Classification with hyperparameter tuning

Aim: Show classification with different strategies for the tuning and evaluation of the classifier

- 1. simple holdout
- 2. cross validation on training set, then score on test set

NB: You should not interpret those experiments as a way to find the *best* evaluation method, but simply as examples of *how* to do the evaluation.

If you look at the final report, both methods are meant for increasing evaluation reliability, method 2 is the more reliable, but it requires several repetitions for cross validation, therefore, if the learning method is expensive, it requires long processing time. If, due to intrinsic variation caused by random sampling, it turns out that method 1 gives higher accuracy, this means simply that the forecast towards generalisation is less reliable.

Workflow

- download the data
- drop the useless data
- separe the predicting attributes X from the class attribute y
- split X and y into training and test
- part 1 single run with default parameters
 - initialise an estimator with the chosen model generator
 - fit the estimator with the training part of X
 - show the tree structure
 - part 1.1
 - o predict the y values with the fitted estimator and the train data
 - o compare the predicted values with the true ones and compute the accuracy on the training set
 - part 1.2

- o predict the y values with the fitted estimator and the test data
 - o compare the predicted values with the true ones and compute the accuracy on the test set
- part 2 compute accuracy with cross validation
 - prepare the structure to hold the accuracy data for the multiple runs
 - repeat for all the values of the parameter
 - o initialise an estimator with the current parameter value
 - o compute the accuracy with cross validation and store the value
 - find the parameter value for the top accuracy
 - fit the estimator with the entire X
 - show the resulting tree and classification report

The data are already in your folder, use the name winequality-red.csv

```
In [1]:
         import pandas as pd
         import numpy as np
         import matplotlib.pyplot as plt
         import seaborn as sns
         from sklearn import tree
         from sklearn.metrics import accuracy score, classification report, confusion matrix
         from sklearn.model selection import cross val score
         from sklearn.ensemble import BaggingClassifier
         %matplotlib inline
         plt.rcParams['figure.figsize'] = [20, 20]
         random state = 15
         np.random.seed(random state)
         # the random state is reset here in numpy, all the scikit-learn procedure use the numpy random state
         # obviously the experiment can be repeated exactly only with a complete run of the program
         data url = "winequality-red.csv"
         target name = 'quality'
```

Read the data into a dataframe and show the size

```
In [2]:
```

Shape of the input data (1599, 12)

Have a quick look to the data.

- use the .shape attribute to see the size
- use the .head() function to see column names and some data
- use the .hist() method for an histogram of the columns
- use the .unique method to see the class values

In [3]:

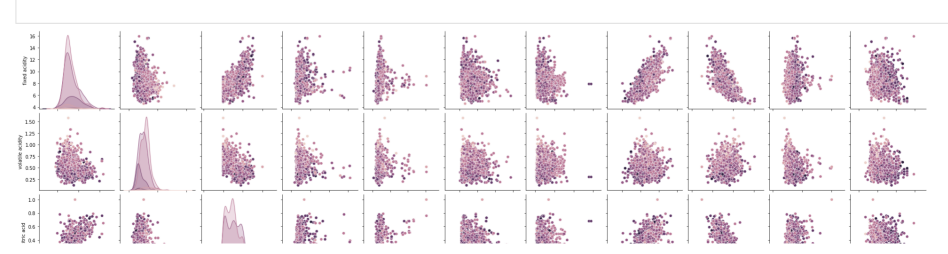
Out[3]:

:	fixed acidity	volatile acidity	citric acid	residual sugar	chlorides	free sulfur dioxide	total sulfur dioxide	density	рН	sulphates	alcohol	quality
0	7.4	0.70	0.00	1.9	0.076	11.0	34.0	0.9978	3.51	0.56	9.4	5
1	7.8	0.88	0.00	2.6	0.098	25.0	67.0	0.9968	3.20	0.68	9.8	5
2	7.8	0.76	0.04	2.3	0.092	15.0	54.0	0.9970	3.26	0.65	9.8	5
3	11.2	0.28	0.56	1.9	0.075	17.0	60.0	0.9980	3.16	0.58	9.8	6
4	7.4	0.70	0.00	1.9	0.076	11.0	34.0	0.9978	3.51	0.56	9.4	5

Use seaborn.pairplot to show the the pairplots of the attributes, using the target as hue

NB: a semicolon at the end of a statement suppresses the <code>Out[]</code>

In [4]:



Print the unique class labels (hint: use the unique method of pandas Series) In [5]: [3 4 5 6 7 8] Split the data into the predicting values X and the class y Drop also the columns which are not relevant for training a classifier, if any The method "drop" of dataframes allows to drop either rows or columns • the "axis" parameter chooses between dropping rows (axis=0) or columns (axis=1) In [6]: Another quick look to data In [7]: Out[7]: fixed acidity volatile acidity citric acid residual sugar chlorides free sulfur dioxide total sulfur dioxide density pH sulphates alcohol 0 7.4 0.70 0.00 1.9 0.076 11.0 34.0 0.9978 3.51 0.56 9.4 1 7.8 0.88 2.6 25.0 0.68 9.8 0.00 0.098 0.9968 3.20 2 7.8 0.76 2.3 0.9970 3.26 9.8 0.04 0.092 15.0 54.0 0.65 3 11.2 9.8 0.28 0.56 1.9 0.075 17.0 60.0 0.9980 3.16 0.58 7.4 0.70 0.00 1.9 0.076 11.0 34.0 0.9978 3.51 0.56 9.4 In [8]: Out[8]: 5

Name: quality, dtype: int64

2

3

5 6

Prepare a simple model selection: holdout method

- Split X and y in train and test
- Show the number of samples in train and test, show the number of features

In [9]:

There are 1055 samples in the training dataset There are 544 samples in the testing dataset Each sample has 11 features

Part 1

- Initialize an estimator with the required model generator tree.DecisionTreeClassifier(criterion="entropy")
- Fit the estimator on the train data and target

In [10]:

Part 1.1

Let's see how it works on training data

- predict the target using the fitted estimator on the training data

In [11]:

The accuracy on training set is 100.0%

Part 1.2

Let's see how it works on test data, and, comparing with the result on training data, see if you can suspect overfitting

- use the fitted estimator to predict using the test features
- compute the accuracy and store it on a variable for the final summary

- store the maximum depth of the tree, for later use
 - fitted_max_depth = estimator.tree_.max_depth
- store the range of the parameter which will be used for tuning
 - parameter_values = range(1,fitted_max_depth+1)
- print the accuracy on the test set and the maximum depth of the tree

```
In [12]:
```

```
The accuracy on test set is 57.9%

The maximum depth of the tree fitted on X train is 19
```

Part 2 - Tuning with **Cross Validation**

Optimisation of the hyperparameter with **cross validation**. Now we will tune the hyperparameter looping on cross validation with the **training set**, then we will fit the estimator on the training set and evaluate the performance on the **test set**

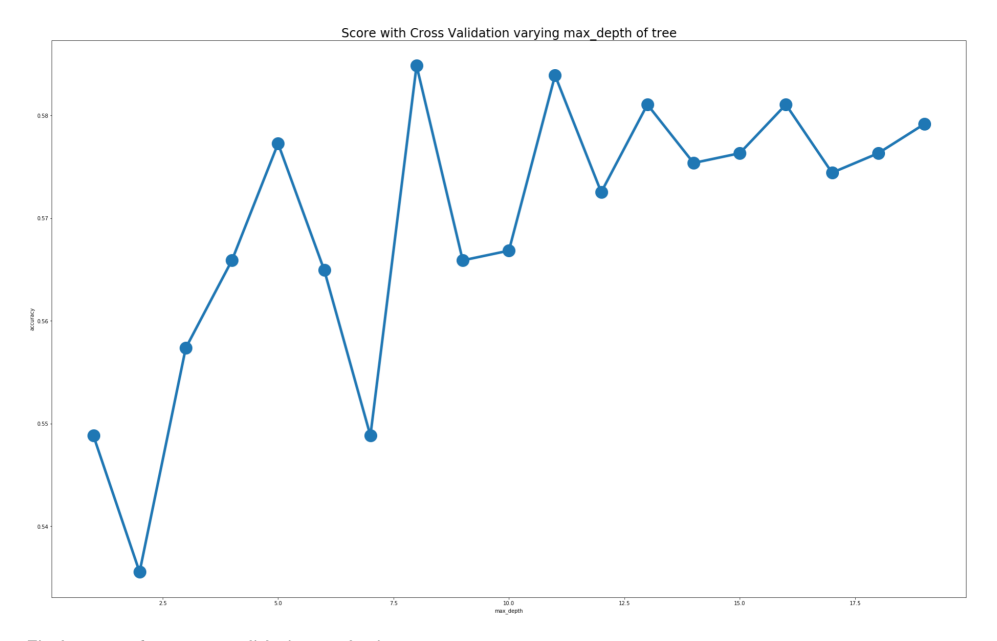
- initialize an empty list for the scores
- loop varying par in parameter_values
 - initialize an estimator with a DecisionTreeClassifier, using par as maximum depth and entropy as criterion
 - compute the score using the estimator on the train part of the features and the target using
 - o cross_val_score(estimator, X_train, y_train, scoring='accuracy', cv = 5)
 - o the result is list of scores
 - compute the average of the scores and append it to the end of the list
- · print the scores

```
In [13]:
```

```
[0.5488151658767773, 0.5355450236966824, 0.5573459715639811, 0.5658767772511849, 0.5772511848341232, 0.5649289099526066, 0.5488151658767773, 0.584834123222749, 0.5658767772511848, 0.5668246445497631, 0.5838862559241706, 0.5725118483412323, 0.581042654028436, 0.5753554502369668, 0.576303317535545, 0.581042654028436, 0.5744075829383886, 0.576303317535545, 0.5791469194312795]
```

Plot using the parameter_values and the list of scores

```
In [14]:
```



Fit the tree after cross validation and print summary

- store the parameter value giving the best score with np.argmax(scores)
- initialize an estimator as a DecisionTreeClassifier, using the best parameter value computed above as maximum depth and entropy as criterion

- fit the estimator using the train part
- use the fitted estimator to predict using the test features
- compute the accuracy on the test and store it on a variable for the final summary
- print the accuracy on the test set and the best parameter value

In [15]:

The accuracy on test set tuned with cross validation is 55.7% with depth 8

Show a more detailed information using the classification_report function of sklearn.metrics, using the true and predicted target values

In [16]:

	precision	recall	f1-score	support
3	0.00	0.00	0.00	5
4	0.00	0.00	0.00	18
5	0.64	0.66	0.65	227
6	0.54	0.57	0.55	223
7	0.42	0.38	0.40	65
8	0.00	0.00	0.00	6
accuracy			0.56	544
macro avg	0.27	0.27	0.27	544
weighted avg	0.54	0.56	0.55	544

- micro: Calculate metrics globally by counting the total true positives, false negatives and false positives.
- macro: Calculate metrics for each label, and find their unweighted mean. This does not take label imbalance into account.
- weighted: Calculate metrics for each label, and find their average weighted by support (the number of true instances for each label). This alters 'macro' to account for label imbalance; it can result in an F-score that is not between precision and recall.

Print the confusion_matrix , using the function of sklearn.metrics

In [17]:

[[0 0 5 0 0 0]

```
[ 0 0 8 9 1 0]
[ 0 7 150 64 6 0]
[ 1 2 67 128 23 2]
[ 0 0 3 36 25 1]
[ 0 0 0 2 4 0]]
```

Final report

Print a summary of the four experiments

Suggested exercises

- try to optimise the parameters "min_impurity_decrease"
- try to optimise using 'gini' instead of 'entropy'
- try to transform the classes: quality<7 transformed to low, quality>=7 transformed to high, then train and optimize with cross-validation, then test
 - try to fit using the parameter class class_weight to balance the classes
 - optimize the f1_score with macro weighting