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Practical Machine Learning Homework 5: Pipeline Optimization

(I) Introduction

The core problem addressed in this report involves optimizing the complete machine learning (ML) pipeline rather than individual components, for the 'winequality-red' dataset. Our focus is on the systematic tweaking of hyperparameters across various stages of preprocessing and model selection, aiming not just for performance but for a deep understanding of the influence and interaction of each component within the pipeline.

(II) Dataset description

The 'winequality-red' dataset comprises data from 1,599 different red wines, characterized by 11 physicochemical features: fixed acidity, volatile acidity, citric acid, residual sugar, chlorides, free sulfur dioxide, total sulfur dioxide, density, pH, sulphates, and alcohol. The dataset targets the wine quality on a scale of 0 to 10. Notably, there are no missing values in the dataset, which simplifies preprocessing steps regarding data completeness.

(III) Experimental setup

\square Programming environment and libraries:

The analysis was conducted using Python, leveraging libraries such as Pandas for data manipulation, Scikit-learn for building ML models and preprocessing, and skopt for advanced hyperparameter optimization.

\square Data preparation and splitting

Data was randomly divided into training (80%) and testing (20%) subsets, using a 'random_state' of 42. Numerical features were subjected to optional logarithmic transformation and standard scaling (to be decided by pipeline optimizer). These steps were encapsulated within a 'ColumnTransformer', ensuring that different transformations apply to appropriate feature types. The ML pipeline optimization task tunes the preprocessing options (e.g., activation of transformations).

\square ML pipeline configuration

The pipeline for the ML integrates:

- Preprocessing: As described above, with components conditionally applied based on optimization results.
- **Feature selection**: **SelectKBest** selects the top 10 features.
- **Classifier**: A placeholder filled by one of the classifiers (Random Forest, Gradient Boosting, KNN) depending on the experiment phase.

\square Hyperparameter optimization strategy

The optimization of hyperparameters was conducted using "Random Search" and "Bayesian Optimization" techniques. Both methods were performed with the following parameters:

- Iterations: 30 random combinations per classifier
- Cross-validation (CV): 5-fold, to evaluate generalizability

Random state: 42, to ensure consistent random sampling

These methods explore a broad parameter space efficiently and investigate multiple configurations to identify the most effective ones. The hyperparameters of the classifiers, the range of these parameters, and the optimized values from both optimization approaches are listed in Tables 1-3.

Table 1. Hyperparameter ranges and optimized values for the Random Forest classifier

Classifier	Parameter	Range/options	Best value (Random Search)	Best value (Bayesian Search)
Random Forest	n_estimators	100 to 1000	838	1000
	max_depth	10 to 100	44	100
	min_samples_leaf	1 to 20	1	1
	log_transform_active	True, False	True	True
	scaler_active	True, False	False	False
	onehot_active	True, False	True	True

Table 2. Hyperparameter ranges and optimized values for the Gradient Boosting classifier

Classifier	Parameter	Range/Options	Best value (Random Search)	Best value (Bayesian Search)
Gradient Boosting	n_estimators	50 to 300	64	203
	learning_rate	0.01 to 0.2	0.1185	0.0228
	max_depth	1 to 10	9	7
	min_samples_leaf	1 to 10	6	1
	log_transform_active	True, False	True	False
	scaler_active	True, False	True	True
	onehot_active	True, False	True	False

Table 3. Hyperparameter ranges and optimized values for the KNN classifier

Classifier	Parameter	Range/Options	Best value (Random Search)	Best value (Bayesian Search)
	n_neighbors	1 to 30	15	21
	weights	Uniform, Distance	Distance	Distance
KNN	algorithm	Auto, Ball Tree, KD Tree, Brute	Ball Tree	Brute
	log_transform_active	True, False	False	True
	scaler_active	True, False	True	True
	onehot_active	True, False	True	True

(IV)Results and observations

The default configurations and the post-optimization results using both Random and Bayesian search methods for each classifier are summarized in Table 4. The illustration of the accuracy values for the test data are also presented in Figure 1.

The results demonstrate that the implementation of pipeline optimization and hyperparameter tuning has led to significant enhancements in model accuracies across all classifiers. Notably, there has been a noteworthy increase in both training and test data accuracies, ranging from 3% to 10% depending on the classifier and optimization technique employed.

For the Random Forest and Gradient Boosting models, the test accuracies were increased by approximately 2-3%. For these two classifiers, Random Search optimization demonstrated a superior performance over Bayesian optimization, which might be attributed to the fact that Random Search explores the hyperparameter space more extensively and is better suited for algorithms with complex and high-dimensional parameter spaces, such as Random Forest and Gradient Boosting. In contrast, Bayesian optimization tends to perform better with algorithms that have smoother, more continuous parameter spaces.

For the KNN classifier, the model initially exhibited poor performance but experienced a remarkable improvement in accuracy (up to 10% on the test data) following the application of Bayesian optimization. This enhancement highlighted the effectiveness of optimization in fine-tuning hyperparameters for ML algorithms.

Table 4. The training and test accuracies before and after optimization of the Random Forest, Gradient Boosting, and KNN classifiers.

Classifier	Default		Random Search		Bayesian Search		
	Test	Training	Test	Training	Test	Training	
Random Forest	63.75%	68.73%	65.94%	70.12%	65.62%	70.48%	
Gradient Boosting	61.56%	65.13%	64.38%	66.89%	62.81%	67.25%	
KNN	56.87%	57.61%	65.31%	66.47%	66.87%	67.82%	

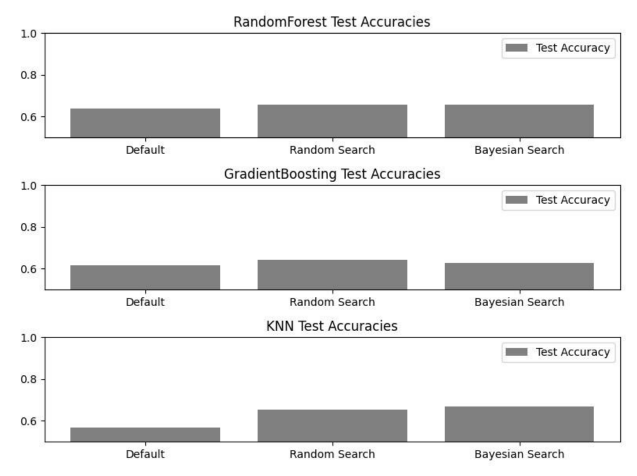


Figure 1: Test accuracies of Random Forest, Gradient Boosting, and KNN classifiers before and after pipeline optimization

(V) Conclusion

This comprehensive experiment demonstrates the significant impact of systematic optimization across the entire ML pipeline. The results also showed that the appropriate hyperparameter optimization technique may vary depending on the nature of the ML model (e.g., Random Search for Random Forest model and Bayesian optimization for the KNN model). By tuning both preprocessing and model parameters together, rather than in isolation, the study provides deeper insights into the interactive effects between the elements of ML pipeline and enhances our understanding of the performance of the predictive models.

(VI) References

- 1) Konstantinos Filippou. (January 2024). Structure Learning and Hyperparameter Optimization Using an Automated Machine Learning (AutoML) Pipeline.
- 2) A Comprehensive Guide on Hyperparameter Tuning and its Techniques (https: analyticsvidhya.com)
- 3) ML Pipeline Architecture Design Patterns(https://neptune.ai/blog/ml-pipeline-architecture-design-patterns)