# **Decision tree learning**

COMS 4771 Fall 2023

**Decision trees** 

#### Decision trees: nested if-then-else statements

- Can be relatively easy to understand (when not too large)
- Can have fast execution time (when not too large)
- Standard learning algorithm has some nice properties

#### Decision trees vs nearest neighbors

- ► Both: try to exploit "local regularity"
- Nearest neighbors: memorize training data
- Decision trees: use training data to carve X into regions
   ...so that, for each region, there is a good constant prediction

Example: iris dataset (using different features)



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### Structure of decision tree (in context of prediction):

- ► Rooted binary tree T
- ► A non-leaf node is associated with a predicate involving single feature
- $\blacktriangleright$  A leaf node is associated with a label from  ${\cal Y}$
- Computing f<sub>T</sub>(x) = prediction of tree T at x: Start at root node
  - If current node is leaf node: return associated label
  - Else if predicate at x is true: recurse on left child
  - Else: recurse on right child



## **Top-down learning algorithm**

Top-down learning algorithm: repeatedly modify tree to reduce its "cost"
Simplest cost function (for classification): training error rate

$$\widehat{\operatorname{err}}[f_T; \mathcal{S}] = \frac{1}{|\mathcal{S}|} \sum_{(x,y) \in \mathcal{S}} \mathbb{1}\{f_T(x) \neq y\}$$

(Classification tree = decision tree for classification problem)

- Initial tree: a single (leaf) node
- Repeat until done: make a modification to tree that reduces the cost the most

Example: iris dataset (using different features)



## Allowed modifications to improve the tree:

Replace a leaf node with a decision stump



How many possible modifications are there?

Example: iris dataset (using different features)



Two steps of top-down algorithm on iris dataset



When to stop modifying the tree? Some options:
Stop when no modification leads to reduction in cost

Stop when # leaves or depth reaches predetermined maximum

Stop when each leaf node is "pure" (i.e., all training examples that "reach" the leaf node have same label or same feature vector)

## Over-fitting training data



Training data	
feature vector	label
(0,0)	0
(0, 1)	1
(1, 0)	1
(1, 1)	0

Decision trees with 1 or 2 leaf nodes make 2 mistakes (Myopic learner does not get past first step)

But the following makes no mistakes:



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#### sklearn.tree.DecisionTreeClassifier

class sklearn.tree.DecisionTreeClassifier(\*, criterion='gini', splitter='best', max\_depth=None, min\_samples\_split=2, min\_samples\_leaf=1, min\_weight\_fraction\_leaf=0.0, max\_features=None, random\_state=None, max\_leaf\_nodes=None, min\_impurity\_decrease=0.0, class\_weight=None, ccp\_alpha=0.0) [source]

A decision tree classifier.

Read more in the User Guide.

Parameters:	<pre>criterion : {"gini", "entropy", "log_loss"}, default="gini" The function to measure the quality of a split. Supported criteria are "gini" for the Gini impurity and "log_loss" and "entropy" both for the Shannon information gain, see Mathematical formulation.</pre>
	splitter : {"best", "random"}, default="best"
	The strategy used to choose the split at each node. Supported strategies are "best" to choose the best split and "random" to choose the best random split.
	max_depth : <i>int, default=None</i>
	The maximum depth of the tree. If None, then nodes are expanded until all leaves are pure or until all leaves contain less than min_samples_split samples.
	min_samples_split : <i>int or float, default=2</i>
	The minimum number of samples required to split an internal node:
	• If int, then consider min_samples_split as the minimum number.
	• If float, then min_samples_split is a fraction and ceil(min_samples_split * n_samples) are the
	minimum number of samples for each split.
	Changed in version 0.18: Added float values for fractions.

min\_samples\_leaf : int or float, default=1

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# **Regression trees**

Regression trees: decision trees for real-valued prediction, (usually) with squared error as loss function

- ▶ Q: How to determine the labels associated with the leaf nodes?
- ► A: Average of labels among examples that "reach" the leaf node

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# Model averaging

Suppose you have many possible predictors  $f_1, f_2, \ldots, f_T$  (or many possible ways to learn a predictor)

- Model selection: try to choose the best one
- Model averaging: combine them into a single predictor by averaging/voting

Simplest form: uniform model averaging

$$f_{\text{avg}}(x) = \frac{1}{T} \sum_{t=1}^{T} f_t(x)$$

(For classification, use majority/plurality vote instead of averaging)

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$$\underbrace{\mathbb{E}\left[(f_{\text{avg}}(X) - Y)^2\right]}_{\text{mse}[f_{\text{avg}}]} = \frac{1}{T} \sum_{t=1}^T \underbrace{\mathbb{E}\left[(f_t(X) - Y)^2\right]}_{\text{mse}[f_t]} - \frac{1}{2T^2} \sum_{s=1}^T \sum_{t=1}^T \mathbb{E}\left[(f_s(X) - f_t(X))^2\right]_{\text{average disagreement}}$$

To generate many "similar" predcitors that may disagree often:

Train each predictor on a different (random) subset of the training data

Popular alternative: Bootstrap resampling of  $S = ((x^{(i)}, y^{(i)}))_{i=1}^n$ Independently sample T new datasets  $S^{(1)}, \ldots, S^{(T)}$ , where

$$\mathcal{S}^{(t)} = ((X^{(t,i)}, Y^{(t,i)}))_{i=1}^n \overset{\text{i.i.d.}}{\sim} \text{Unif}(\mathcal{S})$$

- Differs from "sampling without replacement"
- Some examples in S can appear more than once in  $S^{(t)}$
- Some may not appear at all

- Use bootstrap resampling to generate  $S^{(1)}, \ldots, S^{(T)}$
- ▶ For each t = 1,...,T: Let f<sub>t</sub> = output of learning algorithm on S<sup>(t)</sup>
- Combine f<sub>1</sub>,..., f<sub>T</sub> to form f<sub>avg</sub> using uniform model averaging (Or f<sub>vote</sub> using plurality vote, in case of classification problems)

## Forest cover type $dataset^1$

Problem: Create a program that, given cartographic data about a  $30 \times 30$  meter region of a forest, predict the type of forest cover

- Dataset: "[...] four wilderness areas located in the Roosevelt National Forest of northern Colorado [...] minimal human-caused disturbances [...] forest cover types are more a result of ecological processes rather than forest management practices."
- Classes: spruce/fir (1), lodgepole pine (2), ..., krummholz (7)
- Features (d = 54): elevation, slope, ..., distance to water, distance to roads, ..., amount of shade at 9am, amount of shade at 12pm, ...
- $\blacktriangleright$  Number of training data: 464809; number of test data: 116203

<sup>&</sup>lt;sup>1</sup>https://archive.ics.uci.edu/dataset/31/covertype

#### **Results on cover type**

- Decision tree with trained by top-down algorithm
  - Stopped when all leaf nodes are pure
  - Test error rate: 6.1%
- Bagging + top-down as before (T = 20)
  - $\blacktriangleright$  Individual trees' test error rates: between 7.7% and 8.0%
  - Plurality vote classifier test error rate: 3.5%