Binary Classification with Decision Trees: Identifying Poisonous Mushrooms

Machine Learning and Statistical Learning (Machine Learning module)

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1 Introduction

The aim of this project is to build a binary decision tree from scratch and then test it on a mushroom dataset. The goal is to predict whether a mushroom is poisonous or edible based on numerous attributes.

The first step was to analysed the data, divide it into training and test sets, before proceeding with a data cleaning phase.

The central point of the project was the construction of the decision tree for binary classification. Starting by creating a class for the nodes and then by developing the tree from the fit and prediction functions.

Three methods for the splitting criteria were used: Gini impurity, scaled entropy and squared error. As for the stopping criteria, three were implemented: maximum depth, minimum sample splitting and minimum impurity decrease. These criteria are essential to avoid underfitting or overfitting of the model.

To find the best criteria, the hyperparameters were adjusted, with and without cross-validation, using 0-1 loss as the evaluation metric. This step was essential to optimise the performance of the decision tree.

Finally, a random forest was constructed using the structures previously presented, in order to further study the predictive capabilities of the model and the importance of the features.

2 Data Analysis and Preprocessing

The mushroom dataset includes 61069 hypothetical mushrooms with caps based on 173 species (353 mushrooms per species).[1] Each mushroom is identified as edible or poisonous, and each one is described by the following attributes:

- Classification: Edible or Poisonous
- Cap Diameter (Numerical): Float number in cm
- Cap Shape (Categorical): Bell, Conical, Convex, Flat, Sunken, Spherical, Others
- Cap Surface (Categorical): Fibrous, Grooves, Scaly, Smooth, Shiny, Leathery, Silky, Sticky, Wrinkled, Fleshy, d
- Cap Color (Categorical): Brown, Buff, Gray, Green, Pink, Purple, Red, White, Yellow, Blue, Orange, Black
- Does Bruise or Bleed (Categorical): True, False
- Gill Attachment (Categorical): Adnate, Adnexed, Decurrent, Free, Sinuate, Pores, None

- Gill Spacing (Categorical): Close, Distant, None
- Gill Color (Categorical): Brown, Buff, Gray, Green, Pink, Purple, Red, White, Yellow, Blue, Orange, Black, None
- Stem Height (Numerical): Float number in cm
- Stem Width (Numerical): Float number in mm
- Stem Root (Categorical): Bulbous, Swollen, Club, Cup, Equal, Rhizomorphs, Rooted
- Stem Surface (Categorical): Fibrous, Grooves, Scaly, Smooth, Shiny, Leathery, Silky, Sticky, Wrinkled, Fleshy, None
- Stem Color (Categorical): Brown, Buff, Gray, Green, Pink, Purple, Red, White, Yellow, Blue, Orange, Black, None
- Veil Type (Categorical): Partial, Universal
- Veil Color (Categorical): Brown, Buff, Gray, Green, Pink, Purple, Red, White, Yellow, Blue, Orange, Black, None
- Has Ring (Categorical): True, False
- Ring Type (Categorical): Cobwebby, Evanescent, Flaring, Grooved, Large, Pendant, Sheathing, Zone, Scaly, Movable, None
- Spore Print Color (Categorical): Brown, Buff, Gray, Green, Pink, Purple, Red, White, Yellow, Blue, Orange, Black
- Habitat (Categorical): Grasses, Leaves, Meadows, Paths, Heaths, Urban, Waste, Woods
- Season (Categorical): Spring, Summer, Autumn, Winter

The data was split into training and test sets (80% training, 20% test). To prevent data leakage, only the training set was utilized to clean the dataset.

- Columns with more than 80% missing values were removed:
 - stem-root
 - veil-type
 - veil-color
 - spore-print-color
- Missing values, which were only present for categorical attributes, were replaced using the training set mode. No null values were found for the numerical attributes. However, for stem height and stem width several mushrooms had a value of "0.0". As it was not clear whether these were errors, they were retained. If null values had been present, the median would have been used as a replacement to avoid the influence of outliers.
- Duplicate rows, although few, were removed to reduce the risk of overfitting.
- The classes in the dataset were balanced, so there is no need to apply oversampling or undersampling technique.

No scaling or outlier removal was performed, as decision trees do not require these preprocessing steps.

3 Decision Tree

A decision tree is a machine learning model used for classification and regression tasks. It operates by recursively partitioning the input space into increasingly homogeneous regions, ultimately assigning a predicted class or value to each region. The tree structure consists of interconnected elements called nodes, each representing a decision point or a prediction outcome. [2]

3.1 Structure of a decision tree

At the top of the tree is the root node, which represents the entire data set and is the starting point of the process. The tree expands downwards through a hierarchy of nodes, with parent node branching into child nodes based on specific splitting criteria. This hierarchical structure facilitates a step-by-step division of the data, guided by thresholds or tests at each node, the splitting continues until the final decision or stopping criteria is reached. As this is a binary classification, at each split the decision tree divides the data into two subsets based on the predictor variables. The leaf nodes at the end of each branch represent the final result of the tree.

3.2 How a decision tree works

Decision trees work by recursively partitioning the input space based on selected features, aiming to create subsets of data that are increasingly homogeneous with respect to the target variable. This process is guided by specific splitting criteria, for binary classification tasks the most common are:

• Gini impurity:

$$\psi(p) = 2p(1-p)$$

• Scaled entropy:

$$\psi(p) = -\frac{p}{2}\log_2(p) - \frac{1-p}{2}\log_2(1-p)$$

• Squared impurity:

$$\psi(p) = \sqrt{p(1-p)}$$

At each step, the goal is to reduce the heterogeneity of the groups, meaning that each split should create more homogeneous subgroups compared to the previous level. At the end, the tree's leaf nodes represent highly homogeneous groups that can be assigned to a particular class or output value.

To build the tree, a decision tree algorithm uses recursion. This process begins at the root node, which represents the entire feature space, and recursively divides the data into two child nodes at each step based on the value of a chosen predictor variable. The algorithm evaluates all possible splits at each node and selects the one that maximizes homogeneity or minimizes impurity.

The splitting continues until a stopping criterion is reached, such as:

- Maximum tree depth,
- Minimum number of samples in a node,
- Minimum impurity decrease

3.3 Model Evaluation

The evaluation of decision trees for binary classification involves various performance metrics, which measure the effectiveness of the model in distinguishing between the two classes. These include 0-1 loss, accuracy, precision, recall, F1 score, and the confusion matrix.

Before presenting the formulas for these metrics, some essential terms used in binary classification are:

- True Positives (TP): The number of times the model correctly predicted the positive class.
- False Positives (FP): The number of times the model incorrectly predicted the positive class.
- True Negatives (TN): The number of times the model correctly predicted the negative class.
- False Negatives (FN): The number of times the model incorrectly predicted the negative class.

Now, using these definitions, the most commonly used evaluation metrics can be defined:

• 0-1 Loss is a simple metric that calculates the proportion of incorrect predictions out of the total number of predictions. It is defined as:

$$L_{0-1} = \frac{1}{n} \sum_{i=1}^{n} \ell_i$$

where:

- -n is the total number of predictions,
- $-\ell_i = \mathbb{I}(y_i \neq \hat{y}_i)$ is the loss for the *i*-th sample,
- $-y_i$ is the true label for the *i*-th sample,
- $-\hat{y}_i$ is the predicted label for the *i*-th sample,
- $-\mathbb{I}(\cdot)$ is the indicator function, which returns 1 if $y_i \neq \hat{y}_i$, and 0 otherwise.
- Accuracy measures the proportion of correct predictions out of the total number of predictions. It can be written as:

$$Accuracy = \frac{TP + TN}{TP + TN + FP + FN}$$

Alternatively, accuracy can also be written as:

Accuracy =
$$\frac{1}{n} \sum_{i=1}^{n} \mathbb{I}(y_i = \hat{y}_i)$$

where:

- -n is the total number of predictions,
- $-y_i$ is the true label for the *i*-th sample,
- $-\hat{y}_i$ is the predicted label for the *i*-th sample,
- $-\mathbb{I}(\cdot)$ is the indicator function, which returns 1 if $y_i = \hat{y}_i$, and 0 otherwise.
- Precision is the ratio of correctly predicted positive instances to the total predicted positives. It is defined as:

$$Precision = \frac{TP}{TP + FP}$$

Recall is the ratio of correctly predicted positive instances to the total actual positives. It is defined
as:

$$Recall = \frac{TP}{TP + FN}$$

• F1 Score is the harmonic mean of precision and recall, providing a balance between the two. It is defined as:

$$F1 = 2 \times \frac{\text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}}$$

• Confusion Matrix is a 2x2 table that summarizes the performance of a classification model by showing the counts of TP, FP, TN, and FN:

/	Predicted Positive	Predicted Negative
Actual Positive	TP	FN
Actual Negative	FP	TN

The confusion matrix allows for a detailed analysis of the classification results, providing more insights into model performance beyond accuracy alone.

4 Implementation

This chapter illustrates how the decision tree has been implemented.

4.1 Node Class

The Node class is a central component of the decision tree structure.

4.1.1 Attributes

Each node can have the following attributes:

- feature: The index of the feature used to split the data.
- thr: The threshold value used for the split.
- left: The left child node.
- right: The right child node.
- value: The value assigned to the leaf node.
- categorical: A boolean indicating whether the feature is categorical or numerical.

4.1.2 Methods

The Node class includes the method is_leaf(), which returns True if the node is a leaf node and False otherwise.

4.2 DecisionTree Class

The DecisionTree class is responsible for building and managing the decision tree.

4.2.1 Attributes

The main attributes are:

- max_depth: The maximum depth the tree can reach.
- min_samples_split: The minimum number of samples required to split a node.
- min_impurity_decrease: The minimum decrease in impurity required for a split to occur.
- criterion: The criterion used to evaluate splits (e.g., 'gini', 'scaled_entropy', or 'squared').
- max_feature: The number of features to consider when looking for the best split. (e.g., 'None', 'sqrt', 'log2').

4.2.2 Methods

The main methods are:

• Fit

The decision tree is trained using the fit method, which recursively splits the data. The algorithm begins at the root and continues down the tree by recursively finding the best split at each node. Pseudocode for the fitting method:

Algorithm 1 Recursive Fit

```
    Input: Samples X, target values y, depth depth
    num_sample ← dimension of X
    if depth > max_depth or num_samples < min_samples_split or all samples have the same class then</li>
    return leaf node with the most common class
    end if
    split, gain ← Find Best Split(X, y)
    if gain < min_impurity_decrease then</li>
    return leaf node with the most common class
    end if
    X<sub>left</sub>, y<sub>left</sub>, X<sub>right</sub>, y<sub>right</sub> ← Split Samples(X, y, split)
    left_child ← RecursiveFit(X<sub>left</sub>, y<sub>left</sub>, depth + 1)
    right_child ← RecursiveFit(X<sub>right</sub>, y<sub>right</sub>, depth + 1)
    return node with the split information, left_child, and right_child
```

• Best Split

The best split is determined by evaluating the information gain for each feature and threshold combination. In the implementation of the decision tree for the project, the algorithm first checks whether the feature is categorical or numerical. For numerical features, thresholds on a single feature are set from the 10th to the 90th percentiles for computational efficiency. For categorical features, membership tests using each unique value are applied as a possible split point.

Algorithm 2 Best Split

```
    Input: Samples X, target values y
    for each feature f in X do
    if feature f is numerical then
    split ← percentiles from 10 to 90
    else if feature f is categorical then
    split ← all unique values of f
    end if
    Calculate information gain for each possible split
    end for
    return the split and gain with the highest information gain
```

• Information Gain Calculation

Information gain measures the reduction in impurity achieved by splitting a node. The impurity is calculated based on the criterion chosen during the initialization of the tree (gini, scaled_entropy, or squared).

The general formula for information gain is [2]:

$$IG = \psi(p) - \left(\frac{N_l}{N} \cdot \psi(l) + \frac{N_r}{N} \cdot \psi(r)\right)$$

where:

- -IG is the Information Gain,
- $-\psi(p)$ is the impurity of the parent node,
- $-\psi(l)$ is the impurity of the left child node,
- $-\psi(r)$ is the impurity of the right child node,
- $-N_l$ is the number of samples in the left child node,
- $-N_r$ is the number of samples in the right child node,
- -N is the total number of samples in the parent node.

So, Information Gain is the reduction in impurity achieved by splitting the parent node into child nodes.

• Feature Importance

The importance of a feature is measured by the total information gain it contributes across all nodes where it is used as a splitting criterion. The feature importance is calculated by summing the information gain for each feature and normalizing the values so that the total importance sums to 1.

The feature importance is given by:

$$FI_j = \frac{\sum_{n \in N_j} IG_n}{\sum_{m \in N} IG_m}$$

where:

- FI_i is the importance of feature j,
- $-\sum_{n\in N_j}IG_n$ is the sum of the Information Gain IG for all nodes n where feature j was used for splitting,
- $-\sum_{m\in\mathbb{N}}IG_m$ is the sum of the Information Gain across all nodes m in the entire tree.

So, the feature importance is computed as the ratio of the total Information Gain for feature j to the total Information Gain for all features in the tree.

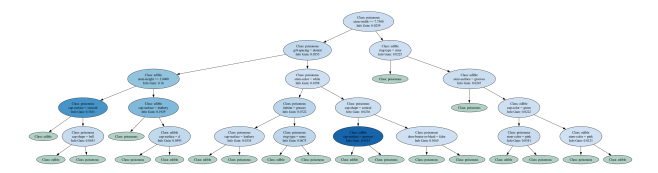


Figure 1: Decision Tree on Mushroom dataset with maximum depth of 5, minimum samples spit of 100 and Gini as criterion. The nodes are colored based on their information gain, with darker shades representing higher information gain. Leaf nodes are shaded differently to distinguish them from decision nodes.

5 Hyperparameter Tuning and Cross-Validation

5.1 Hyperparameter Tuning

The fine-tuning of hyperparameters refers to the process of optimising the hyperparameters of a machine learning model. Hyperparameters are settings that must be specified before training the model and can significantly influence the final performance of the model.

In our decision trees, the hyperparameters are the stopping and splitting criteria. The correct adjustment of these hyperparameters is essential because if the tree grows too large, it can capture noise in the data, resulting in poor generalisation (overfitting). On the other hand, if the tree is too small, it may fail to capture important patterns in the data (underfitting).

Hyperparameters should not be confused with model parameters. Hyperparameters are defined by the user before training, while parameters are values that the model learns automatically during training. An example of a parameter in the Decision Tree is the splitting value associated with each node.

The goal of hyperparameter tuning is to find the optimal balance that maximises the performance of the model on the unseen data.

5.2 Cross-Validation

Cross-validation is a technique used to more reliably assess the performance of a model, particularly when adjusting hyperparameters. It works by splitting the data into several subsets or folds. In the project, k-fold cross-validation was implemented, the data is divided into k parts of equal size. The model is trained k times, each time using k-1 folds as the training set and the remaining folds as the test set. This process is repeated for each fold and the average loss of all iterations is taken as the final evaluation score. [2]

The pseudocodes of the two algorithms:

Algorithm 3 K-Fold

- 1: **Input:** Samples X, Number of fold n-splits
- 2: Inizialize empty list folds
- 3: Shuffle indices of samples
- 4: Determine the size of each fold
- 5: for each fold do
- 6: Assign train and test indices
- 7: Append (train, test) pair to folds
- 8: end for
- 9: return folds

Algorithm 4 Cross-Validation

- 1: **Input:** Samples X, target values y, Number of fold n_splits
- 2: $folds \leftarrow \text{K-Fold}(X, y, n_splits)$
- 3: Initialize empty list losses
- 4: **for** each (train_indices, test_indices) in folds **do**
- 5: Fit model on training set
- 6: $loss \leftarrow compute 0-1 loss on test set$
- 7: Append loss to losses
- 8: end for
- 9: **return** The mean of the *losses*

This technique provides a better estimate of model performance on unseen data. In addition, it allows for a more robust comparison of models or hyperparameter configurations, as it decreases the potential risk of overfitting or underfitting that could result from relying on a single train-test split.

6 Random Forest

Random Forest is an ensemble learning method that combines multiple decision trees to improve predictive performance and reduce overfitting. It consists of building multiple decision trees during training and returning the value that has been predicted multiple times. [2]

6.1 Key Points of Random Forest

- **Bootstrap Sampling**: Each tree is trained on a different bootstrap sample, a random sample with replacement, of the training data, leading to diversity among the trees. [3]
- Feature Randomness: For each split in every tree, a random subset of features is considered for splitting, typically $\sqrt{features}$ or $\log_2(features)$. This process occurs at each node, reducing correlation among trees and improving model robustness.
- Feature Importance: Random Forest provides a reliable measure of feature importance.

6.2 Pseudocode for Random Forest

Algorithm 5 Random Forest

- 1: **Input:** Training data (X, y), number of trees n_trees , max features $max_features$
- 2: Initialize empty list of trees trees
- 3: for i = 1 to n_trees do
- 4: $X_bootstrap, y_bootstrap \leftarrow BootstrapSample(X, y)$
- 5: $tree \leftarrow DecisionTree(X_subset, y_bootstrap, max_features)$
- 6: Append ($tree, X_subset.columns$) to trees
- 7: end for
- 8: return trees

Algorithm 6 Predict Function for Random Forest

- 1: **Input:** Test data X
- 2: Initialize empty array tree_predictions
- 3: for each tree, features in trees do
- 4: $tree_predictions[i] \leftarrow tree.predict(X[features])$
- 5: end for
- 6: Initialize empty array majority_votes of size n_samples
- 7: **for** each sample j in range(n-samples) **do**
- 8: $majority_votes[j] \leftarrow most predicted value for sample j$
- 9: end for
- 10: **return** majority_votes

7 Results

After hyperparameter tuning, both with and without cross-validation, the following parameters were found as the best ones for the decision tree, using 0-1 loss:

Best hyperparameters: {'max_depth': 50, 'min_samples_split': 2, 'min_impurity_decrease': 0.0, 'criterion': 'squared'}, loss: 0.0013334700994973843
Best hyperparameters CV: {'max_depth': None, 'min_samples_split': 10, 'min_impurity_decrease': 0.0, 'criterion': 'gini'}, loss: 0.001148734781476601

Figure 2: Best Hyperparameters found with and without cross-validation and the 0-1 loss evaluated on the validation set

The loss concerns the validation set. From these results, we can observe that the dataset does not suffer of overfitting, as indicated by the similar performance across different validation strategies. The similarity in loss between cross-validation and the initial hyperparameter tuning suggests that the model generalizes well across unseen data, with no significant performance degradation across the different validation folds.

After re-fitting the model with the best hyperparameters found through tuning on the entire training set, the performance of the three splitting criteria was compared on the test set. The results obtained are as follows on the test set:

```
Criterion: gini
0-1 Loss: 0.0015557193154835012
Accuracy: 0.9984442806845165
Precision: 0.9983698873740368
Recall: 0.9988139362490734
F1 Score: 0.9985918624471948
Confusion Matriy_pred:
[[6737 11]
[ 8 5457]]
Criterion: scaled_entropy
0-1 Loss: 0.0018013592074019487
Accuracy: 0.998198640792598
Precision: 0.9974840905727393
Recall: 0.9992587101556709
F1 Score: 0.9983706117612207
Confusion Matriy_pred:
[[6740 17]
    5 5451]]
Criterion: squared
0-1 Loss: 0.0018832391713747645
Accuracy: 0.9981167608286252
Precision: 0.9980735032602253
Recall: 0.9985174203113417
F1 Score: 0.9982954124360779
Confusion Matriy_pred:
[[6735 13]
   10 5455]]
```

Figure 3: Comparison within the three splitting criteria with the best hyperparameters

This comparison further highlights the model's robustness, demonstrating that the choice of splitting criterion has a relatively minor impact on overall performance when using the optimized hyperparameters.

In addition, hyperparameter tuning without Cross-Validation for the random forest has been performed over a small grid.

The feature importances shown in the figure represent those from a Random Forest that achieved perfect results on the test set:

```
0-1 Loss: 0.0
Accuracy: 1.0
Precision: 1.0
Recall: 1.0
F1 Score: 1.0
Confusion Matriy_pred:
[[6745 0]
[ 0 5468]]
```

Figure 4: Results of a Random forest with 30 trees, max_depth of None, min_sample_split of 10 using the Gini criterion and 'sqrt' for max_features.

Regarding the importance of the features the random forest identified the following in descending order:

Feature importance: stem-width: 0.11290158997960267 cap-surface: 0.1122616855002578 gill-color: 0.10350250494969566 gill-attachment: 0.10263590082520457 cap-diameter: 0.07440204725571684 stem-height: 0.07420130606688989 cap-color: 0.07158740937470241 stem-color: 0.06728496418577742 cap-shape: 0.06427840493071298 stem-surface: 0.04478348081206898 gill-spacing: 0.03527509635748149 does-bruise-or-bleed: 0.0319130005021012 ring-type: 0.02977409803213836 habitat: 0.029253904926120792 has-ring: 0.028318124733740478 season: 0.017626481567788378

Figure 5: Feature importance in descending order from a Random Forest model with 100 trees, max_depth of 30, min_sample_split of 10 using the Gini criterion and 'sqrt' for max_features.

This feature importance values provide the factors that have the most significant influence on model predictions, which can be useful for interpretability and further refinement of the model.

References

- [1] Dennis Wagner, D. Heider, and Georges Hattab. Secondary Mushroom. UCI Machine Learning Repository. 2021.
- [2] Shai Shalev-Shwartz and Shai Ben-David. *Understanding Machine Learning: From Theory to Algorithms*. Cambridge, UK: Cambridge University Press, 2014. ISBN: 9781107057135.
- [3] Gareth James et al. An Introduction to Statistical Learning: with Applications in R. New York, NY: Springer, 2013. ISBN: 9781461471387.