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Modelling

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Classification Regression Trees (CART)

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bootstrapping

Building a randon

Building a rando forest

Variable

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Predictor selection

Cubic

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Spatial randon

Data-driven vs.

Data-driven methods for predictive modelling

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February 23, 2022

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Modelling cultures

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Data-driven v

Statistical modelling

- · Statistics starts with data: something we have measured
- Data is **generated** by some (unknown) **mechanism**: input (stimulus) x, output (response) y
- Before analysis this is a black box to us, we only have the data itself
- · Two goals of analysis:
 - **1 Prediction** of future responses, given known inputs
 - Explanation, Understanding of what is in the "black box" (i.e., make it "white" or at least "some shade of grey").

Modelling cultures

Data modelling (also called "model-based")

- · assume an empirical-statistical (stochastic) data model for the inside of the black box. e.g., a functional form such as multiple linear, exponential, hierarchical ...
- · parameterize the model from the data
- · evaluate the model using model diagnostics

Algorithmic modelling (also called "data-driven")

- · find an algorithm that produces v given x
- · evaluate by **predictive** accuracy (note: not internal accuracy)

Reference: Breiman, L. (2001). Statistical Modelina: The Two Cultures (with comments and a rejoinder by the author). Statistical Science, 16(3), 199-231.

https://doi.org/10.1214/ss/1009213726

Explanation vs.

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Explanation vs. prediction

Explanation

- · Testing a causal theory why are things the way they are?
- Emphasis is on correct model specification and coefficient estimation
- Uses conceptual variables based on theory, which are represented by measureable variables

· Prediction

- Predicting new (space, members of population) or future (time) observations.
- · Uses measureable variables only, no need for concepts

Reference: Shmueli, G. (2010). *To Explain or to Predict?* **Statistical Science**, 25(3), 289-310. https://doi.org/10.1214/10-STS330

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Explanation vs. prediction

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Data-driven vs

The expected prediction error (EPE) for a new observation with value \boldsymbol{x} is:

EPE =
$$E\{Y - \hat{f}(x)\}^2$$

= $E\{Y - f(x)\}^2 + \{E(\hat{f}(x)) - f(x)\}^2$
 $+E\{\hat{f}(x) - E(\hat{f}(x))\}^2$
= $Var(Y) + Bias^2 + Var(\hat{f}(x))$

Model variance: residual error with perfect model specification (i.e., noise in the relation)

Bias: mis-specification of the statistical model: $\hat{f}(x) \neq f(x)$

Estimation variance: the result of using a sample to estimate f as $\hat{f}(x)$

Bias/variance tradeoff: explanation vs. prediction

Modelling

Explanation vs. prediction

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Spatial rando forests Explanation Bias should be minimized

 correct model specification and correct coefficients → correct conclusions about the theory (e.g., causual relation)

Prediction Total EPE should be minimized.

- accept some bias if that reduces the estimation variance
- a simpler model (omitting less important predictors) often has better fit to the data

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Data-driven vs

When does an underspecified model better predict than a full model?

- · the data are very noisy (large σ);
- the true absolute values of the left-out parameters are small;
- the predictors are highly correlated; and
- the sample size is small or the range of left-out variables is narrow.

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Data-driven vs

Problems with data modelling

- Mosteller and Tukey(1977): "The whole area of guided regression [an example of, model-based inference] is fraught with intellectual, statistical, computational, and subject matter difficulties."
- It seems we understand nature if we fit a model form, but in fact our conclusions are about the model's mechanism, and not necessarily about nature's mechanism.
- · So, if the model is a poor emulation of nature, the conclusions about nature may be wrong . . .
- · ... and of course the predictions may be wrong we are incorrectly **extrapolating**.

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Data-driven vs.

The philosophy of data-driven methods

- · Also called "statistical learning", "machine learning"
- Build structures to represent the "black box" without using a statistical model
- Model quality is evaluated by predictive accuracy on test sets covering the target population
 - · cross-validation methods can use (part of) the original data set if an independent set is not available

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Data-driven vs

Some data-driven methods

- Covered in this lecture
 - · Classification & Regression Trees (CART) 分类与回归树
 - · Random Forests (RF) 随 机森林
 - Cubist
- Others
 - · Artificial Neural Networks (ANN) 人工神经网络
 - Support Vector Machines
 - Gradient Boosting
- Relevant R packages: https://cran.r-project.org/ web/views/MachineLearning.html

Data-driven (algorithmic) methods

Kev references – texts

- Hastie, T., Tibshirani, R., & Friedman J. H. (2009). The elements of statistical learning data mining, inference, and prediction (2nd ed). New York: Springer. https://doi.org/10.1007%2F978-0-387-84858-7
- James, G., Witten, D., Hastie, T., & Tibshirani, R. (2021). An introduction to statistical learning: with applications in R (second edition). New York: Springer. https://doi.org/10.1007/978-1-0716-1418-1
- · Stanford online course, based on James et al. book: https://www.edx. org/course/statistical-learning?index=product&queryID= d36d87b77d62e1e9ba8218e5f169cf38&position=1
- Kuhn, M., & Johnson, K. (2013). Applied Predictive Modeling (2013) edition). New York: Springer. https://doi.org/10.1007/978-1-4614-6849-3

Trees (CART)
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Key references - papers

- Shmueli, G. (2010). To Explain or to Predict? Statistical Science, 25(3), 289-310. https://doi.org/10.1214/10-STS330
 - Breiman, L. (2001). *Statistical Modeling: The Two Cultures* (with comments and a rejoinder by the author). **Statistical Science**, 16(3), 199-231. https://doi.org/10.1214/ss/1009213726
- Breiman, L. (2001). Random forests. Machine Learning, 45(1), 5-32. https://doi.org/10.1023/A:1010933404324
- Kuhn, M. (2008). Building Predictive Models in R Using the caret Package. Journal of Statistical Software, 28(5), 1-26. https://doi.org/10.18637/jss.v028.i05

Classification & Trees (CART)

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Data-driven vs.

Decision trees 决策树

- · Typical uses in diagnostics (medical, automotive ...)
- · Begin with the full set of possible decisions
- Split into two (binary) subsets based on the values of some decision criterion
- · Each branch has a more limited set of decisions, or at least has more information to help make a decision
- · Continue **recursively** on both branches until there is enough information to make a decision

Engineering Flowchart



Classification & Regression Trees 分类与回归树

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· A type of decision tree; decision is "what is the predicted response, given values of predictors"?

- Aim is to predict the response (target) variable from one or more predictor variables
- If response is categorical (class, factor) we build a classification tree
- · If response is continuous we build a regression tree
- Predictors can be any combination of categorical or continuous

Classification & Regression Trees (CART)

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Data-driven vs.

Advantages of CART

- · A simple model, **no statistical assumptions** other than between/within class variance to decide on splits
 - For example, no assumptions of the distribution of residuals
 - · So can deal with non-linear and threshold relations
- · No need to transform predictors or response variable
- Predictive power is quantified by cross-validation; this also controls complexity to avoid over-fitting

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Data-driven vs

- No model to interpret (although we can see variable importance)
- Predictive power over a population depends on a sample that is representative of that population
- · Quite **sensitive** to the **sample**, even when pruned
- Pruning to a complexity parameter depends on 10-fold cross-validation, which is sensitive to the choice of observations in each fold
- Typically makes only a small number of different predictions ("boxes"), so maps made with it show discontinuities ("jumps")

Modelling

Explanation v prediction Data-driven (algorithmic)

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Model tuning

Data-driven vs.

- rpart: "Recursive partitioning for classification, regression and survival trees. An implementation of most of the functionality of" Breiman, L., Friedman, J. H., Olshen, R. A., & Stone, C. J. (1983). Classification and regression trees. Wadsworth.
- good introduction: vignette("longintro", package='rpart')
- rpart.plot: "Plot rpart models, an enhanced version of plot.rpart

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Data-driven vs

- splitting variable variable to examine, to decide which branch of the tree to follow
 - · root node 根部节点 variable used for first split; overall mean and total number of observations
- · interior node 非叶子节点 splitting variable, value on which to split, mean and number to be split
- · leaf 叶子点 predicted value, number of observations contributing to it
- cutpoint of the splitting variable: value used to decide which branch to follow
- · growing the tree
- · pruning the tree

Classification Regression

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Example regression tree

- Meuse River soil heavy metals dataset
- · Response variable: log(Zn) concentration in topsoil
- · Predictor variables
 - 1 distance to Meuse river (continuous)
 - 2 elevation above sea level (continuous)
 - **3** flood frequency class (categorical, 3 classes)

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Meuse River study area. Sample points, with Zn concentrations as proportional-size circles, shown in Google Earth

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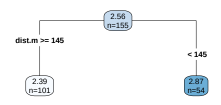
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Example regression tree - first split



Splitting variable: distance to river

Is the point closer or further than 145 m from the river? 101 points *yes*, 54 points *no*.

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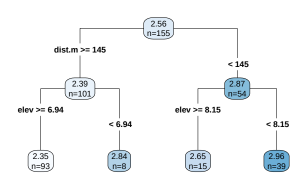
Explanation of first split

- root: average log(Zn) of whole dataset 2.56 log(mg kg⁻¹) fine soil; based on all 155 observations
- · splitting variable at root: distance to river
- · cutpoint at root: 145 m
- · leaves
 - distance < 145 m: 54 observations, their mean is 2.87 log(mg kg⁻¹)
 - · distance \geq 145 m: 101 observations, their mean is 2.39 log(mg kg⁻¹)
 - full dataset has been split into two more homogeneous subsets

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Regression trees

Example regression tree - second split



For both branches, what is the elevation of the point?

Note: this is a coincidence in this case, different splitting variables can be used on different branches.

Regression trees

Explanation of second split

- · interior nodes were leaves after the first split, now 'roots' of subtrees
 - · left: distance \geq 145 m: 101 observations, their mean is 2.39 log(mg kg⁻¹) - note smaller mean on left
 - right: distance < 145 m: 54 observations, their mean is 2.87 log(mg kg⁻¹)
- splitting variable at interior node for < 145 m: elevation
- · cutpoint at interior node for < 145 m: 8.15 m.a.s.l.
- splitting variable at interior node for ≥ 145 m: elevation
- cutpoint at interior node for ≥ 145 m: 6.95 m.a.s.l.
- **leaves** 93, 8, 15, 39 observations; means 2.35, 2.84, 2.65, 2.96 log(mg kg⁻¹)
- These leaves are now more homogeneous than the interior nodes.

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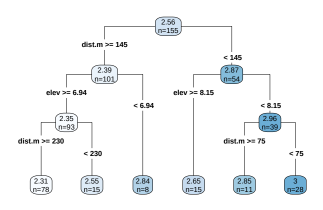
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Data-driven vs.

Example regression tree - third split



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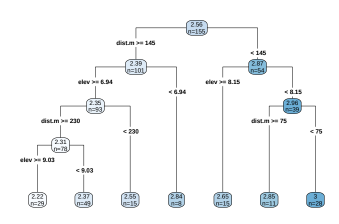
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Example regression tree - fourth split



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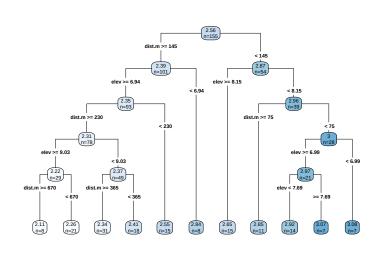
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Example regression tree - fifth split



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Example regression tree - maximum possible splits

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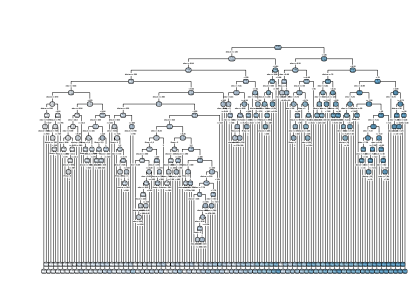
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Data-driven vs

- Take all possible *predictors* and all possible *cutpoints*
- 2 Split the data(sub)set at all combinations
- 3 Compute some **measure of discrimination** for all these i.e., a measure which determine which split is "best"
- 4 Select the predictor/split that most discriminates

Criteria for **continuous** and **categorical** response variables: see next slides

How are splits decided? - Continuous response

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Explanation of prediction of prediction of prediction (algorithmic) methods

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Spatial rando forests Select the predictor/split that most increases between-class variance (this decreases pooled within-class variance):

$$\sum_{\ell} \sum_{i} (y_{\ell,i} - \overline{y_{l}})^{2}$$

- · $y_{\ell,i}$ value i of the target in leaf ℓ
- \cdot $\overline{y_l}$ is the mean value of the target in leaf ℓ

So the set of leaves are **more homogeneous**, on average, than the root.

How are splits decided? - Categorical response

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Data-driven vs

Select the predictor/split that minimizes the *impurity* of the set of leaves:

- Misclassification rate: $\frac{1}{N_m} \sum_{i \in R} I(y_i \neq k(m))$
 - · N_m : number of observations at node m
 - · R_m : the set of observations
 - $\cdot k(m)$ is the majority class; I is the logical T/F function
- Impurity is maximal when all classes have same frequency, and minimal when only one class has any observations in the leaf

So the set of leaves are purer (less confusion), on average, than the root.

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Data-driven
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Regression trees

```
all the possible cutpoints for distance to river
> (distances <- sort(unique(meuse$dist.m)))</pre>
 Γ17
       10
            20
                  30
                       40
                             50
                                  60
                                       70
                                             80
                                                 100
                                                      110
                                                            120
[15]
      160
           170
                 190
                      200
                           210
                                 220
                                      240
                                            260
                                                 270
                                                      280
                                                            290
[29]
      330
                 350
                      360
                            370
                                 380
                                      390
                                           400
                                                      420
                                                            430
            340
                                                 410
[43]
      470
           480
                 490
                      500
                            520
                                 530
                                      540
                                            550
                                                 560
                                                      570
                                                            630
[57]
      690
            710
                 720
                      750
                            760
                                 860 1000
> for (i in 1:nd) {
                      # try them all
  branch.less <- meuse$zinc[meuse$dist.m < distances[i]]</pre>
  branch.more <- meuse$zinc[meuse$dist.m >= distances[i]]
  rss.less <- sum((branch.less-mean(branch.less))^2)
  rss.more <- sum((branch.more-mean(branch.more))^2)
  rss <- sum(rss.less + rss.more)
  results.df[i,2:5] <- c(rss.less, rss.more, rss, 1-rss/tss)
> # find the best split
> ix.r.squared.max <- which.max(results.df$r.squared)</pre>
print(results.df[ix.r.squared.max,])
> print(results.df[ix.r.squared.max.l)
   distance rss.less rss.more
                                     rss r.squared
13
             7127795
                       3030296 10158091
        140
                                          0.510464
> # plot the results
plot(r.squared ~ distance, data=results.df, type="h",
     col=ifelse(distance==d.threshold."red"."grav"))
```

Example split (1)

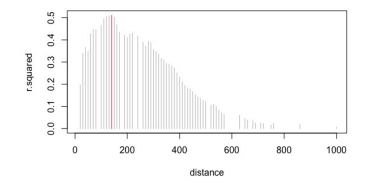
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Data-driven

Example split (2): R^2 vs. cutpoint – distance to river

Try to split the **root node** on this predictor:





Best cutpoint is 140 m; this explains 51% of the total variance

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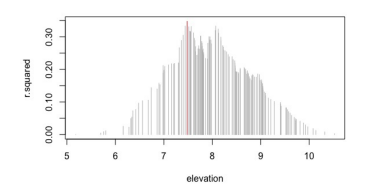
Cubis

Model tuning

forests

Example split (3): R^2 vs. cutpoint – elevation

Try to split the **root node** on this predictor:



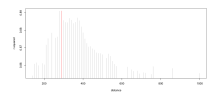
Best cutpoint is 7.48 m.a.s.l.; this only explains 35% of the total variance; so use the distance to river as the first split

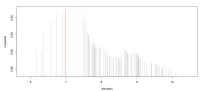
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Regression trees

Example split (4a): left first-level leaf

Try to split the **left first-level leaf** (101 observations):





Best cutpoint is 6.99 m.a.s.l.; this explains 93.0% of the variance in this group. Splitting at 290 m distance would explain 89.1%.

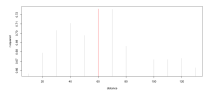
So split this leaf on *elevation* - it becomes an *interior node*

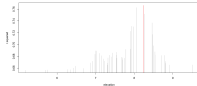
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Regression trees

Example split (4b): right first-level leaf

Try to split the **right first-level leaf** (54 observations):





Best cutpoint is 8.23 m.a.s.l.; this explains 76.6% of the variance in this group. Splitting at 60 m distance would explain 72.6%.

So split on *elevation* – it becomes an *interior node*.

Classification Regression Trees (CART)

Regression trees Sensitivity of Regression Trees Classification trees

Random

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Variable importance Random forests for categorical

Predictor selecti

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Model tunin

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Data-driven vs

Controlling tree complexity

- Fitting a full tree, until there is only one observation per leaf, is always over-fitting to the sample set, and will not be a good predictor of the population.
- · A full tree fits some **noise** as well as **structure**.
- · Can control by the **analyst** or automatically by **pruning** (see below).
- · Analyst can specify:
 - Minimum number of observations in a leaf (fewer: no split is attempted): minsplit
 - Maximum depth of tree: maxdepth
 - Minimum improvement in pooled within-class vs. between-class variance: cp (see below)

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Modelling

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Classification Regression

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Data-driven vs.

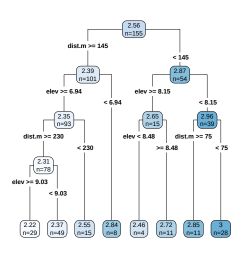
Predicting with the fitted tree

- · A simple 'model' is applied to each leaf:
 - Response variable continuous numeric: mean of observed data in leaf
 - · Categorical variable: most frequent category in leaf
- Value at new location is predicted by running the covariate data down the tree

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Regression trees

Fitted regression tree

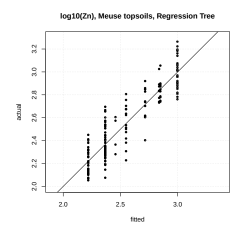


Question: What is the predicted value for a point 100 m from the river and 9 m.a.s.l. elevation?

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Regression trees

Predictions at known points



Note only one prediction per leaf, applies to all points falling in the leaf.

Regression trees

- · The splitting can continue until each calibration observation is in its own leaf
- · This is almost always over-fitting to the current dataset
- · What we want is a tree for the best **prediction**
- · Solution: **grow** a full tree; then **prune** it back to a simpler tree with the best **predictive** power
 - · Similar to using the **adjusted** R^2 to avoid over-fitting a multiple linear regression

Modellin cultures

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Classification Regression Trees (CART)

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Data-driven v

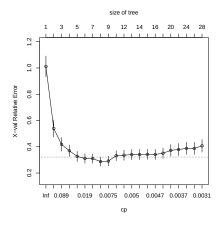
 The cp "complexity parameter" value: Any split that does not decrease the overall lack of fit by a factor of cp is not used.

- Default value is 0.01 (1% increase in R^2)
- · Can be set by the analyst during **growing**
- · Can also be used as a target for **pruning**
- Q: How to decide on the value of cp that gives the best predictive tree?
- A: Use the cross-validation error, also called the out-of-bag error.
 - apply the model to the original data split K-fold (default 10), each time excluding some observations; compare predictions to actual values
 - Note how this fits the philosophy of data-driven approaches: predictive accuracy is the criterion

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Regression trees

X-validation error vs. complexity parameter

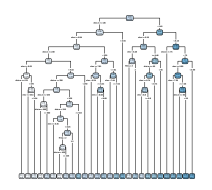


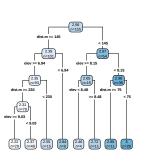
Horizontal line is 1 standard error above the minimum error. Usually choose the largest cp below this; here cp=0.01299 (about 1.3% improvement in R^2).

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Regression trees

Full and pruned trees





Full tree built with cp=0.003 = 0.3%; 27 leaves; pruned to 8 (cp=0.013 = 1.3%)

Interpretation: a noisy dataset if using these two predictors

Classification Regression

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forest

Random forest

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Data-driven v

Variable importance

- Unlike with regression we do not get any coefficient or its standard error for each predictor
- So to evaluate the importance of each predictor we see how much it's used in the tree
 - · simple:
 - \cdot sum of gain in R^2 over all splits based on the predictor
 - complicated;
 - · permute predictor values;
 - · use these to re-build the tree;
 - · compute cross-validation error;
 - the larger the difference, the more important

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Variable importance - example

variable Importance

dist.m 55.5876 elev 38.9996 ffreq 5.4128

Normalized to sum to 100% of the gain in R^2

Distance to river is most important.

Trees (CART)
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Torests

Data-driven ve

Surrogate variables

- What is to be done with observations missing a splitting variable?
 - · e.g., in this case, a new point without a record of its flooding frequency
- Rather than classify the new observation as NA, maybe we can find surrogate variables in the training set: variables that can predict the value of the missing splitting variable at that split.
 - · e.g., maybe *elev* is more useful than random assignment to predict flood frequency.
- These are ranked, and included in the variable importance, even if they are not used in the (pruned) tree.
- Observations missing the split variable are classified using the first surrogate; if that is missing also, the second surrogate, etc.
- · Observations missing the split variable and all surrogates are randomly assigned to one of the branches.

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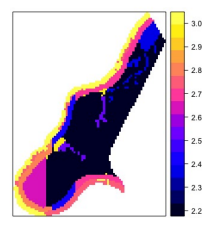
College

Model tunin

Model tuning

Data-driven vs

Map predicted from Regression Tree



This tree: log(Zn) predicted from dist (45% importance); E (17%); soil (15%); N (11%); ffreq. (11%).

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Data-driven vs

Sensitivity of Regression Trees to sample

- **Question**: how sensitive are Regression Trees to the sample?
- Experiment: build trees from random samples of 140 of the 155 observations (only 10% not used!)
 - How different are the optimized trees and the predictive maps?
 - What is the distribution of the optimal complexity parameter and the out-of-bag (predictive) error?

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Modelling

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Explanation

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Predictor selection

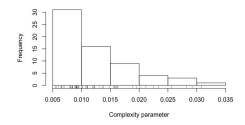
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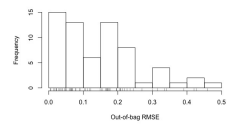
Model tuning

Spatial randor forests

ata-driven vs.

Sensitivity: complexity and out-of-bag error





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Modelling

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Classification

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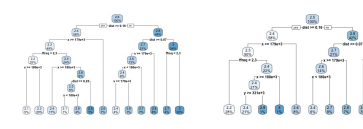
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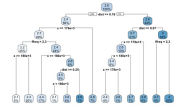
Spatial rando

Data-driven vs

Sensitivity: trees

ffreq = 2,3





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Modellin

Explanation

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Classification Regression

Trees (CART

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Classification

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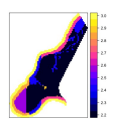
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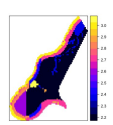
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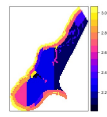
Spatial randor

Data-driven vs

Sensitivity: predictive maps







modelling DGR/罗大维 Modelling cultures

Data-driven

methods for predictive

Regression trees are sensitive to the observations

Classification Regression

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Spatial rand forests · This is a problem!

· Solution: why have one tree when you can have a forest?

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Classification Regression Trees (CART)

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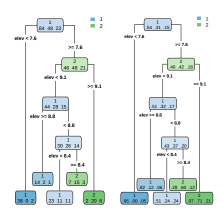
Model tuning

Spatial rando

Data-driven vs

Classification trees

- · Target variable is a categorical variable
- Example (Meuse river): flood frequency class (3 levels) predicted from distance to river and elevation
- Result (pruned): number of observations in each class (left); proportion (right) - note class 3 not predicted!



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Spatial random forests

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Classification & Regression Trees (CART) Regression trees Sensitivity of Regression Trees Classification trees

3 Random forests
Bagging and bootstrapping
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- 4 Cubist
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- 6 Spatial random forests
- Data-driven vs. model-driven methods

Classification Regression Trees (CART)

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Data-driven vs

Random forests - motivation

- · Instead of relying on a *single* (hopefully best) tree, maybe it is better to fit *many* trees.
 - But...how to obtain *multiple* regression trees if we have only *one* data set?
 - Go into field and collect new sample data? too expensive and impractical.
 - Split the dataset and fit trees to the separate parts? Too few observations to build a reliable tree.
 - · **Solution**: Use the *single* sample to generate an *ensemble* (group) of trees; use these together to predict.

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Explanation vs prediction Data-driven (algorithmic)

Classification Regression Trees (CART)

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Torests

"Bag" = a group of samples "in the bag"; others "out-of-bag"

- Suppose we have a large sample that is a good representation of the study area
 - · i.e., sample frequency distribution is close to population frequency distribution
- Generate a new sample is generated by sampling from the sample!

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Explanation of prediction

Data-driven (algorithmic)

Classification Regression

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Data-driven vs

Standard method for sampling in bagging is called **bootstrapping**¹

- · Select same number of points as in sample
- Sample with replacement (otherwise you get the same sample)
- · So some observations are used more than once!
- But, the sample is supposed to represent the population, so these could be values that would have been obtained in a new field sample.

¹ for historical reasons

```
Data-driven
methods
for predictive
modelling
```

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Classification Regression

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forest Variable importance [19] FALSE FALSE

Random forest for categorical

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Model tunin

Spatial rando

Data-driven vs

Sampling with replacement

```
# sample 20 times from (1, 2,... 20) with replacement
> (my.sample <- sample(1:20, 20, replace=TRUE))</pre>
      7 13 5 2 1
                    9 19
                              6
                                 2
                                    9
                                       9 12
                           1
                                             4 11
> sort(my.sample)
 Γ17
                                 9
                                    9
                                       9
                                          9 11 11 12 13 19 20 20
 (1:20) %in% my.sample
                             in bag
      TRUE
            TRUE FALSE
                        TRUE
                              TRUE
                                   TRUE
                                          TRUE FALSE
            TRUE
                  TRUE
                        TRUE FALSE FALSE FALSE FALSE
[19]
      TRUE
            TRUE
> !((1:20) %in% mv.sample)
                            # Out-of-bag
    FALSE FALSE TRUE FALSE FALSE FALSE
                                                TRUF FALSE
[10]
      TRUE FALSE FALSE FALSE
                              TRUE
                                   TRUE
                                          TRUE
                                                TRUE
                                                      TRUE
```

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Model tunin

Data-driven vs

Example: 10 bootstrap samples from the integers 1 ... 20 - sorted

```
b2
           b3 b4 b5
                      b6 b7 b8
                                  b9 b10
        2
    1
                        4
                                         3
                        6
                                         5
                                5
                                       10
                               6
                        8
                                   5
                                       10
6
                      10
                                6
                                   6
                                       11
            8
   11
                      10
                                6
                                       13
            9
   15
                      11
                                       13
   15
           13
               10
                                       13
   16
           15
               10
                      13
                          10
                                   8
                                       14
                                   9
                                       14
                                       14
           16
               14
                                       15
                                       16
                      16
                                       16
               16
                   16
                      18
                                       16
                                       17
                                       19
                                       20
       18 20
                          19 20
              19 19
                      20
                                  20
                                       20
```

Classification Regression Trees (CART)

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forests Bagging and

Bagging and bootstrapping

Building a rando forest Variable

Random fores for categorica

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Cubi

Model tuning

Data-driven vs

Forests with bagging - method

- Fit a full regression tree to each bootstrap sample; do not prune
- Each bootstrap sample results in a **tree** and in a **predicted value** for any combination values of the predictors
- Prediction is the average of the individual predictions from the "forest" of regression trees
- Jumps in predictions are smoothed; more precise predictions

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Explanation prediction Data-driven (algorithmic

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Data-driven vs.

Forest with bagging - limitations

- All predictors are tried at each split, so trees tend to be similar
- Some predictors may never enter into the trees → missing source of diversity
- Solution: random forest variation of bagging two sources of randomness
 - · Random 1: sampling by bagging
 - · Random 2: choice of predictors at each split (see next)

Explanation prediction Data-driven (algorithmic)

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Bagging ar

Building a random forest

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Data-driven v

Random forests

- Multiple samples obtained by bootstrapping, used to build trees (as in bagging)
- · Average predictions over all trees (as in bagging)
- Besides, in each internal node a random subset of splitting variables (predictors) is used
 - · Extra source of diversity among trees
 - Predictors that are "outcompeted" in bagging by stronger competitors may now enter the group of trees

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Explanation prediction Data-driven (algorithmic

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Data-driven vs.

Selecting predictors at each split

- randomForest, ranger parameter mtry: Number of variables randomly sampled as candidates at each split.
 - · ranger default $\lfloor \sqrt{p} \rfloor$, where p is number of possible predictors
 - example: 60 predictors $\rightarrow \lfloor \sqrt{60} \rfloor = \lfloor 7.74 \rfloor = 7$ tried at each split
 - · randomForest default $\lfloor p/3 \rfloor$
 - example: 60 predictors → [60/3] = [20] = 20 tried at each split
- · Can be **tuned**, see below.

Classification Regression

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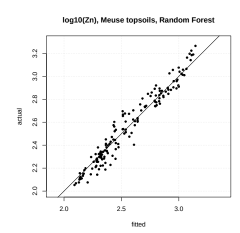
Data-driven vs

Other control parameters

- · number of trees in the forest
 - · ranger parameter min.node.size
 - · randomForest parameter ntree
 - · default = 500
- · minimal node size
 - · ranger parameter min.node.size
 - · randomForest parameter nodesize
 - · default = 5
- (optional) names of variables to always try at each split; weights for sampling of training observations (to compensate for unbalanced samples)

Building a random

forest



Average prediction of many trees, comes close to actual value

Sensitivity of Regression Tree Classification trees

forests

Bagging and

Building a random forest

Variable importance Random forest for categorical

Cubict

Model tunin

forests

Data-driven vs

Out-of-bag ("OOB") evaluation

- · In a bootstrap sample not all samples are present: sampling is with *replacement*.
- · Sample data not in bootstrap sample: **out-of-bag** sample: these were *not* used to build the tree.
- · These data can be used for **evaluation** ("validation"):
 - Use the tree fitted on the bootstrap sample to predict at out-of-bag data, i.e., observations not used in that bootstrap sample.
 - · Compute **squared prediction error** for out-of-bag data.
- This gives a very good estimate of the true prediction error if the sample was representative of the population.

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Explanation v prediction Data-driven

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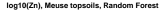
Predictor selection

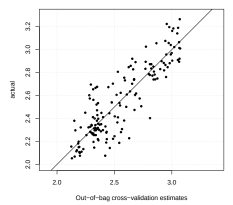
Model tunin

Spatial rando

Data-driven vs

Out-of-bag RF predictions vs. observed





Average prediction of many trees *not* using an observation. Further from actual value; **better estimate of predictive power**

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Modellin

Explanation v prediction Data-driven

Classification Regression

Regression Trees (CART) Regression tree Sensitivity of

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Bagging an bootstrapp

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Predictor selection

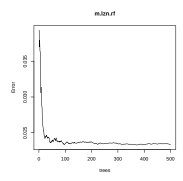
Cubis

Model tuning

forests

How many trees are needed to make a forest?

- Plot mean squared out-of-bag error against number of trees
- · Check whether this is stable
- · If not, increase number of trees



Modelling cultures

Explanation v prediction Data-driven (algorithmic)

Classification Regression Trees (CART)

Sensitivity of Regression Tree Classification

Random forests

Bagging and bootstrapping Building a randor forest

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Cubist

Model tunin

Data-driven v

Importance quantified by permutation accuracy:

- · randomize (permute) values of a predictor
 - · so the predictor can not have any relation with the target
- build a random forest with this randomized predictors and the other (non-randomized) ones
- compute OOB error; compare with OOB error without randomization
 - · the larger the difference, the more important
- Example:

```
% Increase in MSE under randomization ffreq 9.4 dist.m 67.5 elev 54.0
```

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Modellin

Explanation ventorion prediction

Data-driven

Classification

Trees (CART)
Regression trees

Regression 7

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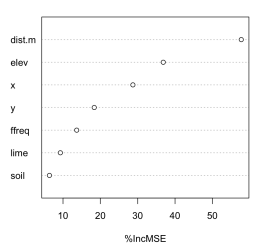
Model tuning

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ata-driven vs.

Variable importance plot





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Modelling

Explanation vs prediction Data-driven (algorithmic)

Classification Regression

Trees (CART)

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Classification trees

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Model tuning

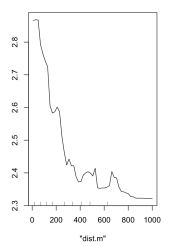
Model tuning

Data-driven vs

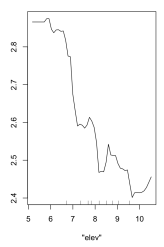
Partial dependence plots

The effect of each variable, with the others held **constant** at their means/most common class.

Partial Dependence on "dist.m"



Partial Dependence on "elev"



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Modellin

Explanation

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Regression Trees (CART

Regression tre Sensitivity of

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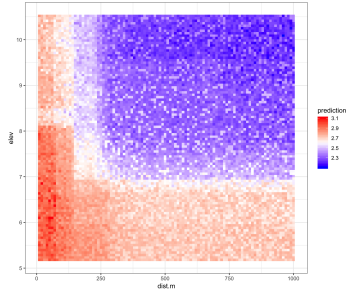
Model tunir

Spatial rando

Data-driven vs.

Two-way partial dependence

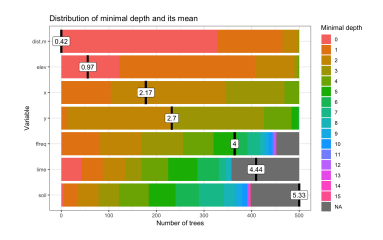
Prediction of the forest for different values of dist.m and elev



DGR/罗大维

Variable importance

Examining the forest - at what depth in the trees are predictors used?



Earlier in tree → most discriminating

Classification Regression

Regression tree Sensitivity of Regression Tree Classification

Random forests

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Predictor selection

Cubis

Model tuning

forests

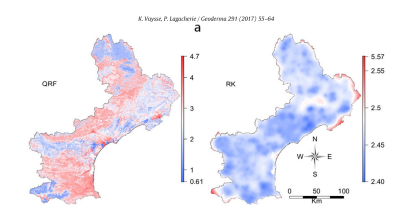
· Recall: RF is built from many trees, each tree makes a

- prediction at each location
- · These are **averaged** to get a "best" predictive map
- However, the set of predictions can be considered a probability distribution of the true value
- · From this we can make a map of any **quantile**, e.g., 5% and 95% confidence limits, or prediction interval width

DGR/罗大维

Variable importance

RF uncertainty vs. RK uncertainty



95% prediction interval for topsoil pH prediction from 2 024 point observations and 18 covariates Languedoc-Roussillon region (F)

DGR/罗大维

Modelling

Explanation v prediction Data-driven (algorithmic)

Classification Regression

Regression tree Sensitivity of Regression Tree Classification trees

Random forests

Bagging and bootstrapping Building a randor

Variable importance

Random forests for categorical

Predictor selecti

Cubis

Model tuni

Spatial rando

Data-driven vs

References for quantile random forests

- Meinshausen, N. (2006). Quantile regression forests. Journal of Machine Learning Research, 7, 983-999.
- Meinshausen, N., & Schiesser, L., 2015. Quantregforest: Quantile Regression Forests. R package. https://cran.r-project.org
- Vaysse, K., & Lagacherie, P. (2017). Using quantile regression forest to estimate uncertainty of digital soil mapping products. Geoderma, 291, 55-64. https://doi.org/10.1016/j.geoderma.2016.12.017

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Modelling

Explanation of prediction Data-driven (algorithmic)

Classification Regression

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Data-driven vs

Random forests for categorical variables

- · Target variable is **categorical**, i.e., a class
 - Example: Meuse river flooding frequency classes (every year, every 2-5 years, rare or none)
- Final prediction is the class predicted by the majority of the regression trees in the forest
- Can also see the probabilty for each class, by predicting with the model with the type="prob" argument to predict.randomForest.

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Modelling

Explanation vs prediction Data-driven (algorithmic)

Classification Regression

Regression Trees (CART)

Sensitivity of Regression Tre Classification

Random

Bagging and bootstrapping Building a rand forest

Random forests for categorical variables

Predictor selection

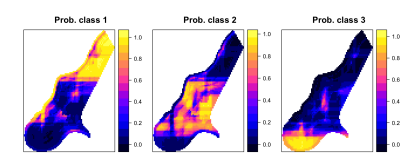
Cubis

Model tuning

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Data-driven vs

Predicted class probabilty



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Modellin

Explanation prediction

Data-driven

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Trees (CART)

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Pandom

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Building a random

forest

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Predictor selection

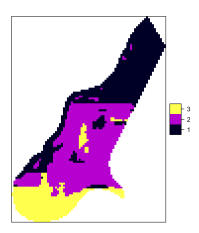
Cubic

Model tu

Spatial randor

Data-driven vs.

Predicted most probable class



Random forests

for categorical variables

Accuracy measures

- naïve agreement: how often a class in the training set is correctly predicted - see with a confusion matrix ("cross-classification")
- · Out-of-bag (OOB) estimate of error rate
- · **Gini impurity**: how often a *randomly chosen* training observation would be incorrectly assignedif it were randomly labeled according to the frequency distribution of labels in the subset.

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Data-driven vs

Cross-classification matrix

A **confusion matrix** (a.k.a. cross-classification matrix) of actual (columns) vs. predicted (rows) classes:

Confusion matrix:

1 2 3 class.error 1 77 7 0 0.08333333

2 3 40 5 0.16666667

3 1 9 13 0.43478261

3 1 9 13 0.434/8261

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Predictor selection

Predictor selection

Problem: large number of possible predictors, can lead to

- Computational inefficiency
- Difficult interpretation of variable importance
- Meaningless good fits, even if using cross-validation²
- · **Solution 1**: expert selection from "known" relations
 - · this is then not pure "data mining" for unsuspected relations
- · **Solution 2**: (semi-)automatic feature selection, see next.

²Wadoux, A. M. J.-C., et al. (2019). A note on knowledge discovery and machine learning in digital soil mapping. European Journal of Soil Science, 71, 133-136. https://doi.org/10.1111/ejss.12909

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Modelling

Explanation vs prediction Data-driven (algorithmic)

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Model tuning

Data-driven vs

Feature selection methods

Wrapper methods: "evaluate multiple models using procedures that add and/or remove predictors to find the optimal combination that maximizes model performance."

- risk of over-fitting
- · high computational load

Filter methods: "evaluate the relevance of the predictors outside of the predictive models and subsequently model only the predictors that pass some criterion"

- does not account for correlation among predictors
- · does not directly assess model performance

Classification Regression Trees (CART)

Sensitivity of Regression Tree Classification trees

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Data-driven vs

Recursive feature elimination

- · A "wrapper" method
- Implemented in caret::rfe "Backwards Feature Selection" function
- Algorithm: "Recursive Feature Elimination (RFE) incorporating resampling"
 - 1 Partition data into training/test sets via resampling
 - 2 Start with **full model**, compute variable importance
 - 3 for each proposed subset size
 - Re-compute model with reduced variable sets
 - Calculate performance profiles using test samples
 - 4 Determine optimum number of predictors

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Predictor selection

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Model tunir

Spatial rando forests

Data-driven v

Reference for feature selection

- \cdot From the documentation of the caret package (§5).
- Feature selection: https://topepo.github.io/caret/ feature-selection-overview.html
- Recursive feature elimination: https://topepo.github.io/caret/ recursive-feature-elimination.html

Cuhist

Modelling cultures

Classification & Regression Trees (CART)

4 Cubist

Model tuning

Cuhist

- · Similar to CART, but instead of **single values** at leaves it creates a multivariate linear regression for the cases in the leaf
- · Advantage vs. CART: predictions are continuous, not discrete values equal to the number of leaves in the regression tree.
 - · Also can be improved with nearest-neighbours, see below
- · Advantage vs. RF: the model can be interpreted, to a certain extent.
- **Disadvantage**: its algorithm is not easy to understand; however its results are generally quite good.

Cuhist

- · "Committees" of models: a sequence of models, where each corrects the errors in the previous one
 - nearest-neighbours adjustment: modify model result at a prediction point from some number of neighbours in feature (predictor) space.

$$\hat{\mathbf{y}}' = \frac{1}{K} \sum_{i=1}^{K} \mathbf{w}_i \left[\mathbf{t}_i + (\hat{\mathbf{y}} - \hat{\mathbf{t}}_i) \right] \tag{1}$$

where t_i is the actual value of the neighbour, \hat{t}_i is its value predicted by the model tree(s), and w_i is the weight given to this neighbour for the adjustment, based on its distance D_i from the target point. These are computed as $w_i = 1/(D_i + 0.5)$ and normalized to sum to one.

```
Data-driven
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```

Cubist

Example cubist model

Rule 1/1: [66 cases, mean 2.288309, range 2.053078 to 2.89098, err 0.1036 if x > 179095, dist > 0.211846then outcome = 2.406759 - 0.32 dist

Rule 1/2: [9 cases, mean 2.596965, range 2.330414 to 2.832509, err 0.1163 if $x \le 179095$. dist > 0.211846

then outcome = -277.415278 + 0.000847 v + 0.56 distRule 1/3: [80 cases, mean 2.772547, range 2.187521 to 3.264582, err 0.157

if dist ≤ 0.211846 then outcome = 2.632508 - 2.1 dist - 2.4e-05 x + 1.4e-05 v

Rule 2/1: [45 cases, mean 2.418724, range 2.10721 to 2.893762, err 0.1822 if $x \le 179826$, ffreq in $\{2, 3\}$

then outcome = 128.701732 - 0.000705 xRule 2/2: [121 cases, mean 2.443053, range 2.053078 to 3.055378, err 0.18

if dist > 0.0703468then outcome = 30.512065 - 0.87 dist - 0.000154 x

Rule 2/3: [55 cases, mean 2.543648, range 2.075547 to 3.055378, err 0.125

if dist > 0.0703468, ffreq = 1 then outcome = $37.730889 - 0.000314 \times - 0.35 \text{ dist} + 6.5e-05 \text{ y}$

Rule 2/4: [34 cases, mean 2.958686, range 2.574031 to 3.264582, err 0.139

if dist ≤ 0.0703468

then outcome = 2.982852 - 0.36 dist

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Modellin

Explanation

Data-driven (algorithmic methods

Classification (Regression

Regression trees

Regression Tre

Random

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Variable

Random forest for categorical

Predictor selection

Cubist

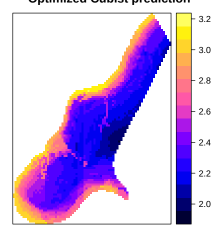
Model tuning

Spatial rando forests

ata-driven vs.

Optimized Cubist prediction

Map predicted by Cubist



Modelling cultures

Classification & Regression Trees (CART)

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Model tuning

Modellin cultures

prediction v prediction Data-driven (algorithmic)

Classification Regression

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Cuhist

Model tuning

Data-driven vs.

 Data-driven models have parameters that control their behaviour and can significantly affect their predictive power.

· CART: complexity parameter

 randomForest: number of predictors to try at each split; minimum number of observations in a leaf; number of trees in the forest

 too many predictors → trees too uniform, loss of diversity; too few → highly-variable trees, poor predictions

 too few observations per leaf to imprecise prediction; too many → over-fitting

 too few trees → sub-optimal model; too many trees → wasted computation

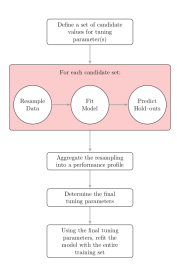
 Cubist: number of committees; number of nearest neighbours

 The model can be tuned to optimize the selection of these.

DGR/罗大维

Model tuning

Model tuning - flow chart



source: Kuhn, M., & Johnson, K. (2013). Applied Predictive Modeling (2013 edition). New York: Springer; figure 4.4

Regression Trees (CART)

Sensitivity of Regression Tre-Classification trees

Random

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Predictor sele

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Spatial randor

ata-driven vs.

Model tuning - algorithm

- For each combination of parameters to be optimized:
 - Split the dataset into some disjunct subsets, for example 10, by random sampling.
 - 2 For each subset:
 - 1 Fit the model with the selected parameters on all but one of the subsets (train subset).
 - 2 Predict at the remaining subset, i.e., the one not used for model building, with the fitted model.
 - 3 Compute the goodness-of-fit statistics of fitting to the test subset
 - e.g., root mean square error (RMSE) of prediction; squared correlation coefficient between the actual and fitted values, i.e., R^2 against a 1:1 line.
 - 3 Average the statistics for the disjunct test subsets.
- 2 Search the table of results for the best results e.g., lowest RMSE, highest R^2 .

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Modelling

Explanation v prediction Data-driven (algorithmic)

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Variable importance Random fores

variables
Predictor selection

Cuhist

Model tuning

ata-driven vs.

Model tuning - R implementation

- · caret "Classification And REgression Training" package
 - Kuhn, M. (2008). Building predictive models in R using the caret package. Journal of Statistical Software, 28(5), 1-26.
 - https://topepo.github.io/caret/index.html
 - · can tune 200+ models; some built-in, some by calling the appropriate package
- method:
 - set up a vector or matrix with the parameter values to test, e.g, all combinations of 1 ... 3 splitting variables to try, and 1 ... 10 observations per leaf
 - 2 run the model for all of these and collect the cross-validation statistics
 - 3 select the best one and build a final model

```
Data-driven
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```

Model tuning

Model tuning example - random forest (1)

```
> ranger.tune <- train(x = preds, y = response, method="ranger",</pre>
              tuneGrid = expand.grid(.mtry = 1:3,
                                        .splitrule = "variance",
                                        .min.node.size = 1:10).
               trControl = trainControl(method = 'cv'))
> print(ranger.tune)
## Resampling: Cross-Validated (10 fold)
   Resampling results across tuning parameters:
##
##
     mtry
           min.node.size
                           RMSE
                                      Rsquared
                                                  MAE
##
     1
            1
                            199.7651
                                      0.8862826
                                                  156, 1662
##
     1
                            200.5215
                                      0.8851154
                                                  156.3225
     1
##
                           200.6421
                                      0.8854146
                                                  156,2801
. . .
##
     3
            8
                           201.9809
                                      0.8793349
                                                  158,7097
     3
##
            9
                           202,9065
                                      0.8781754
                                                  159,7739
##
           10
                            202.5687
                                      0.8788200
                                                  159.5980
##
   RMSE was used to select the optimal model
## Final values: mtrv = 2. min.node.size = 6.
```

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Modellin

Explanation version prediction

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Predictor selection

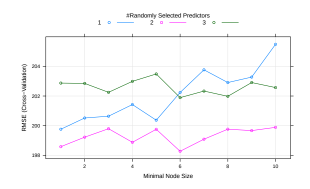
C 11 .

Model tuning

Spatial rando

ata-driven vs

Model tuning example - random forest (2)



Find the minimum RMSE; but favour simpler models (fewer predictors, larger nodes) if not too much difference

```
Data-driven
  methods
for predictive
 modelling
```

Model tuning

Model tuning example - Cubist (1)

```
> cubist.tune <- train(x = all.preds. v = all.resp. method="cubist".</pre>
                      tuneGrid = expand.grid(.committees = 1:12,
                                              .neiahbors = 0:5).
                      trControl = trainControl(method = 'cv'))
```

```
Resampling: Cross-Validated (10 fold)
   Summary of sample sizes: 139, 139, 140, 139, 139, 139, ...
   Resampling results across tuning parameters:
##
##
     committees
                             RMSE
                                        Rsquared
                 neiahbors
                                                   MAF
##
                             0.1898596
                                        0.6678588
                                                    0.1405553
      1
##
                             0.1764705
                                         0.6953460
                                                    0.1189364
##
                             0.1654910
                                         0.7296723
                                                    0.1163660
                             0.1623381
                                        0.7425831
##
                                                    0.1163285
      1
                             0.1631900
                                         0.7453506
                                                    0.1192963
##
. . .
##
                             0.1599994
                                         0.7533962
                                                    0.1139932
     12
##
     12
                 4
                             0.1584434
                                         0.7617762
                                                    0.1153331
##
     12
                             0.1589143
                                         0.7622337
                                                    0.1165942
\##
## RMSE was used to select the optimal model using the smallest value.
```

The final values: committees = 10, neighbors = 4.

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Modellin

Explanation vs prediction Data-driven (algorithmic) methods

Classification Regression Trees (CART)

Sensitivity of Regression Tre Classification

Random

Bagging and bootstrapping Building a random forest

Variable importance Random forests for categorical

Predictor selection

Predictor selection

Model tuning

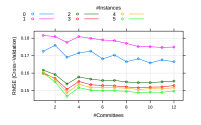
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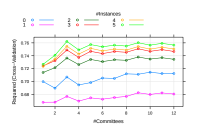
ata-driven v

Model tuning example - Cubist (2)

Criterion: RMSE

Criterion: R²





Adding one neighbour reduces predictive power; adding 2 ...increases it; 3 is close to optimum

Committees improve predictive power; 3 is optimum

Spatial random

forests

Modelling cultures

Classification & Regression Trees (CART)

- Model tuning
- 6 Spatial random forests

Classification Regression Trees (CART)

Regression trees Sensitivity of Regression Tree Classification trees

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Predicto

Cubist

Spatial random

Data-driven vs.

Spatial random forests

- Random forests can use coördinates and distances to geographic features as predictors
 - \cdot e.g., E, N, distance to river, distance to a single point \dots
- · Can also use distances to multiple points as predictors
 - Distance buffers: distance to closest point with some range of values
 - Common approach: compute quantiles of the response variable and one buffer for each
 - Each sample point has a distance to the closest point in each quantile
- This uses separation between point-pairs of different values, but with no model.

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Explanation v prediction

Classification Regression

Regression Trees (CART) Regression tree

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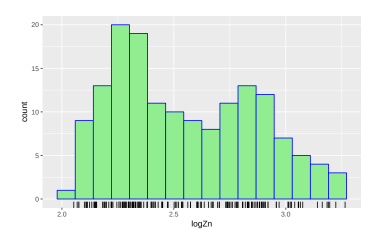
Cubic

Model tuning

Spatial random forests

Data-driven vs

log(Zn) distribution - 16 quantiles



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Modelling

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Classification Regression

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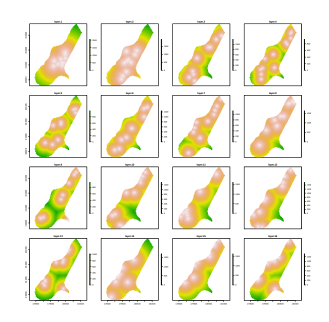
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Data-driven vs.

Distance to closest point in each quantile



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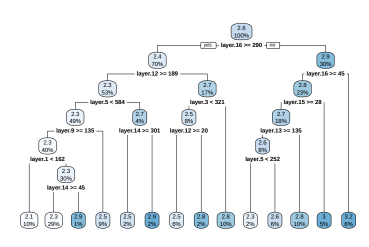
Cubis

Model to

Spatial random forests

Data-driven vs.

Regression tree on 16 distance buffers



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Classification Regression

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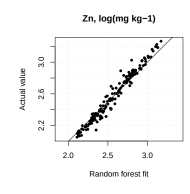
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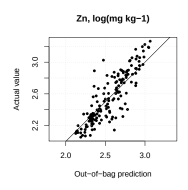
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Data-driven v

Random forest prediction on 16 distance buffers





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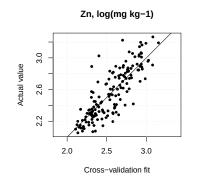
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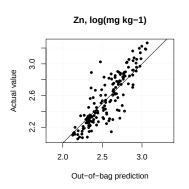
Spatial random

forests

Data-driven v

OOB error vs. OK cross-validation error





OK RF

Note that RF does *not* use any *model* of spatial autocorrelation!

DGR/罗大维

Modellin

Explanation v prediction Data-driven

Classification

Regression Trees (CART

Regression tre Sensitivity of

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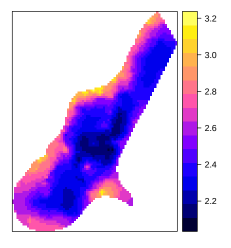
Cubis

Model tu

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Data-driven vs.

Random forest map on 16 distance buffers



Resembles OK map, but no model was used.

DGR/罗大维

Modellir

Explanation vs prediction Data-driven (algorithmic)

Classification Regression

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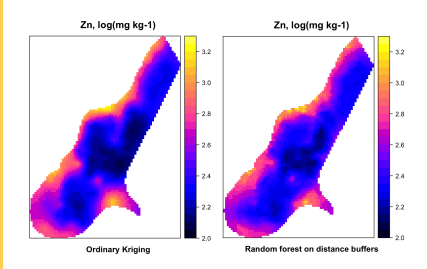
Predictor Self

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Compare with Ordinary Kriging



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Modelling

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Classification & Regression

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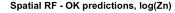
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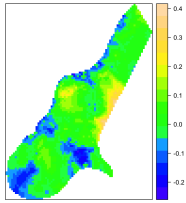
Model tuni

Spatial random forests

Data-driven vs.

Difference spatial RF - OK





DGR/罗大维

Modellin

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Spatial random forests

ata-driven vs.

Reference for spatial random forests

 Hengl, T., Nussbaum, M., Wright, M. N., Heuvelink, G. B. M., & Gräler, B. (2018). Random forest as a generic framework for predictive modeling of spatial and spatio-temporal variables. PeerJ, 6, e5518. https://doi.org/10.7717/peerj.5518

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Modelling

Explanation v prediction Data-driven (algorithmic)

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Classification & Regression Trees (CART) Regression trees Sensitivity of Regression Trees Classification trees

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Data-driven vs.

Conclusion: Data-driven vs. model-based methods

- · Data-driven: main aim is predictive power
 - Individual trees can be interpreted, but forests can not (only can see variable importance, not choice or cutpoints)
- · Model-based: main aim is understanding processes
 - We hope the model is a simplified representation of the process that produced the observations
 - · If the model is correct, predictions will be accurate

Classification Regression Trees (CART)

Sensitivity of Regression Tro Classification

Random

Bagging and bootstrapping Building a random forest

importance
Random forests
for categorical

Predictor selection

Cuhis

Model tunin

Spatial randor forests

Data-driven vs.

Conclusion: limitations

- Data-driven methods depend on their training observations
 - They have no way to extrapolate or even interpolate to unobserved areas in feature space
 - So the observations should cover the entire range of the population
- Model-based methods depend on a correct empirical-statistical model
 - Model is derived from training observations, but many models are possible
 - · Various model-selection techniques
 - Wrong model → poor predictions, incorrect understanding of processes

Modelling

Explanation v prediction Data-driven (algorithmic)

Regression

Trees (CART)
Regression tree

Regression Tree

Random

ваддінд and bootstrapping Building a random

forest

importance Random forests for categorical

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