# Machine learning course – final exercise

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### 1. Algorithm name

#### FastForest

#### 2. Reference

Yates, Darren, and Md Zahidul Islam. "<u>FastForest: Increasing Random Forest Processing Speed While Maintaining Accuracy</u>." arXiv preprint arXiv:2004.02423 (2020).

# 3. Motivation for the algorithm

The algorithm aims to decrease the computation time of the Random Forest algorithm, while keeping its performance at the same level. Often, Random Forest achieves high-quality results. Since the authors' research domain is how to employ AI algorithms in mobile devices, they researched how to decrease the Random Forest runtime during training, to allow it to run in devices with limited resources.

In addition, in applications running in clouds that are being charged by the second (as in AWS), decreasing algorithms' computational time will lead to reduced expanses.

Finally, since the training of the trees can be done in parallel (except for sequential ensemble algorithms, as in the Boosting algorithm), the authors concluded that minimizing the training time per tree will be the best venue for decreasing the overall training time.

### 4. Short Description

### 4.1. Original Random Forest keynotes

The Random Forest algorithm (**RF**) is an ensemble learning method for classification and regression. It uses many decision trees and combines their results in order to predict its task. The trees are not pruned. In order to avoid overfitting it randomizes the data provided to each of its decision trees with:

- 1) Bagging suppose the training samples size is *n*. RF selects a random sample with replacement of size *n* from the training set as an input for each tree in its ensemble
- 2) Subspacing suppose the features number is F. RF selects a subset pf k features, k < F at each split. In scikit-learn implementation  $k = \sqrt{F}$  or  $k = \log_2 F$

#### 4.2. Fast Forest alterations

FastForest (**FF**) uses 3 revised components within the original flow of the Random Forest algorithm, to decrease the training time:

- 1) Half-Subbagging the algorithm randomly chooses half the samples for each tree in the ensemble, so the trees are trained with  $\frac{n}{2}$  of the samples, when the training samples size is n. as mentioned, Random Forests uses Bagging and selects n in random, with replacement
- 2) Logarithmic Split-Point Sampling when testing a numeric feature for the best split value, the method examines only Log<sub>2</sub> of the number of records in that node plus 1 (with non-missing value for that feature). It first sorts all possible values for the feature and then test them in hops so that no more than Log<sub>2</sub> of the number of records in that node plus 1 are considered. In the original Random Forest algorithm

- all unique values of the numeric features were considered as candidates for the split.
- There is no change for nominal features.
- 3) Dynamic Restricted Subspacing when the number of samples reaching a node is  $^1/_8$  of the size of the original training set or less then the number of features selected at the split increases, to allow selecting quality features. In Random Forest the Subspacing value is constant

### 5. Pseudo-Code

### 5.1. For training

```
FastForest - Building the ensemble (fit function)
Input: S - a labeled training set, S := (x_1, y_1), ..., (x_n, y_n)
         T – number of classifiers to train
         F – number of features
  For t=1 to T Do
     // the Half-Subbagging
         S_t \leftarrow \text{random sample with size } \frac{n}{2} \text{ from } S, \text{ with replacement}
2
3
         M_t \leftarrow buildDecisionTree(S_t, M_t, F)
4
         FastForest \leftarrow FastForest + M_t
   End For
    Return FastForest
buildDecisionTree function
Input: S'-a training set passed in the tree build recursion. its size is |S'|
         M_t – a classifier that was trained (decision tree), passes recursively during tree build
         F – number of features
7
     If S'represents a leaf Then
8
         M_t += createLeafWithPrediction()
9
         return M<sub>t</sub>
10 End If
     // the Dynamic Restricted Subspacing
11 If |S'| < 0.125 \times n Then
         k = \log_2\left(F \times \frac{n}{|S|}\right) + 1
12
13 Else
14
         k = \log_2(F) + 1
15 End If
16 A \leftarrow \text{select } k \text{ features from } F \text{ in random}
17 bestSplitInformationGain \leftarrow \infty, bestFeature \leftarrow null
18
     For i = 1 to k Do
19
         splitCandidateInformationGain \leftarrow findSplit(A_i, S)
         If splitCandidateInformationGain > bestSplitInformationGain Then
20
             bestSplitInformationGain \leftarrow splitCandidateInformationGain
21
22
             bestFeature \leftarrow A_i
23
         End If
24 End For
25 newS ← S`remainder after split according to bestFeature
26 M_t += buildDecisionTree(newS, M_t, F)
27 Return M_t
```

Figure 1 FastForest training stage pseudocode

```
findSplit function
Input: A – a feature according to which to split
        S'- a training set at this point
28 bestSplitInformationGain \leftarrow \infty, bestValueToSplit \leftarrow null
29
   If A is nominal Then
30
        return (bestValueforSplit, informationGain)
31 End If
    // the Logarithmic Split-Point Sampling
32 sortedUniqueValues ← find unique values for A in S` and sort them
33 valuesToExamine ← start with sortedUniqueValues[0] till sortedUniqueValues[last]
                          In steps of \frac{sortedUniqueValues[last] - sortedUniqueValues[0]}{}
    For v in valuesToExamine Do
34
35
        informationGain \leftarrow calculateInformationGain(I, S`, v)
        If informationGain > bestSplitInformationGain Then
36
37
             bestSplitInformationGain \leftarrow informationGain
38
             bestValueToSplit \leftarrow v
39
        End If
40 End For
41 Return (bestValueToSplit, bestSplitInformationGain)
```

Figure 2 FastForest training stage pseudocode - continued

## 5.2. For classifying

```
FastForest – predicting
Input: S - a test set, S := (x_1, y_1), ..., (x_a, y_a)
   For s in S Do
2
         votes \leftarrow [0, ..., 0]
                                    // votes size is as the number of classes
2
         For tree in FastForest Do
3
             prediction \leftarrow tree.predict(s)
              votes[prediction] \leftarrow votes[prediction] + 1
4
5
         End For
6
         predictions \leftarrow predictions + argmax(votes)
    End For
    Return predictions
```

Figure 3 FastForest prediction stage pseudocode

# 6. Algorithm Explanation

#### 6.1. For training

The explanation refers to Figure 1 and Figure 2.

In line 1 the FastForest creates a loop to grow the ensemble of *T* decision trees.

In line 2 it produces the Half-Subbagging, the training set for the  $t^{th}$  tree. It randomly selects  $\frac{n}{2}$  samples from the training set and passes it to the function that builds a single decision tree (line 3).

In line 4 the tree that was built,  $M_t$ , is appended to the ensemble of trees of the algorithm.

The next lines describe the *buildDecisionTree* function: it is a recursive function that grows the decision tree. Its input is the training data remained at that stage, the tree grown and the total number of features in the set.

In line 7 the algorithm checks if it reached a stopping criterion for the recursion – whether the training set represents a leaf. This can be true for several reason, among which –

- 1) The training set is pure and has sample of only 1 class in it
- 2) The max\_depth of the tree is achieved
- 3) The minimal number of samples for node is violated

The algorithm then creates a leaf, sets the relevant prediction for that leaf and returns the tree grown so far (lines 8-9).

Otherwise, in lines 11-12, the Subspacing size is calculated – how many features should be considered for the current split. To allow a quality split, the algorithm performs a Dynamic Restricted Subspacing and allows larger number of features when the number of samples in the node is less than  $^1/_8$  of the size of the original training set. In that case the Subspacing size will be  $\log_2\left(F\times\frac{n}{|S^*|}\right)+1$ , where F is the total number of features, n is the total number of samples in the training set and  $|S^*|$  is the number of samples that reached that node. If the condition is not met then the Subspacing size will be  $\log_2(F)+1$  (lines 13-14).

In line 16 the algorithm selects those features in random, and then loops on them and examines each one to determine whether splitting the tree on that feature will result with the best information gain (lines 17-20). The calculation is done with the *findSplit*() function describe below. If the split improved the current best information gain, the algorithm saves the information gain obtained thus far and the best value to split on (lines 21-22).

Once all the features were considered and the best feature and split point discovered, the algorithm creates the new training set according to the split and call the function again with the new parameters.

The *findSplit*() pseudo code is lines 28-41. The input for the function is the feature and the training set at that point. Firstly, the algorithm checks whether the feature in the input is nominal (line 29). If so, then the original code of the Random Forest is called (line 30). Otherwise, the algorithm performs the Logarithmic Split-Point Sampling:

Instead of going over all possible unique values of a numeric feature for a split, the algorithm considers only  $\log_2$  of the number of records with non-missing values reaching that node (32, 33). The algorithm calculates the information gain obtained from splitting on each of these values and returns the best value and information gain obtained (lines 34-41).

### 6.2. For classifying

The FastForest algorithm didn't enhanced the flow of the Random Forest for classifying. The pseudocode of the algorithms is described in Figure 3.

The input for the function is the test set. The algorithm loops on the samples in the set and activated the *T* decision trees predict function on them. It combined all the T predictions and uses a majority vote to output the predicted class per sample in the test set.

#### 7. Illustration

In this section I illustrate the growing of one tree by the FastForest algorithm. The settings of the algorithm are:

- 1. Number of trees to grow (T) = 3 (only one tree will be demonstrated though)
- 2. Max depth of the tree = 3
- 3. Min number of samples in a leaf = 2

I created a synthetic dataset to be used as the training set, based on a simplified version of the iris dataset (Table 1). I'll use the entire dataset for the illustration and will not divide it into train and test sets.

#	f1	f2	f3	f4	class
0	-1.2	0.1	-1.2	-1.3	0
1	-0.5	0.7	-1.2	-1.0	0
2	-0.7	2.4	-1.2	-1.4	0
3	-0.4	2.6	-1.3	-1.3	0
4	-1.1	0.1	-1.2	-1.4	0
5	-1.0	0.3	-1.4	-1.3	0
6	-0.4	1.0	-1.4	-1.3	0
7	-1.1	0.1	-1.2	-1.4	0
8	1.3	0.3	0.5	0.2	1
9	0.6	0.3	0.4	0.3	1
10	1.2	0.1	0.6	0.3	1
11	-0.4	-1.7	0.1	0.1	1
12	0.7	-0.5	0.4	0.3	1
13	-0.1	-0.5	0.4	0.1	1
14	0.5	0.5	0.59	0.5	1
15	-1.1	-0.5	-0.2	-0.2	1
16	0.5	0.3	1.2	1.7	2
17	-0.1	-0.8	0.6	0.3	2
18	1.5	-0.1	1.2	1.7	2
19	0.5	-0.3	1.2	0.2	2
20	0.7	-0.1	1.2	1.7	2
21	2.1	-0.1	0.6	1.7	2
22	-1.1	-1.2	0.4	0.1	2
23	1.7	0.3	0.4	0.2	2

Table 1 A synthetic dataset for the illustration of the FastForest algorithm

Figure 4 illustrates the first step in the algorithm (line 2 in Figure 1). The dataset consists of 24 samples and the algorithm selects half of them in random (Half-Subbagging), per tree. So, the input per tree is a dataset with 12 samples only. We will concentrate on the growing of tree 2, in the first 2 calls for the *buildDecisionTree* unction. The Half-Subbagging result of the illustration example is presented in Table 2.

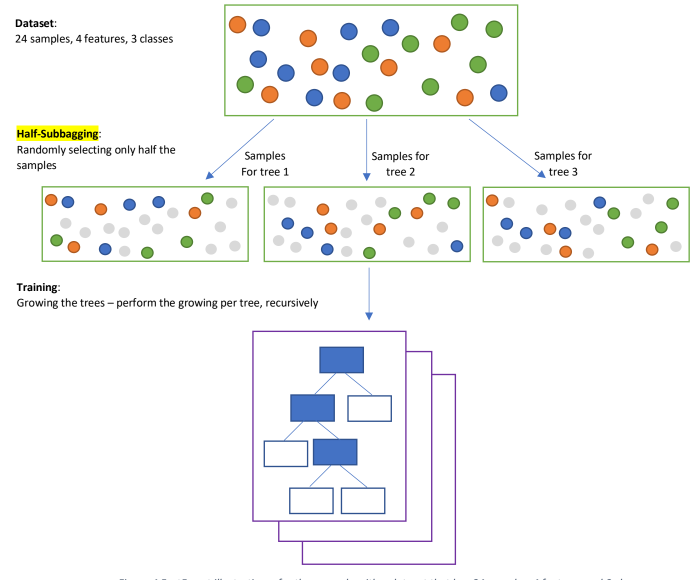


Figure 4 FastForest illustration – for the example with a dataset that has 24 samples, 4 features and 3 classes.

Performing Half-Subbagging on the input and growing 3 trees

Once the Half-Subbagging is completed, the algorithm starts growing the tree. The growing is done recursively and stops once all the data is classified in the leaves. The procedure is illustrated in Figure 5. So, in the first call to the *buildDecisionTree* function (line 7 in Figure 1), the algorithm first checks if the input data can be classified as a leaf. We have 12 samples and they do not meet the conditions for being a leaf – they are not pure, nor the *max\_depth* limitation or the *min\_samples\_per\_leaf* limitation are met.

So, continuing to line 11, the algorithm calculates the Subspacing number: the number of features to select in random from the 4 features in total. Since we are in the first call, the number of samples equals to the total number of samples and the algorithm selects

$$\log_2(4) + 1 \rightarrow 3$$
 features (line 14):  $f_1, f_2, f_3$ .

Then, the algorithm loops over the 3 selected features, and for each finds the best value to split the data on (the *findSplit* function). In a greedy manner, once it discovered the feature

and value that have the best information gain, it splits the data according to them and calls itself again with the new data (line 26).

#	f1	f2	f3	f4	class
0	-1.2	0.1	-1.2	-1.3	0
1	<mark>-0.5</mark>	<mark>0.7</mark>	<mark>-1.2</mark>	<mark>-1.0</mark>	<mark>0</mark>
2	-0.7	2.4	-1.2	-1.4	0
<mark>3</mark>	<mark>-0.4</mark>	<mark>2.6</mark>	<mark>-1.3</mark>	<mark>-1.3</mark>	<mark>O</mark>
<mark>4</mark>	<mark>-1.1</mark>	<mark>0.1</mark>	<mark>-1.2</mark>	<mark>-1.4</mark>	<mark>O</mark>
5	-1.0	0.3	-1.4	-1.3	0
<mark>6</mark>	<mark>-0.4</mark>	<mark>1.0</mark>	<mark>-1.4</mark>	<mark>-1.3</mark>	<mark>O</mark>
7	-1.1	0.1	-1.2	-1.4	0
8	1.3	0.3	0.5	0.2	1
<mark>9</mark>	<mark>0.6</mark>	<mark>0.3</mark>	<mark>0.4</mark>	<mark>0.3</mark>	<mark>1</mark>
10	1.2	0.1	0.6	0.3	1
<mark>11</mark>	<mark>-0.4</mark>	<mark>-1.7</mark>	<mark>0.1</mark>	<mark>0.1</mark>	<mark>1</mark>
<mark>12</mark>	<mark>0.7</mark>	<mark>-0.5</mark>	<mark>0.4</mark>	<mark>0.3</mark>	<mark>1</mark>
13	-0.1	-0.5	0.4	0.1	1
14	0.5	0.5	0.59	0.5	1
<mark>15</mark>	<mark>-1.1</mark>	<mark>-0.5</mark>	<mark>-0.2</mark>	<mark>-0.2</mark>	<mark>1</mark>
16	0.5	0.3	1.2	1.7	2
<mark>17</mark>	<mark>-0.1</mark>	<mark>-0.8</mark>	<mark>0.6</mark>	<mark>0.3</mark>	<mark>2</mark>
<mark>18</mark>	<mark>1.5</mark>	<mark>-0.1</mark>	<mark>1.2</mark>	<mark>1.7</mark>	<mark>2</mark>
19	0.5	-0.3	1.2	0.2	2
<mark>20</mark>	<mark>0.7</mark>	<mark>-0.1</mark>	<mark>1.2</mark>	<mark>1.7</mark>	<mark>2</mark>
21	2.1	-0.1	0.6	1.7	2
<mark>22</mark>	<mark>-1.1</mark>	<mark>-1.2</mark>	<mark>0.4</mark>	<mark>0.1</mark>	<mark>2</mark>
23	1.7	0.3	0.4	0.2	2

Table 2 the Half-Subbagging result for tree 2. The selected rpws from the original dataset are highlighted in yellow

In our dataset all the features are numeric. The *findSplit* function, as presented in the algorithm in Figure 2, first sorts all the values and calculates the split point candidates according to the Logarithmic Split-Point Sampling component. For  $f_1$ , for example, the unique values are: [-1.1 -0.5 -0.4 -0.1 0.6 0.7 1.5], total of 7 different values.

The Logarithmic Split-Point Sampling component takes these values and considers only  $\log_2(|S^*|)$ , where  $|S^*|$  is the number of samples that reached that node, as candidate split values. It takes the first value in the sorted unique values array of the feature and adds to it the hop, which is

$$\frac{last\ sorted\ value-first\ sorted\ value}{\log_2(records\ number)+1}=\ \frac{1.5-(-1.1)}{\log_2(12)+1}=\frac{2.6}{4.6}=0.52.$$

So, the split candidates for  $f_1$  are [-1.1, -0.58, -0.06, 0.46, 0.98], only 5 values.

Then, the algorithm calculates the information gain for splitting according to each of the candidate values and returns the best feature and value to split on. The tables below display the search for the best feature and value in this first call to *buildDecisionTree* function.

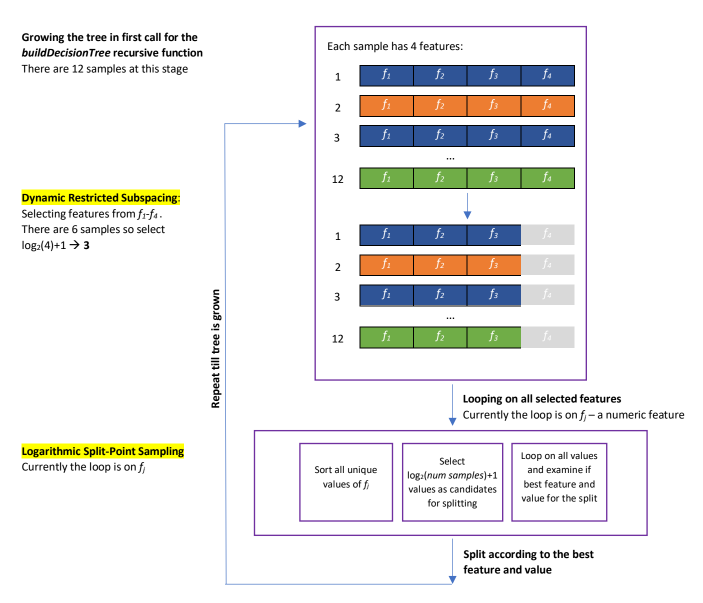


Figure 5 FastForest illustration example continued. It is the first call for the buildDecisionTree function and the input consists of 12 samples. Each sample has 4 features. The algorithm builds the decision tree by selecting the features (Dynamic Restricted Subspacing) and finding the best feature and values to split the tree on. It repeats this procedure till all the dataset is classified in the tree's leaves and returns the final outcome to the FastForest object

the best feature, value and entropy is highlighted in blue in the tables below:

	First call for the <i>buildDecisionTree</i> function - Candidate split feature: f1									
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)						
1	-1.1	∞	1.585	[4,15,22] [1,3,6,9,11,12,17,18,20]						
2	-0.58	1.585	1.585	[4,15,22] [1,3,6,9,11,12,17,18,20]						
3	<del>-0.06</del>	1.585	<b>1.333</b>	[1,3,4,6,11,15,17,22] [9,12,18,20]						
4	0.46	1.333	1.333	[1,3,4,6,11,15,17,22] [9,12,18,20]						
5	0.98	1.333	1.442	[1,3,4,6,9,11,12,15,17,20,22] [18]						

The unique values for  $f_2$  are:

and the output of the Logarithmic Split-Point Sampling component for them is:

[-1.7, -0.84, 0.02, 0.88, 1.74]

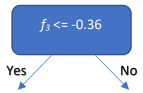
	First call f	or the <i>buildDecision</i> 1	Tree function - Can	didate split feature: f <sub>2</sub>
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)
1	-1.7	1.333 (of feature $f_1$ )	1.441	[11] [1,3,4,6,9,12,15,17,18,20,22]
2	-0.84	1.333	1.476	[11,22] [1,3,4,6,9,12,15,17,18,20]
3	-0.02	1.333	0.875	[11,12,15,17,18,20,22] [1,3,4,6,9]
4	0.88	0.875	1.268	[1,4,9,11,12,15,17,18,20,22] [3,6]
5	1.74	0.875	1.441	[1,4,6,9,11,12,15,17,18,20,22] [3]

# And for $f_3$ :

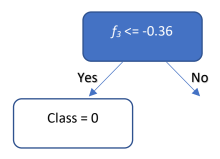
	First call for the <i>buildDecisionTree</i> function - Candidate split feature: f <sub>3</sub>								
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)					
1	-1.4	0.875 (of feature $f_2$ )	1.441	[4] [1,3,6,9,11,12,15,17,18,20,22]					
2	-0.88	0.875	0.667	[1,3,4,6] [9,11,12,15,17,18,20,22]					
3	<del>-0.36</del>	0.667	0.667	[1,3,4,6] [9,11,12,15,17,18,20,22]					
4	0.16	0.667	0.918	[1,3,4,6,11,15] [9, 12,17,18,20,22]					
5	0.68	0.667	1.268	[1,3,4,6,9,11,12,15,17,22] [18,20]					

The output of the first call is to split on  $f_3$  with value -0.36. and now the function calls itself recursively with the split data indices: Indices [1,3,4,6] creates the left sub-tree and [9,11,12,15,17,18,20,22] creates the right sub-tree.

At this stage the algorithm found the root of the tree and it is composed of -



Calling *buildDecisionTree* with the [1,3,4,6] indices creates a leaf node – the data is pure and classifies the samples reaching this node as class 0. The tree now has a leaf:



At this stage the Dynamic Restricted Subspacing takes place again and this time features  $f_1$ ,  $f_2$ ,  $f_4$  are selected.

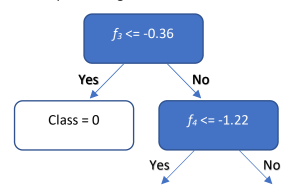
The calculations for the left sub-tree with the [9,11,12,15,17,18,20,22] indices are:

	Building left sub-tree with the <i>buildDecisionTree</i> function - Candidate split feature: f1								
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)					
1	-1.1	∞	1.0	[15,22] [9,11,12,17,18,20]					
2	-0.45	1.0	1.0	[15,22] [9,11,12,17,18,20]					
3	0.2	1.0	1.0	[15,17,22] [9,11,12,18,20]					
4	0.85	1.0	0.862						

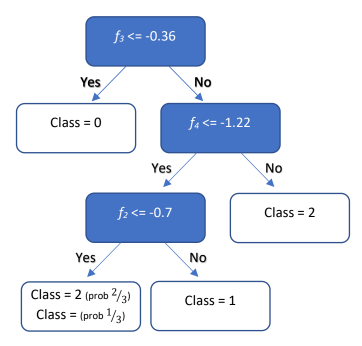
	Building left sub-tree with the <i>buildDecisionTree</i> function - Candidate split feature: <b>f</b> <sub>2</sub>								
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)					
1	<mark>-1.7</mark>	0.862	<mark>0.862</mark>	[11] [9,12,15,17,18,20,22]					
2	1.2	0.862	1.0	[11.22] [9,12,15,17,18,20]					
3	-0.7	0.862	0.951	[11,17,22] [9,12,15,18,20]					
4	-0.2	0.862	0.951	[11, 12,15,17,22] [9,18,20]					

	Building left sub-tree with the <i>buildDecisionTree</i> function - Candidate split feature: fa								
#	Candidate split value	Best gain thus afar	Split entropy	Data indices split (from Table 2)					
1	-0.2	∞	0.862	[12] [9,11,15,17,18,20,22]					
2	0.27	0.862	0.951	[11,15,22] [9,12,17,18,20]					
3	0.75	0.862	0.689	[9,11,12,15,17,22] [18,20]					
4	1.22	0.689	0.689	[9,11,12,15,17,22] [18,20]					

So,  $f_4$  is selected and the tree is proceeding with –



The procedure is repeated until all the data is divided into the tree's leaves or the limitation for the tree's depth or number of samples in the leaf are met. The final outcome for this illustration is:



As can be seen, the tree stops at depth 3 and one of the leaves is not pure.

### 8. Strengths

- 1. FastForest is an optimization for Random Forest. It optimizes the processing time required for training the forest's estimators and in the paper the authors achieved an overall tested speed gain of over 24% compared with Random Forest on 45 datasets (when running on Windows platform).
- 2. It was also compared with 5 other ensemble algorithms: SysFor, Random Subspace, ForestPA, Bagging and Random Committee, and was found to be faster than them as well.
- 3. FastForest was also tested on Android smartphones, on 30 datasets, and outpaced Random Forest on all of them.
- 4. FastForest performance was found to be as good as Random Forest on the datasets tested.

#### 9. Drawbacks

- FastForest is composed of 3 optimization components: the Half Subbagging, the
  Logarithmic Split-Point Sampling and the Dynamic Restricted Subspacing, aiming to
  reduce the processing time. The Logarithmic Split-Point Sampling and the Dynamic
  Restricted Subspacing component is relevant only for numeric features, so when a
  dataset has mainly categorical features then its contribution to the overall result is very
  limited.
- 2. The Logarithmic Split-Point Sampling is not considering the distribution of the unique values of the numeric values and rather assumes a uniform distribution. It divides the range of the values in  $\log_2(\text{number of samples in the node})$  sections and splits the samples accordingly. If the distribution is very skewed then it will miss out many worthy potential candidates. For example: if the unique values of a feature are [0.01, 0.02, 0.03, 0.04, 0.05, 10] and 8 samples reached the node then the hop value will be  $\frac{10-0.1}{\log_2 8+1} \approx 2.5$  and the algorithm will consider these values as split candidates: [2.6, 5.1, 7.6], resulting always in the same split outcome: [0.01, 0.02, 0.03, 0.04, 0.05] and [10]. If the data is normalized then it won't pose as a problem though.
- 3. For very simple datasets, the FastForest algorithm may take longer time for the training of the estimators. Due to the Dynamic Restricted Subspacing component, it may consider more features for splits when compared with the Random Forest algorithm. However, when the dataset is simple then the training time is short at any case.
- 4. When the dataset is extremely imbalanced than the Half-Subbagging may lead the training sets without some of the classes.

### 10. Experimental Results

I conducted the experiments on several computers in parallel, on the Windows platform. The protocol of the experiments—

- 1. Loop on all datasets and
  - a. Read the dataset's csv file
  - Perform minimal and generic preprocessing for the data (more details in 10.1.1)
  - c. Create 10-Fold CV for splitting for training and testing (section 10.1.2). For FastForest and Random Forest do
    - i. Perform 3-Fold CV on the train set to tune the hyper-parameters for each fold created in step 3. Above (section 10.1.3)
    - ii. Create the estimator with the best parameters and predict the class for the test set on the best hyper-parameters
    - iii. Calculate performance measurements (results in 10.2.1)
- 2. Test if the differences between FastForest and Random Forest results are statistically significant (section 10.2.2)
- 3. Build a Meta-learning model with XGBoost and calculate the features importance measures (section 10.2.3)

### 10.1 Implementation notes

The paper's authors did not publish their code for FastForest. My original intention was to derive the Random Forest implementation in scikit-learn package and to override the relevant parts with my implementation of the FastForest ideas.

Random Forest code is written in Python and its estimators are written in Cython in the scikit-learn package. I was able to change the Python code, but couldn't make the estimators code to cythonized. As a result, I implemented a Decision Tree Classifier from scratch, in python, based on examples I found online ([1], [2], [3]). I implemented binary decision trees, as in scikit-learn, and for calculating the information gain I chose the entropy measure.

#### 10.1.1 Datasets preprocessing

The preprocessing was done automatically per dataset and was kept as general and generic as possible.

The classification column (class) was set as the last column of the datasets, except for the 'solar-flare'. The class column appears there as the first column, and I discovered it while analyzing the results from this dataset, and did the change to improve the results.

I filled in missing values with a unique value. I preferred not to drop samples since most datasets are small. I also performed one hot encoding for categorical features and did label encoding for the class column.

Lastly, several datasets have only 1-4 samples of certain class values. I marked these values as extreme minority classes and had several venues for handling –

- 1. Find more data I couldn't find additional relevant datasets online to enrich my data and increase the number of the extreme minority classes
- 2. Perform random oversampling or undersampling most datasets with this issue are quite small, so undersampling is irrelevant. Oversampling is also problematic since there are only 1-4 samples for the process
- 3. Remove the extreme minority classes I chose that venue and dropped the samples with extreme minority classes from the datasets

### 10.1.2 Data split

As instructed, I used 10-fold CV for the splits. Since some dataset were imbalanced I did the splits with stratification, unless the number of the least populated class value was less than the number of folds (10).

In addition, one dataset, 'lenses', was very small, with 24 samples in total, and imbalanced. Consequently, the 10-folds CV failed for it and so I performed only 4-folds instead.

#### 10.1.3 Hyper parameters tuning

I used the RandomizedSearchCV for tuning the algorithms hyper-parameters, as instructed.

For FastForest and Random Forest I tuned -

- 1. n\_estimators the number of trees for the forest to grow. I ran the experiments on several "regular" PCs so I set the options as 5, 15 and 60 trees, to limit the runtime.
- 2. max\_depth the maximal depth of the Decision Tree, 5, 8 or 15. These values should allow growing satisfactory binary trees
- 3. min\_samples\_leaf the minimal number of samples in a leaf node in the Decision Tree. If the number of samples reaching a node were less than this value then a lead node was created. The options were 1, 2, 4

For Random Forest I also tuned -

1. bootstrap – whether to perform bootstrapping or not. For FastForest I always chose half the samples, as required in the algorithm description

- 2. max\_features the number of features to consider for splits (Subspacing). The options were:
  - a. squared root of the number of features in the dataset
  - b. log<sub>2</sub> of the number of features in the dataset

for FastForest I used the Dynamic Restricted Subspacing component, as described in section 6.1.

I chose these 5 parameters since I expected them to have the most meaningful impact on the results. Since I ran the experiments on standard desktop computers I chose the options per parameter to allow the algorithms reasonable performance but with reasonable runtime as well.

#### 10.2 The results

#### 10.2.1 Performance comparison between FastForest and Random Forest

The performance was measured according to several metrics mentioned in the exercise instructions. I used functions implemented in the scikit-learn package, or implemented them myself, specifically –

- 1. Accuracy computed with the accuracy\_score function
- 2. TPR I implemented the calculation by calculating the confusion matrix and FP, FN, TP and TN. Then I calculated the average of the  $\frac{TP}{TP+FN}_{FP}$
- 3. FPR similarly to the calculation of TPR, I calculated  $\frac{FP}{FP+TN}$
- 4. Precision I used the average\_precision\_score function. For multi-classification problems I first binarized the classes and calculated the function in a one-vs-rest manner. Then I calculated the average for the results of all the classes.
- 5. AUC I used the roc\_auc\_score. If the multi-classification problem then I added the *multi\_class* parameter as "ovr" (one-vs-rest) and the *average* parameter as "weighted", since many datasets were imbalanced
- 6. Precision-Recall score I used the average\_precision\_score function with the average parameter as "weighted"
- 7. Training time measured in seconds. I first tuned the hyper parameters and only then measured the training time for the best results
- 8. Inference time measured in seconds and normalized for 1000 samples

The average of each of the performance measurements is in Table 3. We can see the average result per measurement and then its standard deviation in parenthesizes.

	Accuracy	TPR	FPR	Precision	AUC	PR-	Training	Inference
						Curve	Time	Time
Fast	0.7445	0.6584	0.6584	0.723	0.8437	0.7046	1.4197	0.6278
Forest	(0.1898)	(0.2239)	(0.2239)	(0.2189)	(0.1626)	(0.2421)	(1.8446)	(0.5286)
Random	0.6946	0.5996	0.5996	0.6509	0.7844	0.6421	15.2569	0.4512
Forest	(0.2064)	(0.2277)	(0.2277)	(0.2487)	(0.1809)	(0.2515)	(48.164)	(0.4236)

Table 3 Average and std results for the performance measurements per algorithm for the 150 datasets tested

The complete results, per dataset and cross validation run, are in an accompanying file – results\_with\_avg.csv (attaching it here makes the document too long).

We can see that in all the measurements but the inference time, FastForest achieved better average results than the Random Forest algorithm. The measurement in focus, the training time, is significantly lower for FastForest than for Random Forest.

### 10.2.2 Statistical significance

I performed the Mann–Whitney U test to examine the significance of the differences between the results of the two algorithms, as implemented in the Scipy package. The tests' results are summarized in Table 4.

Parameter	FastForest average value	Randome Forest average value	Statistic	p-value
Training time	1.4197	15.2569		
Accuracy	0.7445	0.6946		
AUC	0.8437	0.7844		
Inference time	0.6278	0.4512		

Table 4 Mann-Whitney U test results for the differences between FF and RF

As in the original paper, the training time when applying the FastForest algorithm is statistically significantly lower than the training time of the Random forest algorithm for the 150 datasets tested (p-value ??). It takes on average 10 seconds less to train the algorithm with FastForest, which is ??% faster.

The accuracy and AUC differences also turned out to be statistically significant (p-value ??). in the paper FastForest frequently exceeded Random Forest for classification accuracy as well.

### 10.2.3 Meta-learning model

#### 11. Conclusions

#### 12. Citations

The paper was published on 2020 and was not yet cited by others.

However, the paper's authors performed similar earlier work in [4]. There, they worked on a single Decision Tree and also aspired o decrease the processing time of the tree. When encountering a numeric feature as a candidate for the tree split, they limit the number of values to examine (the Split Point Sampling, or SPS) to up to 20, distributed evenly on the feature's values range. They empirically showed that this method decreased the raining time whilst maintaining the tree's performance similar to a regular Decision Tree.

The FastForest was created to examine SPAARC in an ensemble and to improve it by changing the arbitrary Split Point Sampling value of 20 to a parameter with relevance to the data: the Logarithmic Split-Point Sampling in FastForest takes into account the number of records in each split. It keeps the number of split candidates low when there is a large number of records, and increases the number when the records count decreases.

# 13. Additional references

- [1] Brownlee, J., How to Implement Random Forest From Scratch in Python (2016)
- [2] Kumar, V., Random forests and decision trees from scratch in python (2018)
- [3] Mantey, S., Coding a Decision Tree from Scratch (Python) (2020)
- [4] Yates D., Islam M.Z., Gao J. (2019) SPAARC: A Fast Decision Tree Algorithm. In: Islam R. et al. (eds) Data Mining. AusDM 2018. Communications in Computer and Information Science, vol 996. Springer, Singapore