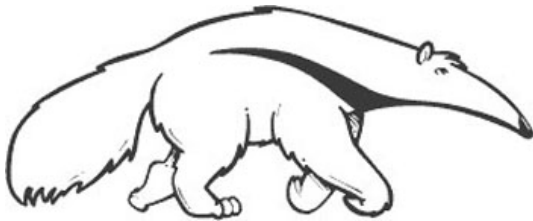


CS178: Machine Learning and Data Mining

Ensembles of Learners

Prof. Alexander Ihler



Ensemble methods

- Why learn one classifier when you can learn many?
- Ensemble: combine many predictors
 - (Weighted) combinations of predictors
 - May be same type of learner or different



Various options for getting help:



“Who wants to be a millionaire?”

Simple ensembles

- “Committees”
 - Unweighted average / majority vote
- Weighted averages
 - Up-weight “better” predictors
 - Ex: Classes: +1 , -1 , weights alpha:
 $\hat{y}_1 = f_1(x_1, x_2, \dots)$
 $\hat{y}_2 = f_2(x_1, x_2, \dots) \quad \Rightarrow \quad \hat{y}_e = \text{sign}(\sum \alpha_i \hat{y}_i)$
...

“Stacked” ensembles

- Train a “predictor of predictors”
 - Treat individual predictors as features

$$\hat{y}_1 = f_1(x_1, x_2, \dots)$$

$$\hat{y}_2 = f_2(x_1, x_2, \dots) \quad \Rightarrow \quad \hat{y}_e = f_e(\hat{y}_1, \hat{y}_2, \dots)$$

...

- Similar to multi-layer perceptron idea
- Special case: binary, f_e linear \Rightarrow weighted vote
- Can train stacked learner f_e on validation data
 - Avoids giving high weight to overfit models

Mixtures of experts

- Can make weights depend on x
 - Weight $\alpha_z(x)$ indicates “expertise”
 - Combine using weighted average (or even just pick largest)

Example

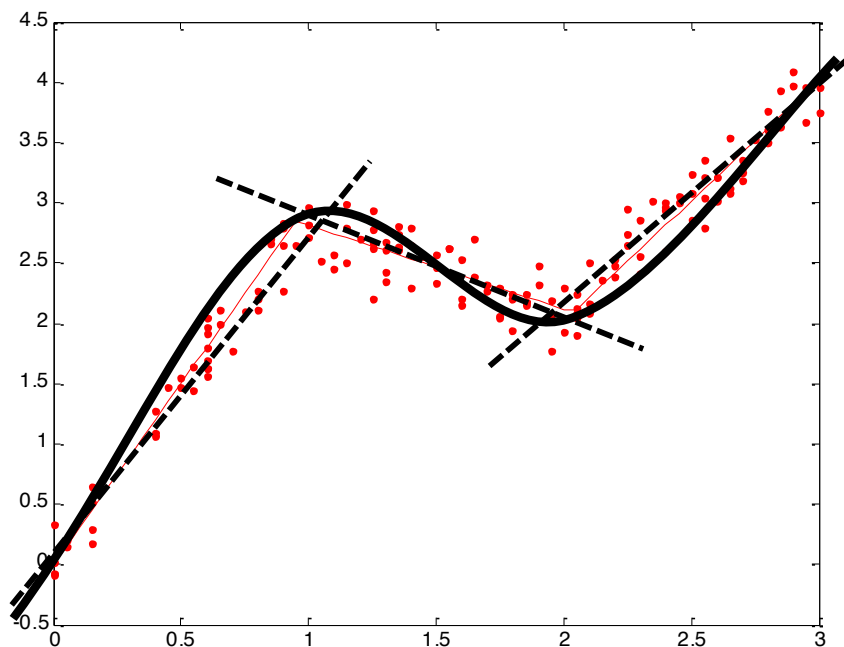
Weighted average:

$$f(x; \omega, \theta) = \sum_z \alpha_z(x; \omega) f_z(x; \theta_z)$$

Weights: (multi) logistic regression

$$\alpha_z(x; \omega) = \frac{\exp(x \cdot \omega^z)}{\sum_c \exp(x \cdot \omega^c)}$$

If loss, learners, weights are all differentiable, can train jointly...



Mixture of three linear predictor experts

Machine Learning

Ensembles: Bagging

Ensembles: Gradient Boosting

Ensembles: Ada Boost

Ensemble methods

- Why learn one classifier when you can learn many?
 - “Committee”: learn K classifiers, average their predictions
- “Bagging” = bootstrap aggregation
 - Learn many classifiers, each with only part of the data
 - Combine through model averaging
- Remember overfitting: “memorize” the data
 - Used test data to see if we had gone too far
 - Cross-validation
 - Make many splits of the data for train & test
 - Each of these defines a classifier
 - Typically, we use these to check for overfitting
 - Could we instead combine them to produce a better classifier?

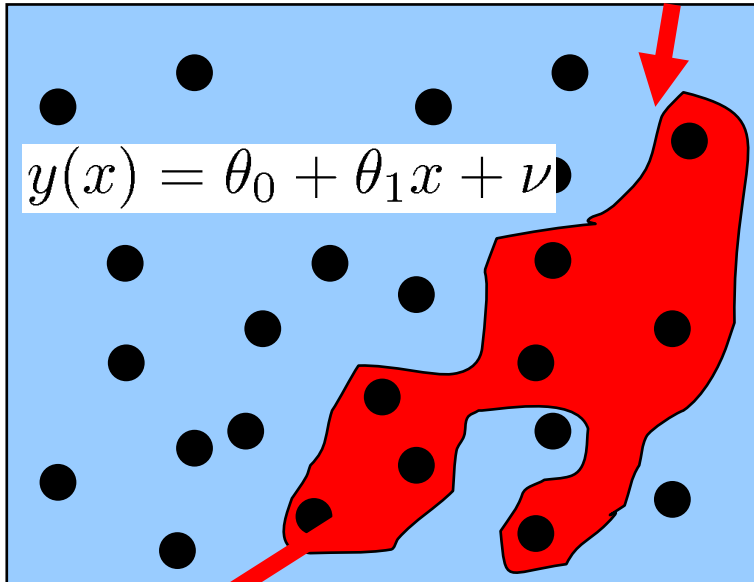
Bagging

- Bootstrap
 - Create a random subset of data by sampling
 - Draw m' of the m samples, with replacement (some variants w/o)
 - Some data left out; some data repeated several times
- Bagging
 - Repeat K times
 - Create a training set of $m' \leq m$ examples
 - Train a classifier on the random training set
 - To test, run each trained classifier
 - Each classifier votes on the output, take majority
 - For regression: each regressor predicts, take average
- Notes:
 - Some complexity control: harder for each to memorize data
 - Doesn't work for linear models (average of linear functions is linear function), but perceptrons OK (linear + threshold = nonlinear)

Bias / variance

“The world”

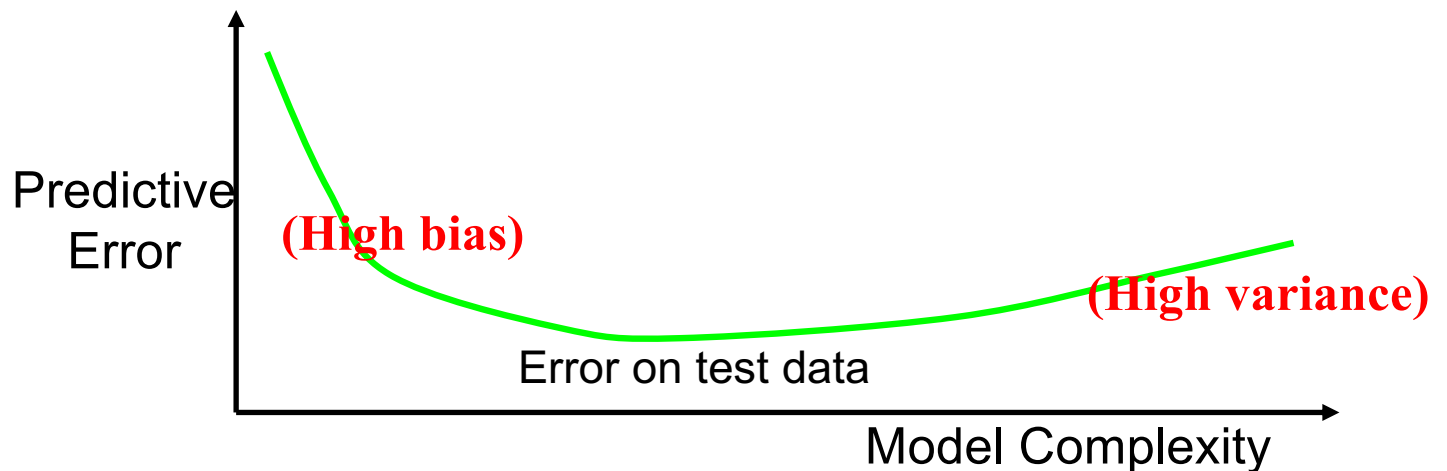
Data we observe



$$y(x) = \theta_0 + \theta_1 x + \nu$$

$$\hat{y}(x) = \hat{\theta}_0 + \hat{\theta}_1 x$$

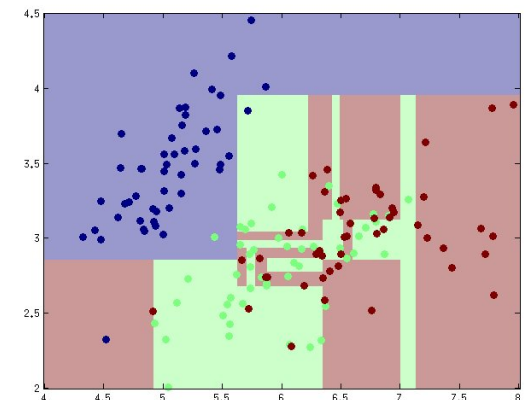
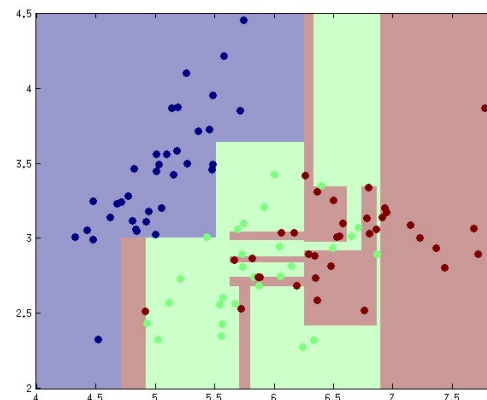
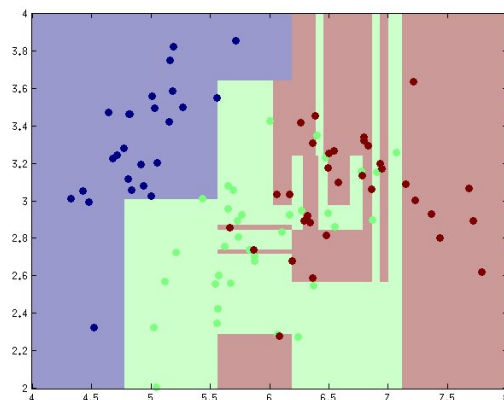
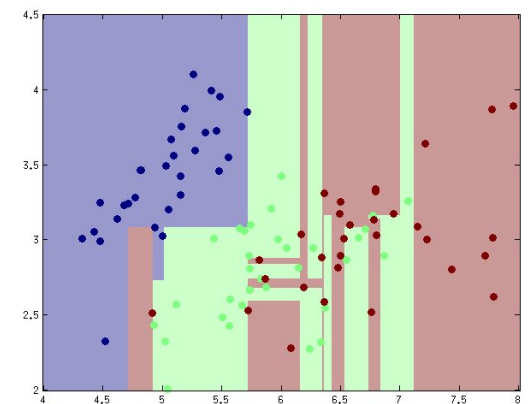
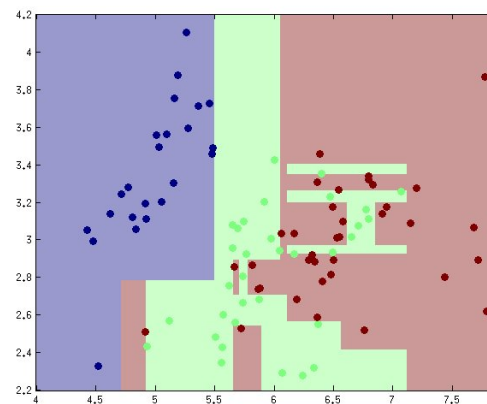
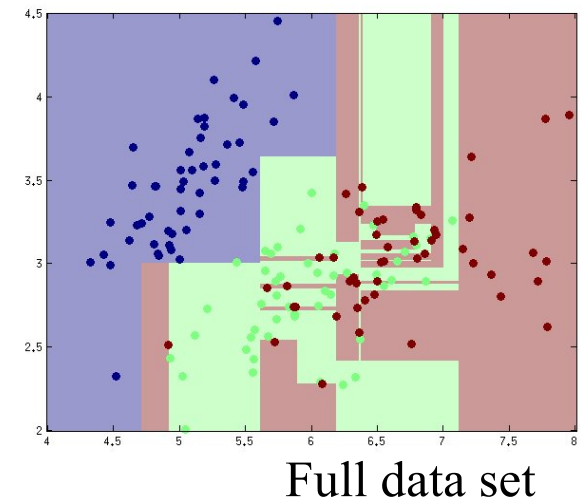
- We only see a little bit of data
- Can decompose error into two parts
 - Bias – error due to model choice
 - Can our model represent the true best predictor?
 - Gets better with more complexity
 - Variance – randomness due to data size
 - Better w/ more data, worse w/ complexity



Bagged decision trees

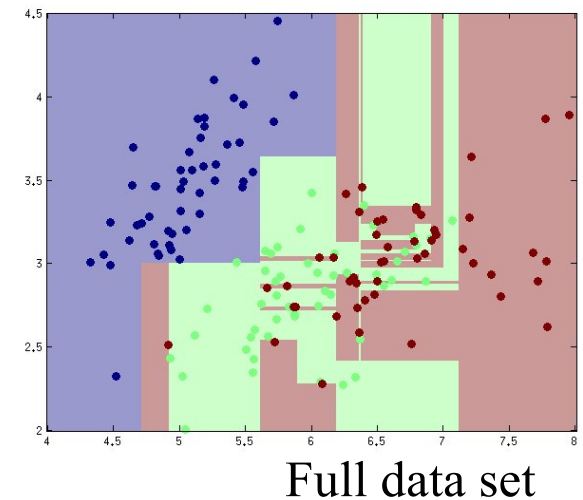
- Randomly resample data
- Learn a decision tree for each
 - No max depth = very flexible class of functions
 - Learner is low bias, but high variance

Sampling:
simulates “equally likely”
data sets we could have
observed instead, &
their classifiers

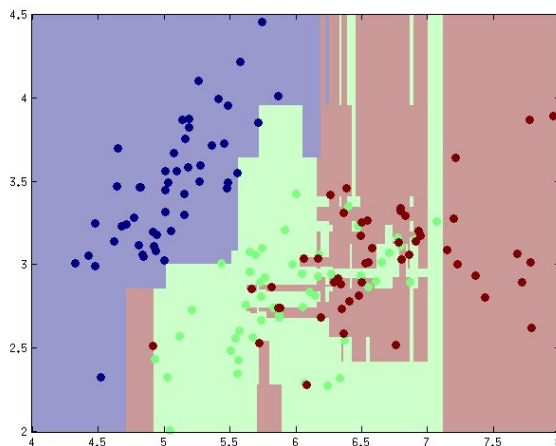


Bagged decision trees

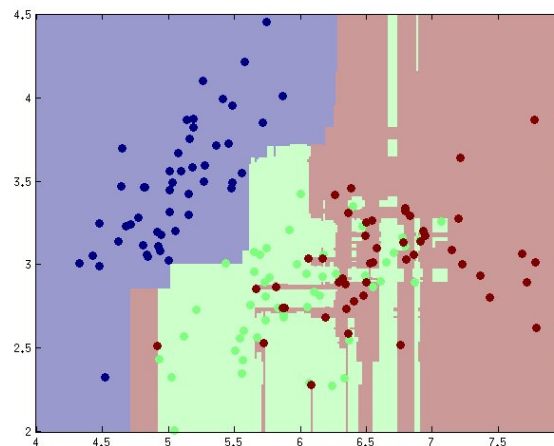
- Average over collection
 - Classification: majority vote
- Reduces memorization effect
 - Not every predictor sees each data point
 - Lowers effective “complexity” of the overall average
 - Usually, better generalization performance
 - Intuition: reduces variance while keeping bias low



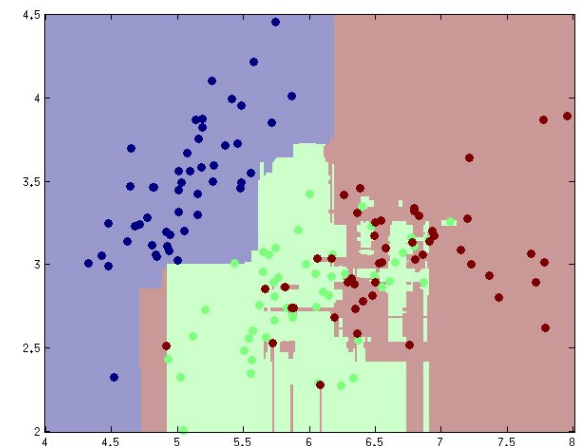
Avg of 5 trees



Avg of 25 trees



Avg of 100 trees



Bagging in Python

```
# Load data set X, Y for training the ensemble...
```

```
m,n = X.shape
```

```
classifiers = [ None ] * nBag
```

```
# Allocate space for learners
```

```
for i in range(nBag):
```

```
    ind = np.floor( m * np.random.rand(nUse) ).astype(int) # Bootstrap sample a data set:
```

```
    Xi, Yi = X[ind,:], Y[ind] # select the data at those indices
```

```
    classifiers[i] = ml.MyClassifier(Xi, Yi) # Train a model on data Xi, Yi
```

```
# test on data Xtest
```

```
mTest = Xtest.shape[0]
```

```
predict = np.zeros( (mTest, nBag) ) # Allocate space for predictions from each model
```

```
for i in range(nBag):
```

```
    predict[:,i] = classifiers[i].predict(Xtest) # Apply each classifier
```

```
# Make overall prediction by majority vote
```

```
predict = np.mean(predict, axis=1) > 0 # if +1 vs -1
```

Random forests

- Bagging applied to decision trees
- Problem
 - With lots of data, we usually learn the same classifier
 - Averaging over these doesn't help!
- Introduce extra variation in learner
 - At each step of training, only allow a (random) subset of features
 - Enforces diversity (“best” feature not available)
 - Keeps bias low (every feature available eventually)
 - Average over these learners (majority vote)

```
# in FindBestSplit(X,Y):  
  for each of a subset of features  
    for each possible split  
      Score the split (e.g. information gain)  
    Pick the feature & split with the best score  
  Recurse on left & right splits
```

Summary

- Ensembles: collections of predictors
 - Combine predictions to improve performance
- Bagging
 - “Bootstrap aggregation”
 - *Reduces* complexity of a model class prone to overfit
 - In practice
 - Resample the data many times
 - For each, generate a predictor on that resampling
 - Plays on bias / variance trade off
 - Price: more computation per prediction

Machine Learning

Ensembles: Bagging

Ensembles: Gradient Boosting

Ensembles: Ada Boost

Ensembles

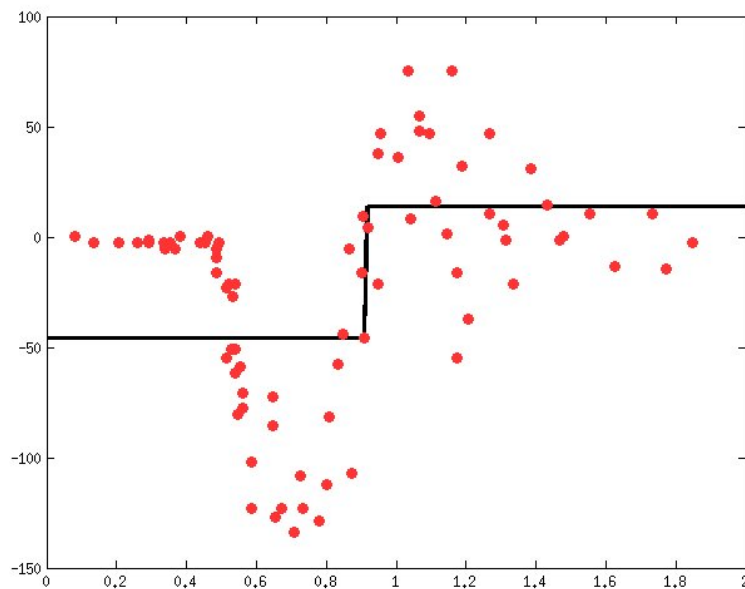
- Weighted combinations of predictors
- “Committee” decisions
 - Trivial example
 - Equal weights (majority vote / unweighted average)
 - Might want to weight unevenly – up-weight better predictors
- Boosting
 - Focus new learners on examples that others get wrong
 - Train learners sequentially
 - Errors of early predictions indicate the “hard” examples
 - Focus later predictions on getting these examples right
 - Combine the whole set in the end
 - Convert many “weak” learners into a complex predictor

Gradient boosting

- Learn a regression predictor
- Compute the error residual
- Learn to predict the residual

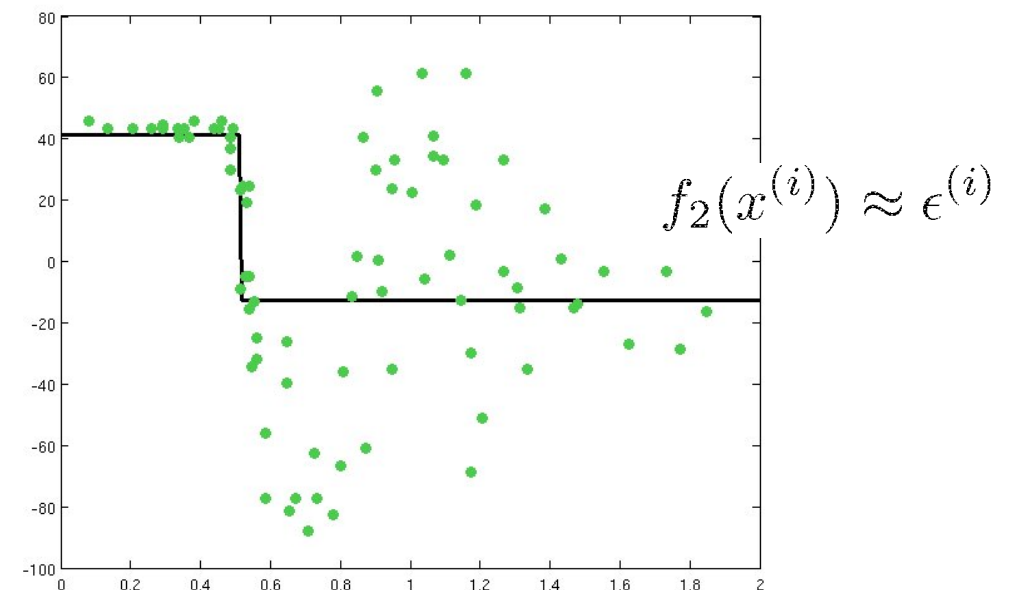
Learn a simple predictor...

$$f_1(x^{(i)}) \approx y^{(i)}$$



Then try to correct its errors

$$\epsilon^{(i)} = y^{(i)} - f_1(x^{(i)})$$



Gradient boosting

- Learn a regression predictor
- Compute the error residual
- Learn to predict the residual

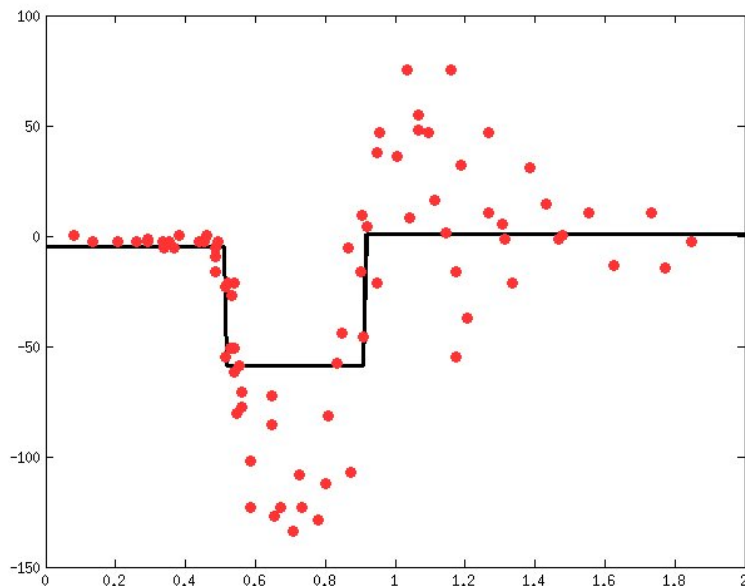
$$f_1(x^{(i)}) \approx y^{(i)}$$

$$\epsilon^{(i)} = y^{(i)} - f_1(x^{(i)})$$

$$f_2(x^{(i)}) \approx \epsilon^{(i)}$$

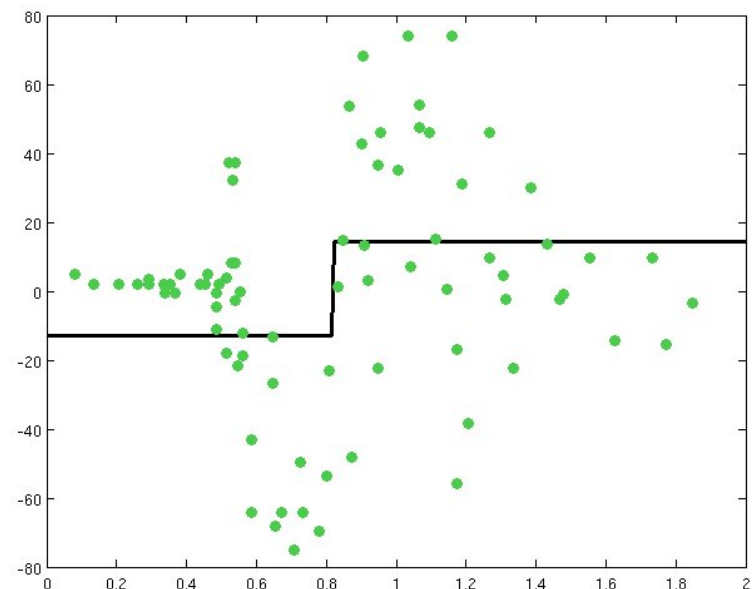
Combining gives a better predictor...

$$\Rightarrow f_1(x^{(i)}) + f_2(x^{(i)}) \approx y^{(i)}$$



Can try to correct its errors also, & repeat

$$\epsilon_2^{(i)} = y^{(i)} - f_1(x^{(i)}) - f_2(x^{(i)}) \quad \dots$$

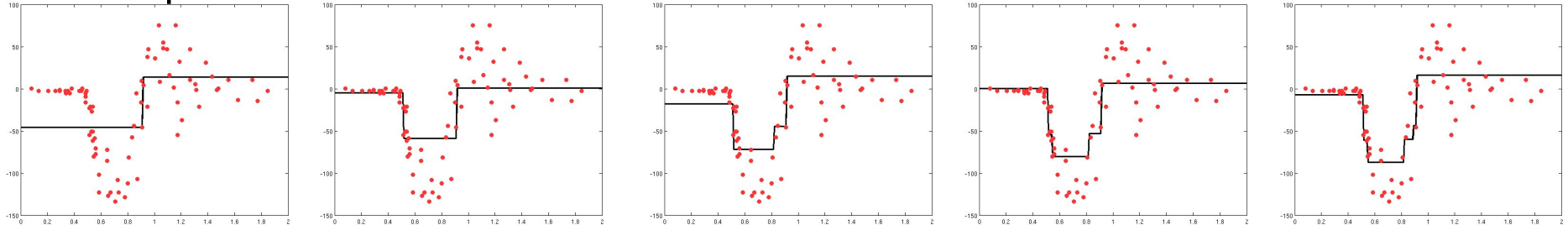


Gradient boosting

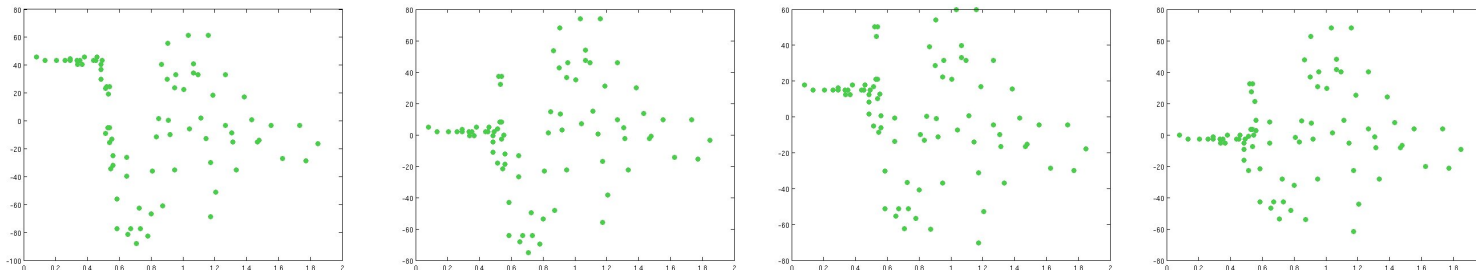
- Learn sequence of predictors
- Sum of predictions is increasingly accurate
- Predictive function is increasingly complex

$$y^{(i)} \approx \sum_z f_z(x^{(i)})$$

Data & prediction function



Error residual



Gradient boosting

- Make a set of predictions $\hat{y}[i]$
- The “error” in our predictions is $J(y, \hat{y})$
 - For MSE: $J(.) = \sum (y[i] - \hat{y}[i])^2$
- We can “adjust” \hat{y} to try to reduce the error
 - $\hat{y}[i] = \hat{y}[i] + \text{alpha } f[i]$
 - $f[i] \approx \nabla J(y, \hat{y}) = (y[i] - \hat{y}[i])$ for MSE
- Each learner is estimating the gradient of the loss function
- Gradient descent: take sequence of steps to reduce J
 - Sum of predictors, weighted by step size alpha

Gradient boosting in Python

```
# Load data set X, Y ...
```

```
learner = [None] * nBoost      # storage for ensemble of models
```

```
alpha = [1.0] * nBoost        # and weights of each learner
```

```
mu = Y.mean()                  # often start with constant "mean" predictor
```

```
dY = Y - mu                    # subtract this prediction away
```

```
for k in range( nBoost ):
```

```
    learner[k] = ml.MyRegressor( X, dY )    # regress to predict residual dY using X
```

```
    alpha[k] = 1.0                # alpha: "learning rate" or "step size"
```

```
    # smaller alphas need to use more classifiers, but may predict better given enough of them
```

```
    # compute the residual given our new prediction:
```

```
    dY = dY - alpha[k] * learner[k].predict(X)
```

```
# test on data Xtest
```

```
mTest = Xtest.shape[0]
```

```
predict = np.zeros( (mTest,) ) + mu      # Allocate space for predictions & add 1st (mean)
```

```
for k in range(nBoost):
```

```
    predict += alpha[k] * learner[k].predict(Xtest) # Apply predictor of next residual & accum
```

Summary

- Ensemble methods
 - Combine multiple classifiers to make “better” one
 - Committees, average predictions
 - Can use weighted combinations
 - Can use same or different classifiers
- Gradient Boosting
 - Use a simple regression model to start
 - Subsequent models predict the error residual of the previous predictions
 - Overall prediction given by a weighted sum of the collection

Machine Learning

Ensembles: Bagging

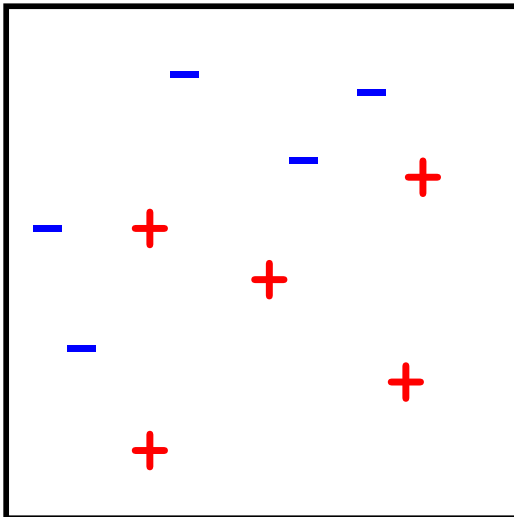
Ensembles: Gradient Boosting

Ensembles: Ada Boost

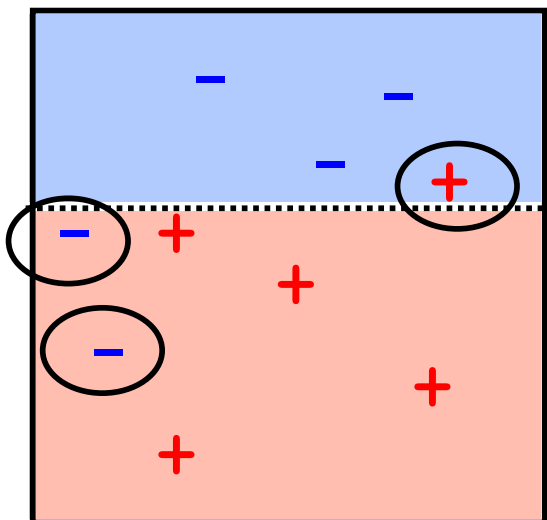
Boosting example

Classes +1, -1

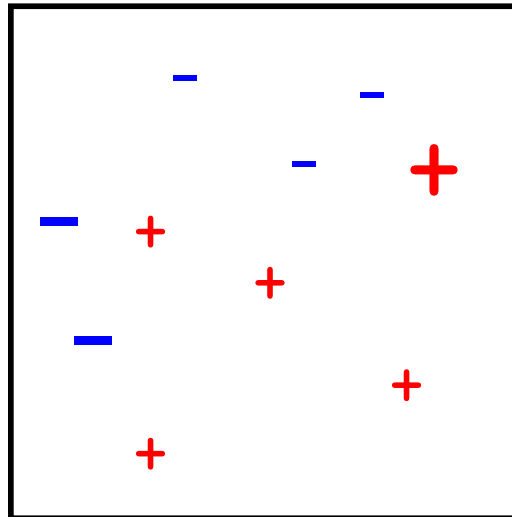
Original data set, D_1



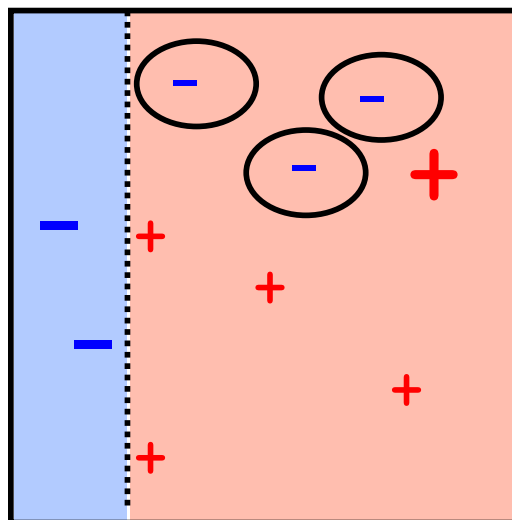
Trained classifier



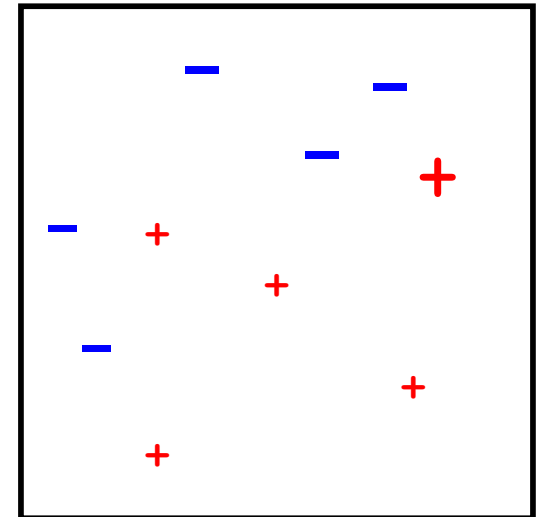
Update weights, D_2



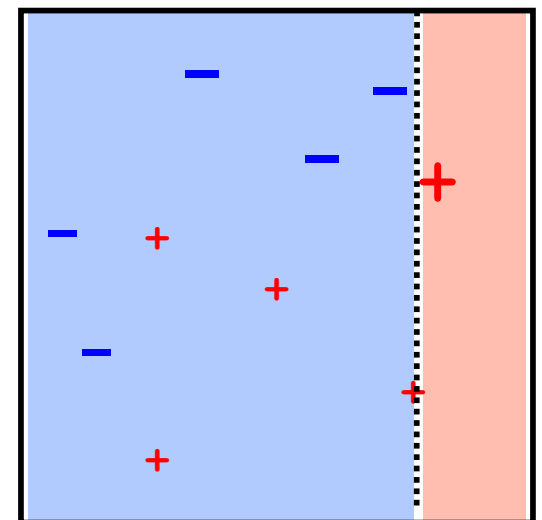
Trained classifier



Update weights, D_3



Trained classifier



Minimizing weighted error

- So far we've mostly minimized unweighted error
- Minimizing weighted error is no harder:

Unweighted average loss:

$$J(\theta) = \frac{1}{m} \sum_i J_i(\theta, x^{(i)})$$

Weighted average loss:

$$J(\theta) = \sum_i w_i J_i(\theta, x^{(i)})$$

For any loss (logistic MSE, hinge, ...)

$$J(\theta, x^{(i)}) = (\sigma(\theta x^{(i)}) - y^{(i)})^2$$

$$J(\theta, x^{(i)}) = \max[0, 1 - y^{(i)} \theta x^{(i)}]$$

To learn decision trees, find splits to optimize *weighted* impurity scores:

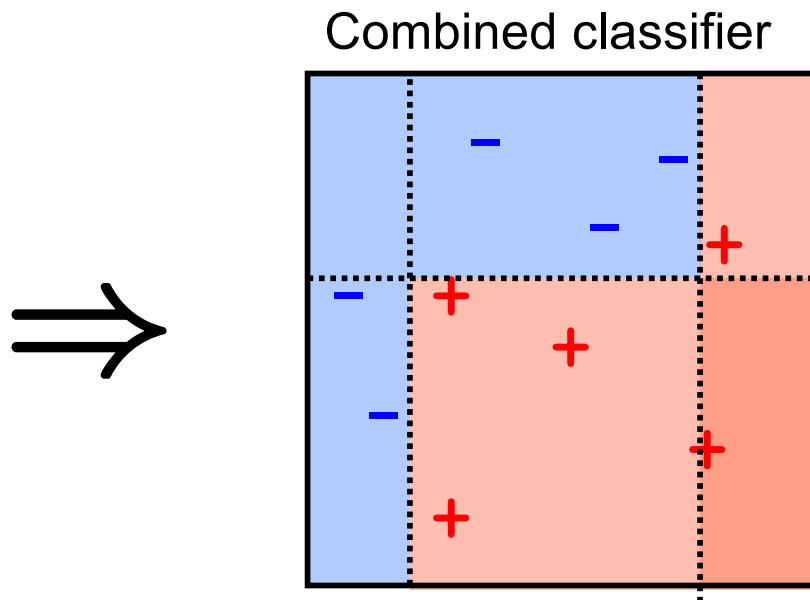
$p(+1)$ = total weight of data with class +1

$p(-1)$ = total weight of data with class -1 $\Rightarrow H(p)$ = impurity

Boosting example

Weight each classifier and combine them:

$$.33 * \begin{array}{|c|} \hline \text{blue} \\ \hline \text{red} \\ \hline \end{array} + .57 * \begin{array}{|c|} \hline \text{blue} \\ \hline \text{red} \\ \hline \end{array} + .42 * \begin{array}{|c|} \hline \text{blue} \\ \hline \text{red} \\ \hline \end{array} \geq 0$$



1-node decision trees
“decision stumps”
very simple classifiers

AdaBoost = “adaptive boosting”

- Pseudocode for AdaBoost

Classes $\{+1, -1\}$

```
# Load data set X, Y ... ; Y assumed +1 / -1
for i in range(nBoost):
    learner[i] = ml.MyClassifier( X, Y, weights=wts ) # train a weighted classifier
    Yhat = learner[i].predict(X)
    e = wts.dot( Y != Yhat ) # compute weighted error rate
    alpha[i] = 0.5 * np.log( (1-e)/e )
    wts *= np.exp( -alpha[i] * Y * Yhat ) # update weights
    wts /= wts.sum() # and normalize them
```

```
# Final classifier:
predict = np.zeros( (mTest,) )
for i in range(nBoost):
    predict += alpha[i] * learner[i].predict(Xtest) # compute contribution of each model
predict = np.sign(predict) # and convert to +1 / -1 decision
```

- Notes

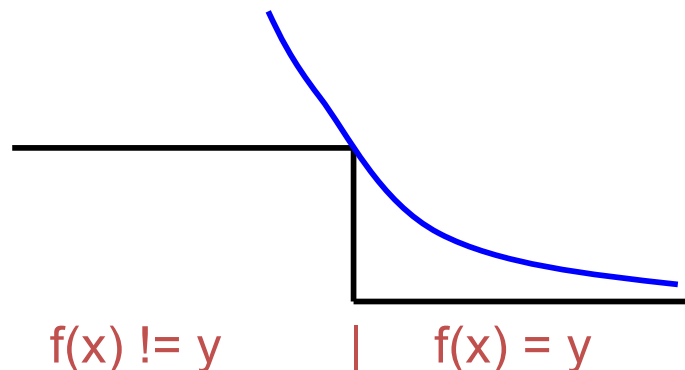
- $e > .5$ means classifier is not better than random guessing
- $Y * Yhat > 0$ if $Y == Yhat$, and weights decrease
- Otherwise, they increase

AdaBoost theory

- Minimizing classification error was difficult
 - For logistic regression, we minimized MSE or NLL instead
 - Idea: low MSE => low classification error
- Example of a surrogate loss function
- AdaBoost also corresponds to a surrogate loss function

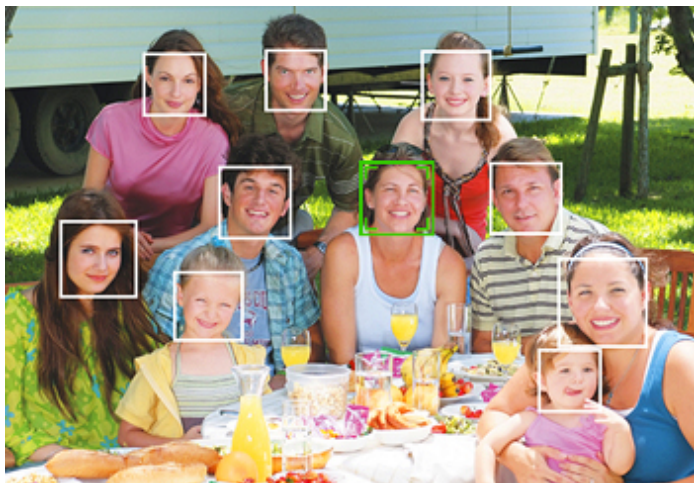
$$C_{ada} = \sum_i \exp[-y^{(i)} f(x^i)]$$

- Prediction is $\hat{y} = \text{sign}(f(x))$
 - If same as y , loss < 1 ; if different, loss > 1 ; at boundary, loss = 1
- This loss function is smooth & convex (easier to optimize)



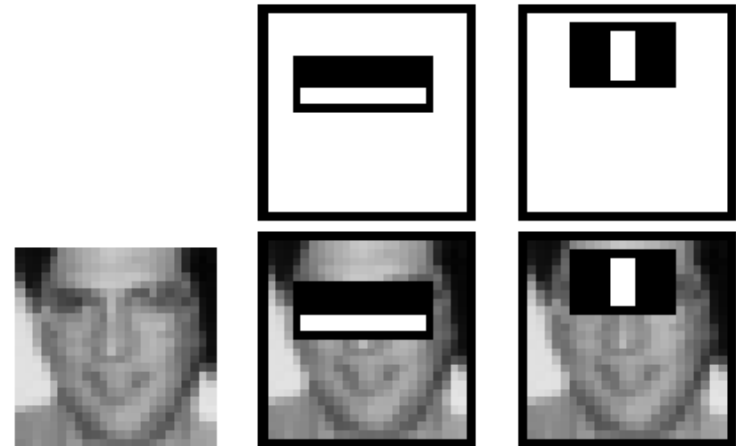
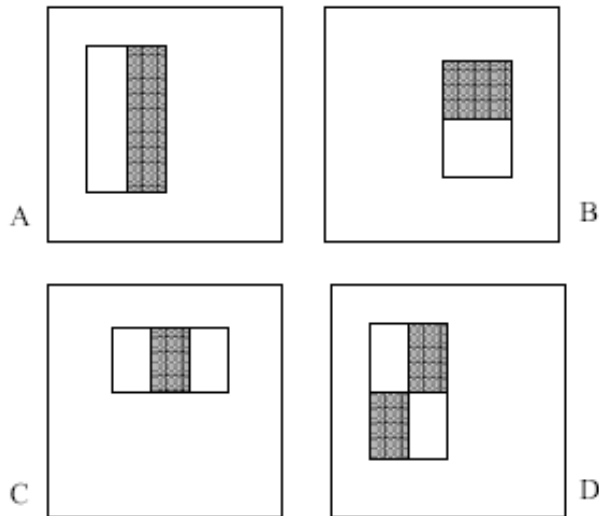
AdaBoost example: Viola-Jones

- Viola-Jones face detection algorithm
- Combine lots of very weak classifiers
 - Decision stumps = threshold on a single feature
- Define lots and lots of features
- Use AdaBoost to find good features
 - And weights for combining as well



Haar wavelet features

- Four basic types.
 - They are easy to calculate.
 - The white areas are subtracted from the black ones.
 - A special representation of the sample called the **integral image** makes feature extraction faster.



Training a face detector

- Wavelets give ~100k features
- Each feature is one possible classifier
- To train: iterate from 1:T
 - Train a classifier on each feature using weights
 - Choose the best one, find errors and re-weight
- This can take a long time... (lots of classifiers)
 - One way to speed up is to not train very well...
 - Rely on adaboost to fix “even weaker” classifier
- Lots of other tricks in “real” Viola-Jones
 - Cascade of decisions instead of weighted combo
 - Apply at multiple image scales
 - Work to make computationally efficient

Summary

- Ensemble methods
 - Combine multiple classifiers to make “better” one
 - Committees, majority vote
 - Weighted combinations
 - Can use same or different classifiers
- Boosting
 - Train sequentially; later predictors focus on mistakes by earlier
- Boosting for classification (e.g., AdaBoost)
 - Use results of earlier classifiers to know what to work on
 - Weight “hard” examples so we focus on them more
 - Example: Viola-Jones for face detection