# **GEOSTAT**

# Mastering Machine Learning for Spatial Prediciton 2

### **Exercises**

Madlene Nussbaum 13 August 2017

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## Preparation

Load needed packages:

```
library(randomForest)
library(geoGAM)
```

Load again the data, select the calibration set and remove missing values in covariates.

```
dim(berne)

## [1] 1052 238

# Continuous response
d.ph10 <- berne[ berne$dataset == "calibration" & !is.na(berne$ph.0.10), ]
d.ph10 <- d.ph10[ complete.cases(d.ph10[13:ncol(d.ph10)]), ]
# covariates start at col 13
l.covar <- names(d.ph10[, 13:ncol(d.ph10)])</pre>
```

## 1 Selection of covariates

For tree based ensemble methods covariate importance can be computed. Based on this measure non-relevant covariates can be excluded and possibly model performance can be increased.

Fit random forest model:

Create the importance plot:

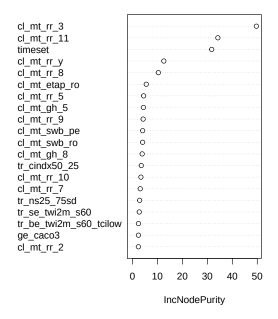
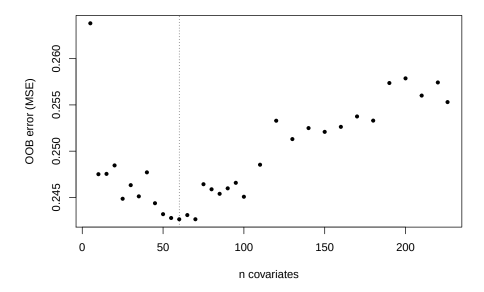


Figure 1: Covariate importance of 20 most important covariates for topsoil pH (before selection).

```
varImpPlot(rf.ph, n.var = 20, main = "")
```

Then, reduce covariates by recursive backward elimination using permuted covariate importance (type = 2 in importance()):

```
# speed up the process by removing 5-10 covariates at a time
s.seq <- sort(c(seq(5, 95, by = 5),
                   seq(100, length(l.covar), by = 10)),
                decreasing = T)
# collect results in list
qrf.elim <- oob.mse <- list()</pre>
# save model and OOB error of current fit
qrf.elim[[1]] <- rf.ph</pre>
oob.mse[[1]] <- tail(qrf.elim[[1]]$mse, n=1)
1.covar.sel <- 1.covar</pre>
# Iterate through number of retained covariates
for( ii in 1:length(s.seq) ){
  t.imp <- importance(qrf.elim[[ii]], type = 2)</pre>
  t.imp <- t.imp[ order(t.imp[,1], decreasing = T),]</pre>
  qrf.elim[[ii+1]] <- randomForest(x = d.ph10[, names(t.imp[1:s.seq[ii]])],</pre>
                                     y = d.ph10$ph.0.10)
  oob.mse[[ii+1]] <- tail(qrf.elim[[ii+1]]$mse,n=1)</pre>
```



**Figure 2:** Path of out-of-bag mean squared error as covariates are removed. Minimum is found at 55 covariates.

## Please continue:

- Optimize  $m_{try}$  before you start the covariate selection (function train, package caret). How much does the OOB error decrease? Are both steps (tuning, selection) worth the effort from a point of view of prediction performance?
- Implement the same covariate selection for gradient boosting with trees as baselearners (package gbm or caret). Do you find the same covariates in the final set? Why do you expect differences?

## 2 Partial dependence plots

Interpretation of the most important covariates can be done by partial dependence plots. But keep in mind that the remaining covariate set might be still multi-collinear, hence covariates might be exchangeable.

```
# select the model with minimum OOB error
rf.selected <- qrf.elim[[ which.min(elim.oob$elim.00Be)]]</pre>
t.imp <- importance(rf.selected, type = 2)</pre>
t.imp <- t.imp[ order(t.imp[,1], decreasing = T),]</pre>
# 4 most important covariates
(t.3 <- names(t.imp[1:4]))
## [1] "cl_mt_rr_3" "cl_mt_rr_11" "timeset"
                                                  "cl_mt_rr_y"
par(mfrow = c(2,2))
# Bug in partialPlot(): function does not allow a variable for the
# covariate name (e. g. x.var = name) in a loop
partialPlot(x = rf.selected,
            pred.data = d.ph10[, names(rf.selected$forest$xlevels)],
            x.var = "cl_mt_rr_3", ylab = "ph [-]", main = "")
partialPlot(x = rf.selected,
            pred.data = d.ph10[, names(rf.selected$forest$xlevels)],
            x.var = "cl_mt_rr_11", ylab = "ph [-]", main = "" )
partialPlot(x = rf.selected,
            pred.data = d.ph10[, names(rf.selected$forest$xlevels)],
            x.var = "timeset", ylab = "ph [-]", main = "" )
partialPlot(x = rf.selected,
            pred.data = d.ph10[, names(rf.selected$forest$xlevels)],
            x.var = "cl_mt_rr_y", ylab = "ph [-]", main = "" )
```

#### Please continue:

- Create partial dependence plots for the boosted trees model (?plot.gbm, plot(.., i.var = ..)). Do you find the same relationships?
- What do you conclude from the plots? Are the plotted covariates good predictors?

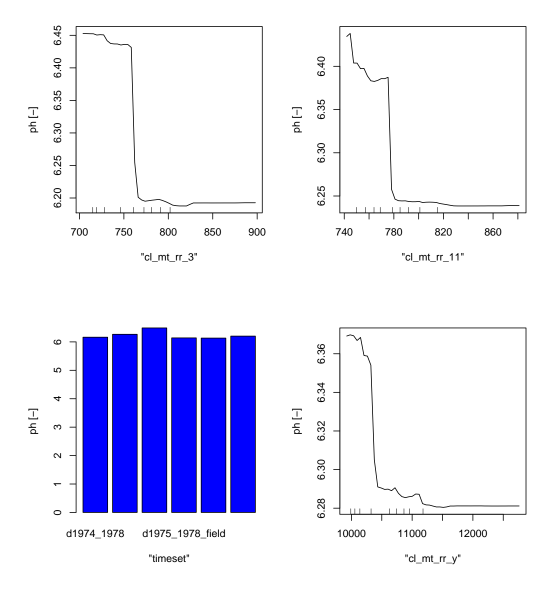
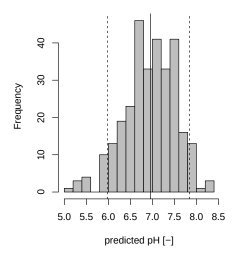


Figure 3: Partial dependence plots for the 4 most important covariates.

# 3 Prediction uncertainty with model-based bootstrapping

When reporting predictions it is important to give prediction uncertainty along with them. For any method not yielding uncertainty estimates form the method itself (e.g. kriging variances) a model-based bootstrap approach can be used.

Compute model based bootstrap with 300 repetitions:



**Figure 4:** Histogram of predictive distribution for one single prediction point, computed by 300 bootstrap repetitions.

Plot for evaluation of prediction intervals (as shown in presentation):

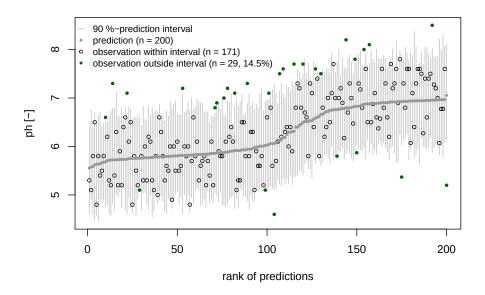
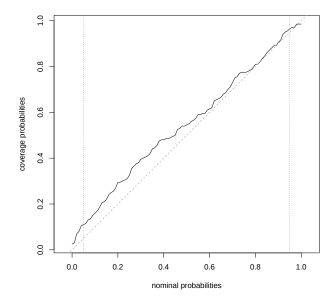


Figure 5: Coverage of 90-prediction intervals computed by model-based boostrap.

```
# Coverage probabilities plot
# create sequence of nominal probabilities
ss < -seq(0,1,0.01)
# compute coverage for sequence
t.prop.inside <- sapply(ss, function(ii){</pre>
  boot.quantile <- t( apply(ph.boot, 1, quantile, probs = c(0,ii) ) )[,2]</pre>
  return( sum(boot.quantile <= d.ph10.val$ph.0.10)/nrow(d.ph10.val) )</pre>
})
plot(x = ss, y = t.prop.inside[length(ss):1],
     type = "1", asp = 1,
     ylab = "coverage probabilities",
     xlab="nominal probabilities" )
# add 1:1-line
abline(0,1, lty = 2, col = "grey60")
# add lines of the two-sided 90 %-prediction interval
abline(v = c(0.05, 0.95), lty = "dotted", col = "grey20")
```

#### Please continue:

- Are you satisfied with the prediction intervals?
- Compute prediction intervals for random forest (Package quantregForest) and create the same plots. How do the intervals of quantile regression forest perform compared to the validation data?
- Create maps of the prediction intervals for the berne.grid data. Is there much spatial structure in the uncertainty? Do quantile regression forest and bootstrapped intervals differ? If yes, why (which assumption)?



**Figure 6:** Coverage probabilities of one-sided prediction intervals computed for the validation data set of topsoil pH of the Berne study area.

## R session information

This document was generated with:

### toLatex(sessionInfo(), locale = FALSE)

- R version 3.4.1 (2017-06-30), x86\_64-pc-linux-gnu
- Running under: Progress Linux 4+ (dschinn-backports)
- Matrix products: default
- BLAS: /usr/lib/libblas/libblas.so.3.7.0
- LAPACK: /usr/lib/lapack/liblapack.so.3.7.0
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: geoGAM 0.1-1, knitr 1.16, randomForest 4.6-12
- Loaded via a namespace (and not attached): codetools 0.2-15, coin 1.2-0, compiler 3.4.1, digest 0.6.12, evaluate 0.10.1, grid 3.4.1, grpreg 3.1-1, highr 0.6, lattice 0.20-34, magrittr 1.5, MASS 7.3-47, Matrix 1.2-7.1, mboost 2.8-0, mgcv 1.8-17, modeltools 0.2-21, multcomp 1.4-6, mvtnorm 1.0-6, nlme 3.1-129, nnls 1.4, parallel 3.4.1, party 1.2-3, quadprog 1.5-5, sandwich 2.3-4, splines 3.4.1, stabs 0.6-2, stats4 3.4.1, stringi 1.1.5, stringr 1.2.0, strucchange 1.5-1, survival 2.40-1, TH.data 1.0-8, tools 3.4.1, zoo 1.8-0