

Supervised learning: Random Forest

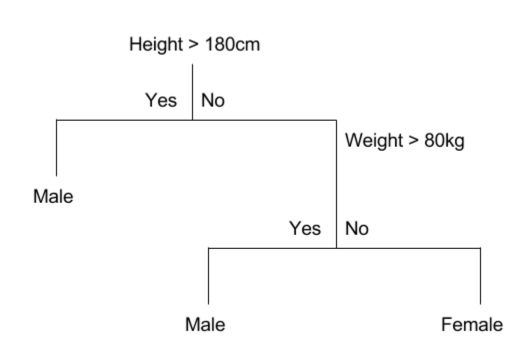
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From decision trees to bagging

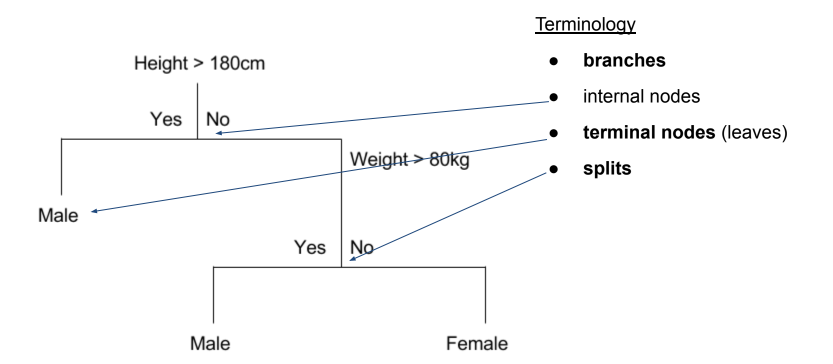




- tree-based methods are simple and useful for interpretation
- can be applied to classification and regression problems
- high variance, usually poor predictive performance
- combining a large number of trees can result in dramatic improvements in prediction accuracy
- bagging, random forest (these methods grow multiple trees which are then combined to yield a single consensus prediction)

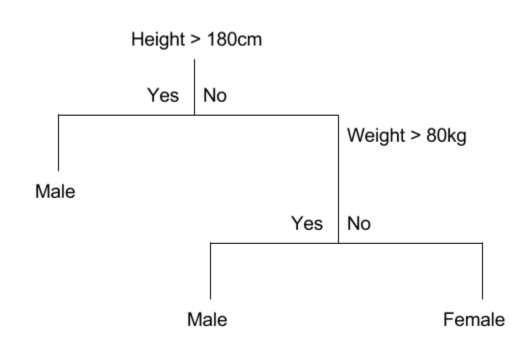
From: https://machinelearningmastery.com/classification-and-regression-trees-for-machine-learning/





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How do we decide on splits?

- min(RSS) for regression trees
- min(Gini index) or min(cross-entropy) for classification trees

$$G = \sum_{k=1}^K p_{mk} (1-p_{mk})$$

- K classes
- p_{mk}: proportion of obs from class k that go to branch m
- G is small if all of the p_{mk} are close to zero or one (node purity)

From: https://machinelearningmastery.com/classification-and-regression-trees-for-machine-learning/



- recursive binary splitting: top-down greedy algorithm for tree-building
 - **top-down**: it begins at the top \rightarrow first variable, first split
 - greedy: at each step the best split is made without considering next steps (splits) in the tree



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continue until stopping criterion is reached (e.g. n. of obs in terminal nodes, reduction in RSS / entropy < threshold etc.)

Decision trees - predictions



- once the tree is trained, **test observations** are run through it and **predicted** by using the **mean** (regression) or **majority vote** (classification) of **training observations** in **that region/node**

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- trees may grow to be very complex → overfitting!
- high threshold as stopping criterion risk of stopping too early ("short trees")
- tree pruning

From: https://www.canstockphoto.com/happy-family-in-garden-28321970.html

Decision trees - pruning



- grow a very large tree, T0
- then **prune it** to obtain a **subtree**
- many subtrees T ⊂ T0
- cost complexity pruning →

$$C = \sum_{m=1}^{|T|} \sum_{i: x_i \in R_m} \left(y_i - \hat{y}_{Rm}
ight)^2 + lpha |T|$$

- |T|: n. terminal nodes in T
- x_i: i_{th} predictor
- R_m: subset of predictor space for m_{th} terminal node
- y_hat_{Rm}: mean of training obs in R_m
- choose T ⊂ T0 so that C is smallest

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Tuning parameter that

controls subtree complexity (trade-off with fit to the data)

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Decision trees - recap



- 1. recursive binary splitting to grow a large tree (training data)
- cost-complexity for tree pruning
- 3. k-fold cross-validation to select α
 - a. repeat steps 1 and 2 on the k-1 training folds
 - b. evaluate prediction accuracy on the kth validation fold
 - c. for each value of α , average over k validation folds
 - d. select α that minimises the average error (max avg accuracy)
- 4. return T (from step 2) that corresponds to selected α

Pros and cons of decision trees



- easy to interpret and explain
- can be displayed graphically
- can easily handle qualitative predictors without the need to create dummy variables
- trees generally compare (very) poorly in terms of predictive accuracy (high variance)



A new resampling method



A resampling method

(make do with what we have)



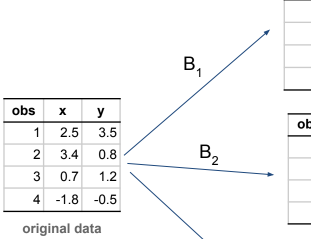
"The Baron had fallen to the bottom of a deep lake. Just when it looked like all was lost, he thought to pick himself up by his own bootstraps."

["The Surprising Adventures of Baron Munchausen" by Rudolph Erich Raspe]



- repeatedly sampling observations from the original data with replacement
- each "bootstrap set" comes from sampling with replacement and is
 the same size as the original data; some observations may appear
 more than once, while some not at all (out-of-bag observations)
- flexible and powerful technique to quantify the uncertainty associated with an estimator or machine learning method (e.g. estimate of the standard error of a coefficient, or a confidence interval around it)





 B_3

obs	X	у	
1	2.5	3.5	
2	3.4	0.8	
2	3.4	0.8	
4	-1.8	-0.5	

obs	х	у
4	-1.8	-0.5
2	3.4	0.8
1	2.5	3.5
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		-	

 Θ_{1}

 Θ_3

$$\Theta = rac{1}{B} \sum_{b=1}^{B} \Theta_i$$

3.4

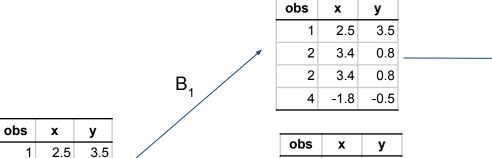
-1.8

original data

8.0 1.2

-0.5





 B_2

 B_3

obs	Х	у
4	-1.8	-0.5
2	3.4	0.8
1	2.5	3.5
4	-1.8	-0.5

(-)	 1	∇^B	Θ_i
O	 \overline{B}	$\angle b=1$	O_i

obs	X	у
1	2.5	3.5
3	0.7	1.2
1	2.5	3.5
4	-1.8	-0.5

You can also estimate the variability of Θ , and its distribution

 Θ_{1}

 Θ_2

 Θ_3



Bagging and Random Forest

Bagging: bootstrap aggregation



 bagging is a general technique to reduce the variance of machine-learning methods

$$Var(\overline{x}) = rac{\sigma^2}{n}$$

averaging reduces the variance (the larger n, the larger the reduction)

Bagging: bootstrap aggregation



 we normally do not have multiple training sets, but we can use bootstrapping to generate them

$$\hat{f}_{bag}(x) = rac{1}{B}\sum_{b=1}^{B}\hat{f}^{b}(x)$$

This is bagging, folks!

Out-of-bag (OOB) observations



- In bootstrapping, typically ¾ of the data are used in each bootstrap replicate → hence, ⅓ is left out → **out-of-bag observations**
- We can use OOB observations to estimate the prediction error (sort of built-in cross-validation)
- This is a valid estimate of the test error, since OOB observations are not used to train the decision tree

Random Forest



- bagging improves over decision trees, but the bootstrapped trees are correlated → overfitting
- Random Forest improves over bagging by decorrelating the bootstrapped trees
- this is done by selecting a random subset of predictors (variables) for each bootstrapped tree
- this way, trees in the forest differ in terms of both observations (sampling with replacement) and variables (random subset)
- typically, ~sqrt(p) predictors are used for classification, ~p/3
 predictors for regression → tuning parameter

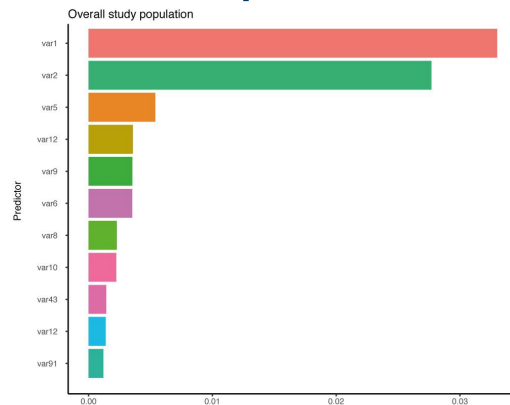
Variable importance



- bagging and RF improve the predictive ability at the expense of model interpretation
- however, one can use the large number of trees to extract variable importance
- this is based on the amount that the RSS (or Gini index) is decreased due to splits over a given predictor, averaged over all B trees. A large value indicates an important predictor

Variable importance





Importance

 ${\color{red} \textbf{source:}} \underline{\textbf{https://stats.stackexchange.com/questions/329995/scale-of-variable-importance-in-randomforest-party-gbm}$

Variable importance



How do we calculate variable importance?

Simple answer:

In the case of RF regression, we can record the total amount that the RSS (residual sum of squares) is decreased due to splits over a given predictor, averaged over all B trees. A large value indicates an important predictor.

Detailed answer:

Once the RF has been trained (and trees with their nodes and splits have been constructed -using OOB observations or testing sets), we can measure the reduction of RSS: each variable is used at some nodes in each tree, and the RSS reduction over such nodes can be summed up and then averaged over the number of trees in the forest.

Alternative method:

Again, after trees have been constructed, OOB observations are passed through each tree twice: i) as they are; ii) after permuting randomly the value for each variable. In both cases the accuracy of prediction is measured. The reduction in the accuracy is then used as a measure of the variable importance (the larger the reduction, the more important the variable)

Variable importance - RSS reduction



- record reduction in RSS or node impurity over all splits involving any given variable
- default variable importance metric in scikitlearn
- average over thousands of trees in the "forest"
- usually rescaled [0,100]
- variables with large average RSS reduction are more important

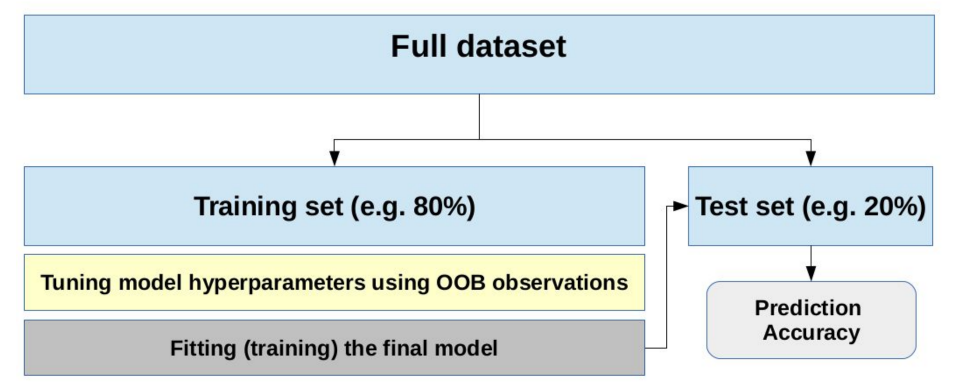
Variable importance - permutations



- for each tree grown in a random forest, get number of correct predictions in OOB data
- random permutation (shuffling) of a predictor's values in the OOB data, then check again the number of correct predictions
- 3. subtract the number of correct predictions in the permuted data from the number of the correct predictions in the original OOB data
- the average of this number over all trees in the forest is the variable importance (can be normalized)
- 5. variables with large differences between permuted and non-permuted data are ranked as more important (a model without that variable gives worse predictions → the variable is important)

Model tuning





Random Forest



- demonstration 8.1
- Exercise 8.1

- → 8.random_forest.Rmd
- → 9.multiclass_random_forest.Rmd