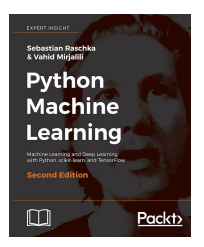
# Introduction to Data Mining (DATS 6103 - 10) Decision Tree and Random Forest

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#### Reference



Picture courtesy of the website of the book code repository and info resource

#### Reference

- This set of slices is an excerpt of the book by Raschka and Mirjalili, with some trivial changes by the creator of the slides
- Please find the reference to and website of the book below:
  - Raschka S. and Mirjalili V. (2017). Python Machine Learning. 2nd Edition.
  - https://sebastianraschka.com/books.html
- Please find the website of the book code repository and info resource below:
  - https://github.com/rasbt/
    python-machine-learning-book-2nd-edition

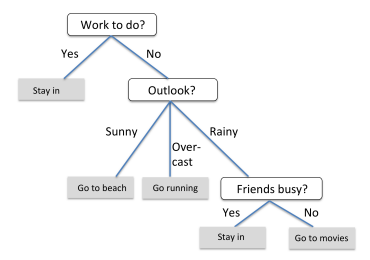
#### Overview

Decision tree

Random forest

#### Interpretability

- An attractive property of decision tree classifier is interpretability
- We can simply read our decision from the tree by asking a sequence of questions
- Figure 1 (see next page) shows an example of a decision tree



#### Attribute-based representations

Examples described by attribute values (Boolean, discrete, continuous, etc.) E.g., situations where I will/won't wait for a table:

Example	Attributes										Target
	Alt	Bar	Fri	Hun	Pat	Price	Rain	Res	Type	Est	WillWait
$X_1$	T	F	F	T	Some	\$\$\$	F	T	French	0–10	T
$X_2$	Τ	F	F	T	Full	\$	F	F	Thai	30–60	F
$X_3$	F	T	F	F	Some	\$	F	F	Burger	0–10	T
$X_4$	Τ	F	T	T	Full	\$	F	F	Thai	10-30	T
$X_5$	Τ	F	T	F	Full	\$\$\$	F	T	French	>60	F
$X_6$	F	T	F	T	Some	\$\$	T	T	Italian	0–10	T
$X_7$	F	T	F	F	None	\$	T	F	Burger	0–10	F
$X_8$	F	F	F	T	Some	\$\$	T	T	Thai	0–10	T
$X_9$	F	T	Т	F	Full	\$	T	F	Burger	>60	F
$X_{10}$	Τ	Т	Т	Τ	Full	\$\$\$	F	Т	Italian	10-30	F
$X_{11}$	F	F	F	F	None	\$	F	F	Thai	0-10	F
$X_{12}$	Τ	T	Τ	T	Full	\$	F	F	Burger	30–60	T

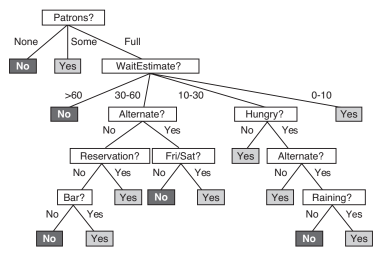
Classification of examples is positive (T) or negative (F)

#### Decision tree learning

Aim: find a small tree consistent with the training examples

Idea: (recursively) choose "most significant" attribute as root of (sub)tree

```
function DTL(examples, attributes, default) returns a decision tree if examples is empty then return default else if all examples have the same classification then return the classification else if attributes is empty then return Mode(examples) else best \leftarrow \texttt{CHOOSE-ATTRIBUTE}(attributes, examples) \\ tree \leftarrow \texttt{a} \text{ new decision tree with root test } best \\ \textbf{for each value } v_i \text{ of } best \text{ do} \\ examples_i \leftarrow \{\texttt{elements of } examples \text{ with } best = v_i\} \\ subtree \leftarrow \texttt{DTL}(examples_i, attributes - best, \texttt{Mode}(examples)) \\ \texttt{add a } \texttt{branch to } tree \text{ with label } v_i \text{ and subtree } subtree \\ \textbf{return } tree
```



## The objective function

Here, our objective function at each split is the information gain:

$$IG(D_p, f) = I(D_p) - \sum_{j=1}^{m} \frac{N_j}{N_p} I(D_j)$$
 where

- ullet f is the feature to perform the split
- ullet  $D_p$  and  $D_j$  are the dataset of the parent p and jth child node
- ullet I is our impurity measure
- $\bullet \ N_p$  and  $N_j$  are the number of samples at the parent and  $j{\rm th}$  child node
- The information gain is simply the difference between the impurity of the parent node and the sum of the child node impurities
  - the lower the child node impurities, the larger the information gain

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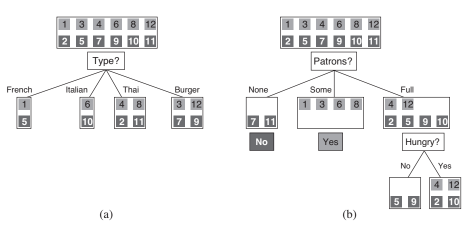
• Q: Shall we maximize or minimize it? Why?

## The objective function

• Here, our objective function at each split is the information gain:

$$IG(D_p, f) = I(D_p) - \sum_{j=1}^{m} \frac{N_j}{N_p} I(D_j)$$

- Q: Shall we maximize or minimize it? Why?
- A: Maximize it
  - generally the lower the impurity of the child nodes  $(I(D_j))$ , the better the classification
  - this is illustrated in Figure 5 (see next page)



#### The most informative feature

- So far, we have been using the concept of the most informative feature (a.k.a., the most significant attribute) without defining it
- Unsurprisingly, the most informative feature at each split is the one that maximizes the objective function

## Binary decision trees

- A special case of decision tree is binary decision tree
- Here, each parent node is split into two child nodes, left and right
- Since the objective function for decision tree is

$$IG(D_p, f) = I(D_p) - \sum_{j=1}^{m} \frac{N_j}{N_p} I(D_j)$$

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$$IG(D_p, f) = I(D_p) - \frac{N_{left}}{N_p} I(D_{left}) - \frac{N_{right}}{N_p} I(D_{right})$$

# Measuring the impurity

- The three commonly used impurity measures
  - Entropy  $(I_H)$
  - ullet Gini impurity  $(I_G)$
  - Classification error  $(I_E)$

#### Entropy

ullet The entropy of node (i.e., feature) t,  $I_H(t)$ , is defined as follows

$$I_H(t) = -\sum_{i=1}^{c} p(i|t) \log_2 p(i|t)$$

- $\bullet$  Here, p(i|t) is the proportion of the samples belonging to class i for a particular node t
- The entropy is 0 if all samples at a node belong to the same class
- The entropy is maximal if we have a uniform class distribution
- For example, in a binary class setting
  - the entropy is 0 if p(i=1|t)=1 or p(i=0|t)=0
  - the entropy is 1 if the classes are distributed uniformly with p(i=1|t)=0.5 and p(i=0|t)=0.5

## Gini impurity

• The gini impurity of node t,  $I_G(t)$ , is defined as follows

$$I_G(t) = \sum_{i=1}^{c} p(i|t)(1 - p(i|t)) = 1 - \sum_{i=1}^{c} p(i|t)^2$$

- The gini impurity is 0 if all samples at a node belong to the same class
- The gini impurity is maximal if we have a uniform class distribution
- For example, in a binary class setting:
  - the gini impurity is 0 if p(i=1|t)=1 or p(i=0|t)=0
  - the gini impurity is 0.5 if the classes are distributed uniformly with p(i=1|t)=0.5 and p(i=0|t)=0.5
- While gini is the default setting for parameter criterion in scikit learn DecisionTreeClassifier, it usually yields very similar results as entropy

#### Classification error

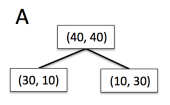
ullet The classification error of node t,  $I_E(t)$ , is defined as follows

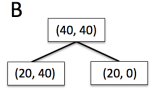
$$I_E(t) = 1 - max\{p(i|t)\}$$

 Compared to entropy and gini impurity, classification error is less sensitive to changes in the class probabilities of the nodes, thus not recommended

## Comparison between the three kinds of criterions

• Considering the following two splitting scenarios





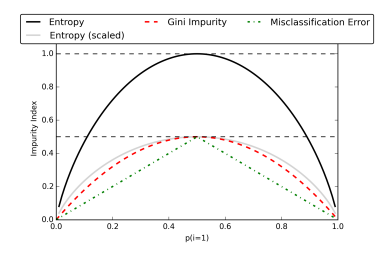
• Which scenario is better?

## Comparison between the three kinds of criterions

Considering the following two splitting scenarios



- Which scenario is better?
  - Entropy would favor scenario B ( $IG_H = 0.31$ ) over A ( $IG_H = 0.19$ )
  - Gini would also favor scenario B ( $IG_G = 0.16$ ) over A ( $IG_H = 0.125$ )
  - ullet Classification error has no preference ( $IG_E=0.25$  for both scenarios)
- A visual comparison between the three is shown in Figure 6 (see next page)



#### Problem of decision tree

- Decision tree tends to go too deep, resulting in overfitting
- Ways to limit the depth in scikit-learn DecisionTreeClassifier
  - max\_depth: the maximum depth of the tree
  - min\_samples\_split: the minimum number of samples required to split an internal node
  - min\_samples\_leaf: the minimum number of samples required to be at a leaf node

#### Random forest

- Random forests have gained huge popularity in application of machine learning
- A random forest can be considered as an ensemble of decision trees
- The idea is to combine weak learners to build a more robust model

## The random forest algorithm

- The random forest algorithm can be summarized in four simple steps
  - ① Draw a random bootstrap sample of size n (randomly choose n samples from the training set with replacement)
  - ② Grow a decision tree from the bootstrap sample. At each node:
    - 2.1 randomly select d features without replacement
    - 2.1 split the node using the feature that provides the best split according to the objective function (e.g., by maximizing the information gain)
  - $\odot$  Repeat steps 1 and 2 k times
  - Aggregate the prediction by each tree to assign the class label by majority vote

#### The hyperparameters

- In most implementations, the value of the hyperparameters, n (in step 1 of the algorithm), d (step 2.1), and k (step 3) are chosen as follows
  - n = the number of samples in the training set
  - $d = \sqrt{m}$  where m is the number of features in the training set
  - the larger k, the better the performance of the random forest classifier, at the expense of an increased computational cost

#### Feature importance

- A unique property of random forest is that it can provide feature importance, which has been widely used in feature selection
- One application is screening the single nucleotide polymorphisms (SNPs)
- Figure 7 (see next page) shows a bar chart of feature importance in Iris dataset

