# **Statistical Machine Learning (BE4M33SSU) Lecture 12: Ensembling**

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# **Overview**

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Topics covered in the lecture:

- Ensemble Methods
- ◆ Bias-Variance Decomposition
- $\blacklozenge$ Bagging
- $\blacklozenge$ Random Forests
- $\blacklozenge$ Boosting and Gradient Boosting
- $\blacklozenge$ Gradient Boosted Trees

#### **Ensemble Methods**

♦ Inspired in Wisdom of the crowd

- (weighted) averaging or taking majority vote
- cancelling effect of noise of individual opinions,
- examples: politics, trial by jury (vs. trial by judge), sports (figure skating, gymnastics), Wikipedia, Quora, Stack Overflow, ...
- $\blacklozenge$ Learning and aggregating multiple predictors
- $\blacklozenge$ Ensemble may be built using single or different types of predictors



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# **Ensembling Approaches**

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- ♦ Bagging (Bootstrap AGGregatING):
	- sample different training sets from the original training set
	- train high variance low bias predictors based on these sets and average them
	- exploits independence between predictors
- $\blacklozenge$ Boosting:
	- sequentially train low variance high bias predictors
	- subsequent predictors learn to fix the mistakes of the previous ones
	- exploits dependence between learners

## **Stacking and Mixture of Experts**







<https://www.commonlounge.com/discussion/9331c0d004704e89bd4d1da08fd7c7bc>

# **Prediction Problem: Expected Risk and Error Decomposition**

Expected risk for data generated by *p*(*x, y*):

$$
R(h) = \mathbb{E}_{(x,y)\sim p} \Big[ \ell(y, h(x)) \Big]
$$

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- The best attainable (Bayes) risk is  $R^* = \inf_{h \in \mathcal{Y}^{\mathcal{X}}} R(h)$
- $\blacklozenge$ The best predictor in  $\mathcal H$  is  $h_{\mathcal H} \in \mathop{\rm Argmin}_{h \in \mathcal H} R(h)$
- $\blacklozenge$ The predictor  $h_m = A(\mathcal{T}^m)$  learned from  $\mathcal{T}^m$  has risk  $R(h_m)$

**Excess error** measures deviation of the learned predictor from the best one:

$$
\underbrace{\left(R(h_m) - R^*\right)}_{\text{excess error}} = \underbrace{\left(R(h_m) - R(h_{\mathcal{H}})\right)}_{\text{estimation error}} + \underbrace{\left(R(h_{\mathcal{H}}) - R^*\right)}_{\text{approximation error}}
$$

#### **Risk Averaged over Datasets**

- ♦ How will our predictor behave when sampling different training sets?
- $\blacklozenge$  We can define the errors considering average over models constructed using all possible datasets  $\mathcal{T}^m$ , i.e.,  $\mathbb{E}_{\mathcal{T}^m} \big[$  $R(h_m)$  $\overline{\phantom{a}}$

♦ The errors can be redefined as:

$$
\underbrace{\left(\mathbb{E}_{\mathcal{T}^m}\left[R(h_m)\right]-R^*\right)}_{\text{excess error}}=\underbrace{\left(\mathbb{E}_{\mathcal{T}^m}\left[R(h_m)\right]-R(h_{\mathcal{H}})\right)}_{\text{estimation error}}+\underbrace{\left(R(h_{\mathcal{H}})-R^*\right)}_{\text{approximation error}}
$$



#### **Predictors Averaged over Datasets**



♦ Let us also define a model averaged over all possible datasets:

$$
g_m(x) = \mathbb{E}_{\mathcal{T}^m} \Big[ h_m(x) \Big]
$$

♦ Unlike individual  $h_m$  models,  $g_m$  has an access to the whole  $p(x,y)$ 

- ♦ Note: in general  $g_m \neq h_H$  due to training algorithm *A* involved in  $h_m$ .
- ♦ Also:  $g_m$  can't be actually evaluated for infinite number of  $\mathcal{T}^m$  datasets

#### **Bias-Variance Decomposition for Regression**

♦ Consider a regression problem with data generated as follows:

$$
y = h^*(x) + \epsilon
$$

where  $\epsilon$  is noise:  $\mathbb{E}[\epsilon] = 0$  and  $\text{Var}(\epsilon) = \sigma^2$ , e.g.,  $\epsilon \sim \mathcal{N}(0, \sigma^2)$ 

Use squared loss:

$$
\ell(y, h(x)) = (h(x) - y)^2
$$

♦ The optimal predictor  $h^*(x)$  has a nonzero risk (for  $\sigma^2 > 0$ ):

$$
R^* = \mathbb{E}_{x,y}\left[\left(h^*(x) - y\right)^2\right] = \mathbb{E}_{\epsilon}\left[\epsilon^2\right] = \text{Var}(\epsilon) = \sigma^2
$$



#### **Bias-Variance Decomposition for Regression 2**

♦ The expected risk for *h<sup>m</sup>* can be decomposed:

$$
R(h_m) = \mathbb{E}_{x,y,\mathcal{T}^m} \left[ \left( h_m(x) - y \right)^2 \right]
$$
  
= ...  
= 
$$
\mathbb{E}_{x,\mathcal{T}^m} \left[ \left( h_m(x) - g_m(x) \right)^2 \right] +
$$
  
variance
$$
+ \mathbb{E}_x \left[ \left( g_m(x) - h^*(x) \right)^2 \right] + \frac{\sigma^2}{\text{noise}}
$$

The error splits into three terms

 $\mathbb{E}_{\mathcal{T}^m} \Big[$ 

- variance: difference of  $h_m$  from the averaged predictor  $g_m$ ,
- $\bullet$  bias $^2$ : difference of the averaged predictor  $g_m$  from the optimal one,
- **noise**: irreducible determined by data



#### **Excess Error vs. Bias and Variance**

The excess error is defined as:

$$
\mathbb{E}_{\mathcal{T}^m}\Big[R(h_m)\Big]-R^*
$$

 $\blacklozenge$  As  $R^* = \sigma^2$  we get:

$$
\mathbb{E}_{\mathcal{T}^m} \Big[ R(h_m) \Big] - R^* = \mathbb{E}_x \Big[ \Big( g_m(x) - h^*(x) \Big)^2 \Big] + \mathbb{E}_{x, \mathcal{T}^m} \Big[ \Big( h_m(x) - g_m(x) \Big)^2 \Big]
$$
  
variance

 $\blacklozenge$ Compare

- **bias**<sup>2</sup> vs. approximation error,
- **variance** vs. estimation error
- averaged model  $g_m$  vs. best predictor  $h_{\mathcal{H}}$

$$
\begin{array}{c}\n\textcircled{\color{blue}0}\text{m p} \\
\hline\n\end{array}
$$

#### **Derivation of the Bias-Variance Decomposition**







#### **Derivation of the Bias-Variance Decomposition 2**

We get:

$$
\mathbb{E}_{\mathcal{T}^m} \Big[ R(h_m) \Big] = \mathbb{E}_{x, \mathcal{T}^m} \Big[ \Big( h_m(x) - g_m(x) \Big)^2 \Big] + \mathbb{E}_{x, y} \Big[ \Big( g_m(x) - y \Big)^2 \Big]
$$
  
= 
$$
\text{Var}_{x, \mathcal{T}^m} \Big( h_m(x) \Big) + \mathbb{E}_{x, y} \Big[ \Big( g_m(x) - y \Big)^2 \Big]
$$

Note that the second term does not depend on  $\mathcal{T}^m$ .



Let us continue with the second term:

$$
\mathbb{E}_{x,y}\left[\left(g_m(x) - y\right)^2\right] = \mathbb{E}_{x,\epsilon}\left[\left(g_m(x) - h^*(x) - \epsilon\right)^2\right]
$$

$$
= \mathbb{E}_{x,\epsilon}\left[\left(g_m(x) - h^*(x)\right)^2 + \epsilon^2 - 2\epsilon\left(g_m(x) - h^*(x)\right)\right]
$$

$$
= \mathbb{E}_x\left[\left(g_m(x) - h^*(x)\right)^2\right] + \mathbb{E}_\epsilon\left[\epsilon^2\right]
$$

$$
-2\mathbb{E}_{x,\epsilon}\left[\epsilon\left(g_m(x) - h^*(x)\right)\right]
$$

$$
= \underbrace{\mathbb{E}_x\left[\left(g_m(x) - h^*(x)\right)^2\right]}_{\text{bias}^2} + \underbrace{\sigma^2}_{\text{noise}}
$$

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#### **Pointwise Bias-Variance**



We can express the bias and variance as function of *x* by not integrating over in expected values

$$
\mathbb{E}_{y|x,\mathcal{T}^m} \left[ \ell(y, h_m(x)) \right] = \mathbb{E}_{y|x,\mathcal{T}^m} \left[ \left( h_m(x) - y \right)^2 \right]
$$

$$
= \underbrace{\text{Var}_{\mathcal{T}^m} \left( h_m(x) \right)}_{\text{variance(x)}} + \underbrace{\left( g_m(x) - h^*(x) \right)^2}_{\text{bias(x)}^2} + \underbrace{\sigma(x)^2}_{\text{noise}}
$$

#### **Bias-Variance: Example**

♦ Polynomial regression with a varying degree of polynomial



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<span id="page-15-0"></span>Figure 4.2: Bias-variance decomposition of the expected generalization error Gilles Louppe: Understanding Random Forests: From Theory to Practice, 2014

# **Decision/Regression Trees**



- $\overline{\mathbf{S}}$ .  $\overline{\mathbf{S}}$ original training data as well as mutually exclusive subsets of the input Nodes at the same level correspond to mutually exclusive subsets of the  $\blacklozenge$ space  $\mathcal{X}$ 
	- $\blacklozenge$ Inner node further splits its subset



<span id="page-16-1"></span><span id="page-16-0"></span>Hastie et al.: The Elements of Statistical Learning, 2009

#### **Decision/Regression Trees (contd.)**

- $\blacktriangledown$  Training set:  $\mathcal{T}^m = \{(\bm{x}_i, y_i) | i = 1, \ldots, m\}$ ,  $\bm{x}_i = (x_{i1}, x_{i2}, \ldots, x_{ip})$
- ♦ Input space split into regions defined in leaves:  $R_r$ ,  $r \in \{1, \ldots, M\}$
- We can model *region responses* by constants  $c_r, \ r \in \{1, \ldots, M\}$  but other possibilities, e.g., linear regression are possible
- ♦ Prediction:

$$
h(\boldsymbol{x}) = \sum_{r=1}^M c_r[\boldsymbol{x} \in R_r]
$$

 $\blacklozenge$  For sum of squares *loss function*  $\sum_{i=1}^{m}(y_i-h(\boldsymbol{x}_i))^2$  we set the responses to be the averages over regions:

$$
\hat{c}_r = \frac{1}{|S_r|} \sum_{(\boldsymbol{x}_i, y_i) \in S_r} y_i \qquad \text{(see seminar)}
$$

 $\mathbf{w}$ here  $S_r = \{(\boldsymbol{x}_i, y_i): (\boldsymbol{x}_i, y_i) \in \mathcal{T}^m \land \boldsymbol{x}_i \in R_r\}$ 



# **Greedy Learning of Decision/Regression Trees**

- ♦ How many distinct decision trees with *p* Boolean attributes for binary classification?
	- at least as many as boolean functions of *p* attributes
	- $\bullet$  = number of distinct truth tables with  $2^p$  rows:  $2$ 2*p*
	- For 6 Boolean attributes at least 18,446,744,073,709,551,616 trees!
- ◆ Learning is NP-complete: [Hyafil and Rivest 1976]
- $\blacklozenge$ We need heuristics ⇒ **greedy approach**
- $\blacklozenge$ Recursively choose the "most important" attribute to find a small tree consistent with the training data
- $\blacklozenge$ Split points:
	- **nominal attribute**: try all possibilities
	- **ordinal/continuous attribute**: try attribute values based on all training data samples or their subset



#### **Regression Trees: Which Attribute to Split?**

- The "most important" attribute for regression trees would be the one, for which the split reduces the loss (sum of squared errors) by the greatest amount
- $\blacklozenge$ We have:

$$
h(\boldsymbol{x}) = \sum_{r=1}^M c_r[\boldsymbol{x} \in R_r]
$$

 $\blacklozenge$ Consider splitting attribute *j* and split point *s*, we split an original region *R* into a pair of half-planes for an ordinal (e.g., continuous) attribute:

$$
R_L(j,s) = \{ \boldsymbol{x} | \boldsymbol{x} \in R \land x_j \leq s \} \text{ and } R_R(j,s) = \{ \boldsymbol{x} | \boldsymbol{x} \in R \land x_j > s \}
$$

similarly for a nominal attribute:

$$
R_L(j,s) = \{ \boldsymbol{x} | \boldsymbol{x} \in R \land x_j = s \} \text{ and } R_R(j,s) = \{ \boldsymbol{x} | \boldsymbol{x} \in R \land x_j \neq s \}
$$

 $\blacklozenge$ Denote the corresponding subsets of of  $\mathcal{T}^m$  as  $S_L$  and  $S_R$ 



## **Regression Trees: Which Attribute to Split? (contd.)**



$$
\min_{c_L} \sum_{(\bm{x}_i, y_i) \in S_L(j,s)} (y_i - c_L)^2 + \min_{c_R} \sum_{(\bm{x}_i, y_i) \in S_R(j,s)} (y_i - c_R)^2
$$

for  $(\boldsymbol{x}_i, y_i) \in S$  and  $S = S_L \cup S_R$ 

Inner minimizations (region response values) are solved by averaging tree outputs per region:

$$
\hat{c}_L = \frac{1}{|S_L(j,s)|} \sum_{(\bm{x}_i, y_i) \in S_L(j,s)} y_i \text{ and } \hat{c}_R = \frac{1}{|S_R(j,s)|} \sum_{(\bm{x}_i, y_i) \in S_R(j,s)} y_i
$$

 $\blacklozenge$ Root node:  $S = \mathcal{T}^m$ 



# **Tree Learning Algorithm**



```
BUILD-TREE(S)
 i = \text{IMPUTY}(S) // e.g., the squared loss
 2 \hat{i}, \hat{j}, \hat{s}, \hat{S}_L, \hat{S}_R = 0, 0, 0, \emptyset, \emptyset // current best kept in these
 3 for j \in \{1, \ldots, p\} // iterate over attributes
 4 for s \in \text{SPLIT-POINTS}(S, j) // iterate over all split points
 5 S_L, S_R = \text{SPLIT}(S, j, s)6 i_L = \text{IMPUTY}(S_L)7 i_R = \text{IMPUTY}(S_R)8 if i_L + i_R < \hat{i} and |S_L| > 0 and |S_R| > 0<br>9 \hat{i}, \hat{j}, \hat{s}, \hat{S}_L, \hat{S}_R = (i_L + i_R), j, s, S_L, S_R(\hat{i}, \hat{j}, \hat{s}, \hat{S}_L, \hat{S}_R = (i_L + i_R), j, s, S_L, S_R10 if \hat{i} < i11 N_L = \text{BULD-TREE}(\hat{S}_L)12 N_R = \text{BULD-TREE}(\hat{S}_R)13 return DECISION-NODE(\hat{j}, \hat{s}, N_L, N_R)14 else return LEAF-NODE(S)
```
- 
- 
- 
- 

### **Bias and Variance of Decision Trees**



- ◆ Small changes of training data lead to big differences in final trees
- ♦ Decision trees grown deep enough have typically:
	- low bias
	- high variance
	- ⇒ **overfitting**
- $\blacklozenge$ Idea: average multiple models to reduce variance while (happily) not increasing bias much

#### **Averaging Models**



♦ Define bagging model *b* as an average of *K* component models:

$$
b(x) = \frac{1}{K} \sum_{i=1}^{K} h_m^{(i)}(x)
$$

trained using a set of i.i.d. datasets of size  $m\colon\mathcal{D}^{m}=\{\mathcal{T}_{1}^{m},\ldots,\mathcal{T}_{K}^{m}\}$ so  $h_{m}^{(1)}(x)$  is trained using  $\mathcal{T}_{1}^{m}$  $\mathcal{T}^m_1$ ,  $h^{(2)}_m(x)$  using  $\mathcal{T}^m_2$  $2^m$ , etc.

 $\blacklozenge$ Note that  $b(x)$  approximates the *averaging model*:

$$
g_m(x) = \mathbb{E}_{\mathcal{T}^m} \Big[ h_m(x) \Big]
$$

 $\blacklozenge$  We can define the *averaging model* for  $b(x)$  as well:

$$
g_m^B(x) = \mathbb{E}_{\mathcal{D}^m} \left[ b(x) \right]
$$

#### **Averaging Models: Bias**



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bias
$$
x^2 = (g_m^B(x) - h^*(x))^2
$$
  
\n
$$
= \left(\mathbb{E}_{\mathcal{D}^m} \left[b(x)\right] - h^*(x)\right)^2
$$
  
\n
$$
= \left(\mathbb{E}_{\mathcal{D}^m} \left[\frac{1}{K} \sum_{i=1}^K h_m^{(i)}(x)\right] - h^*(x)\right)^2
$$
  
\n
$$
= \left(\frac{1}{K} \sum_{i=1}^K \mathbb{E}_{\mathcal{T}_i^m} \left[h_m^{(i)}(x)\right] - h^*(x)\right)^2
$$
  
\n
$$
= \left(\mathbb{E}_{\mathcal{T}^m} \left[h_m(x)\right] - h^*(x)\right)^2 = \left(g_m(x) - h^*(x)\right)^2
$$

#### **Averaging Models: Variance**



 $\blacklozenge$  For uncorrelated component models  $h_m^{(i)}(x)$ :

$$
\operatorname{Var}_{\mathcal{D}^m}\left(b(x)\right) = \operatorname{Var}_{\mathcal{D}^m}\left(\frac{1}{K} \sum_{i=1}^K h_m^{(i)}(x)\right)
$$

$$
= \frac{1}{K^2} \sum_{i=1}^K \operatorname{Var}_{\mathcal{T}_i^m}\left(h_m^{(i)}(x)\right) = \frac{1}{K} \operatorname{Var}_{\mathcal{T}^m}\left(h_m(x)\right)
$$

which is a great improvement based on the very **strong** assumption **There is no improvement for maximum correlation, i.e., for all** component models equal:  $h_m^{(i)}(x) = h_m(x)$  for  $i = 1, \ldots, K$ , we get:

$$
\text{Var}_{\mathcal{D}^m}\Big(b(x)\Big) = \text{Var}_{\mathcal{D}^m}\left(\frac{1}{K}\sum_{i=1}^K h_m^{(i)}(x)\right) = \text{Var}_{\mathcal{T}^m}\Big(h_m(x)\Big)
$$

⇒ we need to train **uncorrelated** (diverse) component models while **keeping their bias reasonably low**

# **Bootstrapping**



- $\blacklozenge$  In practice we have only a single training dataset  $\mathcal{T}^m$
- ♦ Bootstrapping is a method producing datasets  $\mathcal{T}^m_i$  $\tilde{I}_i^m$  for  $i=1,\ldots K$  by sampling  $\mathcal{T}^m$  uniformly with *replacement*
- ◆ Bootstrap datasets have the same size as the original dataset  $|\mathcal{T}^m_i| = |\mathcal{T}^m|$
- $\blacklozenge$   $\mathcal{T}^m_i$  $\tilde{t}^m_i$  is expected to have the fraction  $1-\frac{1}{e}$  $\frac{1}{e} \approx 63.2\%$  of unique samples from  $\mathcal{T}^m$ , others are duplicates (see seminar)

# **Bagging**



- ♦  $Bagging = Bootstrap AGGregating [Breiman 1994]:$ 
	- 1. Use bootstrapping to generate *K* datasets
	- 2. Train a model  $h^{(i)}_m(x)$  on each dataset  $\mathcal{T}^m_i$ *i*
	- 3. Average the models getting the bagging model *b*(*x*)
- $\blacklozenge$ When decision trees are used as the models ⇒ **random forests**
- $\blacklozenge$ Low bias is achieved by growing the trees to maximal depth
- Trees are decorrelated by:
	- training each tree on a different bootstrap dataset
	- randomization of split attribute selection

# **Random Forest Algorithm**



- 1. For  $i = 1...K$ :
	- (a) draw a bootstrap dataset  $\mathcal{T}^m_i$  $\mathcal{T}^m_i$  from  $\mathcal{T}^m$ ,  $|\mathcal{T}^m_i| = |\mathcal{T}^m| = m$
	- (b) grow a tree  $h_m^{(i)}$  using  $\mathcal{T}_i^m$  $\tilde{C}^m_i$  by recursively repeating the following, until the minimum node size  $n_{\text{min}}$  is reached:
		- i. select *k* attributes at random from the *p* attributes
		- ii. pick the best attribute and split-point among the *k*
		- iii. split the node into two daughter nodes
- 2. Output ensemble of trees  $b(x)$  averaging  $h^{(i)}_m(x)$  (regression) or selecting a majority vote (classification)
	- $\blacklozenge$ Node size  $n_{\sf min}$  is the number of the training dataset samples associated with the node, limits tree depth

# **Out-of-Bag (OOB) Error**



- ♦ Cheap way of generalization error assessment for bagging
- ♦ Bagging produces bootstrapped sets T *m*  $\mathcal{T}^m_1, \mathcal{T}^m_2$  $\tau_2^m, \ldots \mathcal{T}_K^m$ *K*
- For each  $(\boldsymbol{x}_i, y_i) \in \mathcal{T}^m$  select only trees which were not trained on this sample:  $H_i = \{h^{(j)}_m \mid (\boldsymbol{x}_i, y_i) \notin \mathcal{T}^m_i\}$ *j* }
- Average only the OOB trees in  $H_i$  when evaluating error for  $(\boldsymbol{x}_i, y_i)$
- $\blacklozenge$ Replacement for K-fold cross-validation

#### **Feature Importance**

- ♦ Random forests allow easy evaluation of feature importances
- ♦ Mean Decrease Impurity (MDI):
	- set  $f_i = 0$  for all attributes  $j = 1, \ldots, p$
	- traverse all trees processing all internal nodes
	- for each node having a split attribute *j* add its impurity decrease multiplied by the proportion of the node size to *f<sup>j</sup>*

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- $\blacklozenge$ Mean Decrease Accuracy (MDA), permutaion importance:
	- evaluate the forest using OOB
	- do the same with permuted values of an attribute *j*
	- watch decrease in accuracy: low decrease means unimportant feature

## **Random Forest Summary**



- ♦ Easy to use method: robust w.r.t. parameter settings (*K*, node size)
- ♦ While *statistical consistency* is proven for decision trees (both regression and classification) we have only proofs for simplified versions of random forests [Breiman, 1984]
- ♦ Related methods: boosted trees

# **Boosting**

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- Sequentially train weak learners/predictors low variance high bias
- ♦ Subsequent predictors fix the mistakes of the previous ones reducing bias
- ♦ Methods discussed here:
	- Forward Stagewise Additive Modeling
	- Gradient Boosting Machine
	- Gradient Boosted Trees
	- AdaBoost



### **Forward Stagewise Additive Modeling (FSAM)**

1. Initialize  $f_0(x) = 0$ 

2. For  $k=1$  to  $K$ :

(a) Find

$$
(\beta_k, \theta_k) = \underset{\beta, \theta}{\text{argmin}} \sum_{i=1}^m \ell\Big(y_i, f_{k-1}(x_i) + \beta b(x_i; \theta)\Big)
$$

 $\mathsf{where} \,\, b(x_i; \theta_k)$  is the *basis function* and  $\beta_k$  the corresponding coefficient (b) Set  $f_k(x) = f_{k-1}(x) + \beta_k b(x; \theta_k)$ 

3. Return  $h_m(x) = f_K(x)$ 

# **FSAM and Gradient Descent**



FSAM update looks very similar to the gradient descent one:

$$
f_k(x) = f_{k-1}(x) + \beta_k b(x; \theta_k)
$$

- ♦ Just think of
	- $\beta_k \approx$  step size (learning rate)
	- $\bullet \ \ b(x_i; \theta_k) \approx \text{the negative of gradient}$

# **FSAM for Squared Loss**

♦ Once again, consider regression with the squared loss:

$$
\ell(y, f(x)) = (y - f(x))^2
$$

♦ For FSAM we get:

$$
\ell(y_i, f_k(x_i)) = \ell(y_i, f_{k-1}(x_i) + \beta_k b(x_i; \theta_k))
$$
  
=  $(y_i - f_{k-1}(x_i) - \beta_k b(x_i; \theta_k))^2$   
=  $(r_{ik} - \beta_k b(x_i; \theta_k))^2$ 

where  $r_{ik} = y_i - f_{k-1}(x_i)$  is the *residual* of the current model for the *i*-th sample

- $\blacklozenge$ The task of FSAM is to fit the model  $\beta_k b(x_i; \theta_k)$  to match the residuals
- $\blacklozenge$ The method is sometimes called the *least-squares boosting*



### **Gradient Boosting for Regression**

 $\blacklozenge$ In case of regression with squared loss we minimize:

$$
\mathcal{L} = \sum_{i=1}^{m} \ell(y_i, f(x_i)) = \sum_{i=1}^{m} \frac{1}{2} (y_i - f(x_i))^2,
$$

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which is same as minimization of the empirical risk

 We can treat  $f(x_1), f(x_2), \ldots, f(x_m)$  as parameters and take derivatives:

$$
\frac{\partial \mathcal{L}}{\partial f(x_i)} = \frac{\partial \left(\sum_{j=1}^m \ell(y_j, f(x_j))\right)}{\partial f(x_i)} = \frac{\partial \ell(y_i, f(x_i))}{\partial f(x_i)}
$$

$$
= f(x_i) - y_i = -r_i
$$

The least-squares boosting hence takes steps in the negative gradient direction where  $r_i = -\frac{\partial \mathcal{L}}{\partial f(x_i)}$ *∂f*(*xi*)

 $\blacklozenge$ This approach can be generalized for any differentiable loss function!

# **Gradient Boosting Machine**



2. For  $k = 1$  to  $K$ :

(a) Compute:

$$
\boldsymbol{g}_k = \left[\frac{\partial \ell(y_i, f_{k-1}(x_i))}{\partial f_{k-1}(x_i)}\right]_{i=1}^m
$$

(b) Fit a regression model *b*(·; *θ*) to −*g<sup>k</sup>* using squared loss:

$$
\theta_k = \operatorname*{argmin}_{\theta} \sum_{i=1}^{m} [(-\boldsymbol{g}_k)_i - b(x_i; \theta)]^2
$$

(c) Choose a fixed step size  $\beta_k = \beta > 0$  or use line search:

$$
\beta_k = \operatorname*{argmin}_{\beta > 0} \sum_{i=1}^m \ell\Big(y_i, f_{k-1}(x_i) + \beta b(x_i; \theta_k)\Big)
$$

(d) Set  $f_k(x) = f_{k-1}(x) + \beta_k b(x; \theta_k)$ 

3. Return  $h_m(x) = f_K(x)$ 



#### **Multinominal Classification: Gradient Boosting Machine**

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- Train one GBM per target class
- ♦ Use softmax to get probability distribution
- ♦ Use multinominal cross-entropy as the loss

#### **Gradient Boosted Trees**



- ♦ Gradient Boosting Tree is GBM where all weak learners are decision or regression trees
- ♦ Use limit on depth/number of leaves/node size for the weak learners  $\Rightarrow$ high bias
- ♦ Often single-level tree: decision stump
- ♦ Meta-parameters such as *K* (number of trees) and *β* (learning rate) have to be found using cross validation
- $\blacklozenge$ Model is built sequentially (unlike random forests)
- $\blacklozenge$ Highly optimized algorithms based on Gradient Boosting Trees:
	- XGBoost, LightGBM
	- parallelization, scalability, regularization

#### **AdaBoost M1**

Binary classifier:  $\mathcal{Y} = \{-1, 1\}$ 

- 1. Initialize the weights  $w_i = 1/m$  for  $i = 1, 2, \ldots m$
- 2. For  $k = 1$  to  $K$ :

(a) Fit a classifier  $f_k(x; \theta_k)$  to the training data using loss weighted by  $w_i$ :

$$
\theta_k = \operatorname*{argmin}_{\theta} \sum_{i=1}^{m} w_i [y_i \neq f_k(x_i; \theta)]
$$

(b) Compute the weighted error rate

$$
\epsilon_k = \frac{\sum_{i=1}^m w_i [y_i \neq f_k(x_i; \theta_k)]}{\sum_{i=1}^m w_i}
$$

(c) Compute the scaling coefficient  $\alpha_k = \log((1 - \epsilon_k)/\epsilon_k)$ (d) Set  $w_i \leftarrow w_i \cdot \exp(\alpha_k \cdot [y_i \neq f_k(x_i; \theta_k)])$  for  $i = 1, 2, \ldots m$ 

3. Return 
$$
h_m(x) = \text{sign}\left[\sum_{k=1}^K \alpha_k f_k(x; \theta_k)\right]
$$



#### **AdaBoost is FSAM: the Loss**

♦ Claim: AdaBoost is FSAM using the exponential loss

> $-1.0$   $-0.5$  0.0 0.5 1.0  $yf(x)$  $\Omega$  $(\beta_k, \theta_k) = \text{argmin}$ *β,θ*  $\sum$ *m i*=1  $\ell$  $\sqrt{ }$  $y_i, f_{k-1}(x_i) + \beta b(x_i; \theta)$  $\setminus$  $=$  argmin *β,θ*  $\sum$ *m i*=1  $\exp\Big(-y_i$  $\sqrt{ }$  $f_{k-1}(x_i) + \beta b(x_i; \theta)$  $\bigwedge$  $=$  argmin *β,θ*  $\sum$ *m i*=1  $w_i^{(k)}$  $\sum_{i}^{(k)} \exp\left(-y_i \beta b(x_i; \theta)\right)$  $\setminus$ *,*

 $\ell(y, f(x)) = \exp(-yf(x))$ 

 $w_i^{(k)} \triangleq \exp(-y_if_{k-1}(x_i))$ 

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0/1 loss hinge

[exponential](#page-51-0)

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2

3



#### **AdaBoost is FSAM II: Fitting the Classifier**



$$
(\beta_k, \theta_k) = \underset{\beta, \theta}{\operatorname{argmin}} \sum_{i=1}^m w_i^{(k)} \exp\left(-y_i \beta b(x_i; \theta)\right)
$$
  
= 
$$
\underset{\beta, \theta}{\operatorname{argmin}} \left[e^{-\beta} \sum_{y_i = b(x_i; \theta)} w_i^{(k)} + e^{\beta} \sum_{y_i \neq b(x_i; \theta)} w_i^{(k)}\right]
$$
  
= 
$$
\underset{\beta, \theta}{\operatorname{argmin}} \left[e^{-\beta} \sum_{i=1}^m w_i^{(k)} + \underbrace{(e^{\beta} - e^{-\beta})}_{>0 \text{ for } \beta > 0} \sum_{i=1}^m w_i^{(k)} [y_i \neq b(x_i; \theta)]\right]
$$

For any *β >* 0 we can minimize *θ* separately:

$$
\theta_k = \operatorname*{argmin}_{\theta} \sum_{i=1}^{m} w_i^{(k)} [y_i \neq b(x_i; \theta)] \quad \text{(same as AdaBoost 2(a))}
$$







◆ Let's minimize

$$
(e^{\beta} - e^{-\beta}) \sum_{i=1}^{m} w_i^{(k)} [y_i \neq b(x_i; \theta_k)] + e^{-\beta} \sum_{i=1}^{m} w_i^{(k)}
$$

with respect to *β*

 $\blacklozenge$ 

$$
(e^{\beta_k} + e^{-\beta_k}) \sum_{i=1}^{m} w_i^{(k)} [y_i \neq b(x_i; \theta_k)] - e^{-\beta_k} \sum_{i=1}^{m} w_i^{(k)} = 0
$$

$$
(e^{\beta_k} + e^{-\beta_k}) \epsilon_k - e^{-\beta_k} = 0
$$

where 
$$
\epsilon_k = \frac{\sum_{i=1}^m w_i [y_i \neq b(x_i; \theta_k)]}{\sum_{i=1}^m w_i}
$$
 as in *Adaboot* 2(b)  
Solving for  $\beta_k$ :

$$
\beta_k = \frac{1}{2} \log \frac{1 - \epsilon_k}{\epsilon_k}
$$

 $\blacklozenge$ Define  $\alpha_k \triangleq 2\beta_k$  and compare to AdaBoost 2(c)

#### **AdaBoost is FSAM IV: the Weight Update**

\n- We have 
$$
w_i^{(k)} = e^{-y_i f_{k-1}(x_i)}
$$
 and  $f_k(x) = f_{k-1}(x) + \beta_k b(x; \theta_k)$  so:
\n- $w_i^{(k+1)} = e^{-y_i \left(f_{k-1}(x_i) + \beta_k b(x_i; \theta_k)\right)} = w_i^{(k)} \cdot e^{-y_i \beta_k b(x_i; \theta_k)}$
\n- Finally  $-y_i b(x_i; \theta_k) = 2 \cdot [y_i \neq b(x_i; \theta_k)] - 1$  gives the weight update:
\n- $w_i^{(k+1)} = w_i^{(k)} \cdot e^{\alpha_k \left[y_i \neq b(x_i; \theta_k)\right]} \cdot e^{-\beta_k}$
\n

 $\epsilon$  corresponding to *AdaBoost 2(d)* up to the factor  $e^{-\beta_k}$  which is same for all weights and hence has no effect



















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