Tree Learning

16s1: COMP9417 Machine Learning and Data Mining

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March 28, 2017

Acknowledgements

Material derived from slides for the book "Machine Learning" by T. Mitchell McGraw-Hill (1997) http://www-2.cs.cmu.edu/~tom/mlbook.html

Material derived from slides by Andrew W. Moore http:www.cs.cmu.edu/~awm/tutorials

Material derived from slides by Eibe Frank http://www.cs.waikato.ac.nz/ml/weka

Material derived from slides for the book "Machine Learning" by P. Flach Cambridge University Press (2012) http://cs.bris.ac.uk/~flach/mlbook

Aims

This lecture will enable you to describe decision tree learning, the use of entropy and the problem of overfitting. Following it you should be able to:

define the decision tree representation

list representation properties of data and models for which decision trees are appropriate

reproduce the basic top-down algorithm for decision tree induction (TDIDT)

define entropy in the context of learning a Boolean classifier from examples

describe the inductive bias of the basic TDIDT algorithm define overfitting of a training set by a hypothesis

describe developments of the basic TDIDT algorithm: pruning, rule generation, numerical attributes, many-valued attributes, costs, missing values

describe regression and model trees

Brief History of Decision Tree Learning Algorithms

late 1950's – Bruner et al. in psychology work on modelling *concept acquisition*

early 1960s – Hunt et al. in computer science work on *Concept Learning Systems (CLS)*

late 1970s – Quinlan's *Iterative Dichotomizer 3 (ID3)* based on CLS is efficient at learning on then-large data sets

early 1990s – ID3 adds features, develops into C4.5, becomes the "default" machine learning algorithm

late 1990s - C5.0, commercial version of C4.5 (available from SPSS and www.rulequest.com)

current - widely available and applied; influential techniques

Why use decision trees?

Decision trees are probably the single most popular data mining tool

Easy to understand

Easy to implement

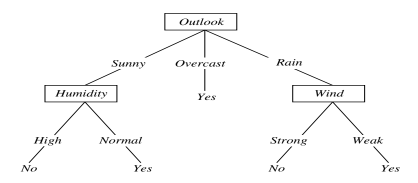
Easy to use

Computationally cheap (efficient, even on big data)

There are some drawbacks, though — e.g., high variance

They do *classification*, i.e., predict a categorical output from categorical and/or real inputs, or *regression*

Decision Tree for PlayTennis



A Tree to Predict C-Section Risk

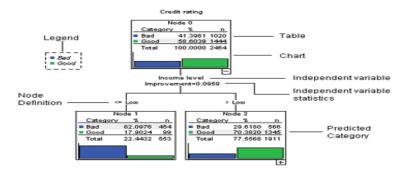
Learned from medical records of 1000 women

Negative examples are C-sections

```
[833+,167-] .83+ .17-

Fetal_Presentation = 1: [822+,116-] .88+ .12-
| Previous_Gsection = 0: [767+,81-] .90+ .10-
| | Primiparous = 0: [399+,13-] .97+ .03-
| | Primiparous = 1: [368+,68-] .84+ .16-
| | | Fetal_Distress = 0: [334+,47-] .88+ .12-
| | | | Birth_Weight < 3349: [201+,10.6-] .95+ .05-
| | | | Birth_Weight >= 3349: [133+,36.4-] .78+ .22-
| | | Fetal_Distress = 1: [34+,21-] .62+ .38-
| Previous_Gsection = 1: [55+,35-] .61+ .39-
Fetal_Presentation = 2: [3+,29-] .11+ .89-
Fetal_Presentation = 3: [8+,22-] .27+ .73-
```

Decision Tree for Credit Rating



Decision Trees

Decision tree representation:

Each internal node tests an attribute

Each branch corresponds to attribute value

Each leaf node assigns a classification

How would we represent the following expressions?

$$\land, \lor, \mathsf{XOR}$$

 $(A \land B) \lor (C \land \neg D \land E)$

M of N

Decision Trees

$X \wedge Y$

```
X = t:
| Y = t: true
| Y = f: no
X = f: no
```

$X \vee Y$

Decision Trees

2 of 3

```
| Y = t: true
| Y = f:
| | Z = t: true
| | Z = f: false
X = f:
| Y = t:
| | Z = t: true
| | Z = f: false
| Y = f: false
```

So in general decision trees represent a *disjunction of conjunctions* of constraints on the attributes values of instances.

When are Decision Trees the Right Model?

With Boolean values for the instances ${\bf X}$ and class Y, the representation adopted by decision-trees allows us to represent Y as a Boolean function of the ${\bf X}$

Given d input Boolean variables, there are 2^d possible input values for these variables. Any specific function assigns Y=1 to some subset of these, and Y=0 to the rest

Any Boolean function can be trivially represented by a tree. Each function assigns Y=1 to some subset of the 2^d possible values of \mathbf{X} . So, for each combination of values with Y=1, have a path from root to a leaf with Y=1. All other leaves have Y=0

When are Decision Trees the Right Model?

This is nothing but a re-representation of the truth-table, and will have 2^d leaves. More compact trees may be possible, by taking into account what is common between one or more rows with the same Y value

But, even for Boolean functions, there are some functions for which compact trees may not be possible (the parity and majority functions are examples)

In general, although possible in principle to express any Boolean function, our search and prior restrictions may not allow us to find the correct tree in practice.

BUT: If you want readable models that combine logical tests with a probability-based decision, then decision trees are a good start

When to Consider Decision Trees?

Instances described by a mix of numeric features and discrete attribute—value pairs

Target function is discrete valued (otherwise use regression trees)

Disjunctive hypothesis may be required

Possibly noisy training data

Interpretability is an advantage

Examples are extremely numerous, including:

Equipment or medical diagnosis

Credit risk analysis

Modeling calendar scheduling preferences

etc.

Top-Down Induction of Decision Trees (TDIDT)

Main loop:

 $A \leftarrow$ the "best" decision attribute for next node

Assign A as decision attribute for node

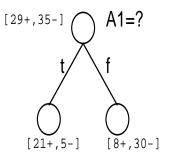
For each value of A, create new descendant of node

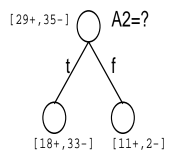
Sort training examples to leaf nodes

If training examples perfectly classified, Then STOP, Else iterate over new leaf nodes

Essentially this is the "ID3" algorithm (Quinlan, 1986) — the first efficient symbolic Machine Learning algorithm.

Which attribute is best?





Bits

You are watching a set of independent random samples of X You observe that X has four possible values

$$P(X = A) = \frac{1}{4} \mid P(X = B) = \frac{1}{4} \mid P(X = C) = \frac{1}{4} \mid P(X = D) = \frac{1}{4}$$

So you might see: BAACBADCDADDDA...

You transmit data over a binary serial link. You can <code>encode</code> each reading with two bits (e.g. A = 00, B = 01, C = 10, D= 11)

01000010010011101100111111100...

Someone tells you that the probabilities are not equal

$$P(X = A) = \frac{1}{2} \mid P(X = B) = \frac{1}{4} \mid P(X = C) = \frac{1}{8} \mid P(X = D) = \frac{1}{8}$$

It's possible ...

 \dots to invent a *coding* for your transmission that only uses 1.75 bits on average per symbol. How ?

Someone tells you that the probabilities are not equal

$$P(X = A) = \frac{1}{2} \mid P(X = B) = \frac{1}{4} \mid P(X = C) = \frac{1}{8} \mid P(X = D) = \frac{1}{8}$$

It's possible ...

... to invent a *coding* for your transmission that only uses 1.75 bits per symbol on average. How ?

Α	0		
В	10		
С	110		
D	111		

(This is just one of several ways)

Suppose there are three equally likely values

$$P(X = A) = \frac{1}{3} \mid P(X = B) = \frac{1}{3} \mid P(X = C) = \frac{1}{3}$$

Here's a naïve coding, costing 2 bits per symbol

Α	00
В	01
С	10

Can you think of a coding that would need only 1.6 bits per symbol on average?

Suppose there are three equally likely values

$$P(X = A) = \frac{1}{3} \mid P(X = B) = \frac{1}{3} \mid P(X = C) = \frac{1}{3}$$

Using the same approach as before, we can get a coding costing 1.6 bits per symbol on average ...

Α	0
В	10
С	11

This gives us, on average $\frac{1}{3}\times 1$ bit for A and $2\times\frac{1}{3}\times 2$ bits for B and C, which equals $\frac{5}{3}\approx 1.6$ bits.

Is this the best we can do?

Suppose there are three equally likely values

$$P(X = A) = \frac{1}{3} \mid P(X = B) = \frac{1}{3} \mid P(X = C) = \frac{1}{3}$$

From information theory, the optimal number of bits to encode a symbol with probability p is $-\log_2 p$...

So the best we can do for this case is $-\log_2\frac{1}{3}$ bits for each of A, B and C, or 1.5849625007211563 bits per symbol

General Case

Suppose X can have one of m values $\ldots V_1, V_2, \ldots V_m$

$$P(X = V_1) = p_1 \mid P(X = V_2) = p_2 \mid \dots \mid P(X = V_m) = p_m$$

What's the smallest possible number of bits, on average, per symbol, needed to transmit a stream of symbols drawn from X's distribution? It's

$$H(X) = -p_1 \log_2 p_1 - p_2 \log_2 p_2 - \dots - p_m \log_2 p_m$$

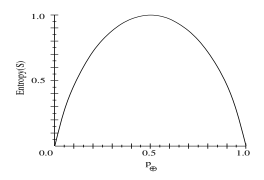
=
$$-\sum_{j=1}^m p_j \log_2 p_j$$

H(X) =the *entropy* of X

General Case

"High entropy" means X is very uniform and boring "Low entropy" means X is very varied and interesting

Entropy



Where:

 ${\cal S}$ is a sample of training examples

 p_{\oplus} is the proportion of positive examples in S

 p_{\ominus} is the proportion of negative examples in S

Entropy

Entropy measures the "impurity" of S

$$Entropy(S) \equiv -p_{\oplus} \log_2 p_{\oplus} - p_{\ominus} \log_2 p_{\ominus}$$

A "pure" sample is one in which all examples are of the same class.

Entropy

Entropy(S) =expected number of bits needed to encode class (\oplus or \ominus) of randomly drawn member of S (under the optimal, shortest-length code)

Why?

Information theory: optimal length code assigns $-\log_2 p$ bits to message having probability p.

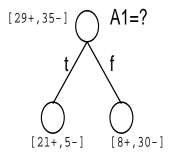
So, expected number of bits to encode \oplus or \ominus of random member of S:

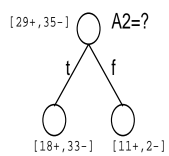
$$p_{\oplus}(-\log_2 p_{\oplus}) + p_{\ominus}(-\log_2 p_{\ominus})$$

$$Entropy(S) \equiv -p_{\oplus} \log_2 p_{\oplus} - p_{\ominus} \log_2 p_{\ominus}$$

Gain(S,A) = expected reduction in entropy due to sorting on A

$$Gain(S, A) \equiv Entropy(S) - \sum_{v \in Values(A)} \frac{|S_v|}{|S|} Entropy(S_v)$$





$$Gain(S, A1) = Entropy(S) - \left(\frac{|S_t|}{|S|}Entropy(S_t) + \frac{|S_f|}{|S|}Entropy(S_f)\right)$$

$$= 0.9936 - \left(\frac{26}{64}\left(-\frac{21}{26}\log_2\left(\frac{21}{26}\right) - \frac{5}{26}\log_2\left(\frac{5}{26}\right)\right)\right) + \left(\frac{38}{64}\left(-\frac{8}{38}\log_2\left(\frac{8}{38}\right) - \frac{30}{38}\log_2\left(\frac{30}{38}\right)\right)\right)$$

$$= 0.9936 - \left(0.2869 + 0.4408\right)$$

$$= 0.2658$$

$$Gain(S, A2) = 0.9936 - (0.7464 + 0.0828)$$

= 0.1643

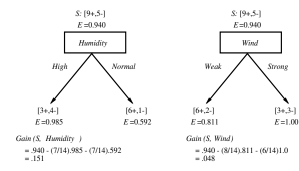
So we choose A1, since it gives a larger expected reduction in entropy.

Training Examples

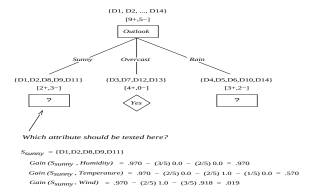
Day	Outlook	Temperature	Humidity	Wind	PlayTennis
D1	Sunny	Hot	High	Weak	No
D2	Sunny	Hot	High	Strong	No
D3	Overcast	Hot	High	Weak	Yes
D4	Rain	Mild	High	Weak	Yes
D5	Rain	Cool	Normal	Weak	Yes
D6	Rain	Cool	Normal	Strong	No
D7	Overcast	Cool	Normal	Strong	Yes
D8	Sunny	Mild	High	Weak	No
D9	Sunny	Cool	Normal	Weak	Yes
D10	Rain	Mild	Normal	Weak	Yes
D11	Sunny	Mild	Normal	Strong	Yes
D12	Overcast	Mild	High	Strong	Yes
D13	Overcast	Hot	Normal	Weak	Yes
D14	Rain	Mild	High	Strong	No

Information gain once more

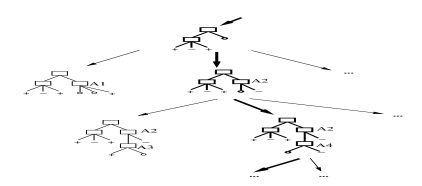
Which attribute is the best classifier?



Information gain once more



Hypothesis Space Search by ID3



Hypothesis Space Search by ID3

This can be viewed as a graph-search problem

Each vertex in the graph is a decision tree

Suppose we only consider the two-class case ($\omega = \omega_1$ or ω_2), and all the features x_i are Boolean, so each vertex is a binary tree

A pair of vertices in the graph have an edge if the corresponding trees differ in just the following way: one of the leaf-nodes in one vertex has been replaced by a non-leaf node testing a feature that has not appeared earlier (and 2 leaves)

This is the full space of all decision trees (is it?). We want to search for a single tree or a small number of trees in this space. How should we do this?

Hypothesis Space Search by ID3

Usual graph-search technique: greedy or beam search, starting with the vertex corresponding to the "empty tree" (single leaf node)

Greedy choice: which one to select? The neighbour that results in the greatest increase in $P(D \vert T)$

How?

Suppose T is changed to T'. Simply use the ratio of P(D|T')/P(D|T)

Most of the calculation will cancel out: so, we will only need to do the local computation at the leaf that was converted into a non-leaf node

RESULT: set of trees with (reasonably) high posterior probabilities given D: we can now use these to answer questions like $P(y'=\omega_1|\ldots)$? or even make a decision or a classification that $y'=\omega_1$, given input data ${\bf x}$

Hypothesis Space Search by ID3

Hypothesis space is complete! (contains all finite discrete-valued functions w.r.t attributes)

Target function surely in there...

Outputs a single hypothesis (which one?)

Can't play 20 questions...

No back tracking

Local minima...

Statistically-based search choices

Robust to noisy data...

Inductive bias: approx "prefer shortest tree"

Inductive Bias in ID3

Note H is the power set of instances X

 \rightarrow Unbiased?

Not really...

Preference for short trees, and for those with high information gain attributes near the root

Bias is a $\ensuremath{\textit{preference}}$ for some hypotheses, rather than a $\ensuremath{\textit{restriction}}$ of hypothesis space H

an incomplete search of a complete hypothesis space *versus* a complete search of an incomplete hypothesis space (as in learning conjunctive concepts)

Occam's razor: prefer the shortest hypothesis that fits the data

Occam's Razor

William of Ockham (c. 1287-1347)

Entities should not be multiplied beyond necessity

Why prefer short hypotheses?

Argument in favour:

- Fewer short hypotheses than long hypotheses
- ightarrow a short hyp that fits data unlikely to be coincidence
- ightarrow a long hyp that fits data might be coincidence

Occam's Razor

Argument opposed:

There are many ways to define small sets of hypotheses

e.g., all trees with a prime number of nodes that use attributes beginning with "Z"

What's so special about small sets based on size of hypothesis??

Look back to linear classification lecture to see how to make this work using Minimum Description Length (MDL)

Why does overfitting occur?

Greedy search can make mistakes. We know that it can end up in local minima — so a sub-optimal choice earlier might result in a better solution later (*i.e.* pick a test whose posterior gain (or information gain) is less than the best one

But there is also another kind of problem. We know that training error is an optimistic estimate of the true error of the model, and that this optimism increases as the training error decreases

We will see why this is the case later (lectures on Evaluation) Suppose we have two models h_1 and h_2 with training errors e_1 and e_2 and optimism o_1 and o_2 . Let the true error of each be $E_1=e_1+o_1$ and $E_2=e_2+o_2$

If $e_1 < e_2$ and $E_1 > E_2$, then we will say that h_1 has overfit then training data

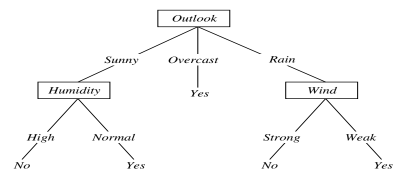
So, a search method based purely on training data estimates may end overfitting the training data

Overfitting in Decision Tree Learning

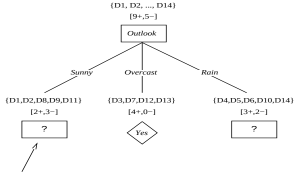
Consider adding noisy training example #15:

Sunny, Hot, Normal, Strong, PlayTennis = No

What effect on earlier tree?



Overfitting in Decision Tree Learning



Which attribute should be tested here?

$$S_{Sunny} = \{D1,D2,D8,D9,D11\}$$

 $Gain (S_{Sunny}, Humidity) = .970 - (3/5) 0.0 - (2/5) 0.0 = .970$
 $Gain (S_{Sunny}, Temperature) = .970 - (2/5) 0.0 - (2/5) 1.0 - (1/5) 0.0 = .570$
 $Gain (S_{Sunny}, Wind) = .970 - (2/5) 1.0 - (3/5) .918 = .019$

Overfitting in General

Consider error of hypothesis h over

training data: $error_{train}(h)$

entire distribution \mathcal{D} of data: $error_{\mathcal{D}}(h)$

Definition

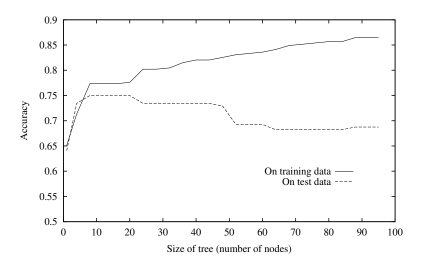
Hypothesis $h \in H$ overfits training data if there is an alternative hypothesis $h' \in H$ such that

$$error_{train}(h) < error_{train}(h')$$

and

$$error_{\mathcal{D}}(h) > error_{\mathcal{D}}(h')$$

Overfitting in Decision Tree Learning



How can we avoid overfitting? Pruning

pre-pruning stop growing when data split not statistically significant
post-pruning grow full tree, then remove sub-trees which are
overfitting

Post-pruning avoids problem of "early stopping"

How to select "best" tree:

Measure performance over training data?

Measure performance over separate validation data set ?

MDL: minimize size(tree) + size(misclassifications(tree)) ?

Pre-pruning

Usually based on statistical significance test

Stops growing the tree when there is no statistically significant association between any attribute and the class at a particular node

Most popular test: chi-squared test

ID3: chi-squared test plus information gain

Only statistically significant attributes were allowed to be selected by information gain procedure

Early stopping

Pre-pruning may suffer from early stopping: may stop the growth of tree prematurely

Classic example: XOR/Parity-problem

No individual attribute exhibits a significant association with the class

Target structure only visible in fully expanded tree

Prepruning won't expand the root node

But: XOR-type problems not common in practice

And: pre-pruning faster than post-pruning

Post-pruning

Builds full tree first and prunes it afterwards

Attribute interactions are visible in fully-grown tree

Problem: identification of subtrees and nodes that are due to chance effects

Two main pruning operations:

Subtree replacement

Subtree raising

Possible strategies: error estimation, significance testing, MDL principle

We examine two methods: Reduced-error Pruning and Error-based

Pruning

Reduced-Error Pruning

Split data into training and validation set

Do until further pruning is harmful:

Evaluate impact on validation set of pruning each possible node (plus those below it)

Greedily remove the one that most improves validation set accuracy

Reduced-Error Pruning

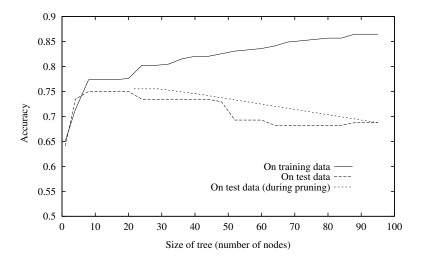


Good produces smallest version of most accurate subtree

Not so good

reduces effective size of training set

Effect of Reduced-Error Pruning



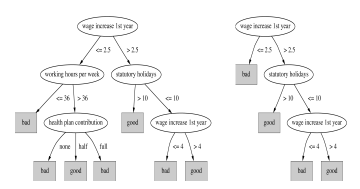
Error-based pruning (C4.5 / J48 / C5.0)

```
Quinlan (1993) describes the successor to ID3 – C4.5
    many extensions - see below
    post-pruning using training set
    includes sub-tree replacement and sub-tree raising
    also: pruning by converting tree to rules
    commercial version - C5.0 - is widely used
         RuleQuest.com
         now free
    Weka version – J48 – also widely used
```

Pruning operator 1: Sub-tree replacement

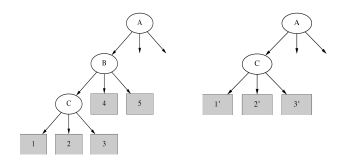
Bottom-up:

tree is considered for replacement once all its sub-trees have been considered



Pruning operator 2: Sub-tree raising

Deletes node and redistributes instances - more complicated, slow



Goal is to improve estimate of error on unseen data using all and only data from training set

But how can this work?

Make the estimate of error **pessimistic**!

Apply pruning operation if this does not increase the estimated error C4.5's method: using upper limit of standard confidence interval derived from the training data

Standard Bernoulli-process-based method

Note: statistically motivated, but not statistically valid

However: works well in practice!



The error estimate for a tree node is the weighted sum of error estimates for all its subtrees (possibly leaves).

Upper bound error estimate e for a node (simplified version):

$$e = f + Z_c \cdot \sqrt{\frac{f \cdot (1 - f)}{N}}$$

f is actual (empirical) error of tree on examples at the tree node

N is the number of examples at the tree node

 \mathcal{Z}_c is a constant whose value depends on *confidence* parameter c

C4.5's default value for confidence c=0.25

If c = 0.25 then $Z_c = 0.69$ (from standardized normal distribution)

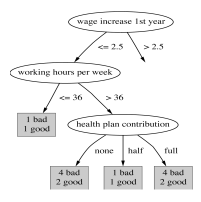
How does this method implement a pessimistic error estimate ?

What effect will the c parameter have on pruning ?

As
$$c \uparrow$$
, $z \downarrow$

See example on next slide (note: values not calculated using exactly the above formula)





health plan contribution: node measures $f=0.36,\,e=0.46$ sub-tree measures:

none: f = 0.33, e = 0.47

 $\mathsf{half:}\ f=0.5,\ e=0.72$

full: f = 0.33, e = 0.47

sub-trees combined 6:2:6 gives 0.51 sub-trees estimated to give $\it greater$ error so prune away

Rule Post-Pruning



This method was introduced in Quinlan's C4.5

Convert tree to equivalent set of rules

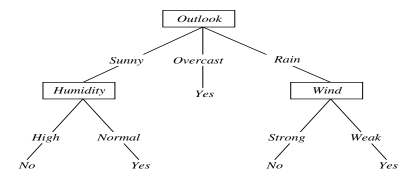
Prune each rule independently of others

Sort final rules into desired sequence for use

For: simpler classifiers, people prefer rules to trees

Against: does not scale well, slow for large trees & datasets

Converting A Tree to Rules



$$\begin{array}{ll} \mathsf{IF} & (Outlook = Sunny) \wedge (Humidity = High) \\ \mathsf{THEN} & PlayTennis = No \\ \mathsf{IF} & (Outlook = Sunny) \wedge (Humidity = Normal) \\ \mathsf{THEN} & PlayTennis = Var. \\ \end{array}$$

PlayTennis = YesTHEN

Rules from Trees (Rule Post-Pruning)

Rules can be simpler than trees but just as accurate, e.g., in C4.5Rules:

path from root to leaf in (unpruned) tree forms a *rule*i.e., tree forms a *set of rules*can simplify rules independently by deleting conditions
i.e., rules can be generalized while maintaining accuracy
greedy rule simplification algorithm

drop the condition giving lowest estimated error (as for pruning)
continue while estimated error does not increase

Rules from Trees

Select a "good" subset of rules within a class (C4.5Rules):

goal: remove rules not useful in terms of accuracy

find a subset of rules which minimises an MDL criterion

trade-off accuracy and complexity of rule-set

stochastic search using simulated annealing

Sets of rules can be ordered by class (C4.5Rules):

order classes by increasing chance of making *false positive* errors set as a default the class with the most training instances not covered by any rule

Continuous Valued Attributes



Decision trees originated for **discrete** attributes only. Now: continuous attributes.

Can create a discrete attribute to test continuous value:

Temperature = 82.5

 $(Temperature > 72.3) \in \{t, f\}$

Usual method: continuous attributes have a binary split

Note:

discrete attributes – one split exhausts all values continuous attributes – can have many splits in a tree

Continuous Valued Attributes

Splits evaluated on all possible split points

More computation: n-1 possible splits for n values of an attribute in training set

Fayyad (1991)

sort examples on continuous attribute find midway boundaries where class changes, e.g. for Temperature $\frac{(48+60)}{2}$ and $\frac{(80+90)}{2}$

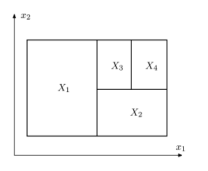
Choose best split point by info gain (or evaluation of choice)

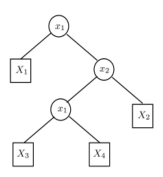
Note: C4.5 uses actual values in data

Temperature:	40	48	60	72	80	90
PlayTennis:	No	No	Yes	Yes	Yes	No

Continuous Valued Attributes

Dyadic decision trees — split each variable in turn at interval mid-point:





[&]quot;Algorithms for optimal dyadic decision trees". Hush, D & Porter, R. (2010)

Attributes with Many Values

Problem:

If attribute has many values, Gain will select it Why? more likely to split instances into "pure" subsets Maximised by singleton subsets Imagine using Date = March 28, 2017 as attribute High gain on training set, useless for prediction

Attributes with Many Values



One approach: use GainRatio instead

$$GainRatio(S, A) \equiv \frac{Gain(S, A)}{SplitInformation(S, A)}$$

$$SplitInformation(S, A) \equiv -\sum_{i=1}^{c} \frac{|S_i|}{|S|} \log_2 \frac{|S_i|}{|S|}$$

where S_i is subset of S for which A has value v_i

Attributes with Many Values

Why does this help?

sensitive to how broadly and uniformly attribute splits instances actually the entropy of S w.r.t. values of A

i.e., the information of the partition itself

i.e., the information of the partition itself

therefore higher for many-valued attributes, especially if mostly uniformly distributed across possible values

Attributes with Costs

Consider

medical diagnosis, BloodTest has cost \$150 robotics, $Width_from_1ft$ has cost 23 sec.

How to learn a consistent tree with low expected cost?

Attributes with Costs

One approach: replace gain by

Tan and Schlimmer (1990)

$$\frac{Gain^2(S,A)}{Cost(A)}.$$

Nunez (1988)

$$\frac{2^{Gain(S,A)} - 1}{(Cost(A) + 1)^w}$$

where $w \in [0,1]$ determines importance of cost

Attributes with Costs

Key idea: evaluate gain *relative to* cost, so prefer decision trees using lower-cost attributes.

More recently

Domingos (1999) – MetaCost, a meta-learning wrapper approach uses ensemble learning method to estimate probabilities decision-theoretic approach

General problem: class (misclassification) costs, instance costs, \dots SEE5 / C5.0 can use a misclassification cost matrix.

Can give false positives a different cost to false negatives

Unknown Attribute Values

What if some examples missing values of A?

Use training example anyway, sort through tree. Here are 3 possible approaches

If node n tests A, assign most common value of A among other examples sorted to node n

assign most common value of \boldsymbol{A} among other examples with same target value

assign probability p_i to each possible value v_i of A

assign fraction p_i of example to each descendant in tree

Note: need to classify new (unseen) examples in same fashion

Windowing

Early implementations – training sets too large for memory

As a solution ID3 implemented windowing:

- select subset of instances the window
- 2. construct decision tree from all instances in the window
- 3. use tree to classify training instances not in window
- 4. if all instances correctly classified then halt, else
- 5. add selected misclassified instances to the window
- 6. go to step 2

Windowing retained in C4.5 because it can lead to *more accurate* trees. Related to *ensemble learning*.

Non-linear Regression with Trees

Despite some nice properties of Neural Networks, such as generalization to deal sensibly with unseen input patterns and robustness to losing neurons (prediction performance can degrade gracefully), they still have some problems:

Back-propagation is often difficult to scale – large nets need lots of computing time; may have to be partitioned into separate modules that can be trained independently, e.g. NetTalk, DeepBind

Neural Networks are not very *transparent* – hard to understand the representation of what has been learned

Possible solution: exploit success of tree-structured approaches in ML

Regression trees

Differences to decision trees:

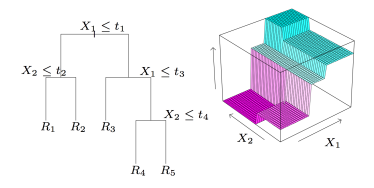
Splitting criterion: minimizing intra-subset variation Pruning criterion: based on numeric error measure Leaf node predicts average class values of training instances reaching that node

Can approximate piecewise constant functions

Easy to interpret

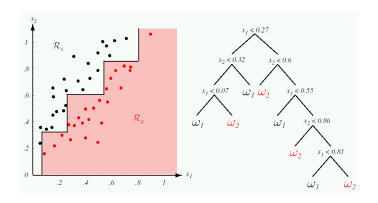
More sophisticated version: model trees

A Regression Tree and its Prediction Surface



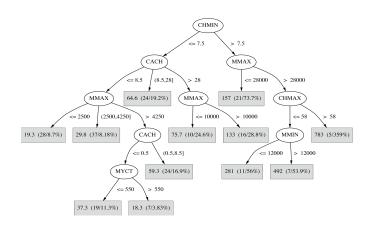
"Elements of Statistical Learning" Hastie, Tibshirani & Friedman (2001)

Axis-parallel Splitting



"Pattern Classification" Duda, Hart, and Stork, (2001)

Regression Tree on CPU dataset



Tree learning as variance reduction

The variance of a Boolean (i.e., Bernoulli) variable with success probability \dot{p} is $\dot{p}(1-\dot{p})$, which is half the Gini index. So we could interpret the goal of tree learning as minimising the class variance (or standard deviation, in case of $\sqrt{\mathrm{Gini}}$) in the leaves.

In regression problems we can define the variance in the usual way:

$$Var(Y) = \frac{1}{|Y|} \sum_{y \in Y} (y - \overline{y})^2$$

If a split partitions the set of target values Y into mutually exclusive sets $\{Y_1,\ldots,Y_l\}$, the weighted average variance is then

$$Var(\{Y_1, \dots, Y_l\}) = \sum_{j=1}^l \frac{|Y_j|}{|Y|} Var(Y_j) = \dots = \frac{1}{|Y|} \sum_{y \in Y} y^2 - \sum_{j=1}^l \frac{|Y_j|}{|Y|} \overline{y}_j^2$$

The first term is constant for a given set Y and so we want to maximise the weighted average of squared means in the children.

Learning a regression tree

Imagine you are a collector of vintage Hammond tonewheel organs. You have been monitoring an online auction site, from which you collected some data about interesting transactions:

#	Model	Condition	Leslie	Price
1.	B3	excellent	no	4513
2.	T202	fair	yes	625
3.	A100	good	no	1051
4.	T202	good	no	270
5.	M102	good	yes	870
6.	A100	excellent	no	1770
7.	T202	fair	no	99
8.	A100	good	yes	1900
9.	E112	fair	no	77

Learning a regression tree

From this data, you want to construct a regression tree that will help you determine a reasonable price for your next purchase.

There are three features, hence three possible splits:

```
\begin{aligned} \mathsf{Model} &= [\mathsf{A}100, \mathsf{B}3, \mathsf{E}112, \mathsf{M}102, \mathsf{T}202] \\ &= [1051, 1770, 1900][4513][77][870][99, 270, 625] \\ \mathsf{Condition} &= [\mathsf{excellent}, \mathsf{good}, \mathsf{fair}] \\ &= [1770, 4513][270, 870, 1051, 1900][77, 99, 625] \\ \mathsf{Leslie} &= [\mathsf{yes}, \mathsf{no}] \quad [625, 870, 1900][77, 99, 270, 1051, 1770, 4513] \end{aligned}
```

The means of the first split are 1574, 4513, 77, 870 and 331, and the weighted average of squared means is $3.21 \cdot 10^6$. The means of the second split are 3142, 1023 and 267, with weighted average of squared means $2.68 \cdot 10^6$; for the third split the means are 1132 and 1297, with weighted average of squared means $1.55 \cdot 10^6$. We therefore branch on Model at the top level. This gives us three single-instance leaves, as well as three A100s and three T202s.

Learning a regression tree

For the A100s we obtain the following splits:

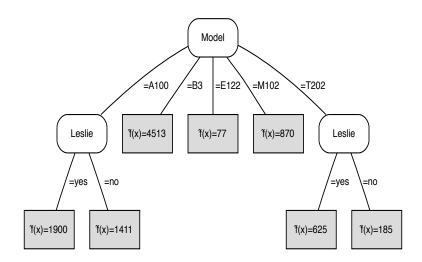
```
 \begin{array}{ll} {\sf Condition} = [{\sf excellent}, {\sf good}, {\sf fair}] & [1770][1051, 1900][] \\ {\sf Leslie} = [{\sf yes}, {\sf no}] & [1900][1051, 1770] \\ \end{array}
```

Without going through the calculations we can see that the second split results in less variance (to handle the empty child, it is customary to set its variance equal to that of the parent). For the T202s the splits are as follows:

```
Condition = [excellent, good, fair] [][270][99, 625]
Leslie = [yes, no] [625][99, 270]
```

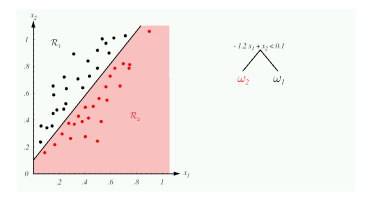
Again we see that splitting on Leslie gives tighter clusters of values. The learned regression tree is depicted on the next slide.

A regression tree



A regression tree learned from the Hammond organ dataset.

Splitting on Linear Combinations of Features



"Pattern Classification" Duda, Hart, and Stork, (2001)

Model trees

Like regression trees but with linear regression functions at each node Linear regression applied to instances that reach a node after full tree has been built

Only a subset of the attributes is used for LR

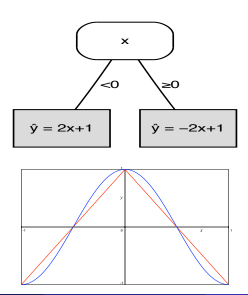
Attributes occurring in subtree (+maybe attributes occurring in path to the root)

Fast: overhead for Linear Regression (LR) not large because usually only a small subset of attributes is used in tree

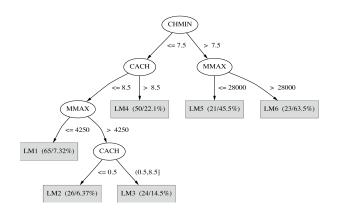
Two uses of features

Suppose we want to approximate $y=\cos\pi x$ on the interval $-1\leq x\leq 1$. A linear approximation is not much use here, since the best fit would be y=0. However, if we split the x-axis in two intervals $-1\leq x<0$ and $0\leq x\leq 1$, we could find reasonable linear approximations on each interval. We can achieve this by using x both as a splitting feature and as a regression variable (next slide).

A small model tree



Model Tree on CPU dataset



Smoothing

Naïve prediction method – output value of LR model at corresponding leaf node

Improve performance by *smoothing* predictions with *internal* LR models

Predicted value is weighted average of LR models along path from root to leaf

Smoothing formula: $p' = \frac{np+kq}{n+k}$ where

- p' prediction passed up to next higher node
- p prediction passed to this node from below
- q value predicted by model at this node
- n number of instances that reach node below
- k smoothing constant

Same effect can be achieved by incorporating the internal models into the leaf nodes

Building the tree

Splitting criterion: standard deviation reduction

$$SDR = sd(T) - \sum_{i} \frac{|T_i|}{|T|} \times sd(T_i)$$

where T_1, T_2, \ldots are the sets from splits of data at node.

Termination criteria (important when building trees for numeric prediction):

Standard deviation becomes smaller than certain fraction of sd for full training set (e.g. 5%)

Too few instances remain (e.g. less than four)

Pruning the tree

Pruning is based on estimated absolute error of LR models Heuristic estimate:

$$\frac{n+v}{n-v} \times \mathsf{average_absolute_error}$$

where n is number of training instances that reach the node, and v is the number of parameters in the linear model

LR models are pruned by greedily removing terms to minimize the estimated error

Model trees allow for heavy pruning: often a single LR model can replace a whole subtree

Pruning proceeds bottom up: error for LR model at internal node is compared to error for subtree

Discrete (nominal) attributes

Nominal attributes converted to binary attributes and treated as numeric

Nominal values sorted using average class value for each one For k-values, k-1 binary attributes are generated

the ith binary attribute is 0 if an instance's value is one of the first i in the ordering, 1 otherwise

Best binary split with original attribute provably equivalent to a split on one of the new attributes

Pseudo-code for M5prime

Four methods:

Main method: MakeModelTree()

Method for splitting: split()
Method for pruning: prune()

Method that computes error: subtreeError()

Note: linear regression method is assumed to perform attribute subset selection based on error

```
MakeModelTree()
MakeModelTree(instances)
   SD = sd(instances)
   for each k-valued nominal attribute
      convert into k-1 synthetic binary attributes
   root = newNode
   root.instances = instances
   split(root)
   prune(root)
   printTree(root)
```

```
split()
split(node)
   if sizeof(node.instances) < 4 or
         sd(node.instances) < 0.05*SD
      node.type = LEAF
   else
      node.type = INTERIOR
      for each attribute
         for all possible split positions of the attribute
            calculate the attribute's SDR
      node.attribute = attribute with maximum SDR
      split(node.left)
      split(node.right)
```

```
prune()
prune(node)
{
   if node = INTERIOR then
        prune(node.leftChild)
        prune(node.rightChild)
        node.model = linearRegression(node)
        if subtreeError(node) > error(node) then
            node.type = LEAF
}
```

Summary – decision trees

Decision tree learning is a practical method for many classifier learning tasks — a "Top 10" data mining algorithm

TDIDT family descended from ID3 searches complete hypothesis space - the hypothesis is there, somewhere...

Uses a search or *preference* bias, search for optimal tree is, in general, not tractable

Overfitting is inevitable with an expressive hypothesis space and noisy data, so pruning is important

Decades of research into extensions and refinements of the general approach, e.g., for numerical prediction, logical trees

Often the "try-first" machine learning method in applications, illustrates many general issues

Performance can be improved with use of "ensemble" methods

Summary – regression and model trees

Regression trees were introduced in CART

Quinlan proposed the M5 model tree inducer

M5': slightly improved version that is publicly available (M5Pin Weka is based on this)

Quinlan also investigated combining instance-based learning with M5

CUBIST: Quinlan's rule learner for numeric prediction

www.rulequest.com

Interesting comparison: Neural nets vs. model trees — both do non-linear regression

other methods also can learn non-linear models