Pipeline Optimization Report

Soudabeh Bolouri

Introduction:

This exercise involves a comprehensive approach to predicting wine quality utilizing multiple ML algorithms including SVM, Random Forest, and KNN. This experiment optimizes the entire ML approach, including any preprocessing, rather than optimizing just one component. Using nested cross-validation, we were able to determine which model generalized the data the best. The goal was to assess the impact of hyperparameter tuning on the performance of three different ML classifiers. The experiment involved comparing the default accuracy of these models with their post-tuning accuracy.

Dataset Description:

The dataset that we utilized in this exercise is the "Wine Quality" dataset, precisely the "white" wine version. It contains data on different features of white wines, including fixed acidity, volatile acidity, citric acid, residual sugar, chlorides, free sulfur dioxide, total sulfur dioxide, density, PH, sulphates, and alcohol. The dataset includes a totality of 4,898 rows and 12 features. The "quality" of the wine is the target variable, which we desire to predict. Also, we did not find any missing values in the dataset.

Experimental Setup:

We used the Python programming language for this exercise. First, we need to import our dataset "winequality-white.csv" using Panda. In the next step, the dataset is preprocessed to ensure its suitability for model training. Missing values are replaced with the mean of the column via an imputer (this is a general case for implementing pipelines even though there are no missing values for this specific dataset). As a way of standardizing features, it uses the Standard Scaler to remove the mean and scale to unit variance. Then feature scaling and one-hot encoding are performed using **StandardScaler** and **OneHotEncoder** respectively.

Four classifiers are assessed in this analysis:

- Support Vector Machine (SVM)
- Random Forest
- K-Nearest Neighbors (KNN)

The hyperparameters and their values to be searched for each model are as follows:

SVM:

Hyperparameter	Values	Description
classifierC	[1, 10, 100]	This parameter controls the trade-off
		between maximizing the margin and
		minimizing classification error.
classifier_gamma	[0.01, 0.1, 'scale']	This parameter defines the influence of a
		single training example's distance,
		affecting the reach of the kernel
classifier_kernel	['rbf']	Kernel specifies the type of kernel used in
		the SVM algorithm, determining the
		decision boundary shape in the feature
		space.

Random Forest:

Hyperparameter	Values	Description
classifier_n_estimators	[5, 20, 50, 100]	number of trees in the rand
		om forest
classifier_max_features	['auto', 'sqrt']	number of features in
		consideration at every split
classifier_max_depth	[int(x) for x in np.linspace(10,	maximum number of levels
	[120, num = 12)]	allowed in each decision tre
		e
classifier_min_samples_split	[2, 6, 10]	minimum sample number t
		o split a node
classifier_min_samples_leaf	[1, 3, 4]	minimum sample number
		that can be stored in a leaf
		node
classifier_bootstrap	[True, False]	method used to sample data
		points

KNN:

Hyperparameter	Values	Description
classifier_n_neighbors	np.arange(2, 30, 1)	Number of neighbors to consider. It de
		fines the value of 'k' in the KNN algori
		thm.
classifier_weights	['uniform', 'distance']	The weight function used in prediction

The hyperparameters for preprocessing steps (**imputer** strategies for both numerical and categorical features) were also included in the tuning process.

Nested cross-validation was implemented with an outer 5-fold Stratified K-Fold and an inner 3-fold Stratified K-Fold. The inner loop involved hyperparameter tuning using **RandomizedSearchCV** with 10 iterations, optimizing the entire pipeline, including preprocessing and the classifier. The outer loop assessed the optimized model's performance.

Results:

The following accuracies were observed before and after tuning:

Random Forest:

Default Accuracy: 0.6809Tuned Accuracy: 0.6801

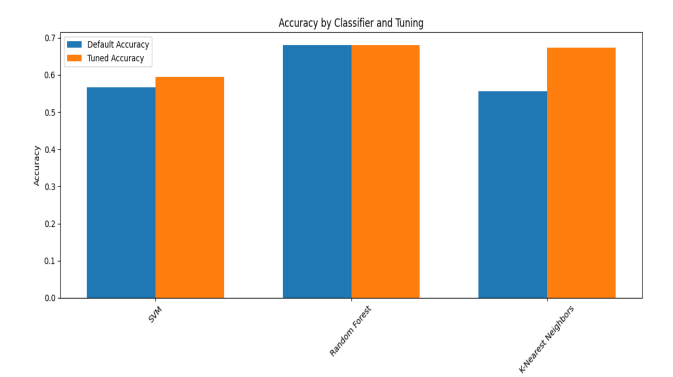
K-Nearest Neighbors (KNN):

Default Accuracy: 0.5570Tuned Accuracy: 0.6744

SVM:

Default Accuracy: 0.5666Tuned Accuracy: 0.5945

Based on the bellow bar chart showcasing the Comparison of Model Performance Before and After Hyperparameter Tuning, we can interpret the results as follows:



Tuning does not significantly improve the performance of the Random Forest model. With its default hyperparameters, the Random Forest model already achieves a high degree of accuracy (0.6809), which indicates it is quite robust from the beginning. After tuning, the accuracy slightly decreases to 0.6801.

Models

By tuning hyperparameters, KNN shows an impressive improvement; the default accuracy of 0.5570 is significantly increased to 0.6744 after tuning. A marked improvement in model performance may have been attributed to the optimization of parameters like neighbor number and weighting method.

For the SVM model, the tuning process results in a moderate increase in accuracy, from 0.5666 to 0.5945.

References:

- [1] https://towardsdatascience.com/step-by-step-tutorial-of-sci-kit-learn-pipeline-62402d5629b6
- [2] https://medium.com/mlearning-ai/how-to-use-sklearns-pipelines-to-optimize-your-analysis-b6cd91999be
- [3] https://scikit-learn.org/stable/modules/grid_search.html
- [4] https://towardsdatascience.com/hyper-parameter-tuning-and-model-selection-like-a-movie-star-a884b8ee8d68
- [5] https://towardsdatascience.com/nested-cross-validation-hyperparameter-optimization-and-model-selection-5885d84acda
- [6] https://machinelearningmastery.com/nested-cross-validation-for-machine-learning-with-python/
- [7] https://www.kaggle.com/code/arjunprasadsarkhel/simple-random-forest-with-hyperparameter-tuning
- [8] https://www.kaggle.com/code/funxexcel/p2-logistic-regression-hyperparameter-tuning
- [9] https://medium.com/@agrawalsam1997/hyperparameter-tuning-of-knn-classifier-a32f31af25c7