP-DM process, we successfully built a predictive model for wine quality. The final **weighted averaging ensemble**, calibrated using

Isotonic Regression, achieved an $\mathbf{R^2}$ score of 0.524. Despite initial challenges in preprocessing and model optimization, the methodology and solutions proposed helped create a robust model suitable for real-world applications in wine production.