

# report-homework-1

November 9, 2019

## 1 Dataset

The following machine learning exercises are applied on [sklearn wine dataset](#) that is taken from [UCI wine dataset](#). Before to go deep into the analysis it's useful for the reader to understand how wine dataset is composed, that is it might be helpful to give a look at its features, features distribution, cardinalities, kind of labels that are in the dataset and so on, since it's a crucial task in a data analysis. We can get a brief description of the dataset in the following table.

Features	Min	Max	Mean	SD
Alcohol	11.0	14.8	13.0	0.8
Malic Acid	0.74	5.80	2.34	1.12
Ash	1.36	3.23	2.36	0.27
Alcalinity of Ash	10.6	30.0	19.5	3.3
Magnesium	70.0	162.0	99.7	14.3
Total Phenols	0.98	3.88	2.29	0.63
Flavanoids:	0.34	5.08	2.03	1.00
Nonflavanoid Phenols	0.13	0.66	0.36	0.12
Proanthocyanins	0.41	3.58	1.59	0.57
Colour Intensity	1.3	13.0	5.1	2.3
Hue	0.48	1.71	0.96	0.23
OD280/OD315 of diluted wines	1.27	4.00	2.61	0.71
Proline	278	1680	746	315

- Missing Attribute Values: None
- Class Distribution: class\_0 (59), class\_1 (71), class\_2 (48)

From this brief description we can see that the dataset is composed by 13 continuous features and there are 3 class labels that represent 3 different wine categories. For the following exercises we will use a 2D representation of the dataset, in particular we will use only the first two features (i.e. Alcohol and Malic Acid). It could be interesting to see the distribution of these features, among all classes and one per class.

```
[2]: import sklearn as sk
from sklearn import datasets
import seaborn as sns
import pandas as pd
import matplotlib.pyplot as plt
```

```

raw_data = sk.datasets.load_wine()
features = pd.DataFrame(data=raw_data['data'],columns=raw_data['feature_names'])
data = features
data['target']=raw_data['target']
data['class']=data['target'].map(lambda ind: raw_data['target_names'][ind])
sns.distplot(data['alcohol'],kde=0)
plt.title("Alcohol distribution among all classes")
plt.show()

for i in data.target.unique():
    sns.distplot(data['alcohol'][data.target==i],
                  kde=1,label='{}'.format(i))

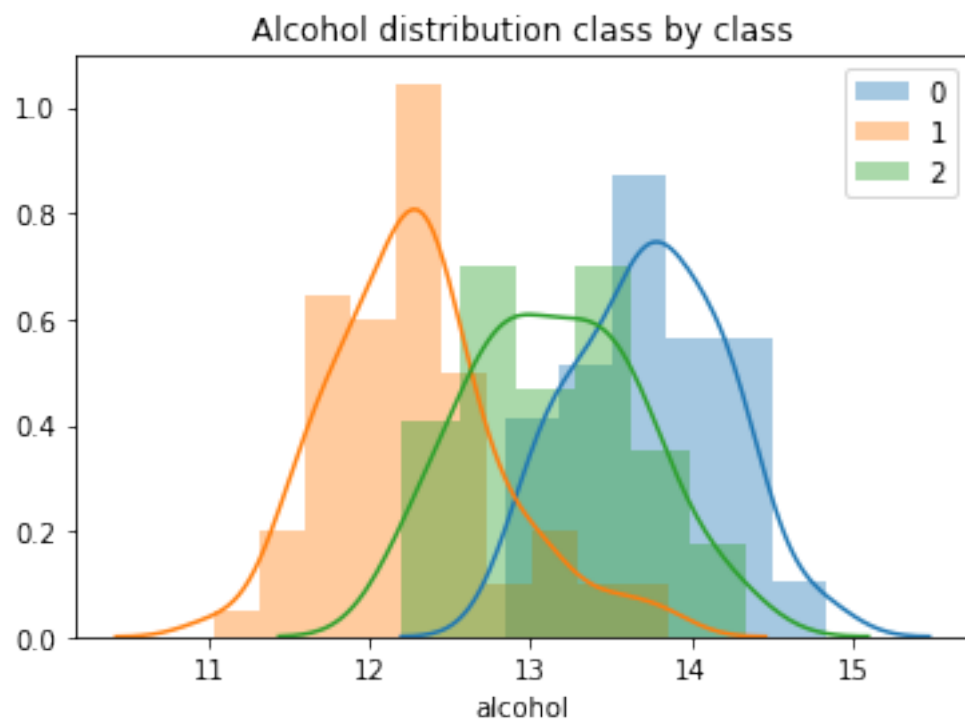
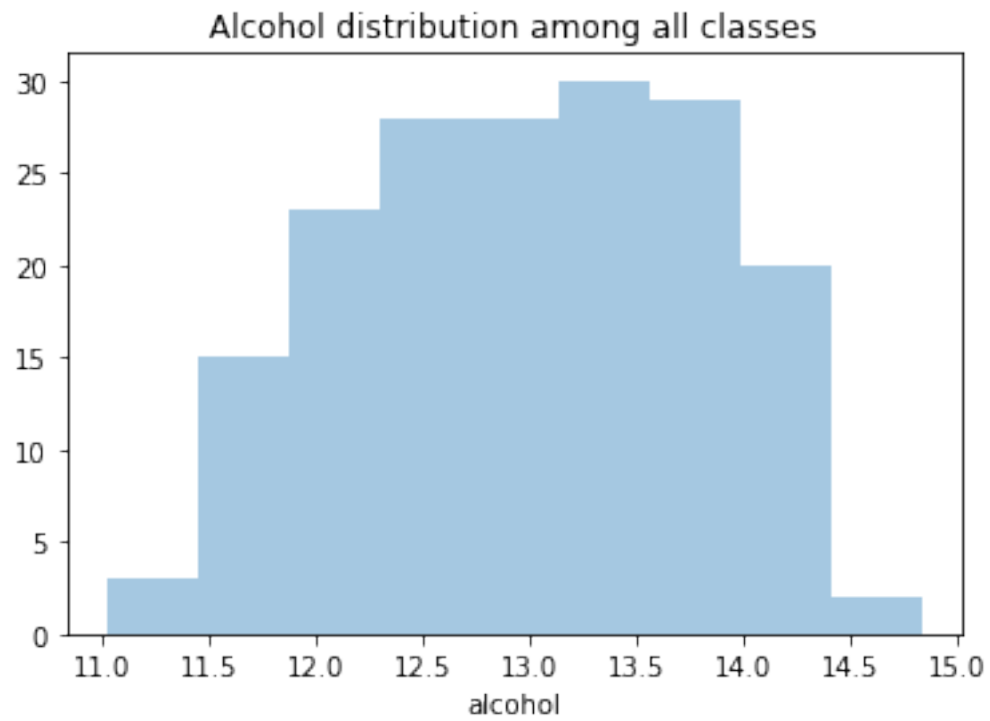
plt.title("Alcohol distribution class by class")
plt.legend()
plt.show()

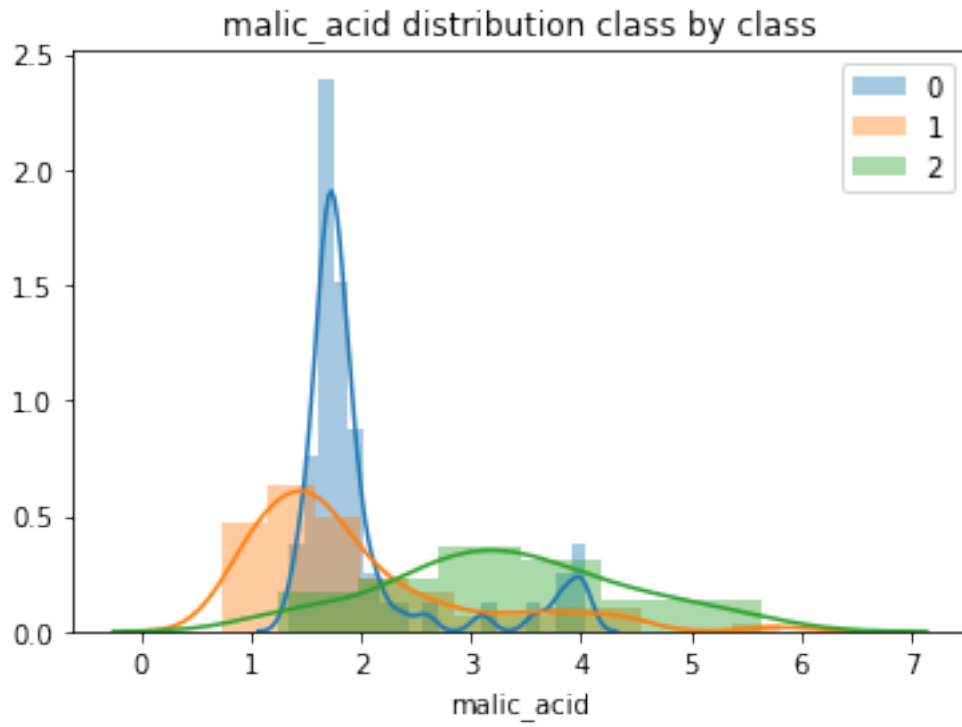
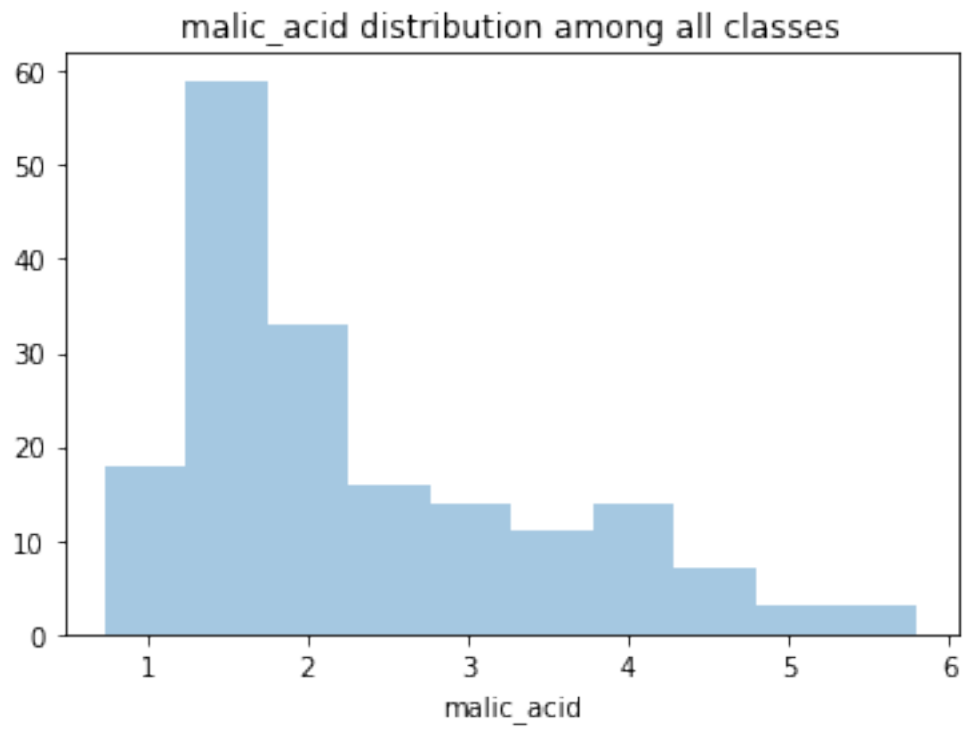
sns.distplot(data['malic_acid'],kde=0)
plt.title("malic_acid distribution among all classes")
plt.show()

for i in data.target.unique():
    sns.distplot(data['malic_acid'][data.target==i],
                  kde=1,label='{}'.format(i))

plt.title("malic_acid distribution class by class")
plt.legend()
plt.show()

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