Plan

In this lecture we review several Classical Machine Learning models.

These also allow us to introduce some concepts that are useful for many other models.

- Decision Trees
 - Simple Trees
 - Random Forests
 - introduce
 - Ensembles
 - Bootstrapping, Bagging

- Boosting
- Support Vector Machines
 - Margin Loss

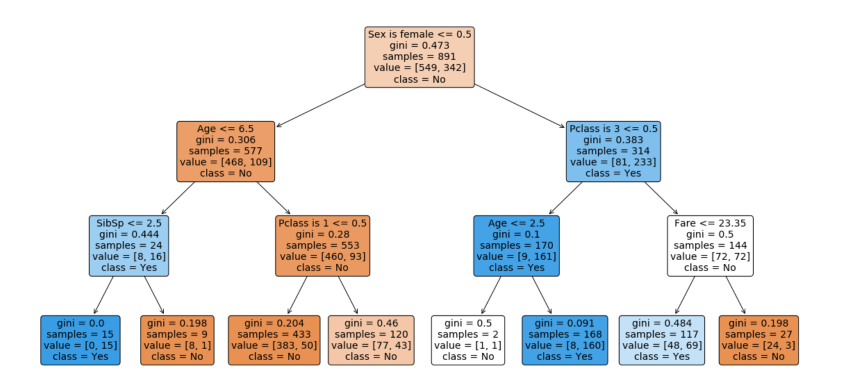
Decision Tree Terminology

In contrast to other models (e.g., Logistic Regression), which had some mathematical basis, Decision Trees will see very operational.

Let's dive in with an example: a Decision Tree to solve the Titanic Surival Classification task.

As usual we let ${\cal C}$ denote the set of distinct categories/classes (possible targets) for our Classification task.

```
In [4]: th = dthelp.TitanicHelper()
    ret = th.make_titanic_png(max_depth=3)
    if hasattr(ret, "fname"):
        Image(filename=ret["fname"] + ".png")
```



Nodes

- Each box is called a node
- There are two types of nodes
 - Those with no arrows exiting (called a *leaf* or *terminal* node)
 - Those with arrows exiting (called an *interior* or *non-terminal* node)
- The single node with no arrow entering is called the *root* node

Let U be the universe of possible examples, i.e, \mathbf{x}, \mathbf{y} pairs

A subset of the universe of examples associated

Edges

- An arrow, which is directed, is called an edge
 - The node from which an edge exits is called a *parent* node
 - The node to which an edge is directed is called a *child* node

Edges connect only a parent to a child.

The edges thus define an acyclic graph

Labels

A non-leaf node is labelled with a True/False question/test

- The test is evaluated on an example, from the universe of examples
- The left child of the node is associated with a True evaluation of the test
- The right child of the node is associated with a False evaluation of the test

A leaf node is labelled with one category/class in C

Let S be the set (universe) of all possible examples: \mathbf{x} , \mathbf{y} pairs.

- There is a subset of S associated with each node: $S_{
 m n}$
- ullet The test at node f n partitions $S_{f n}$ into disjoint subsets

$$egin{array}{lll} S_{
m n} &=& L_{
m n} \cup R_{
m n} \ \phi &=& L_{
m n} \cap R_{
m n} \ L_{
m n} &=& \{(\mathbf{x^{(i)}},\mathbf{y^{(i)}})|(\mathbf{x^{(i)}},y^{(i)}) \in S_{
m n}, ext{ "True" answer to question} \} \ R_{
m n} &=& \{(\mathbf{x^{(i)}},\mathbf{y^{(i)}})|(\mathbf{x^{(i)}},y^{(i)}) \in S_{
m n}, ext{ "False" answer to question} \} \end{array}$$

That is, the question partitions $S_{\rm n}$ into "left" and right subsets $L_{\rm n}, R_{\rm n}$ depending on the answer.

Example: Decision Tree for Titanic Survival

Let's illustrate using a Decision Tree for the Titanic Survival Classification task.

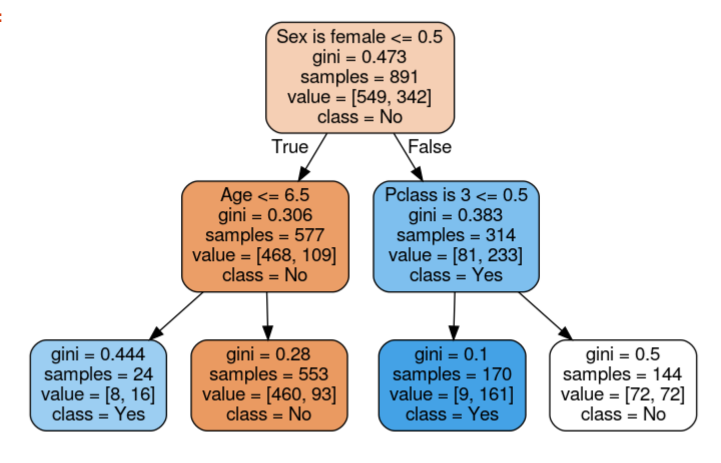
- High level
- We will subsequently explain the exact details for the labels and colors

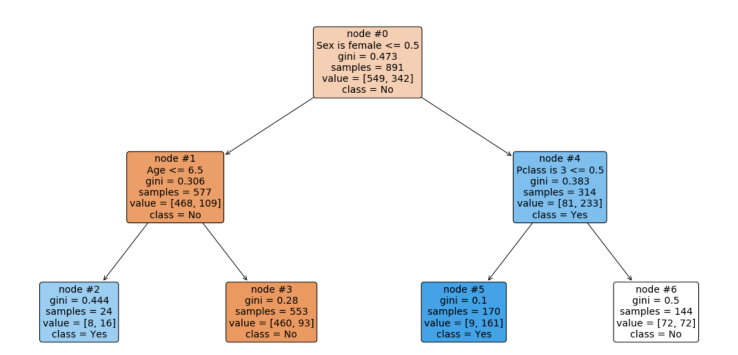
```
In [5]: th = dthelp.TitanicHelper()

ret = th.make_titanic_png(max_depth=2, node_ids=True)
Image(filename=ret["fname"] + ".png")

fig_titanic2 = ret["plt"]["fig"]
```

Out[5]:





- Root node #0
 - ullet $S_{\#0}=S$ is the entire universe of examples
 - Labelled with question: "Is **x** non Female ?"
- Left child #1
 - ullet $S_{\#1}=$ subset of $S_{\#0}$ that are `Male'
- Right child #4
 - ullet $S_{\#4}=$ subset of $S_{\#0}$ that are `Female'

- Left child #2
 - ullet $S_{\#2}$ subset of $S_{\#1}$ with Age ≤ 6.5
 - Is a leaf
 - Labeled with class Yes (i.e., Survive)
 - \circ Corresponding to the subset of entire universe S that are Males aged no more than 6.5 years
- Right child #3
 - $lacksquare S_{\#3}$ subset of $S_{\#1}$ with Age >6.5
 - Is a leaf
 - Labeled with class No (i.e., did not Survive)

Prediction

Given a test example (with features \mathbf{x})

- ullet Apply the sequence of questions to ${f x}$
 - Evaluate the test of the current node on x
 - Depending on the evaluation
 - Evaluate the test of the Left/Right child of the current node
- This defines a path to a leaf node
- Prediction \hat{y} is the class label of the leaf

This makes prediction in Decision Trees very fast.

Return to parent notebook

Training: a first look at the algorithm

Feature encoding

Before describing the algorithm we enumerate the features we will use

• Numeric: Age, SibSp, Parch, Fare

• Categorical: Sex, Pclass

The categorical features will be One Hot Encoded

- The Sex feature is replaced by two binary features: Is Female, Is Male
- The Pclass feature is replaced by three binary features Is Class 1, Is Class 2, Is Class 3

So don't expect to see a test like Sex == Male?

• Instead: Is Male == 1?

The training algorithm

We use the training examples as the universe of examples

$$S_{ ext{train}} = \langle \mathbf{X}, \mathbf{y}
angle = [\mathbf{x^{(i)}}, \mathbf{y^{(i)}} | 1 \leq i \leq m]$$

to build the tree recursively.

Let's start with the root node

- Let $n_{root}\mbox{ denote the root node}$
- Associate the entire set of training examples $S_{
 m train}$ with $S_{
 m root} = S_{
 m train}$

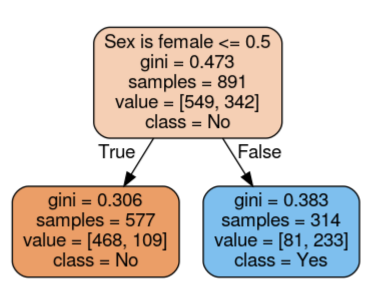
We construct a node n by the procedure Construct (n):

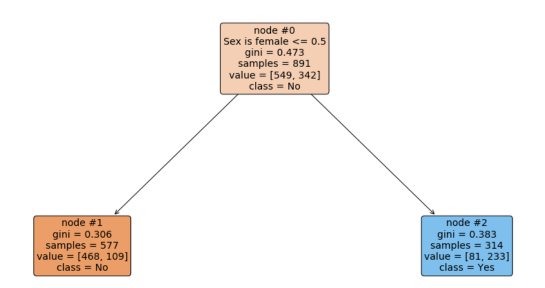
- Use S_{n} to label n with a test/question
- The test splits $S_{\rm n}$ into
 - $L_{
 m n}$: the subset of $S_{
 m n}$ consisting of examples where the test is True
 - $R_{\rm n}$: the subset of $S_{\rm n}$ consisting of examples where the test is False
- if $L_{\rm n}$ is not empty:
 - ullet create a new node \mathbf{n}_L as the left child or \mathbf{n}
 - Construct(n_L)
- ullet if $R_{
 m n}$ is not empty:
 - create a new node n_R as the right child of n
 - Construct (n_R)

To illustrate, let's apply Construct($n_{\rm root}\,$) :

```
In [6]: th = dthelp.TitanicHelper()
    ret = th.make_titanic_png(max_depth=1, node_ids=True)
    Image(filename=ret["fname"] + ".png")
```

Out[6]:





- Root node: #0
 - $S_{\#0}$ is the entire training set
 - samples = 891: This is number of examples in the training set
 - $\circ~$ The examples of $S_{\#0}$ split into 549 No, 342 Yes: `values = [549, 342]'
 - Labelled with question: "Is **x** non Female?"

- Left child of root: #1
 - samples = 577: This is the subset of $S_{\#0}$ (training set) consisting of the 577 Male examples
 - The examples of $S_{\#1}$ split into 468 No, 109 Yes: `values = [468, 109]'
- Right child of root #2:
 - ullet samples = 314: This is the subset of $S_{\#0}$ (training set) consisting of the 577 Female examples
 - The examples of $S_{\#2}$ split into 81 No, 233 Yes: `values = [468, 109]'

At this point, the left and right children are both leaf nodes

- Left child of root: #1
 - Labelled with prediction "No": class = No
- Right child of root: #2
 - Labelled with prediction "Yes": class = Yes

"No" is colored orange "Yes" is colored blue

Digression

The question

"Is
$$x$$
 non Female?" -- encoded via the test: Sex is female ≤ 0.5

is a bit contorted.

This is an artifact of the feature Sex being replaced by binary features "Is Female", "Is Male".

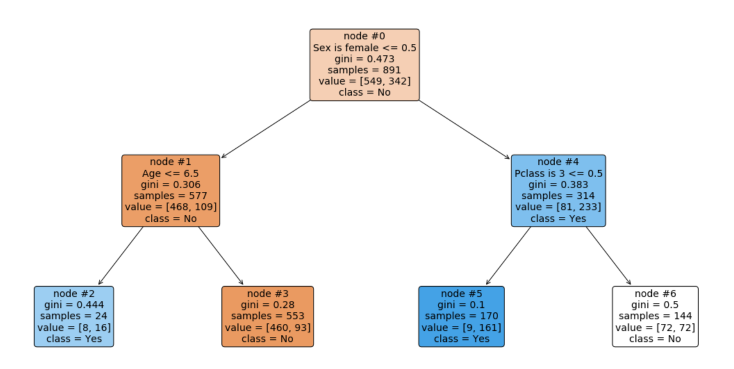
represents Is Female == 0, hence my translation to "non Female".

If the categorical variable had more than 2 classes, this test would appear less contorted.

We can now recursively apply <code>Construct(#1)</code> and <code>Consruct(#2)</code>

• Note that the numbering of the nodes changes

```
In [7]: th = dthelp.TitanicHelper()
    ret = th.make_titanic_png(max_depth=2, node_ids=True)
```



With the tree now depth 2, we have 4 leaf nodes.

If we were to continue this procedure indefinitely

- We would eventually have "pure" leaf nodes
 - All examples in the node are in the same class
 - Further splitting would not change the predicted class
 - The algorithm would stop

Return to parent notebook

Training: a deeper look at the algorithm

Encoding the test

The test evaluated at node n is a comparison

- Of a feature \mathbf{x}_j
- With a threshold value $t_{\mathrm{n},j}$

Thus we can represent the test at n as the pairs $(\mathbf{x}_j, t_{\mathrm{n},j})$.

Deciding the j and $t_{n,j}$ at a node ${\mathbf n}$ will be at the heart of the algorithm.

The threshold

Consider feature \mathbf{x}_j .

We restrict the possible threshold values V_j for comparison with \mathbf{x}_j to

ullet The distinct values of ${f x}_j$ in the training set

$$V_j = \{\mathbf{x}_i^{(\mathbf{i})} | 1 \le i \le m\}$$

This is true for both numeric and categorical features \mathbf{x}_{j} .

Note

- A variant uses the *mid-point* between distinct values
- The labelling of questions in our diagram always uses the comparison

$$\mathbf{x}_j \leq \text{midpoint value}$$

• So categorical tests look like

Is Female
$$\leq 0.5$$

which is equivalent to

Is Female
$$== 0$$

Choosing the test

There are

- A finite number (*n*) of features
- A finite number $\vert \vert V_j \vert \vert$ of distinct values for the threshold

How do we choose the test $(\mathbf{x}_j, t_{\mathrm{n},j})$ with which to label a non-leaf node ?

<u>sklearn manual (https://scikit-learn.org/stable/modules/tree.html#mathematical-formulation)</u>

Let's re-write our initial algorithm to give us a little more flexibility:

Initialization:

- $\bullet \ \mbox{Let} \ n_{root}$ denote the root node
- Associate the entire set of training examples $S_{
 m train}$ with $S_{
 m root} = S_{
 m train}$

We construct a sub-tree rooted at node ${
m n}$ by the procedure $\ {
m split}$ (${
m n}, S_{
m n}$) :

- if we can split $S_{
 m n}$
 - ullet Determine the question that "best" splits $S_{
 m n}$ into $L_{
 m n}, R_{
 m n}$
 - lacksquare Create a child node n_L with corresponding examples L_n
 - ullet Create a child node n_R with corresponding examples R_n
 - $\mathsf{split}(\mathrm{n}_L, L_\mathrm{n})$
 - $\mathsf{split}(\mathrm{n}_R, R_\mathrm{n})$

The high level description was vague on

- \bullet how do we define the "best" split of the examples at node n ?
- what does "if we can split"a node mean?

We answer each in turn.

Measuring the quality of a split

In order to determine the "best" split, we need a metric of the quality of the split.

We start with measuring the "randomness" of a node n

- ullet The examples S_n may correspond to different classes, each with its own frequency.
- So S_n induces a probability distribution on the class labels.
- We need a metric that measures the randomness of this distribution at node n

If we had a metric of randomness, we can define our quality metric of a split as the difference between

- the randomness of S_n
- ullet the weighted (by size) randomness of nodes L_n, R_n

We call this metric the information gain achieved by the split.

The best split would be the one which maximizes the information gain.

Any ideas for a metric of randomness of a distribution?

Entropy!

This a a very good measure to use.

In the interest of showing alternatives, we continue with a different choice.

Gini.

Gini score

For node n:

ullet Let $p_{n,c}$ be the fraction of S_n examples with class $c\in C$

$$p_{n,c} = rac{count_{n,c}}{|S_n|}$$

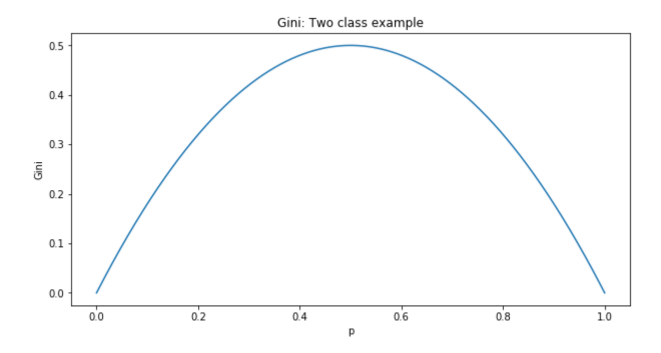
Then the Gini score (metric of randomness) of node ${\bf n}$ is defined as

$$G_n=1-\sum_{c\in C}{p_{n,c}}^2$$

 G_n is called the **impurity** of node ${f n}$

Note so we will try to minimize *impurity* (just as we would minimize entropy) Goal is to have pure nodes, i.e., all observations in node ${\bf n}$ are in same target class. Here's what Gini looks like as a function of $p_{n,c}$ for binary C

```
In [8]: gh = dthelp.GiniHelper()
    _ =gh.plot_Gini()
```



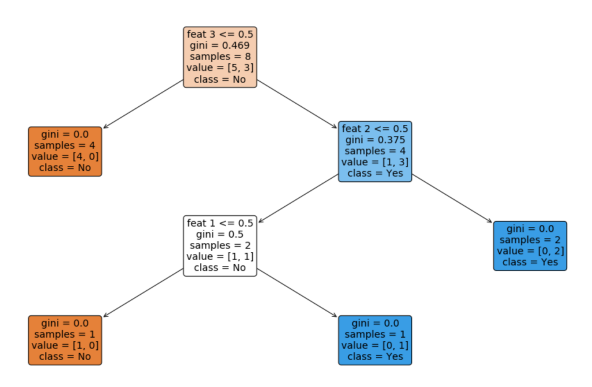
Just like Entropy, Gini impurity

- is minimized by a pure distribution (of either class)
- is maximized by an equally balanced distribution

Gini illustration

It will be easiest to illustrate with a toy example having only categorical features.

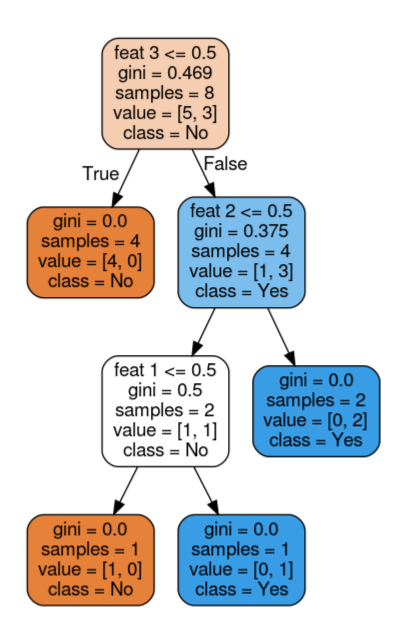
```
In [9]: gh = dthelp.GiniHelper()
    _ = gh.make_logicTree_png()
```





Out[10]:

	feat 1	feat 2	feat 3	target
0	0	0	0	0
1	0	0	1	0
2	0	1	0	0
3	0	1	1	1
4	1	0	0	0
5	1	0	1	1
6	1	1	0	0
7	1	1	1	1





```
In [11]: | gini_lt = gh.gini(df_lt, gh.target_name_lt, gh.feature_names_lt, noisy=True)
         print("\n\nMethod returns ", gini_lt)
         Gini, by hand:
         Count by target:
         0
              5
         Name: target, dtype: int64
         Frequency by target:
              0.625
              0.375
         Name: target, dtype: float64
         1 - sum(freq**2) = 0.469
         Method returns 0.46875
```



```
In [12]: | df_right = df_lt[ df_lt["feat 3"] > 0.5 ]
         gh.gini( df_right, gh.target_name_lt, gh.feature_names_lt, noisy=True)
         Gini, by hand:
         Count by target:
              3
         Name: target, dtype: int64
         Frequency by target:
              0.75
              0.25
         Name: target, dtype: float64
         1 - sum(freq**2) = 0.375
Out[12]: 0.375
```

The "best" (j, k) split: From Gini of children, to Cost for split at parent

Let node n

- ullet have child nodes n_L, n_R
- ullet S_n denote the set of examples corresponding to node ${f n}$
- ullet have Gini impurity score G_n

We can associate a Cost with the choice of splitting node ${f n}$ with question $(j,t_{{f n},j})$:

$$\mathrm{Cost}_{j,t_{\mathrm{n},j}}(S_{\mathrm{n}}) = rac{m_L}{(m_L+m_R)}G_L + rac{m_R}{(n_L+m_R)}G_R$$

$$egin{aligned} \bullet & ext{where } m_L \ &= | \ |L_{
m n}| \ |, m_R \ &= | \ |R_{
m n}|| \end{aligned}$$

That is, the Cost of splitting $S_{
m n}$ on $X_j \leq t_{n,j}$ is

• the weighted sum of the Gini's of the partitions created.

Finally:

• the best split $(j,t_{\mathrm{n},j})$ for node n is the one that minimizes the Cost $j,t_{\mathrm{n},j}=\operatorname*{argmin}_{j,t_{\mathrm{n},j}}\operatorname*{Cost}_{j,t_{\mathrm{n},j}}$

- $1 \leq j \leq n$
- $ullet t_{\mathrm{n},j} \in V_j$

This is the split that maximizes Information Gain (since G_n is constant, relative to the choices)



Out[13]: 0.1875

So split on (3,0) (feature "feat 3", threshold 0) gives the minimum cost.

That explains the split at the root.

What does "if we can split" a node mean

Time to answer our second vague statement: is there a time when we can't/shouldn't split node ${\bf n}$

When can't we split $S_{
m n}$?

- $|S_{\mathrm{n}}|=0$
 - an empty child, which we ignore (n.b., whose sibling is pure)
- $S_{
 m n}$ is pure

When shouldn't we split $S_{ m n}$?

One obvious case

• when the Information Gain of all possible splits is negative

There are some less obvious cases related to the Performance Measure of our out of sample test set.

If we don't restrict the answer to "if we can split"

- Then we will eventually have leaf nodes that are all pure.
- That's good, but it's also possible to have a single example corresponding to a leaf node
- Overfitting!
 - memorize training: each leaf memorizes an example

Prediction $\hat{\mathbf{y}}_n$ for node \mathbf{n}

We need to label a *leaf* node n with a category $c \in C$.

This will be the prediction $\hat{y}^{(i)}$ that will be made for test example $\mathbf{x}^{(i)}$.

We do this by choosing

- The class c
- ullet That occurs most frequently in $S_{
 m n}$, the set of training examples associated with node m n

$$egin{aligned} count_{ ext{n},c} &= ||\{i|(\mathbf{x^{(i)}},\mathbf{y^{(i)}}) \in S_{ ext{n}}, \mathbf{y^{(i)}} = c\}|| \ \hat{\mathbf{y}}_{ ext{n}} &= rgmax_{c \in C} \end{aligned}$$

Note

Our diagrams use the same logic for labelling non-leaf nodes with a class.

Prediction $\hat{\mathbf{y}}_n$ for node \mathbf{n}

Let's count the number of examples in S_n for each class $c \in C$

$$count_{n,c} = |\{i|(\mathbf{x^{(i)}},\mathbf{y^{(i)}}) \in S_n, \mathbf{y^{(i)}} = c\}|$$

Then our prediction for a node $\mathbf n$ (not just leaf nodes) is the class c with the greatest count.

$$\hat{\mathbf{y}}_n = rgmax count_{n,c} \ c \in C$$

Return to parent notebook

Decision Tree Regression

A Classifier, like the Decision Tree, that partitions examples can be modified to solve a Regression task.

We simply need to modify

- The category label assigned to a leaf node
- The measure used for the quality of a split

A category label is associated with each leaf node ${\bf n}$

ullet Classification: label is target category occurring with highest frequency in the examples in $S_{
m n}$

$$\hat{ extbf{y}}_n = rgmax_{c \in C} count_{n,c}$$

• Regression: label is average of the targets of the examples in S_n \$\$ \hat{y}\node{n} = \dfrac{1}{|| S\node{n} ||}\sum\limits{\scriptstyle (\x^\ip, \y^\ip) \in S\node{n}}y^{(i)}



The quality of the split of S_n into L_n (size m_L) and R_n (size m_R)

 Classification: minimize weighted impurity (or entropy) of the subsets created by split

$$\mathrm{Cost}_{j,t_{\mathrm{n},j}}(S_{\mathrm{n}}) \;\;=\;\; rac{m_L}{(m_L+m_R)}G_L + rac{m_R}{(n_L+m_R)}G_R$$

where

$$G_s = ext{impurity/entropy of set s}, s \in L, R$$

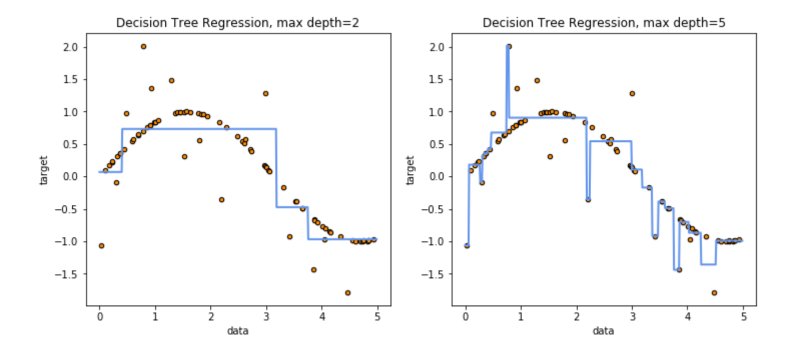
Regression: minimize the MSE of the subsets created by split \$\$ \begin{array}[III]\
\text{Cost}{j, t{\node{n},j}}(S_\node{n}) & = & \dfrac{m_L}{(m_L + m_R)}\text{MSE}_L + \dfrac{m_R}{(m_L + m_R)}\text{MSE}_R \ \text{where } \
\text{MSE}_s & = & \text{MSE of set s}, s \in {L,R}\

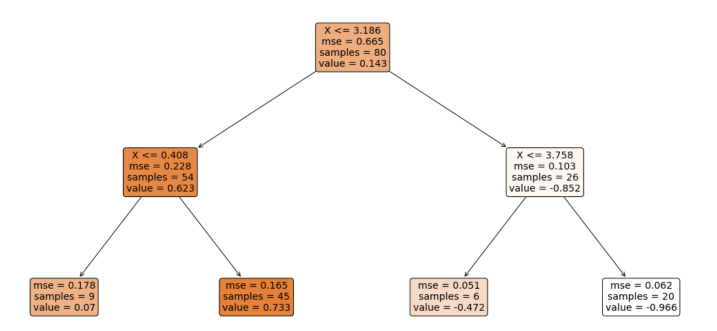
\end{array} \$\$

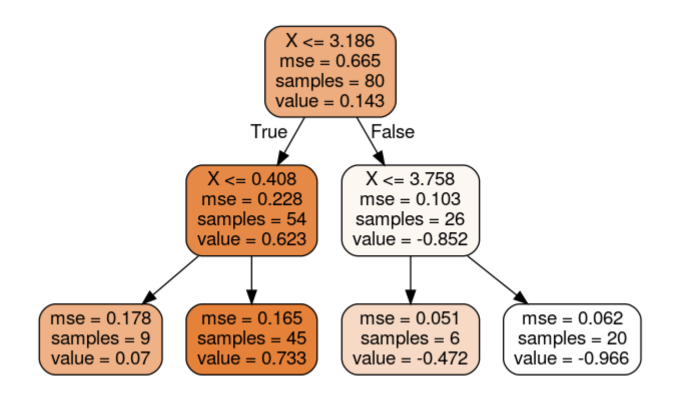
Return to parent notebook

Overfitting example

In [14]: rh = dthelp.RegressionHelper()
 _ =rh.make_plot()







Return to parent notebook

Hyper parameters for Decision Trees

Hyper parameters to control overfitting

You can combat overfitting with several parameters

- max depth: maximum depth of tree
- min_samples: minimum size (no. of observations) to split a node
- min_samples_leaf: minimum number of samples for a leaf

Other hyper parameters

Recall that our comparator $t_{n,j}$ for feature \mathbf{x}_j was drawn from V_j

ullet set of distinct values of ${f x}_j$ in the training set

This can be quite large. Perhaps defining a smaller number of threshholds may work even better.

• The choice of threshholds is a hyper paramter

Return to parent notebook

```
In [15]: print("Done")
```

Done