# Trees, Bagging, and Boosting

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April 14, 2019

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## Today's lecture

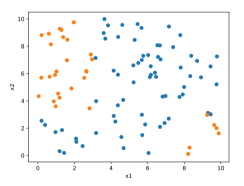
- Our first inherently non-linear classifier: decision trees.
- Ensemble methods: bagging and boosting.

### **Decision Trees**

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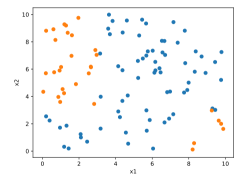
# Motivating example in 2d

• Partition data into different (axis-aligned) regions recursively



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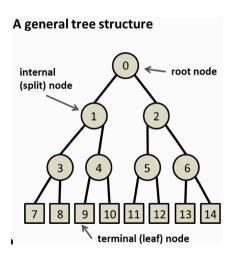
#### Classification flowchart



Is this a linear or non-linear classifier?

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#### Decision trees setup



#### We'll only consider

- binary trees (vs multiway trees where nodes can have more than 2 children)
- each node contains a subset of data points
- decisions at each node involve only a single feature (i.e. input coordinate)
- for continuous variables, splits always of the form

$$x_i \leqslant t$$

 for discrete variables, partitions values into two groups

From Criminisi et al. MSR-TR-2011-114, 28 October 2011.

# Regularization of decision trees

- What will happen if we keep splitting the data?
  - Every data point will be in its own region—overfitting.
- When to stop splitting? (control complexity of the hypothesis space)
  - Limit number of total nodes.
  - Limit number of terminal nodes.
  - Limit tree depth.
  - Require minimum number of data points in a terminal node.
  - Backward pruning the approach of CART (Breiman et al 1984):
    - **Q** Build a really big tree (e.g. until all regions have  $\leq 5$  points).
    - Prune the tree back greedily all the way to the root, assessing performance on validation.

Goal Find a tree that minimize the task loss (e.g., squared loss) within a given complexity.

Problem Finding the optimal binary tree is computationally intractable.

Solution Greedy algorithm.

- Find the best split (according to some criteria) for a non-terminal node (initially the root)
- Add two children nodes
- Repeat until a stopping criterion is reached (e.g., max depth)

### Evaluate splits

Let's think about what makes a good split.

Which one is better?

Split 1 
$$R_1:8+/2 R2:2+/8-$$
  
Split 2  $R_1:6+/4 R2:1+/9-$ 

Which one is better?

Split 1 
$$R_1:8+/2-R2:2+/8-$$
  
Split 2  $R_1:6+/4-R2:0+/10-$ 

In general, we want to produce pure nodes, i.e. close to single-class node.

#### Misclassification error in a node

Let's formalize things a bit.

- Consider classification case:  $\mathcal{Y} = \{1, 2, ..., K\}$ .
- What's in a node?
  - Let node m represent region  $R_m$ , with  $N_m$  observations
  - Denote proportion of observations in  $R_m$  with class k by

$$\hat{\rho}_{mk} = \frac{1}{N_m} \sum_{\{i: x_i \in R_m\}} 1(y_i = k).$$

• Predict the majority class in node m:

$$k(m) = \arg\max_{k} \hat{p}_{mk}.$$

Misclassification rate in node m:

$$1-\hat{p}_{mk(m)}$$
.

# Node Impurity Measures

#### How to quantify impurity?

• Three measures of **node impurity** for leaf node *m*:

Misclassification error

$$1-\hat{p}_{mk(m)}$$
.

Gini index

$$\sum_{k=1}^{K} \hat{p}_{mk} (1 - \hat{p}_{mk})$$

Entropy / Information gain

$$-\sum_{k=1}^K \hat{p}_{mk} \log \hat{p}_{mk}.$$

• Gini index and entropy work well in practice.

# Impurity of a split

A potential split produces two nodes,  $R_L$  and  $R_R$ . How do we score it?

- Suppose we have  $N_L$  points in  $R_L$  and  $N_R$  points in  $R_R$ .
- Let  $Q(R_L)$  and  $Q(R_R)$  be the node impurity measures for each node.
- Then find split that minimizes the weighted average of node impurities:

$$\frac{N_L Q(R_L) + N_R Q(R_R)}{N_L + N_R}$$

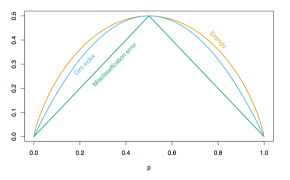
#### Example:

$$R_1:8+/2 R_2:1+/4-$$

What's the weighted misclassification rate?

### Two-Class Node Impurity Measures

Consider binary classification. Let p be the relative frequency of class 1.



Misclassification error is not strictly concave thus may not guarantee improvement over the parent node.

## Finding the Split Point

How to find a split point that minimizes a given impurity measure?

- Consider splitting on the j'th feature  $x_i$ .
- If  $x_{j(1)}, \ldots, x_{j(n)}$  are the sorted values of the j'th feature,
  - we only need to check split points between adjacent values
  - traditionally take split points halfway between adjacent values:

$$s_j \in \left\{ \frac{1}{2} \left( x_{j(r)} + x_{j(r+1)} \right) \mid r = 1, \dots, n-1 \right\}.$$
  $n-1 \text{ splits}$  (1)

• Enumerate d features and n-1 split points for each feature.

#### Regression trees

Predict the mean value of a node

$$k(m) = \operatorname{mean}(y_i \mid x_i \in R_m). \tag{2}$$

- Squared loss as the node impurity measure.
- Everything else remains the same as classification trees.

# Categorical features

- For a categorical feature, we split its values into two groups.
- Given a set of categories of size k, how many distinct splits? (its power set)
- Finding the optimal split is intractable in general.
- Approximations

Numeric encoding Randomly assign a number to each category

- Binary classification: proportion of class 0
- Regression: mean of targets of examples in the category, i.e.
   mean encoding

One-hot encoding May grow imbalanced trees, e.g., left-branching Binary encoding Robust to large cardinality

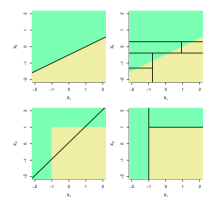
- Statistical issues with categorical features
  - If a category has a very large number of categories, we can overfit.
  - Extreme example: Row Number could lead to perfect classification with a single split.

### Interpretability

- Trees are certainly easier to explain than other classifiers.
- Can be used to discover non-linear features.
- Small trees seem interpretable. For large trees, maybe not so easy.
- Approximate neural network decision boundaries to gain interpretability
  - Wu M, Hughes M, Parbhoo S, Zazzi M, Roth V, Doshi-Velez F. Beyond Sparsity: Tree Regularization of Deep Models for Interpretability. Association for the Advancement of Artificial Intelligence (AAAI). 2018

#### Trees vs linear models

Trees have to work much harder to capture linear relations.



#### Review

#### Decision trees:

- Non-linear classifier that recursively partitions the input space.
- Non-metric: make no use of geometry, i.e. no inner-product or distances.
- Non-parametric: make no assumption of the data distribution.

#### Pros:

- Simple to understand.
- Interpretable, feature selection for free.

#### Cons:

- Poor linear modeling.
- Unstable / high variance, tend to overfit.  $\rightarrow$  Next, how to fix this.

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Bagging and Random Forests

# Recap: statistic and point estimator

- Observe data  $\mathcal{D} = (x_1, x_2, \dots, x_n)$  sampled i.i.d. from a parametric distribution  $p(\cdot \mid \theta)$ .
- A statistic  $s = s(\mathcal{D})$  is any function of the data.
  - E.g., sample mean, sample variance, histogram, empirical data distribution
- A statistic  $\hat{\theta} = \hat{\theta}(\mathcal{D})$  is a **point estimator** of  $\theta$  if  $\hat{\theta} \approx \theta$ .

#### Review questions

In frequentist statistics.

- Is  $\theta$  random?
- Is  $\hat{\theta}$  random?
- Is the function  $s(\cdot)$  random?

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## Recap: bias and variance of an estimator

- Statistics are random, so they have probability distributions.
- The distribution of a statistic is called a sampling distribution.
- The standard deviation of the sampling distribution is called the standard error.
- What are some parameters of the sampling distribution we might be interested in?

$$\begin{array}{c} \mathsf{Bias} \ \mathsf{Bias}(\hat{\theta}) \stackrel{\mathrm{def}}{=} \mathbb{E}\left[\hat{\theta}\right] - \theta. \\ \mathsf{Variance} \ \mathsf{Var}(\hat{\theta}) \stackrel{\mathrm{def}}{=} \mathbb{E}\left[\hat{\theta}^2\right] - \mathbb{E}^2\left[\hat{\theta}\right]. \end{array}$$

- Is bias and variance random?
  - Neither bias nor variance depend on a specific sample  $\mathcal{D}_n$ . We are taking expectation over  $\mathcal{D}$ .
- Why do we care about variance?
  - $\hat{\theta}(\mathcal{D}) = x_1$  is an unbiased estimator of the mean of a Gaussian, but would be farther away from  $\theta$  than the sample mean.

#### Variance of a Mean

Using a single estimate may have large standard error

- $\bullet \ \ \mathsf{Let} \ \hat{\theta}(\mathcal{D}) \ \mathsf{be} \ \mathsf{an} \ \mathsf{unbiased} \ \mathsf{estimator} \colon \ \mathbb{E} \left\lceil \hat{\theta} \right\rceil = \theta \text{, } \mathsf{Var}(\hat{\theta}) = \sigma^2.$
- We could use a single estimate  $\hat{\theta} = \hat{\theta}(\mathcal{D})$  to estimate  $\theta$ .
- The standard error is  $\sqrt{\mbox{Var}(\hat{\theta})} = \sigma.$

#### Average of estimates has smaller standard error

- Consider a new estimator that takes the average of i.i.d.  $\hat{\theta}_1, \dots, \hat{\theta}_n$  where  $\hat{\theta}_i = \hat{\theta}(\mathcal{D}^i)$ .
- Average has the same expected value but smaller standard error:

$$\mathbb{E}\left[\frac{1}{n}\sum_{i=1}^{n}\hat{\theta}_{i}\right] = \theta \quad \text{Var}\left[\frac{1}{n}\sum_{i=1}^{n}\hat{\theta}_{i}\right] = \frac{\sigma^{2}}{n}$$
(3)

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# Averaging Independent Prediction Functions

Let's apply averaging to reduce variance of prediction functions.

- Suppose we have B independent training sets from the same distribution  $(\mathcal{D} \sim p(\cdot \mid \theta))$ .
- Learning algorithm (estimator) gives B prediction functions:  $\hat{f}_1(x), \hat{f}_2(x), \dots, \hat{f}_B(x)$
- Define the average prediction function as:

$$\hat{f}_{\text{avg}} \stackrel{\text{def}}{=} \frac{1}{B} \sum_{b=1}^{B} \hat{f}_{b} \tag{4}$$

- What's random here?
  - The B independent training sets are random, which gives rise to variation among the  $\hat{f}_b$ 's.
- Concept check: What's the distribution of  $\hat{f}$  called? What do we know about the distribution?

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## Averaging reduce variance of predictions

• The average prediction on  $x_0$  is

$$\hat{f}_{avg}(x_0) = \frac{1}{B} \sum_{b=1}^{B} \hat{f}_b(x_0).$$

- $\hat{f}_{avg}(x_0)$  and  $\hat{f}_b(x_0)$  have the same expected value, but
- $\hat{f}_{avg}(x_0)$  has smaller variance (see 3):

$$\operatorname{Var}(\hat{f}_{\mathsf{avg}}(x_0)) = \frac{1}{B} \operatorname{Var}\left(\hat{f}_1(x_0)\right)$$

• Problem: in practice we don't have B independent training sets...

# The Bootstrap Sample

How do we simulate multiple samples when we only have one?

- A **bootstrap sample** from  $\mathcal{D}_n = (x_1, ..., x_n)$  is a sample of size n drawn with replacement from  $\mathcal{D}_n$ .
- Some elements of  $\mathcal{D}_n$  will show up multiple times, and some won't show up at all.

How similar are the bootstrap samples?

- Each  $x_i$  has a probability of  $(1-1/n)^n$  of not being selected.
- Recall from analysis that for large n,

$$\left(1 - \frac{1}{n}\right)^n \approx \frac{1}{e} \approx .368. \tag{5}$$

• So we expect ~63.2% of elements of  $\mathcal{D}_n$  will show up at least once.

# The Bootstrap Method

#### Definition

A **bootstrap method** is when you simulate having B independent samples from P by taking B bootstrap samples from the sample  $\mathfrak{D}_n$ .

- Given original data  $\mathcal{D}_n$ , compute B bootstrap samples  $D_n^1, \ldots, D_n^B$ .
- For each bootstrap sample, compute some function

$$\phi(D_n^1), \ldots, \phi(D_n^B)$$

- Work with these values as though  $D_n^1, \ldots, D_n^B$  were i.i.d. samples from P.
- Amazing fact: This is often very close to what we'd get with independent samples from *P*.

### Independent vs Bootstrap Samples

- Want to estimate  $\alpha = \alpha(P)$  for some unknown P and some complicated  $\alpha$ .
- Point estimator  $\hat{\alpha} = \hat{\alpha}(\mathcal{D}_{100})$  for samples of size 100.
- Histogram of  $\hat{\alpha}$  based on
  - 1000 independent samples of size 100, vs
  - 1000 bootstrap samples of size 100

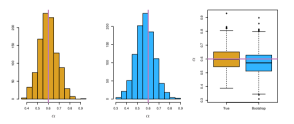


Figure 5.10 from ISLR (Springer, 2013) with permission from the authors: G. James, D. Witten, T. Hastie and R. Tibshirani.

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# Side note: Bootstrap in Practice

We can use bootstrap to get error bars in a cheap way.

- Suppose we have an estimator  $\hat{\theta} = \hat{\theta}(\mathcal{D}_n)$ .
- To get error bars, we can compute the "bootstrap variance".
  - Draw B bootstrap samples.
  - Compute sample variance of  $\hat{\theta}(\mathcal{D}_n^1), \ldots, \hat{\theta}(\mathcal{D}_n^B)$ ...
  - Could report

$$\hat{\theta}(\mathfrak{D}_n) \pm \sqrt{\mathsf{Bootstrap Variance}}$$

#### Ensemble methods

#### Key ideas:

- Averaging i.i.d. estimates reduces variance without making bias worse.
- Can use bootstrap to simulate multiple data samples.

#### **Ensemble methods:**

- Combine outputs from multiple models.
  - Same learner on different datasets: ensemble + bootstrap = bagging.
  - Different learners on one dataset: they may make similar errors.
- Parallel ensemble: models are built independently, e.g., bagging
- Sequential ensemble: models are built sequentially, e.g., boosting
  - Try to add new learners that do well where previous learners lack

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# Bagging

- Draw B bootstrap samples  $D^1, \ldots, D^B$  from original data  $\mathfrak{D}$ .
- Let  $\hat{f}_1, \hat{f}_2, \dots, \hat{f}_B$  be the prediction functions from training on  $D^1, \dots, D^B$ , respectively.
- The bagged prediction function is a combination of these:

$$\hat{f}_{\mathsf{avg}}(x) = \mathsf{Combine}\left(\hat{f}_1(x), \hat{f}_2(x), \dots, \hat{f}_B(x)\right)$$

- How might we combine
  - prediction functions for regression?
  - binary class predictions?
  - binary probability predictions?
  - multiclass predictions?

# Out-of-Bag Error Estimation

- Each bagged predictor is trained on about 63% of the data.
- Remaining 37% are called out-of-bag (OOB) observations.
- For ith training point, let

$$S_i = \{b \mid D^b \text{ does not contain } i\text{th point}\}.$$

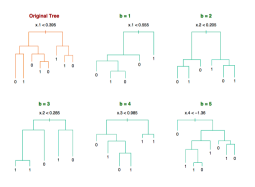
• The OOB prediction on  $x_i$  is

$$\hat{f}_{OOB}(x_i) = \frac{1}{|S_i|} \sum_{b \in S_i} \hat{f}_b(x_i).$$

- The OOB error is a good estimate of the test error.
- OOB error is similar to cross validation error both are computed on training set.

# Bagging Classification Trees

• Input space  $\mathfrak{X}=\mathsf{R}^5$  and output space  $\mathfrak{Y}=\{-1,1\}$ . Sample size n=30.



- Each bootstrap tree is quite different: different splitting variable at the root
- **High variance**: high degree of model variability from small perturbations of the training data.
- Conventional wisdom: Bagging helps most when base learners are relatively unbiased but has high variance / low stability 

  decision trees.

#### Variance of a Mean of Correlated Variables

#### Recall the motivating principle of bagging:

• For  $\hat{\theta}_1, \dots, \hat{\theta}_n$  i.i.d. with  $\mathbb{E}\left[\hat{\theta}\right] = \theta$  and  $\mathrm{Var}\left[\hat{\theta}\right] = \sigma^2$ ,

$$\mathbb{E}\left[\frac{1}{n}\sum_{i=1}^{n}\hat{\theta}_{i}\right] = \mu \qquad \operatorname{Var}\left[\frac{1}{n}\sum_{i=1}^{n}\hat{\theta}_{i}\right] = \frac{\sigma^{2}}{n}.$$

- What if  $\hat{\theta}$ 's are correlated?
- Suppose  $\forall i \neq j$ ,  $\mathsf{Corr}(\hat{\theta}_i, \hat{\theta}_i) = \rho$  . Then

$$\operatorname{Var}\left[\frac{1}{n}\sum_{i=1}^{n}\hat{\theta}_{i}\right] = \rho\sigma^{2} + \frac{1-\rho}{n}\sigma^{2}.$$

• For large n, the  $\rho \sigma^2$  term dominates – limits benefit of averaging.

## Correlation between bootstrap samples

- Averaging  $\hat{f}_1, \ldots, \hat{f}_B$  reduces variance if they're based on i.i.d. samples from  $P_{\mathfrak{X} \times \mathfrak{Y}}$
- Bootstrap samples are
  - independent samples from the training set, but
  - are not independent samples from  $P_{X \times Y}$ .
- This dependence limits the amount of variance reduction we can get.
- Solution: reduce the dependence between  $\hat{f}_i$ 's by randomization.

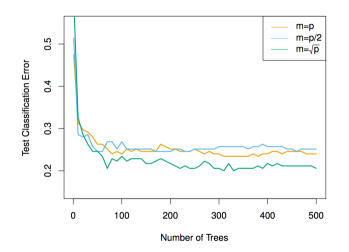
#### Random Forest

#### Key idea

Use bagged decision trees, but modify the tree-growing procedure to reduce the dependence between trees.

- Build a collection of trees independently (in parallel).
- When constructing each tree node, restrict choice of splitting variable to a randomly chosen subset of features of size *m*.
  - Avoid dominance by strong features.
- Typically choose  $m \approx \sqrt{p}$ , where p is the number of features.
- Can choose m using cross validation.

#### Random Forest: Effect of *m* size



From An Introduction to Statistical Learning, with applications in R (Springer, 2013) with permission from the authors: G. James, D. Witten, T. Hastie and R. Tibshirani.

- Usual approach is to build very deep trees—low bias but high variance
- Ensembling many models reduces variance
  - Motivation: Mean of i.i.d. estimates has smaller variance than single estimate.
- Use bootstrap to simulate many data samples from one dataset
  - $\implies$  Bagged decision trees
- But bootstrap samples (and the induced models) are correlated.
- Bagging seems to work better when we are combining a diverse set of prediction functions.
  - ⇒ random forests (randomized tree building)

Boosting

#### Overview

- Bagging Reduce variance of a low bias, high variance estimator by ensembling many estimators trained in parallel.
- Boosting Reduce the error rate of a high bias estimator by ensembling many estimators trained in sequential.
  - A weak/base learner is a classifier that does slightly better than chance.
  - Weak learners are like "rules of thumb":
    - "Viagra" ⇒ spam
    - From a friend  $\implies$  not spam
  - Key idea:
    - Each weak learner focuses on different examples (reweighted data)
    - Weak learners have different contributions to the final prediction (reweighted classifier)

## AdaBoost: Setting

- Binary classification:  $\mathcal{Y} = \{-1, 1\}$
- Base hypothesis space  $\mathcal{H} = \{h : \mathcal{X} \to \{-1, 1\}\}.$
- Typical base hypothesis spaces:
  - Decision stumps (tree with a single split)
  - Trees with few terminal nodes
  - Linear decision functions

# Weighted Training Set

Each base learner is trained on weighted data.

- Training set  $\mathcal{D} = ((x_1, y_1), \dots, (x_n, y_n)).$
- Weights  $(w_1, ..., w_n)$  associated with each example.
- Weighted empirical risk:

$$\hat{R}_n^w(f) \stackrel{\text{def}}{=} \frac{1}{W} \sum_{i=1}^n w_i \ell(f(x_i), y_i)$$
 where  $W = \sum_{i=1}^n w_i$ 

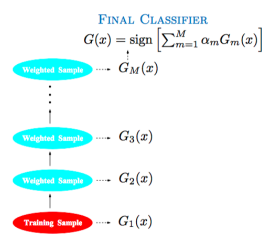
Examples with larger weights have more influence on the loss.

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## AdaBoost - Rough Sketch

- Training set  $\mathcal{D} = ((x_1, y_1), \dots, (x_n, y_n)).$
- Start with equal weight on all training points  $w_1 = \cdots = w_n = 1$ .
- Repeat for m = 1, ..., M:
  - Find base classifier  $G_m(x)$  that tries to fit weighted training data (but may not do that well)
  - Increase weight on the points  $G_m(x)$  misclassifies
- So far, we've generated M classifiers:  $G_1, \ldots, G_M : \mathcal{X} \to \{-1, 1\}$ .

#### AdaBoost: Schematic



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# AdaBoost - Rough Sketch

- Training set  $\mathcal{D} = \{(x_1, y_1), \dots, (x_n, y_n)\}.$
- Start with equal weight on all training points  $w_1 = \cdots = w_n = 1$ .
- Repeat for m = 1, ..., M:
  - Base learner fits weighted training data and returns  $G_m(x)$
  - Increase weight on the points  $G_m(x)$  misclassifies
- Final prediction  $G(x) = \operatorname{sign}\left[\sum_{m=1}^{M} \alpha_m G_m(x)\right]$ . (recall  $G_m(x) \in \{-1,1\}$ )
- What are desirable  $\alpha_m$ 's?
  - nonnegative
  - larger when  $G_m$  fits its weighted  $\mathcal{D}$  well
  - smaller when  $G_m$  fits weighted  $\mathfrak D$  less well

# Adaboost: Weighted Classification Error

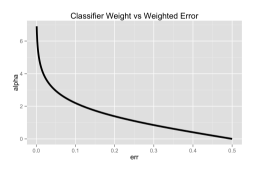
- Weights of base learners depend on their performance. How to evaluate each base learner?
- In round m, base learner gets a weighted training set.
  - Returns a base classifier  $G_m(x)$  that minimizes weighted 0-1 error.
- The weighted 0-1 error of  $G_m(x)$  is

$$\operatorname{err}_m = \frac{1}{W} \sum_{i=1}^n w_i \mathbb{1}(y_i \neq G_m(x_i))$$
 where  $W = \sum_{i=1}^n w_i$ .

• Notice:  $err_m \in [0, 1]$ .

## AdaBoost: Classifier Weights

• The weight of classifier  $G_m(x)$  is  $\alpha_m = \ln\left(\frac{1 - \text{err}_m}{\text{err}_m}\right)$ .



- Higher weighted error ⇒ lower weight
- When is  $\alpha_m < 0$ ?

## Adaboost: Example Reweighting

- We train  $G_m$  to minimize weighted error, and it achieves err<sub>m</sub>.
- Then  $\alpha_m = \ln\left(\frac{1 \text{err}_m}{\text{err}_m}\right)$  is the weight of  $G_m$  in final ensemble.

We want the base learner to focus more on examples misclassified by the previous learner.

- Suppose  $w_i$  is weight of example i before training:
  - If  $G_m$  classfies  $x_i$  correctly, then  $w_i$  is unchanged.
  - Otherwise, w<sub>i</sub> is increased as

$$w_i \leftarrow w_i e^{\alpha_m}$$

$$= w_i \left( \frac{1 - \operatorname{err}_m}{\operatorname{err}_m} \right)$$

• For  $err_m < 0.5$  (weak learner), this always increases the weight.

## AdaBoost: Algorithm

Given training set  $\mathcal{D} = \{(x_1, y_1), \dots, (x_n, y_n)\}.$ 

- 1 Initialize observation weights  $w_i = 1, i = 1, 2, ..., n$ .
- 2 For m = 1 to M:
  - Base learner fits weighted training data and returns  $G_m(x)$
  - 2 Compute weighted empirical 0-1 risk:

$$\operatorname{err}_m = \frac{1}{W} \sum_{i=1}^n w_i \mathbb{1}(y_i \neq G_m(x_i))$$
 where  $W = \sum_{i=1}^n w_i$ .

- Compute classifier weight:  $\alpha_m = \ln\left(\frac{1 \text{err}_m}{\text{err}_m}\right)$ .
- Update example weight:  $w_i \leftarrow w_i \cdot \exp\left[\alpha_m \mathbf{1}(y_i \neq G_m(x_i))\right]$
- **3** Return voted classifier:  $G(x) = \text{sign} \left[ \sum_{m=1}^{M} \alpha_m G_m(x) \right]$ .

### AdaBoost with Decision Stumps

• After 1 round:

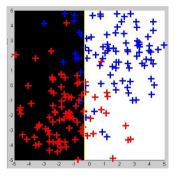


Figure: Plus size represents weight. Blackness represents score for red class.

### AdaBoost with Decision Stumps

• After 3 rounds:

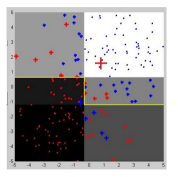


Figure: Plus size represents weight. Blackness represents score for red class.

### AdaBoost with Decision Stumps

• After 120 rounds:

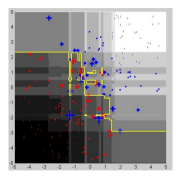
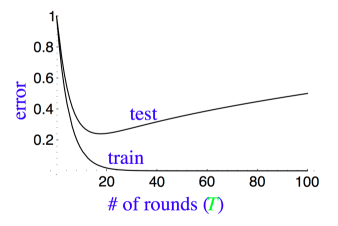


Figure: Plus size represents weight. Blackness represents score for red class.

# Typical Train / Test Learning Curves

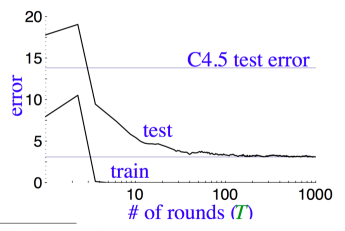
• Might expect too many rounds of boosting to overfit:



From Rob Schapire's NIPS 2007 Boosting tutorial.

### Learning Curves for AdaBoost

- In typical performance, AdaBoost is surprisingly resistant to overfitting.
- Test continues to improve even after training error is zero!



From Rob Schapire's NIPS 2007 Boosting tutorial.

### Summary

- Shallow decision tree + boosting
  - "best off-the-shelf classifier in the world"—Leo Brieman
  - Used in the first successful real-time face detector (Viola and Jones, 2001)
  - XGBoost: very popular in competitions
- Next week
  - What is the objective function of Adaboost?
  - Generalize to other loss functions.