## Day 2, Practical 2

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### Super learner

1. Load the following libraries (install them if necessary).

```
library(riskRegression)
library(nnls)
library(foreach)
library(SuperLearner)
library(ranger)
library(randomForest)
library(randomForestSRC)
library(rms)
library(data.table)
```

- 2. Download the pph data from the course website. The pph data are saved in rds (R Data Single) format.
  - Load the data into R (readRDS).
  - Split the data into two data sets:
    - pph09 contains only data from calendar year 2009 (Year==2009)
    - pph contains data from the previous years (Year!=2009).
  - The outcome variable is planned cesarian section at second delivery: plannedCS (1="yes", 0="no"). Calculate the probability of a planned cesarian section at second delivery by calendar year. Is there a calendar time trend?
  - The predictor variables are the following characteristics of the first delivery:
    - MotherAge
    - PrevPPHbin
    - PrevArgumented
    - PrevPraeecl
    - PrevAbruptio

- PrevCS
- PrevRetained
- PrevInduced

Calculate the median age and the probabilities of the other risk factors at second delivery by calendar year. Are there calendar time trends?

- 3. Fit the following models in the data pph (before 2009):
  - logistic regression (glm) with additive effects of all predictor variables.
  - generalized additive model (gam) with additive effects of all predictor variables where the effect of MotherAge is modeled by a smoothing spline.
  - Random forest (rfsrc) with 200 trees (to make the fit faster) and otherwise default parameters.

Then predict the probabilities of a planned cesarian section for the 2009 data. Scatterplot the predicted probabilities against each other. Calculate the average Brier scores in the 2009 data.

- 4. Create level-one data
  - data splitting: 10-fold cross-validation
    - split the data pph (before 2009) at random into 10 non-overlapping subsets (called folds in what follows) with roughly same size.
  - in a foreach::foreach loop (with argument .combine set to "rbind")
     across the folds do:
    - fit all models (i.e., train all learners) from step 3 in the data which excludes the current fold
    - predict the probabilities of planned cesarian section for the subjects in the current fold
    - combine the resulting matrix with the observed outcome values (plannedCS) of the current fold into a data frame: the level-1 data corresponding to this fold.
  - the result of the foreach loop are the level-1 data obtained with 10-fold cross-validation. There are 4 columns where the first is the observed outcome (plannedCS) and the remaining are the cross-validated predicted probabilities of the three learners. The level-1 data are used in the following to construct super learners.
- 5. Calculate the following super learners:
  - discrete super learner (manual programming)
  - Polley's default: non-negative least squares (package nnls)

• Breiman's suggestion: shrinkage (e.g., package: rms::lrm, set penalty to 0.2)

Then scatterplot the predicted probabilities of the 3 super learners against each other.

#### 6. Use the SuperLearner package:

- from the learning data extract the outcome column (argument Y) and construct the design matrix which contains the "dummy coding" of the predictor values (argument X), e.g., with model.matrix.
- specify the following super learner libraries:
  - SL.mean
  - SL.glm
  - SL.gam
  - SL.glmnet
- Consider the coefficients of the nnls fit: object\$coef
- Plot the predicted values: object\$SL.predict against the manually computed super learner from step 5.

### 7. Compare the average Brier scores of:

- all single models from step 3
- the super learner model from step 6.
- the single library results of the super learner from step 6.

#### 8. Monte-Carlo error

- Check the program and mark all lines with random seed dependence.
- Set the seed (set.seed) to control the randomness and re-run the whole program.
- Visualize the seed dependence of the super learner obtained with the SuperLearner package (step 6) by scatterplotting the predicted values in the 2009 data for two different seeds.

### 9. Tune the random forest parameters

- create a list of strong random forest learners by varying the ranger parameters mtry (values 1,3,7 and minimum node size (values 20,50,100). See section 10 of the R package vignette: Guide-to-SuperLearner.
- run the SuperLearner by adding the tuned forest to the libraries of step 7, check the coefficients ...

# References

- 1. David H Wolpert. Stacked generalization. Neural Networks, 5(2):241-259, 1992.
- 2. Leo Breiman. Stacked regressions. Machine Learning, 24(1):49–64, 1996.
- 3. Mark J Van der Laan, Eric C Polley, and Alan E Hubbard. Super learner. Statistical Applications in Genetics and Molecular Biology, 6(1), 2007.