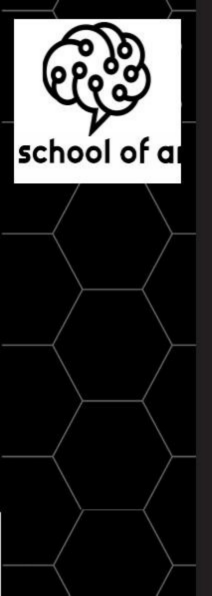


#### **Ensemble Learning: Random Forests**

**AAA-Python Edition** 



#### Plan

- 1- Random Forest
- 2- Extreme Random Forest
- 3- Balanced Random Forest
- 4- Feature Importance
- 5- Grid Search
- 6- Balancing by resampling



# **L- Random Forest**

#### Concept

A random forest is :

Bagging or pasting.

+

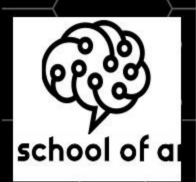
Decision Trees: with random split

==

**Bagging** with **Decision Trees** as classifier. The trees The trees use a random subset of features at each split

==

An **ensemble learning method** composed only of Decision Trees. They are trained on different random subsets of training data (with or without replacement). To split a node, the trees select the best features from a random subset of features.



# - Random Forest

#### Random forest in sklearn: method 1

 Using a Bagging Classifier: since a random forest is a special case, of a bagging method. We can use a Bagging Classifier with decision trees as estimator.

Confusion matrix =
[[11 0 0]
[ 0 10 0]
[ 0 1 16]]
score= 0.97

By specifying the splitter= "random", at each split, the chosen feature will be the "best random feature": the best feature from a random subset of features.



# **L- Random Forest**

#### Random forest in sklearn: method 2

 Using the random forest classifier class: in sklearn, the size of the samples used in the trees is the same as the original data size.

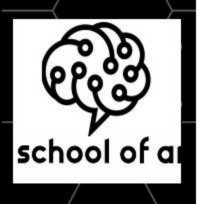
Same parameters used in the bagging classifier for the previous example.

[ 0 1 16]]

score= 0.97

There is no "splitter" parameter. Almost all other default parameters are the same. The only one that differs is : max\_features which is here set by default to "auto". The only parameter we had to specify is (to be adequate with the tree of the previous example) is : max\_depth

```
max_features ="auto" means that:
max_features = sqrt(numb_features)
```



## z- Extreme Random Forest

#### Concept

- In Extreme random forests, the thresholds used in the decision trees are selected randomly, instead of choosing the best one.
- They are also called "Extremely randomized Trees" or "Extra Trees".
- In sklearn, they are: "ExtraTreesClassifier" and "ExtraTreesRegeressor"

• "Extra**Trees**Classifier" are random forests, but there is in sklearn "Extra**Tree**Classifier" which is a decision tree and not a forest.



## z- Extreme Random Forest

#### Example

```
Confusion matrix =
[[11 0 0]
[ 0 10 0]
[ 0 1 16]]
score= 0.97
```

```
from sklearn.ensemble import ExtraTreesClassifier

myERF2= ExtraTreesClassifier(n_estimators=500, n_jobs=-1,max_depth=4,oob_score=True,bootstrap=True)
myERF2.fit(x_train, y_train)
y_pred = myERF2.predict(x_test)

print("Confusion matrix = \n",confusion_matrix(y_test,y_pred))
print("score=".np.round(myERF2.score(x_test.v_test).2))
```



## 2- Extreme Random Forest

#### Out of bag scores

- The bag refers to the training samples selected and used by a predictor in a bagging method.
- The out of bag, are the remaining samples.
- For each predictor, it is possible to compute its score using its out of bag samples.
- At the end of the training, the average of all these scores will represent the "out of bag score" of the learning method.
- In sklearn, you have to specify: "oob\_score =True" as parameter of the bagging method.



## 5- balanced Random Forest

#### Concept

- Sometimes, in classification, the representation of your classes in the training data is "**unbalanced**". For example, in a binary classification, you can have a big number of samples belonging to the first class. And, only few samples related to the other one.
- In this case, the learning method will not be able to classify correctly.
- To avoid this situation, scikit-learn allow us to define a parameter called: class\_weight, with a special value: "balanced" ==> it will attribute bigger weights for less present classes.
- The formula is as follow: n\_samples / (n\_classes \* np. incount(y))

  Count number of occurrences of each value in



## 3- Balanced Random Forest

#### Example without balancing

```
from pandas import DataFrame as DF, Series as S
import pandas as pd
myData = pd.read_csv("AAA-Ped-Week5/A3P-w5-data_imbalance.txt", header = -1)
x2= myData.iloc[:,[0,1]].values
y2= myData.iloc[:,2].values
```

```
0 1 2
0 5.66 6.77 1
1 4.40 5.05 0
2 3.52 4.73 1
```

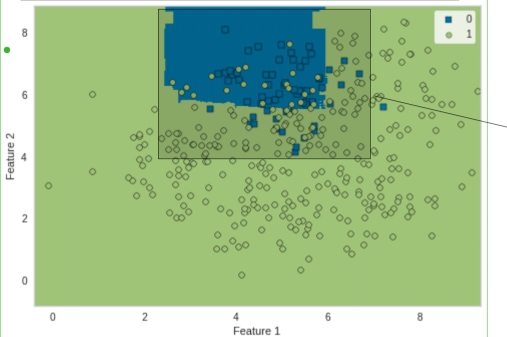
We will use **yellowbrick** library to visualize our classification boundaries.

Loading the data, and extracting **x** and **y** values



## 3- Balanced Random Forest

#### Example without balancing (suite)



The classes are not balanced

We see that some samples belonging to class 0 are predicted to be in class 1 and vice versa.

pr	ecision	recall	f1-score	support
Θ	0.65	0.66	0.66	62
1	0.93	0.93	0.93	313



#### Same example with balancing

```
myBRF2= RandomForestClassifier(n estimators=100, n jobs=-1,oob score=True, max depth=4,
                                     class weight="balanced")
myVis = DecisionViz(myBRF2, "Balanced Learning", features = ["Feature 1", "Feature 2"],
                        classes = [0.1])
                                                    Balanced classes
myVis.fit(x2_train,y2_train)
myVis.draw(x2_test, y2_test)
myVis.poof(outpath="balanced.png")
```

```
    We see clearly that the

  decision boundaries
  changed: the area of
  class 0 is bigger now
```

0.92

0.84

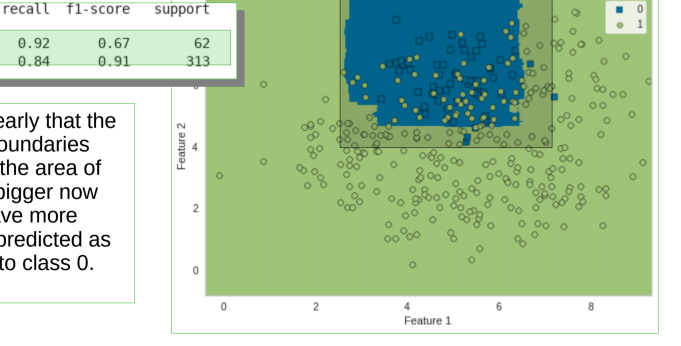
precision

0.53

0.98

 But, we have more classes 1 predicted as belonging to class 0.

[By Amina Delali]





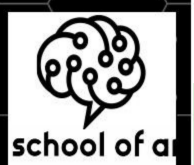
## 4- Grid Search

#### Concept

- GridSearch is a tool used to test a learning model with a list of dictionaries containing different parameters.
- We will use it to find the best parameters for a Random Forest method. We will apply it on the already used iris data.

```
print("\nThe obtained scores are:")
for params, mean, ally in myClassifier.grid_scores_:
    print(params, '-->', np.round(mean, 3))
print("The best parameters are :", myClassifier.best_params_)
```

First the first dictionary we have: 9 combinations of parameters. For the second one we have: 12 combinations.



## 4- Grid Search

#### Example results

```
{'min samples leaf': 1, 'n estimators': 20} ==> 0.964
{'min samples leaf': 1, 'n estimators': 40} ==> 0.964
{'min samples leaf': 1, 'n estimators': 60} ==> 0.964
{'min samples leaf': 2, 'n estimators': 20} ==> 0.973
{'min samples leaf': 2, 'n estimators': 40} ==> 0.973
{'min samples leaf': 2, 'n estimators': 60} ==> 0.964
{'min_samples_leaf': 3, 'n_estimators': 20} ==> 0.973
{'min samples leaf': 3, 'n estimators': 40} ==> 0.973
{'min samples leaf': 3, 'n estimators': 60} ==> 0.964
{'max depth': 1, 'n estimators': 10} ==> 0.946
{'max depth': 1, 'n estimators': 20} ==> 0.955
{'max depth': 1, 'n estimators': 30} ==> 0.964
{'max depth': 1, 'n estimators': 40} ==> 0.955
{'max depth': 2, 'n estimators': 10} ==> 0.955
{'max depth': 2, 'n estimators': 20} ==> 0.964
{'max depth': 2, 'n estimators': 30} ==> 0.964
{ 'max depth': 2, 'n estimators': 40} ==> 0.973
{ 'max depth': 3, 'n_estimators': 10} ==> 0.964
{'max depth': 3, 'n estimators': 20} ==> 0.964
{'max depth': 3, 'n estimators': 30} ==> 0.964
{| max depth|: 3. | n estimators|: 40} ==> 0.964
```

- We can see that augmenting the number of estimators doesn't necessary enhance the results.
- Same observation about max\_depth value

The best parameters are : {'min\_samples\_leaf': 2, 'n\_estimators': 20}



## 4- Grid Search

#### Best result visualization

We will display the confusion matrix using yellowbrick.

```
from yellowbrick.classifier import ConfusionMatrix
myEstimator= RandomForestClassifier(min_samples_leaf=2,n_estimators= 20)
myVisualizer = ClassificationReport(myEstimator, classes=["setosa","versicolour","virginica"],
                                                   support=True)
 myVisualizer.fit(x train,y train)
myVisualizer.score(x_test,y_test)
myVisualizer.poof(outpath="Confusion_
                                                                               RandomForestClassifier Classification Report
                                                                        0.909
                                                                                         0.909
                                                                                                           0.909
                                                                                                                               11
                                                          virginica
               The confusion
                                                                                                                                                  0.6
               matrix is
                                                                        0.923
                                                                                         0.923
                                                                                                           0.923
                                                                                                                               13
                                                        versicolour
               printed as a
                                                                                                                                                  0.4
               "heat-map":
               the best
                                                                                                                                                  02
               values are the
                                                                        1.000
                                                                                          1.000
                                                                                                                               14
                                                                                                           1.000
                                                           setosa
               darkest, and
               vice-versa.
[By Amina Delali]
```



## lmpor Featur 1

#### Concept

- The features in a data set have not the same importance.
- In a decision tree, this importance can be deduced from the distance between the appearance of a feature in a decision node and the root.
- In a random forest, the average of the distances corresponding to each tree will represent the feature's importance.
- Scikit-learn implements this method.

```
# we will use the previous trained estimator

for feature, importance in zip(myIris["feature names"], myEstimator.feature_importances_):
    print(feature,'==>', np.round(importance,5))

We can see clearly
    that the most important features
    petal length (cm) => 0.07181
    sepal length (cm) => 0.3492
    petal length (cm) => 0.57557

mportant features
    are: "petal length"
    and "petal width"
```

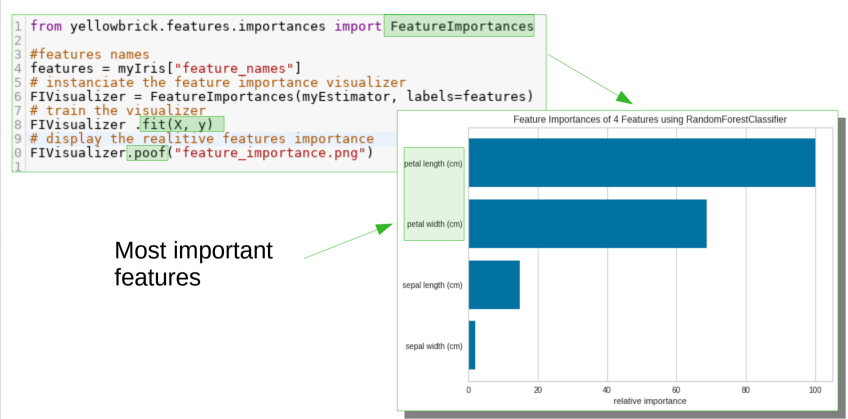


# - Feature Importance

2

#### Visualization of the features importance

Again, we will use yellowbrick.





# Feature Importance

L

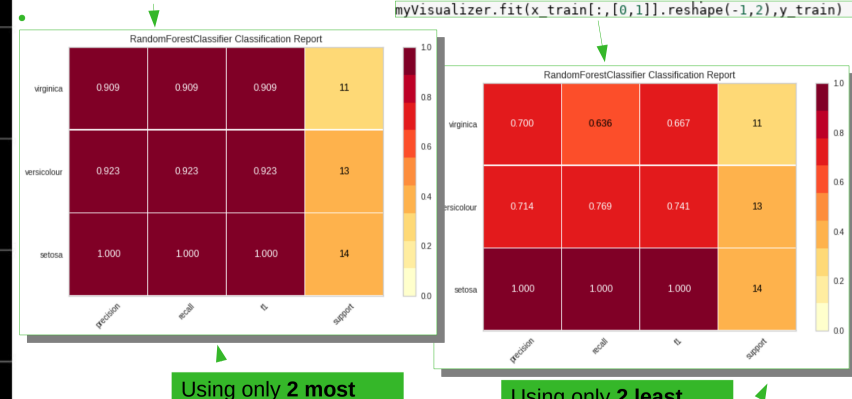
#### Comparison of results

myVisualizer.fit(x\_train[:,[2,3]].reshape(-1,2),y\_train)

**important** features

all the features)

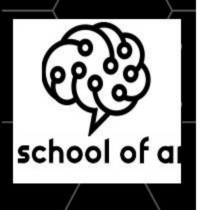
(same result as using



[By Amina Delali]

Using only 2 least important features (same result as using all the features)

18



### - Balancing y resampling

#### Concept

- An other way to obtain a balanced training set, is to remove samples from the majority class. Which is called: undersampling. Or, to add new samples belonging to the less represented class. Which is called: over-sampling.
- There is a variety of resampling techniques. The library **imbalanced-learn** library implements some of them.
- For a RandomForest classifier, it applies: random undersampling technique with different strategies.
- The default one is to: resample all classes but the minority class;



### o- Balancing by resampling

#### Example

 In this example, we combine the 2 libraries: yellowbrick and imbalanced-learn

```
print(classification_report(y2_test,myBRF3.predict(x2_test)))
```

support	f1-score	recall	precision		
72	0.64	0.89	0.50	0	
303	0.87	0.79	0.97	1	
375	0.81	0.81	0.81	avg	micro
375	0.75	0.84	0.73		macro
375	0.82	0.81	0.88		weighted

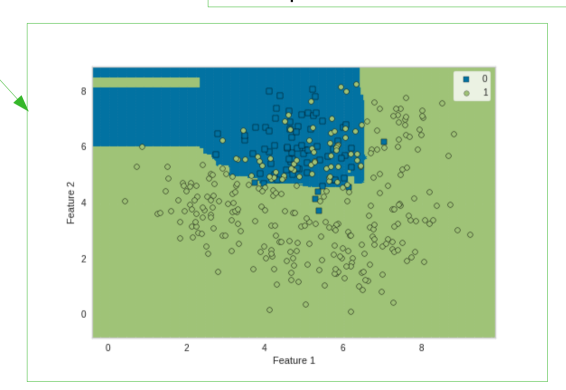


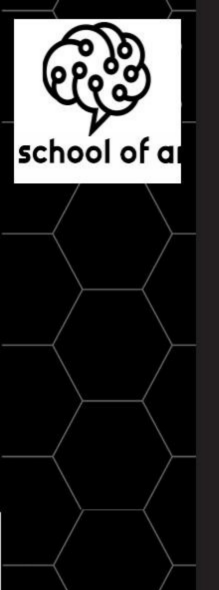
### - Balancıng y resampling

#### Example (suite)



We can see that the **class 0** region obtained is different than the one obtained using sckit-learn weighting technique





#### References

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## Thank you!

FOR ALL YOUR TIME