

Lecture 12: Tree-Based Algorithms

Applied Machine Learning

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Part 1: Decision Trees

We are now going to see a different way of defining machine models called *decision trees*.

Review: Components of A Supervised Machine Learning Problem

At a high level, a supervised machine learning problem has the following structure:

| Training Dataset + | - Learning Algorithm | \rightarrow Predictive Model |
|-----------------------|-------------------------------------|--------------------------------|
| | | |
| Attributes + Features | Model Class + Objective + Optimizer | |

The UCI Diabetes Dataset

To explain what is a decision tree, we are going to use the UCI diabetes dataset that we have been working with earlier.

Let's start by loading this dataset.

```
In [4]: import numpy as np
import pandas as pd
%matplotlib inline
import matplotlib.pyplot as plt
plt.rcParams['figure.figsize'] = [12, 4]
from sklearn import datasets
# Load the diabetes dataset
diabetes = datasets.load_diabetes(as_frame=True)
print(diabetes.DESCR)
```

.. diabetes dataset:

Diabetes dataset

Ten baseline variables, age, sex, body mass index, average blood pressure, and six blood serum measurements were obtained for each of n = 442 diabetes patients, as well as the response of interest, a guantitative measure of disease progression one year after baseline.

Data Set Characteristics:

:Number of Instances: 442

:Number of Attributes: First 10 columns are numeric predictive values

:Target: Column 11 is a quantitative measure of disease progression one ye ar after baseline

:Attribute Information:

| - | age | age in years |
|---|-----|---|
| - | sex | |
| _ | bmi | body mass index |
| - | bp | average blood pressure |
| - | s1 | tc, T-Cells (a type of white blood cells) |
| - | s2 | ldl, low-density lipoproteins |
| _ | e J | hdl high_density linonroteins |

| | 55 | <u>тат</u> , | magn-accord tabobroccano |
|---|------------|--------------|-----------------------------|
| _ | s4 | tch, | thyroid stimulating hormone |
| _ | s 5 | ltg, | lamotrigine |
| _ | s6 | glu, | blood sugar level |

Note: Each of these 10 feature variables have been mean centered and scaled by the standard deviation times `n_samples` (i.e. the sum of squares of each column totals 1).

Source URL: https://www4.stat.ncsu.edu/~boos/var.select/diabetes.html

For more information see:

Bradley Efron, Trevor Hastie, Iain Johnstone and Robert Tibshirani (2004) "L east Angle Regression," Annals of Statistics (with discussion), 407-499. (https://web.stanford.edu/~hastie/Papers/LARS/LeastAngle_2002.pdf)

We can also look at the data directly.

```
In [2]: # Load the diabetes dataset
diabetes_X, diabetes_y = diabetes.data, diabetes.target
# create a binary risk feature
diabetes_y_risk = diabetes_y.copy()
diabetes_y_risk[:] = 0
diabetes_y_risk[diabetes_y > 150] = 1
# Print part of the dataset
diabetes_X.head()
```

| Out[2]: | | age | sex | bmi | bp | s1 | s2 | s3 | s4 | s5 | s6 |
|---------|---|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| | 0 | 0.038076 | 0.050680 | 0.061696 | 0.021872 | -0.044223 | -0.034821 | -0.043401 | -0.002592 | 0.019908 | -0.017646 |
| | 1 | -0.001882 | -0.044642 | -0.051474 | -0.026328 | -0.008449 | -0.019163 | 0.074412 | -0.039493 | -0.068330 | -0.092204 |
| | 2 | 0.085299 | 0.050680 | 0.044451 | -0.005671 | -0.045599 | -0.034194 | -0.032356 | -0.002592 | 0.002864 | -0.025930 |
| | 3 | -0.089063 | -0.044642 | -0.011595 | -0.036656 | 0.012191 | 0.024991 | -0.036038 | 0.034309 | 0.022692 | -0.009362 |
| | 4 | 0.005383 | -0.044642 | -0.036385 | 0.021872 | 0.003935 | 0.015596 | 0.008142 | -0.002592 | -0.031991 | -0.046641 |

Decision Trees: Intuition

Decision tress are machine learning models that mimic how a human would approach this problem.

- 1. We start by picking a feature (e.g., age)
- 2. Then we *branch* on the feature based on its value (e.g, age > 65?)
- 3. We select and branch on one or more features (e.g., is it a man?)
- 4. Then we return an output that depends on all the features we've seen (e.g., a man over 65)

Decision Trees: Example

Let's first see an example on the diabetes dataset.

We will train a decision tree using it's implementation in sklearn.





Decision Rules

Let's now define a decision tree a bit more formally. The first important concept is that of a rule.

• A decision rule *r* : *X* → {true, false} is a partition of the feature space into two disjoint regions, e.g.:

$$r(x) = \begin{cases} \text{true} & \text{if } x_{\text{bmi}} \le 0.009\\ \text{false} & \text{if } x_{\text{bmi}} > 0.009 \end{cases}$$

- Normally, a rule applies to only one feature or attribute x_j of x.
- If x_j is continuous, the rule normally separates inputs x_j into disjoint intervals
 −∞, c], (c, ∞).

Decision Regions

The next important concept is that of a decision region.

- A decision region R ⊆ X is a subset of the feature space defined by the application of a set of rules r₁, r₂, ..., r_m and their values v₁, v₂, ..., v_m ∈ {true, false}, i.e.:
 R = {x ∈ X | r₁(x) = v₁ and ... and r_m(x) = v_m}
- For example, a decision region in the diabetes problem is:

 $R = \{x \in \mathcal{X} \mid x_{bmi} \le 0.009 \text{ and } x_{bp} > 0.004\}$

Decision Trees: Definition

A decision tree is a model $f : \mathcal{X} \to \mathcal{Y}$ of the form $f(x) = \sum_{R \in \mathcal{R}} y_R \mathbb{I}\{x \in R\}.$

- The $[\{\cdot\}$ is an indicator function (one if $\{\cdot\}$ is true, else zero) and values $y_R \in \mathcal{Y}$ are the outputs for that region.
- The set \mathcal{R} is a collection of decision regions. They are obtained by *recursive binary splitting*.
- The rules defining the regions \mathcal{R} can be organized into a tree, with one rule per internal node and regions being the leaves.

In [4]: plot_tree(clf, feature_names=diabetes_X.columns[:4], impurity=False) print('')



We can also illustrate decision trees via this figure from Hastie et al.





The illustrations are as follows:

- Top left: regions that cannot be represented by a tree
- Top right: regions that can be represented by a tree
- Bottom left: tree generating the top right regions
- Bottom right: function values assigned to the regions

Pros and Cons of Decision Trees

Decision trees are important models in machine learning

- They are highly interpretable.
- Require little data preparation (no rescaling, handle continuous and discrete features).

Their main disadvantages are that:

- If they stay small and interpretable, they are not as powerful.
- If they are large, they easily overfit and are hard to regularize.



Part 2: Learning Decision Trees

We saw how decision trees are represented. How do we now learn them from data?

Recall: Decision Trees

A decision tree is a model $f: \mathcal{X} \to \mathcal{Y}$ of the form

$$f(x) = \sum_{R \in \mathcal{R}} y_R \mathbb{I}\{x \in R\}.$$

- The $[\{\cdot\}$ is an indicator function (one if $\{\cdot\}$ is true, else zero) and values $y_R \in \mathcal{Y}$ are the outputs for that region.
- The set \mathcal{R} is a collection of decision regions. They are obtained by *recursive binary splitting*.
- The rules defining the regions \mathcal{R} can be organized into a tree, with one rule per internal node and regions being the leaves.

In [5]: plot_tree(clf, feature_names=diabetes_X.columns[:4], impurity=False) print('')



We can also illustrate decision trees via this figure from Hastie et al.





Learning Decision Trees

At a high level, decision trees are grown by adding nodes one at a time.

```
def build_tree(tree, data):
    whlie tree.is_complete() is False:
        region, region_data = tree.get_region()
        new_rule = split_region(region_data)
        tree.add_rule(region, new_rule)
```

Most often, we build the tree until it reaches a maximum number of nodes. The crux of the algorithm is in split_region.

Learning New Decision Rules

How does the split_region function choose new rule r? Given a dataset $\mathcal{D} = \{(x^{(i)}, y^{(i)} \mid i = 1, 2, ..., n\}$, we greedily choose the rule that achieves the dataset split with the lowest possible loss.

This can be written as the following optimization problem:



where L is a loss function over a subset of the data flagged by the rule and \mathcal{U} is the set of possible rules.

What is the set of possible rules? When x has continuous features, the rules have the following form:

$$r(x) = \begin{cases} \text{true} & \text{if } x_j \leq t \\ \text{false} & \text{if } x_j > t \end{cases}$$

for a feature index j and threshold $t \in \mathbb{R}$.

When x has categorial features, rules may have the following form:

$$r(x) = \begin{cases} \text{true} & \text{if } x_j = t_k \\ \text{false} & \text{if } x_j \neq t_k \end{cases}$$

for a feature index j and possible value t_k for x_j .

Objectives for Trees: Regression

What loss functions might we want to use? In regression, it is common to minimize the L2 error between the data and the single best prediction we can make on this data:

$$L(\mathcal{D}) = \sum_{(x,y)\in\mathcal{D}} (y - average-y(\mathcal{D}))^2.$$

If this was a leaf node, we would predict average-y(D), the average y in the data. The above loss measures the resulting squared error.



where $p_{\text{true}}(r) = \text{average-y}(\{(x, y) \mid (x, y) \in \mathcal{D} \text{ and } r(x) = \text{true}\})$ and $p_{\text{false}}(r) = \text{average-y}(\{(x, y) \mid (x, y) \in \mathcal{D} \text{ and } r(x) = \text{false}\})$ are the average predictions on each part of the data split.

Objectives for Trees: Classification

In classification, we may similarly use the misclassification rate

$$L(\mathcal{D}) = \sum_{(x,y)\in\mathcal{D}} \mathbb{I} \{ y = \text{most-common-y}(\mathcal{D}) \}.$$

If this was a leaf node, we would predict most-common-y(D), the most common class y in the data. The above loss measures the resulting misclassification error.

Other losses that can be used include the entropy or the gini index. These all optimize for a split in which different classes do not mix.

Other Practical Considerations

A few additional comments on the above training procedure;

- Nodes are added to until tree reaches a maximum depth or the leaves can't be split anymore.
- In practice trees are also often pruned in order to reduce overfitting.
- There exist alternative algorithms, including ID3, C4.5, C5.0. See Hastie et al. for details.

Algorithm: Classification and Regression Trees (CART)

- **Type**: Supervised learning (regression and classification).
- Model family: Decision trees.
- **Objective function**: Squared error, mislcassification error, Gini index, etc.
- **Optimizer**: Greedy addition of rules, followed by pruning.



Part 3: Bagging

Next, we are going to see a general technique to improve the performance of machine learning algorithms.

We will then apply it to decision trees to define an improved algorithm.

Review: Overfitting

Overfitting is one of the most common failure modes of machine learning.

- A very expressive model (a high degree polynomial) fits the training dataset perfectly.
- The model also makes wildly incorrect prediction outside this dataset, and doesn't generalize.

Recall this example, in which we take random samples around a true function.

```
In [5]: from sklearn.pipeline import Pipeline
from sklearn.preprocessing import PolynomialFeatures
from sklearn.linear_model import LinearRegression

def true_fn(X):
    return np.cos(1.5 * np.pi * X)

np.random.seed(2)
n_samples = 40
X = np.sort(np.random.rand(n_samples))
y = true_fn(X) + np.random.randn(n_samples) * 0.1
X_test = np.linspace(0, 1, 100)
plt.plot(X_test, true_fn(X_test), label="True function")
plt.scatter(X, y, edgecolor='b', s=20, label="Samples")
```

Out[5]: <matplotlib.collections.PathCollection at 0x12c905b00>



Fitting High-Degree Polynomials

Let's see what happens if we fit a high degree polynomial to random samples of 20 points from this dataset.

```
In [9]: n_plots, X_line = 3, np.linspace(0,1,20)
plt.figure(figsize=(14, 5))
for i in range(n_plots):
    ax = plt.subplot(1, n_plots, i + 1)
    random_idx = np.random.randint(0, 30, size=(30,))
```

```
X_random, y_random = X[random_idx], y[random_idx]
polynomial_features = PolynomialFeatures(degree=6, include_bias=False)
linear_regression = LinearRegression()
pipeline = Pipeline([("pf", polynomial_features), ("lr", linear_regression)]
)
pipeline.fit(X_random[:, np.newaxis], y_random)
ax.plot(X_line, true_fn(X_line), label="True function")
ax.plot(X_line, pipeline.predict(X_line[:, np.newaxis]), label="Model")
ax.scatter(X_random, y_random, edgecolor='b', s=20, label="Samples", alpha=0
.2)
ax.set_xlim((0, 1))
ax.set_ylim((-2, 2))
ax.legend(loc="best")
ax.set_title('Random sample %d' % i)
```



High-Variance Models

Each small subset of the data that we train on results is a very different model.

An algorithm that has a tendency to overfit is also called *high-variance*, because it outputs a predictive model that varies a lot if we slightly perturb the dataset.

Bagging: Bootstrap Aggregation

The idea of *bagging* is to reduce model variance by averaging many models trained on random subsets of the data.

```
for i in range(n_models):
    # collect data samples and fit models
    X_i, y_i = sample_with_replacement(X, y, n_samples)
    model = Model().fit(X_i, y_i)
    ensemble.append(model)

# output average prediction at test time:
y test = ensemble.average prediction(x test)
```

The data samples are taken with replacement and known as bootstrap samples.

Bagged Polynomial Regression

Let's apply bagging to our polynomial regression problem.

We are going to train a large number of polynomial regressions on random subsets of the dataset of points that we created earlier.

We start by training an ensemble of bagged models.

```
In [8]: n models, n subset = 10000, 30
        ensemble, Xs, ys = [], [], []
         for i in range(n models):
            # take a random subset of the data
            random idx = np.random.randint(0, 30, size=(n subset,))
            X random, y random = X[random idx], y[random idx]
            # train a polynomial regression model
            polynomial features = PolynomialFeatures(degree=6, include bias=False)
            linear regression = LinearRegression()
            pipeline = Pipeline([("pf", polynomial features), ("lr", linear regression)]
         )
            pipeline.fit(X random[:, np.newaxis], y random)
            # add it to our set of bagged models
            ensemble += [pipeline]
            Xs += [X random]
            ys += [y random]
```

Let's visualize the prediction of the bagged model on each random dataset sample and compare to predictions from an un-bagged models.





Extensions of Bagging

There exist a few closely related techniques to bagging.

- Pasting is when samples are taken without replacement.
- Random features are when we randomly sample the features.
- Random patching is when we do both of the above.

Summary: Bagging

Bagging is a general technique that can be used with high-variance ML algorithms.

It averages predictions from multiple models trained on random subset of the data.



Part 4: Random Forests

Next, let's see how bagging can be applied to decision trees. This will also provide us with a new algorithm.

Review: Bagging

The idea of *bagging* is to reduce model variance by averaging many models trained on random subsets of the data.

```
for i in range(n_models):
    # collect data samples and fit models
    X_i, y_i = sample_with_replacement(X, y, n_samples)
    model = Model().fit(X_i, y_i)
    ensemble.append(model)

# output average prediction at test time:
y test = ensemble.average prediction(y test)
```

The data samples are taken with replacement and known as bootstrap samples.

Review: Decision Trees

A decision tree is a model $f : \mathcal{X} \to \mathcal{Y}$ of the form $f(x) = \sum_{R \in \mathcal{R}} y_R \mathbb{I}\{x \in R\}.$

- The $[\{\cdot\}$ is an indicator function (one if $\{\cdot\}$ is true, else zero) and values $y_R \in \mathcal{Y}$ are the outputs for that region.
- The set \mathcal{R} is a collection of decision regions. They are obtained by *recursive binary splitting*.
- The rules defining the regions \mathcal{R} can be organized into a tree, with one rule per internal node and regions being the leaves.

We can also illustrate decision trees via this figure from Hastie et al.





Classification Dataset: Iris Flowers

Let's now look at the performance of decision trees on a new dataset, Iris flowers.

It's a classical dataset originally published by <u>R. A. Fisher</u> (<u>https://en.wikipedia.org/wiki/Ronald_Fisher</u>) in 1936. Nowadays, it's widely used for demonstrating machine learning algorithms.

```
In [10]: import numpy as np
import pandas as pd
from sklearn import datasets
# Load the Iris dataset
iris = datasets.load_iris(as_frame=True)
print(iris.DESCR)
```

```
.. iris dataset:
Iris plants dataset
**Data Set Characteristics:**
   :Number of Instances: 150 (50 in each of three classes)
   :Number of Attributes: 4 numeric, predictive attributes and the class
   :Attribute Information:
       - sepal length in cm
       - sepal width in cm
       - petal length in cm
       - petal width in cm
       - class:
              - Iris-Setosa
              - Iris-Versicolour
              - Iris-Virginica
   :Summary Statistics:
   _____ ____ ____ ____ ____ ____ ____
                  Min Max
                                   SD
                                        Class Correlation
                            Mean
   _____ ____ ____ ____ ____ ____ ____
   sepal length: 4.3 7.9 5.84
                                  0.83 0.7826
                  2.0 4.4 3.05 0.43 -0.4194
   sepal width:
                  1.0 6.9 3.76 1.76
                                       0.9490 (high!)
   petal length:
   petal width:
                  0.1 2.5 1.20 0.76
                                          0.9565
                                                (high!)
   ______ ____ ____ ____ ____ ____ ____
   :Missing Attribute Values: None
   :Class Distribution: 33.3% for each of 3 classes.
   :Creator: R.A. Fisher
   :Donor: Michael Marshall (MARSHALL%PLU@io.arc.nasa.gov)
   :Date: July, 1988
The famous Iris database, first used by Sir R.A. Fisher. The dataset is take
```

```
n
```

from Fisher's paper. Note that it's the same as in R, but not as in the UCI Machine Learning Repository, which has two wrong data points.

This is perhaps the best known database to be found in the pattern recognition literature. Fisher's paper is a classic in the field an d is referenced frequently to this day. (See Duda & Hart, for example.) The data set contains 3 classes of 50 instances each, where each class refers to а type of iris plant. One class is linearly separable from the other 2; the latter are NOT linearly separable from each other. .. topic:: References - Fisher, R.A. "The use of multiple measurements in taxonomic problems" Annual Eugenics, 7, Part II, 179-188 (1936); also in "Contributions to Mathematical Statistics" (John Wiley, NY, 1950). - Duda, R.O., & Hart, P.E. (1973) Pattern Classification and Scene Analys is. (Q327.D83) John Wiley & Sons. ISBN 0-471-22361-1. See page 218. - Dasarathy, B.V. (1980) "Nosing Around the Neighborhood: A New System Structure and Classification Rule for Recognition in Partially Exposed Environments". IEEE Transactions on Pattern Analysis and Machine Intelligence, Vol. PAMI-2, No. 1, 67-71. - Gates, G.W. (1972) "The Reduced Nearest Neighbor Rule". IEEE Transacti

ons

on Information Theory, May 1972, 431-433.

| Out[11]: | | sepal length (cm) | sepal width (cm) | petal length (cm) | petal width (cm) | target |
|----------|---|-------------------|------------------|-------------------|------------------|--------|
| | 0 | 5.1 | 3.5 | 1.4 | 0.2 | 0 |
| | 1 | 4.9 | 3.0 | 1.4 | 0.2 | 0 |
| | 2 | 4.7 | 3.2 | 1.3 | 0.2 | 0 |
| | 3 | 4.6 | 3.1 | 1.5 | 0.2 | 0 |
| | 4 | 5.0 | 3.6 | 1.4 | 0.2 | 0 |

```
In [12]: # Plot also the training points
p1 = plt.scatter(iris_X.iloc[:, 0], iris_X.iloc[:, 1], c=iris_y, s=50, cmap=plt.
cm.Paired)
plt.xlabel('Sepal Length')
plt.ylabel('Sepal Width')
plt.legend(handles=p1.legend_elements()[0], labels=['Setosa', 'Versicolour', 'Vi
rginica', 'Query'], loc='lower right')
```

Out[12]: <matplotlib.legend.Legend at 0x12d881978>



Decision Trees on the Flower Dataset

Let's now consider what happens when we train a decision tree on the Iris flower dataset.

The code below will be used to visualize predictions from decision trees on this dataset.

In []: # https://scikit-learn.org/stable/auto_examples/neighbors/plot_classification.ht
 ml
 from sklearn.tree import DecisionTreeClassifier
 from matplotlib.colors import ListedColormap

```
import warnings
warnings.filterwarnings("ignore")
def make grid(X):
    # Plot the decision boundary. For that, we will assign a color to each
    # point in the mesh [x min, x max]x[y min, y max].
    x \min, x \max = X.iloc[:, 0].min() - 0.1, X.iloc[:, 0].max() + 0.1
    y \min_{x} y \max_{x} = X.iloc[:, 1].min() - 0.1, X.iloc[:, 1].max() + 0.1
    xx, yy = np.meshgrid(np.arange(x min, x max, 0.02),
                         np.arange(y min, y max, 0.02))
    return xx, yy, x min, x max, y min, y max
def make 2d preds(clf, X):
    xx, yy, x min, x max, y min, y max = make grid(X)
    Z = clf.predict(np.c [xx.ravel(), yy.ravel()])
    Z = Z.reshape(xx.shape)
    return Z
def make 2d plot(ax, Z, X, y):
    # Create color maps
    cmap light = ListedColormap(['orange', 'cyan', 'cornflowerblue'])
    cmap bold = ListedColormap(['darkorange', 'c', 'darkblue'])
    xx, yy, x min, x max, y min, y max = make grid(X)
    # Put the result into a color plot
    ax.pcolormesh(xx, yy, Z, cmap=plt.cm.Paired)
    # Plot also the training points
    ax.scatter(X.iloc[:, 0], X.iloc[:, 1], c=y, cmap=plt.cm.Paired, edgecolor='k
', s=50)
    ax.set xlabel('Sepal Length')
   ax.set ylabel('Sepal Width')
    ax.set xlim(xx.min(), xx.max())
    ax.set ylim(yy.min(), yy.max())
```

We may now train and visualize a decision tree on this dataset.



Two Problems With Decision Trees

We see two problems with the output of the decision tree on the Iris dataset:

- The decision boundary between the two classes is very non-smooth and blocky.
- The decision tree overfits the data and the decision regions are highly fragmented.

High-Variance Decision Trees

When the trees have sufficiently high depth, they can quickly overfit the data.

Recall that this is called the *high variance* problem, because small perturbations of the data lead to large changes in model predictions.

Consider the perofmrance of a decision tree classifier on 3 random subsets of the data.





Random Forests

In order to reduce the variance of the basic decision tree, we apply bagging -- the variance reduction technique that we have seen earlier.

We refer to bagged decision trees as Random Forests.

Instantiating our definition of bagging with decision trees, we obtain the following pseudocode definition of random forests:

```
for i in range(n_models):
    # collect data samples and fit models
    X_i, y_i = sample_with_replacement(X, y, n_samples)
    model = DecisionTree().fit(X_i, y_i)
    random_forest.append(model)
```

```
# output average prediction at test time:
y_test = random_forest.average_prediction(y_test)
```

We may implement random forests in python as follows:

```
In [121]: np.random.seed(1000)
n_models, n_flowers, n_subset = 300, iris_X.shape[0], 10
random_forest = []
for i in range(n_models):
    # sample the data with replacement
    random_idx = np.random.randint(0, n_flowers, size=(n_subset,))
    X_random, y_random = iris_X.iloc[random_idx, :2], iris_y[random_idx]
    # train a decision tree model
    clf = DecisionTreeClassifier()
    clf.fit(X_random, y_random)
    # append it to our ensemble
    random forest += [clf]
```

Random Forests on the Flower Dataset

Consider now what happens when we deploy random forests on the same dataset as before.

Now, each prediction is the average on the set of bagged decision trees.

```
In [122]: # Visualize predictions from a random forest
ax = plt.gca()
# compute average predictions from all the models in the ensemble
X_all, y_all = iris_X.iloc[:,:2], iris_y
Z_list = []
for clf in random_forest:
    Z_clf = make_2d_preds(clf, X_all)
    Z_list += [Z_clf]
Z_avg = np.stack(Z_list, axis=2).mean(axis=2)
# visualize predictions
make_2d_plot(ax, np.rint(Z_avg), X_all, y_all)
```



The boundaries are much more smooth and well-behaved.

Algorithm: Random Forests

- **Type**: Supervised learning (regression and classification).
- Model family: Bagged decision trees.
- **Objective function**: Squared error, mislcassification error, Gini index, etc.
- **Optimizer**: Greedy addition of rules, followed by pruning.

Pros and Cons of Random Forests

Random forests remain a popular machine learning algorithm:

- They require little data preparation (no rescaling, handle continuous and discrete features, work well for classification and regression).
- They are often quite accurate.

Their main disadvantages are that:

- They are not interpretable.
- They do not work with unstructured data (images, audio).