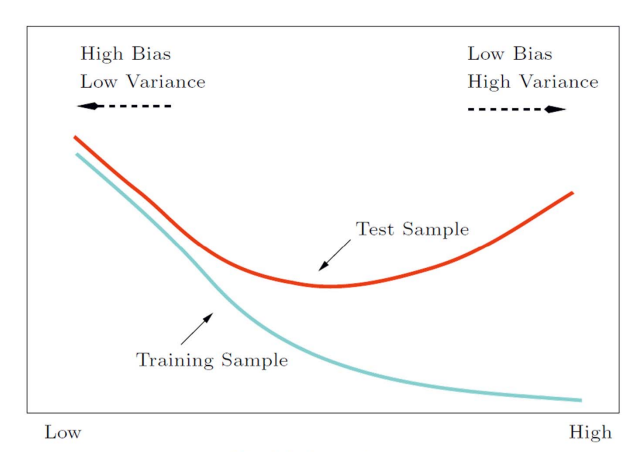


## So Far...

- Our goal (supervised learning)
  - Naïve Bayesian classifier
  - Linear Regression
  - Logistic Regression
  - SVM
  - Perceptron
  - Neural Network
  - k Nearest Neighbor
  - Decision Tree
  - Single Classifier
- Can the classifiers be combined to achieve better performance?
  - Two heads are better than one



# Prediction Error



Model Complexity

$$EPE(f) = (bias)^2 + variance + noise$$

## **Ensemble Methods & Random Forest**

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## The Bagging Algorithm

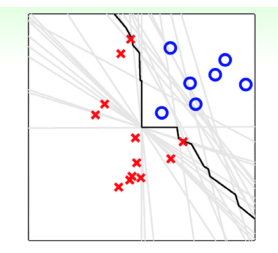
- ► The name Bagging came from the abbreviation of *Bootstrap Aggregating* [Breiman, 1996d], implies the two key ingredients of Bagging are bootstrap and aggregation.
  - Bootstrap sampling: re-sample *N* examples from D uniformly with replacement—can also use arbitrary *N'* instead of original *N*
  - **Aggregating**: adopts the most popular strategies for aggregating the outputs of the base learners, that is, *voting* for classification and *averaging* for regression.



## The Bagging Algorithm

Pocket is a modified Perceptron Learning Algorithm.

Bagging 25 Pocket Models. The max iteration for each Pocket Algorithm is 1000.



 $T_{\mathsf{POCKET}} = 1000; \, T_{\mathsf{BAG}} = 25$ 

- very diverse  $g_t$  from bagging
- proper non-linear boundary after aggregating binary classifiers
- bagging works reasonably well if base algorithm sensitive to data randomness



## Why Bagging Works

Let f denote the ground-truth function and h(x) denote a learner trained from the bootstrap distribution  $D_{bs}$ . The aggregated learner generated by Bagging is

$$H(\mathbf{x}) = E_{D_{bs}}[h(\mathbf{x})]$$

▶ With simple algebra and the inequality  $(E[X])^2 \le E[X^2]$ , we have

$$(f - H(\mathbf{x}))^2 \le E_{D_{bs}} \left[ (f - h(\mathbf{x}))^2 \right]$$

Thus, by integrating both sides over the distribution, we can get that the mean-squared error of H(x) is smaller than that of h(x) averaged over the bootstrap sampling distribution



## Why Bagging Works: Bias & Variance

$$\left\{E_D(f(\boldsymbol{x};D)) - E(y|\boldsymbol{x})\right\}^2 + E_D\left\{\left[f(\boldsymbol{x};D) - E_D(f(\boldsymbol{x};D))\right]^2\right\}$$

Let f denote the ground-truth function and h(x) denote a learner trained from the bootstrap distribution  $D_{bs}$ . The aggregated learner generated by Bagging is

$$H(\mathbf{x}) = E_{D_{hs}}[h(\mathbf{x})]$$

- ▶ Bias:  $E_D[h(x)] f \approx H(x) f$
- Variance:
  - Assume there are T bs samples, if data is i. i. d. and each variance is  $\sigma^2$ . The ensemble variance is  $\frac{1}{T}\sigma^2$

$$H(x) = E_{D_{bs}}[h(x)] = \frac{1}{T} \sum_{i=1}^{T} h_i(x)$$

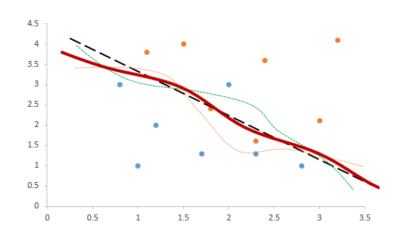
$$Var(H(x)) = Var\left(\frac{1}{T}\sum_{i}^{T}h_{i}(x)\right) = \frac{1}{T^{2}}Var\left(\sum_{i=1}^{T}h_{i}(x)\right) = \frac{1}{T^{2}}\sum_{i=1}^{T}Var(h_{i}(x)) = \frac{1}{T}\sigma^{2}$$

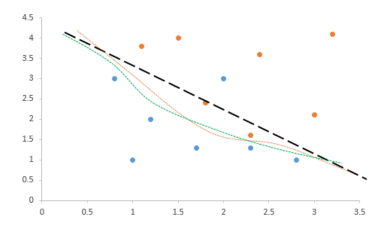
• In reality, Bootstrap Sampled data is i.d. (not necessarily independent) with correlation  $\rho$ 

$$\rho\sigma^2 + \frac{1-\rho}{T}\sigma^2$$



# Why Bagging Works

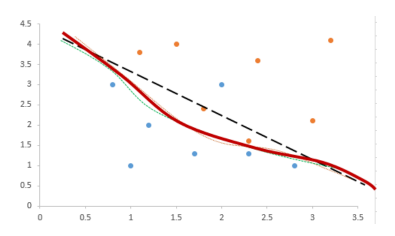




Why?

If Bootstrap Sampled data is i.d. with correlation  $\rho$ :

$$\rho\sigma^2 + \frac{1-\rho}{T}\sigma^2$$





## Why Bagging Works

- ▶ Performance improvement brought by Bagging is large when the base learner is unstable(large variance).
  - The base learner should aware the little change, sometimes overfitting is allowable.
- ▶ Thus, Friedman and Hall [2007] concluded that Bagging can reduce the variance of higher-order components, yet not affect the linear components. This implies that Bagging is better applied with highly nonlinear learners.



## Which Classifier is a good choice for base learner?

- Naïve Bayesian classifier
- Perceptron
- Linear Regression
- Logistic Regression
- SVM
- Neural Network
- k Nearest Neighbor
- Decision Tree

Model with low bias benefits from bagging

- Decision Tree!
  - Non-linear classifier
  - Easy to use and interpret
  - Can perfectly fit to any training data (overfitting) with high test error. (zero bias, high variance)



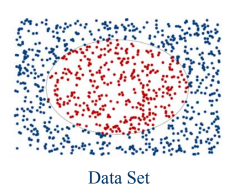
#### **Random Forest**

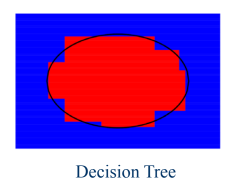
- Random Forest, which uses decision tree as basic unit in bagging, is an ensemble model.
- Why Random Forest?
  - Advantages of Decision Tree:
    - Handling missing value
    - Robust to outliers in input space
    - Fast
    - Good Interpretability
  - Limitations of Decision Tree:
    - Low accuracy, low bias with high variance and easy to overfit
  - Ensemble to maintain advantages while increasing accuracy



## **Random Forest**

- Bagging sample is a random process.
  - Bootstrap sample process: random sampling he given
     N samples with replacement
  - Repeat T times and create new T training sets
  - Get T models
- Aggregating all models will fuzzy up the decision boundary, which help reduce the variance, and prevent the *one man rule* danger

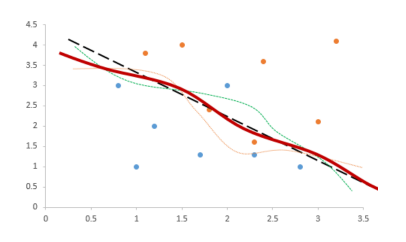


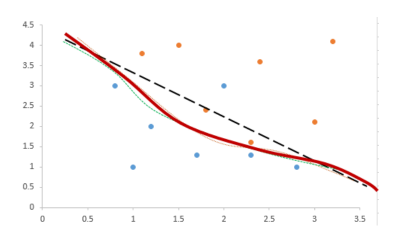






## How to increase the variation of each tree?







#### **Random Forest**

- ► The main difference between the method of RF and Bagging is that RF incorporate randomized feature selection at each split step.
- At each split step: Get feature subsets f from the whole F
  - More efficient in building trees
    - each time we only pick the best feature from size(f) rather than size(F).
      - We often let size(f) =  $\sqrt{k}$  in classification and k/3 in regression
  - Each tree is not as good as DT, but work better when they are combined.



#### **Construct Random Forest**

- Let N be the number of trees and K be the feature subset size.
- For each N iterations: (building a tree in forest)
  - 1. Select a new bootstrap sample from original training set
  - 2. Growing a tree...
  - 3. At each internal node, randomly select K features from ALL features and then, determine the best split in ONLY the K features.
  - 4. Do not pruning
  - 5. Until Test Error never decrease (here means Validation error in RF)
- At last, overall prediction as the average( or vote) from N trees.



#### 3 Rules of RF makes the learners more Diverse.

Random forest need basic learner aware the little change, sometimes overfit is allowable.

► Each time, basic learner doesn't learn from all data, but from Random bootstrap sampled data.

Basic learner doesn't use all features, but Random select some features.



## Other ways to generate base learner?

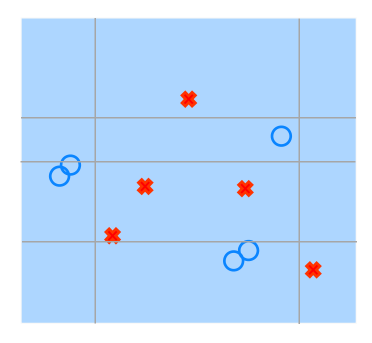
- ► The base learner should have low bias? Not necessary
- ▶ The base learner should have high variance? What does this mean?
  - Different base learners should be different.
  - Different base learners should have different result.

$$\min_{f} \left\{ \sum_{i=1}^{n} \ell(f, \mathbf{x}_i, y_i) + \lambda R(f) \right\}$$

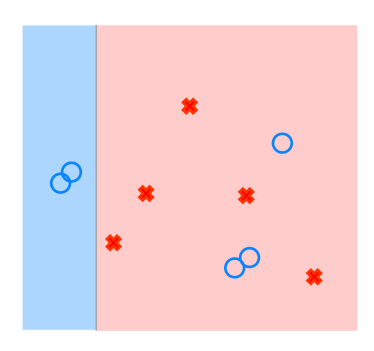
$$\min_{f} \left\{ \sum_{i=1}^{n} w_i \ell(f, \mathbf{x}_i, y_i) + \lambda R(f) \right\}$$

- Boosting
  - Training base learners by assigning different weights to the samples

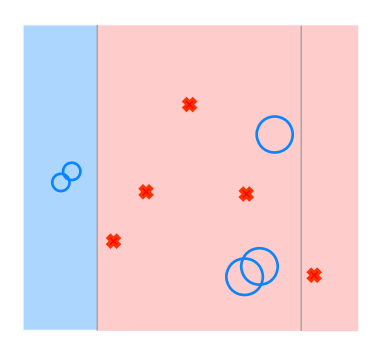




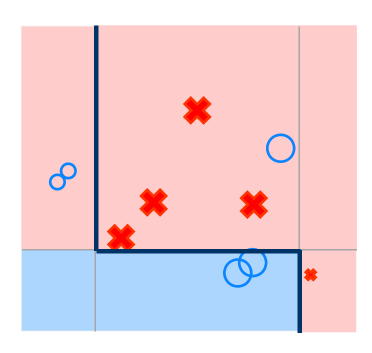




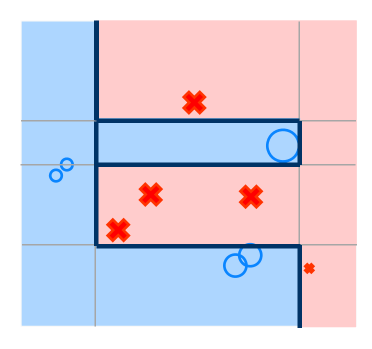




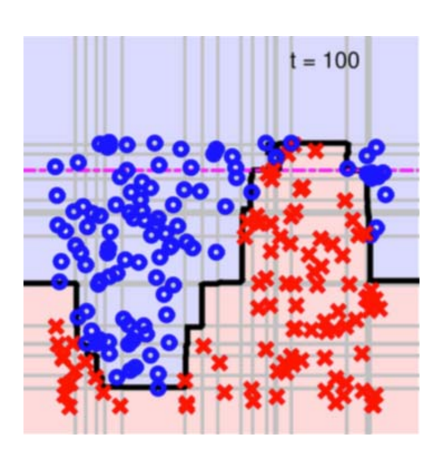














## Observation

- In order to come up an algorithm, we need to answer two underlying questions:
  - Question 1: How to change the weights of samples so that misclassified samples get more weight
  - Question 2: How to combine base learners in final phase



## AdaBoost

▶ 1. Initialize the data weighting coefficients  $\mathbf{w}$  by setting  $w_n^{(1)} = 1/N$  for n = 1, ..., N.

- 2. For m = 1, ..., M:
  - (a) Fit a classifier  $y^{(m)}(x)$  to the training data by minimizing the weighted error function

$$J_{m} = \sum_{n=1}^{N} w_{n}^{(m)} I(y^{(m)}(x_{n}) \neq t_{n})$$



#### AdaBoost

(b) Evaluate the Errorrate:

$$\epsilon_m = \frac{\sum_{n=1}^{N} w_n^{(m)} I(y^{(m)}(x_n) \neq t_n)}{\sum_{n=1}^{N} w_n^{(m)}}$$

and then use this to evaluate

$$\alpha_m = \ln \frac{1 - \epsilon_m}{\epsilon_m}$$

 $\alpha_m = \ln \frac{1 - \epsilon_m}{\epsilon_m}$  Log odds, smaller the error rates, bigger this value.

(c) the data weighting coefficients 
$$w_n^{(m+1)} = w_n^{(m)} \exp\{\alpha_m I(y^{(m)}(\boldsymbol{x}_n) \neq t_n)\} = \begin{cases} w_n^{(m)} \frac{1 - \epsilon_m}{\epsilon_m}, & \text{if } y^{(m)} \text{ makes error} \\ w_n^{(m)}, & \text{otherwise} \end{cases}$$

Answer to question 1, How to change the weights of samples so that misclassified samples get more weight.



## AdaBoost

▶ 3. Make predictions using the final model, which is given by

$$Y_M(x) = \operatorname{sign}\left(\sum_{m=1}^M \alpha_m y^{(m)}(x)\right)$$

Question 2: How to Combine base learners in final phase

weighting coefficients  $\alpha_m$  give greater weight to the more accurate classifiers when computing the overall output



# **Insights Behind Adaboost**

▶ In each round, the algorithm try to get a different base learner, so that model diversity is achieved



## Re-weight for More Diverse Base Learner

$$g_{m} \leftarrow \min_{h \in H} (\sum_{n=1}^{N} w_{n}^{(m)} I(y_{n}^{(m)} \neq t_{n}))$$

$$g_{m+1} \leftarrow \min_{h \in H} (\sum_{n=1}^{N} w_{n}^{(m+1)} I(y_{n}^{(m+1)} \neq t_{n}))$$

if  $g_m$  'not good' for  $w^{(m+1)} \Rightarrow g_{m+1}$  diverse from  $g_m$ 

**Idea: construct**  $w^{m+1}$  to make  $g_m$  random-like

$$\frac{\sum_{n=1}^{N} w_n^{(m+1)} I(y_n^{(m)} \neq t_n))}{\sum_{n=1}^{N} w_n^{(m+1)}} = \frac{1}{2} \frac{\sum_{n=1}^{N} w_n^{(m+1)} I(y_n^{(m)} \neq t_n))}{\sum_{n=1}^{N} w_n^{(m+1)} I(y_n^{(m)} \neq t_n)) + \sum_{n=1}^{N} w_n^{(m+1)} I(y_n^{(m)} = t_n))} = \frac{1}{2}$$

Solve this equation will give us 
$$w_n^{(m+1)} = w_n^{(m)} \sqrt{\frac{1-\epsilon_m}{\epsilon_m}}$$

$$w_n^{(m+1)} = w_n^{(m)} \exp\left\{-\frac{1}{2}\alpha_m t_n y^{(m)}(\boldsymbol{x}_n)\right\} \qquad w_n^{(m+1)} = w_n^{(m)} \sqrt{\frac{\epsilon_m}{1-\epsilon_m}}$$

$$w_n^{(m+1)} = w_n^{(m)} \sqrt{\frac{\epsilon_m}{1-\epsilon_m}}$$

which is the same as update equation in Adaboost Algorithm



# **Insights Behind Adaboost**

▶ In each round, the algorithm try to get a different base learner, so that model diversity is achieved

 Adaboost can be see as a sequential optimization process of an additive model under exponential error



# **Optimization of Exponential Loss**

Consider the exponential error function defined by

$$E = \sum_{n=1}^{N} \exp\{-t_n f_m(\boldsymbol{x}_n)\}\$$

where  $f_m(x)$  is a classifier defined in terms of a linear combination of base classifiers  $y_l(x)$  of the form  $f_m(x) = \frac{1}{2} \sum_{l=1}^{m} \alpha_l y_l(x)$ 

and  $t_n \in \{-1, 1\}$  are the training set target values. Our goal is to minimize E with respect to both the weighting coefficients  $\alpha_l$  and the parameters of the base classifiers  $y_l(x)$ .

Instead of doing a global error function minimization, however, we shall suppose that the base classifiers  $y_1(x), \dots, y_{m-1}(x)$  are fixed, as are their coefficients  $\alpha_1, \dots, \alpha_{m-1}$ , and so we are minimizing only with respect to  $\alpha_m$  and  $y_m(x)$ .



# **Optimization of Exponential Loss**

$$E = \sum_{n=1}^{N} \exp\{-t_n f_m(\mathbf{x}_n)\} \qquad f_m(\mathbf{x}) = \frac{1}{2} \sum_{l=1}^{m} \alpha_l y_l(\mathbf{x})$$

• the base classifiers  $y_1(x), \dots, y_{m-1}(x)$  are fixed, as are their coefficients  $\alpha_1, \dots, \alpha_{m-1}$ ,

$$= \sum_{n=1}^{N} \exp \left\{ -t_n f_{m-1}(\mathbf{x}_n) - \frac{1}{2} t_n \alpha_m y_m(\mathbf{x}_n) \right\}$$

$$= \sum_{n=1}^{N} \exp \left\{ -t_n f_{m-1}(\mathbf{x}_n) \right\} \exp \left\{ -\frac{1}{2} t_n \alpha_m y_m(\mathbf{x}_n) \right\}$$

$$= \sum_{n=1}^{N} w_n^{(m)} \exp \left\{ -\frac{1}{2} t_n \alpha_m y_m(\mathbf{x}_n) \right\}$$

$$w_n^{(m+1)} = w_n^{(m)} \exp\left\{-\frac{1}{2}t_n \alpha_m y_m(\mathbf{x}_n)\right\} \qquad t_n y_m(\mathbf{x}_n) = 1 - 2I(y^{(m)}(\mathbf{x}_n) \neq t_n)$$

$$w_n^{(m+1)} = w_n^{(m)} \exp\left\{-\frac{\alpha_m}{2}\right\} \exp\left\{\alpha_m I(y^{(m)}(\mathbf{x}_n) \neq t_n)\right\}$$



## **Optimization of Exponential Loss**

If we denote by  $T_m$  the set of data points that are correctly classified by  $y_m(x)$ , and if we denote the remaining misclassified points by  $M_m$ , then we can in turn rewrite the error function in the form.

$$E = e^{-\frac{\alpha_m}{2}} \sum_{n \in T_m} w_n^{(m)} + e^{\frac{\alpha_m}{2}} \sum_{n \in M_m} w_n^{(m)}$$

$$= \left(e^{\frac{\alpha_m}{2}} - e^{-\frac{\alpha_m}{2}}\right) \sum_{n=1}^N w_n^{(m)} I(y_m(x_n) \neq t_n) + e^{-\frac{\alpha_m}{2}} \sum_{n=1}^N w_n^{(m)}$$

$$J_m = \sum_{n=1}^N w_n^{(m)} I(y^{(m)}(x_n) \neq t_n)$$

When we minimize this with respect to  $y_m(x)$ , we see that the second term is constant, and so this is equivalent to the procedure in Adaboost Algorithm 2-(a) and we get  $\varepsilon_m$  which is the same as in Adaboost. Similarly, minimizing with respect to  $\alpha_m$ , we get the  $\alpha_m$  the same as  $\alpha_m$  in Adaboost.



# Loss functions for boosting

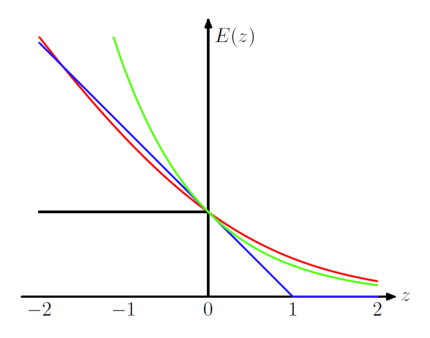
► The exponential loss function that is minimized by the AdaBoost algorithm is

$$E = \sum_{n=1}^{N} \exp\{-t_n f_m(\boldsymbol{x}_n)\}\$$

- Why these loss function?
- What properties do this loss function have?
- Can we choose other loss functions?
- ▶ Those questions opens the door to a wide range of boosting-like algorithms, including multiclass extensions, by altering the choice of loss function. It also motivates the extension to regression problems



# **Comparing to Other Loss Function**



Comparing exponential loss, logistic loss, and hinge loss

Error function grows exponentially with |ty(x)|, it penalizes large negative values of ty(x), thus be much less robust to outliers or misclassified data



## Summary

- They are two different Ensemble Paradigms
  - Boosting is sequential ensemble methods, where the base learners are generated sequentially.
  - Bagging is parallel ensemble methods: where the base learners are generated in parallel.
  - Boosting exploit the *dependence* between the base learners, since the overall performance can be boosted in a residual-decreasing way.
  - Bagging exploit the *independence* between the base learners, since the error can be reduced dramatically by combining independent base learners.