# Committees, ensembles.

# Petr Pošík

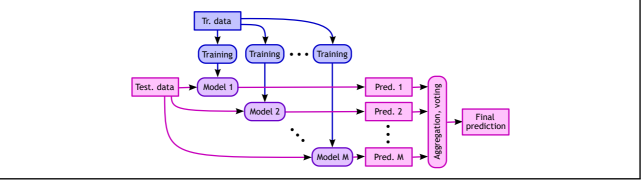
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# Introduction

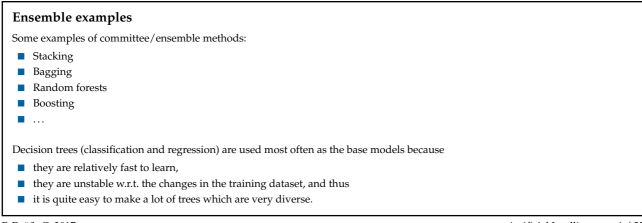
## Ensemble a.k.a committee

- ML model composing multiple different models to obtain better predictive performance than could be obtained from any of the constituent models.
- A way to compensate for poor learning algorithms by performing a lot of extra computations.
- Ensembles tend to yield better results when there is a significant diversity among the models (Intuition: averaging reduces variance).
- Individual ensamble/committee methods differ in the way they create individual *models different from each other*.
- Use different kinds of models, or models unstable w.r.t. a change in the training data.



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## Aggregation

The final aggregation of results of individual models is usually done by

- (weighted) *voting* of individual models for *classification* problems,
- (weighted) *averaging* of individual models for *regression* problems,
- or by other techniques.

#### Stacking

- Assume we have M different models  $h_m$  created for the same modeling task, each being a function  $h_m(x)$  of the input features x.
- The predictions of these models,  $h(x) = (h_1(x), \dots, h_M(x))$ , may be considered new features extracted from the data set (basis expansion).
- We can thus train a higher-level classification/regression model  $h_{\text{stack}}$  as a function of these new features, i.e.  $h_{\text{stack}}(h)$  (sometimes together with the original features, i.e.  $h_{\text{stack}}(x, h)$ ).
- For classification, logistic regression is often used as  $h_{\text{stack}}$ .
- For regression, multiple linear regression is often used as  $h_{\text{stack}}$  with the constraint on the weights  $w_i$  such that  $\sum w_i = 1$  and  $w_i > 0 \forall i$ .
- An obvious way to estimate the weights w as  $w^* = \arg \min_{w} \sum_{i=1}^{|T|} L\left(y_i, \sum_{m=1}^{M} w_m h_m(x_i)\right)$ , however, can result in overfitting; this is solved by LOO cross-validation, i.e. using the estimate  $w^* = \arg \min_{w} \sum_{i=1}^{|T|} L\left(y_i, \sum_{m=1}^{M} w_m \hat{h}_m^{-i}(x_i)\right)$ , where  $\hat{h}_m^{-i}$  is a predictor obtained by training on data excluding  $(x_i, y_i)$ , i.e. at the price of high-computational demands.

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# Bagging

## Bootstrapping

- A general statistical technique for assessing the accuracy of parameter estimates and for hypotheses testing.
- It relies on *many repetitions* and *random sampling with replacement*.

Example: Assume we want to estimate the average height of all the people in the world. How to do that?

- Cannot measure the whole population, measure just a sample of *N* people.
- Using this sample, we can obtain a (single) point estimate of the average population height:  $\hat{h} = \frac{1}{N} \sum_{i=1}^{N} h_i$ .
- We also need some measure of uncertainty/variability of this estimate. How to do that?
- Use "classic" statistics: compute the sample variance  $\hat{s}_h^2$  and compute the variance of the estimate as  $\hat{s}_{\hat{t}}^2 = \frac{\hat{s}_h^2}{N}$ , or:
- Use *bootstrapping*:
  - 1. Repeat *M* times  $(M = 10^2, ..., 10^6)$ :
    - Create a bootstrap sample from the original dataset.
    - Compute *b*th estimate of the statistic (here average) from the bootstrap sample.
  - 2. Now you have a histogram of the estimates (here averages), from which you can estimate the mean, variance, ... of the sampling distribution.

Similar process works for many other estimators.

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## **Bootstrap sample**

Assume we have a dataset *T* with *N* items. What is the **bootstrap sample**  $T^b$ ?

- A pertubed version of the original dataset *T*.
- Each item of  $T^b$  was chosen uniformly with replacement from the original dataset T. Usually,  $|T| = N = |T^b|$ .
- Some items of *T* are copied to *T<sup>b</sup>* more than once. Some items are not copied at all.

How many unique elements of T are present in  $T^b$  (on average)?

- Probability that a particular item will not be chosen in one particular pick:  $1 \frac{1}{N}$
- Probability that a particular item will not be chosen in any of *N* picks:  $\left(1 \frac{1}{N}\right)^N$

The expected number of items that will not be copied to a bootstrap sample:  $N\left(1-\frac{1}{N}\right)^N \approx Ne^{-1} = N \cdot 0.368$ 

• The expected number of unique elements copied from  $T: N\left(1 - \left(1 - \frac{1}{N}\right)^N\right) \approx N\left(1 - e^{-1}\right) = N \cdot 0.632$ 

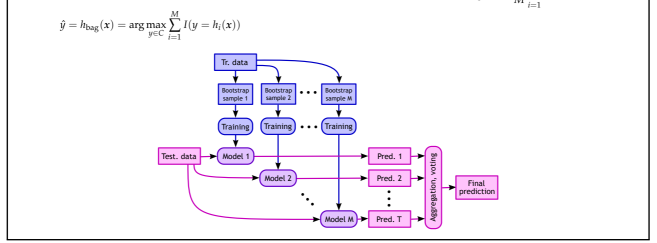
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## Bagging a.k.a. Bootstrap aggregation

- Uses bootstrap to improve the estimate or the prediction itself.
- Aggregating results of several models reduces variance and prevents overfitting.
- Algorithm:
  - 1. Create *M* bootstrap samples  $T^i$  from training data T (i = 1, ..., M).
  - 2. Build a model  $h_i$  on each bootstrap sample  $T^i$ .

3. Construct final model by averaging/voting the predictions of individual models:  $\hat{y} = h_{\text{bag}}(x) = \frac{1}{M} \sum_{i=1}^{M} h_i(x)$ , resp.



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## Features

Bagging

- leads to improvements for unstable procedures (artificial neural networks, classification and regression trees, etc.), but
- it can mildly degrade the performance of stable methods such as K-nearest neighbors.
- Thanks to bootstrapping, it can provide not only predictions, but also estimates of uncertainty of those predictions.

Estimate of prediction error (out-of-bag error):

- Around 37 % of training examples are not part of a bootstrap sample; they are called OOB (out of bag).
- We can predict the model response for each training sample  $x_i$  using only the models that did not have  $x_i$  in their bootstrap sample.
- We can average these predicted responses (regression) or can take a majority vote (classification) to get a single "OOB prediction" for the each observation.
- OOB predictions then can be used to compute OOB estimate of the error.
- With *M* sufficiently large, OOB error is virtually equivalent to leave-one-out cross-validation error.

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# **Random forests**

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## Random forest (RF)

An ensamble method using set of decision trees (i.e. forest):

- Trees that are grown very deep tend to learn highly irregular patterns: they overfit their training sets, i.e. have low bias, but very high variance.
- RF perform averaging of multiple deep decision trees, trained on different parts of the same training set, with the goal of reducing the variance.

RF combine

- bagging, and
- random subspace method (see below).

Predictions are computed using voting/averaging.

To train a single tree, RF algorithm

- creates a bootstrap sample of the training data (bagging), and
- uses a modified tree-learning algorithm which considers only *a random subset of input features* at each candidate split in the learning process ("feature bagging"; this further decorrelates the resulting trees). Suggestions:
  - Classification: consider  $\sqrt{D}$  features at each split.
  - Regression: consider *D*/3 features at each split, use minimum node size of 5.
- In *ExtraTrees* (extremely randomized trees), instead of searching for the locally optimal split for each variable, a random value is used for the split.

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## **RF** features

Estimate of prediction uncertainty and OOB error:

See bagging.

#### Variable importance:

- 1. Grow the forest. Compute OOB error for each data point averaged over the whole forest.
- 2. To measure the importance of *j*th variable, permute its values, and compute OOB error on this perturbed dataset. Compute the difference of the estimates before and after permutation.
- 3. The larger the difference, the larger the importance of variable *j*.

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# Boosting

# Boosting

Hypothesis Boosting Problem

If there exists an efficient algorithm able to create *weak classifiers* (i.e. classifiers only slightly better than random guessing), does it also mean that there is an efficient algorithm able to build *strong classifiers* (i.e. classifiers with an arbitrary precision)?

No constraint on the algorithm.

Most (not all) Boosting algorithms

- sequentially learn weak classifiers using weighted training set (using information from previous trees),
- construct the final strong classifier as a weighted sum of the weak classifiers,
- assign the weights to individual weak learners depending on their accuracy,
- re-weight the training data for another round of the weak learner,
- differ in the way how they weight the training data and/or the individual weak classifiers.

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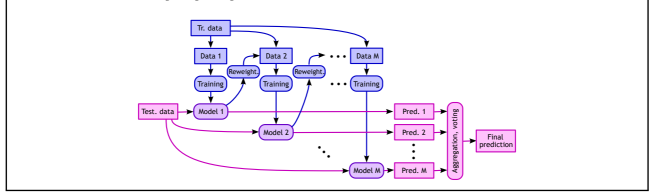
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# AdaBoost (informally)

## AdaBoost

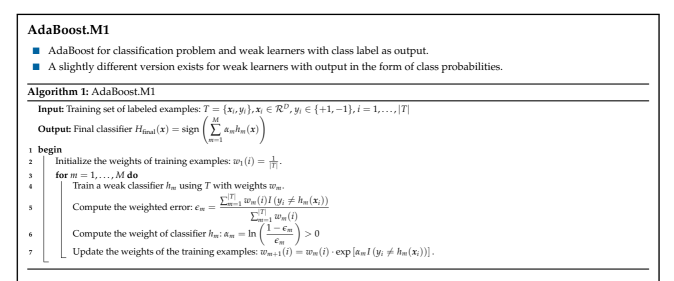
Training data:

- In each iteration t = 1, ..., M, it uses different weights  $w_t(i)$  of the training examples  $x_i$ .
- Misclassified examples get a larger weight for the next iteration.
- The resulting classifier:
  - Weighted voting.
  - More accurate models get larger weight.



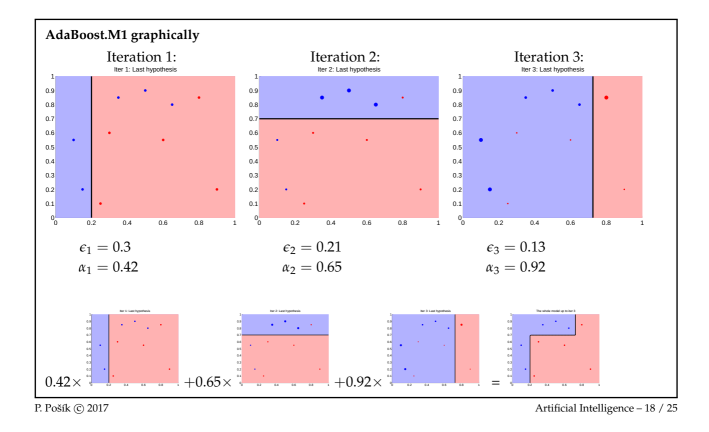
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# AdaBoost: remarks

The training error:

- Let  $\gamma_t = 0.5 \epsilon_t$  be the improvement of the *t*-th model over a random guess.
- Let  $\gamma = \min_t \gamma_t$  be the minimal improvement, i.e. the difference of error of all models  $h_t$  compared to the error of random guessing is at least  $\gamma$ , i.e.

 $\forall t: \gamma_t \geq \gamma > 0.$ 

It can be shown that the training error

 $\operatorname{Err}_{\operatorname{Tr}}(H_{\operatorname{final}}) \leq e^{-2\gamma^2 M}$ 

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## Forward stagewise additive modeling

Boosting tries to solve the following optimization problem:  $f^* = \arg \min_{f \in \mathcal{F}} \sum_{i=1}^{|T|} L(y_i, f(x_i))$  Finding the optimal  $f^*$  is hard; we shall

tackle it sequentially:

Algorithm 2: Forward stagewise additive modeling (FSAM)

1 begin Initialize  $f_0(x) = 0$ . 2 for  $m = 1, \ldots, M$  do 3 Compute  $(\alpha_m, \theta_m) = \arg \min_{\alpha, \theta} \sum_{i=1}^{|T|} L(y_i, f_{m-1}(\mathbf{x}_i) + \alpha h(\mathbf{x}_i; \theta)).$ 4 Set  $f_m(\mathbf{x}) = f_{m-1}(\mathbf{x}) + \alpha_m h(\mathbf{x}; \theta_m)$ 5

AdaBoost.M1 is equivalent to FSAM using the **exponential loss function**  $L(y, f(x)) = \exp(-y \cdot f(x))$ .

$$\begin{aligned} (\alpha_m, \theta_m) &= \arg\min_{\alpha, \theta} \sum_{i=1}^{|I|} \exp\left[-y_i \left(f_{m-1}(\mathbf{x}_i) + \alpha h(\mathbf{x}_i; \theta)\right)\right] \\ &= \arg\min_{\alpha, \theta} \sum_{i=1}^{|T|} w_m(i) \exp\left[-y_i \alpha h(\mathbf{x}_i; \theta)\right], \end{aligned}$$

where  $w_m(i) = \exp(-y_i f_{m-1}(x_i))$  depend neither on  $\alpha_m$  nor  $\theta_m$  and can be regarded as weights of training examples, which change each iteration. AdaBoost.M1 then follows from minimization of the last expression.

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#### L2Boosting

Suppose we need to solve regression problem with squared error loss (L2).

■ Then at step *m* we have:

$$\begin{split} L(y_i, f_m(\mathbf{x}_i)) &= L(y_i, f_{m-1}(\mathbf{x}_i) + \alpha_m h(\mathbf{x}_i; \theta_m)) = \\ &= (y_i - f_{m-1}(\mathbf{x}_i) - \alpha_m h(\mathbf{x}_i; \theta_m))^2 = \\ &= (r_{im} - \alpha_m h(\mathbf{x}_i; \theta_m))^2, \end{split}$$

where we define  $r_{im} = y_i - f_{m-1}(x_i)$  to be the current residual of the model for *i*th data point.

**B** By fitting each weak model  $h_m$  to the residuals  $r_{im}$ , the *m*th model  $f_m$  learns to correct its predecessor  $f_{m-1}$ .

- Observation: the residuals  $r_{im} = y_i f_{m-1}(x_i)$  are negative gradients of the squared error loss function  $\frac{1}{2}(y f(x))^2$ .
- The algorithm can be viewed as a gradient descent in the space of functions.
- The generalization of
  - FSAM using exponential loss (AdaBoost.M1) and
  - FSAM using L2 loss (L2Boosting)

for a general differentiable loss function L is called Gradient Boosting Machine.

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# Gradient Boosting Algorithms Algorithm 3: Gradient boosting Input: Training set of labeled examples: $T = \{x_i, y_i\}, i = 1, ..., |T|$ , a differentiable loss function L(y, f(x)), number of iterations M. Output: Final model $f_M(x)$ . 1 begin 2 Initialize model with constant value: $f_0(x) = \arg \min_{\gamma} \sum_{i=1}^{|T|} L(y_i, \gamma)$ 3 for m = 1, ..., M do 4 Compute pseudo-residuals $r_{im} = -\frac{\partial L(y_i, f(x_i))}{\partial f(x_i)} \Big|_{f(x) = f_{m-1}(x)}$ for all i = 1, ..., |T|.

5 Fit model  $h_m(\mathbf{x})$  to pseudo-residuals, i.e. use training set  $\{(\mathbf{x}_i, y_i)\}_{i=1}^{|T|}$ . 6 Compute multiplier  $\alpha_m$  by solving the following 1D opt. problem:  $\alpha_m = \arg\min_{\alpha} \sum_{k=1}^{|T|} L(y_i, f_{m-1}(\mathbf{x}_i) + \alpha h_m(\mathbf{x}_i))$ .

Update the model:  $f_m(\mathbf{x}) = f_{m-1}(\mathbf{x}) + \alpha_m h_m(\mathbf{x})$ 

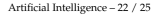
By plugging in different loss functions, we can construct different boosting variants like

AdaBoost,

- L2Boost,
- LogitBoost,
- etc.

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# **Further considerations**

Choosing the number of models *M*:

- The optimal value usually found by tracking the error on validation set.
- Often, we do not bother; we just set it sufficiently high (several hundreds). Boosting can overfit, but is quite resistant to it.

## Shrinkage:

• Often, the so-called shrinkage is applied, i.e. only a small part of the *m*th model is used:

 $f_m(\mathbf{x}) = f_{m-1}(\mathbf{x}) + \nu \alpha_m h(\mathbf{x}; \theta_m),$ 

where  $\nu \in (0, 1)$ , often  $\nu \approx 0.1$ , is the so-called *learning rate*.

Learning is slowed down; it requires more models to be added to the model, providing a configuration trade-off between the number of trees and learning rate.

## Stochastic gradient boosting

- It is possible to subsample the training data set and use only a subset of it to train each model.
- Subsample examples as in boosting (but without replacement).
- Subsample features as in random forests.
- It further *prevents overfitting*, *speeds up learning* of individual models, and gives chance to *compute out-of-bag error* estimates.

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## Summary

## Competencies

After this lecture, a student shall be able to ...

- describe the basic principle behind all committee/ensemble methods;
- list and conceptually compare several methods to achieve diversity among models trained on the same data, and know which
  of these methods are used in which ensemble algorithms;
- explain the purpose and the basic principle of stacking;
- explain how a bootstrap sample is created from the available data, and describe its properties;
- describe features of bagging;
- explain how to compute out-of-bag error estimate when using bagging;
- explain the principle of random forests and describe their difference to bagging with trees;
- explain how to compute a score of variable importance using random forest;
- explain the hypothesis boosting problem, and define a weak and a strong classifier in this context;
- explain the basic principle of AdaBoost.M1 algorithm;
- relate the training error of the AdaBoost algorithm to the number of constituent models and to the errors of individual models;
- describe the relations of AdaBoost.M1, L2Boost, and Gradient Boosting.

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