COMPSCI762: Foundations of Machine Learning Ensembles

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Motivation







This week we will cover



Motivation

Ensembles

Averaging

Random Forests

AdaBoost

XGBoost

Partially based on Slides from University of British Columbia

Ensembles

Ensembles



- Ensembles are classifiers that have classifiers as input
- With great names:
 - Averaging
 - Boosting
 - Bootstrapping
 - Bagging
 - Cascading
 - Random Forests
 - Stacking
- Ensemble methods often have higher accuracy than the input classifiers
- How would you build an ensemble classifier?

Ensembles



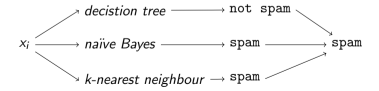
- Remember the fundamental trade-off
 - 1. E_{train} : How small you can make the training error vs.
 - 2. E_{approx} : How well training error approximates the test error
- Goal of ensemble methods is that the ensemble classifier
 - Does much better on one of these than the individual classifiers
 - Doesn't do too much worse on the other
- This suggests two types of ensemble methods
 - 1. **Boosting**: Improves training error of classifiers with high E_{train}
 - 2. Averaging: Improves approximation error of classifiers with high E_{approx}

Averaging





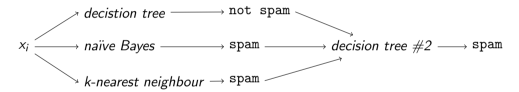
- Input to averaging is the predictions of a set of models, for example
 - Decision trees make one prediction
 - Naïve Bayes makes another prediction
 - KNN makes another prediction
- Simple model averaging:
 - Take the mode of the predictions (or average probabilities if probabilistic)



Stacking



- A common variation of averaging is stacking
 - Fit another classifier that use the predictions as features



- How does the training set of *decision tree #2* look like?
- Averaging or stacking often performs better than individual models
 - Typically used by Kaggle winners
 - E.g., the Netflix \$1M user-rating competition winner was a stacked classifier

Why does averaging work?



- Consider 3 binary classifiers, each independently correct with probability 0.8
- With simple averaging, the ensemble is correct if we have at least 2 right:

$$P(\text{all 3 right}) = 0.8^3$$
 = 0.512
 $P(\text{2 right, 1 wrong}) = 3 * 0.8^2(1 - 0.8)$ = 0.384
 $P(\text{1 right, 2 wrong}) = 3 * (1 - 0.8)^2 0.8$ = 0.096
 $P(\text{all 3 wrong}) = (1 - 0.8)^3$ = 0.512

- So the ensemble is right with a probability of 0.896 (=0.512+0.384)
- Note:
 - For averaging to work, classifiers need to be at least somewhat independent
 - You also want the probability of being right to be > 0.5, otherwise it will do much worse
 - Probabilities also shouldn't be to different (otherwise, it might be better to take most accurate)

Why does averaging work?



- Consider a set of classifiers that makes these predictions:
 - Classifier 1: spam
 - Classifier 2: spam
 - Classifier 3: spam
 - Classifier 4: not spam
 - Classifier 5: spam
 - Classifier 6: not spam
 - Classifier 7: spam
 - Classifier 8: spam
 - Classifier 9: spam
 - Classifier 10: spam
- I these independently get 80% accuracy, the mode will be close to 100%
 - In practice errors won't be completely independent due to noise in the labels

Why can averaging work?



- Why can averaging lead to better results?
- Consider classifiers that overfit (like deep decision trees)
 - If they all overfit in exactly the same way, averaging does nothing Why?
- But if they make **independent** errors
 - Probability that average is wrong can be lower than for each classifier
 - Less attention to specific overfitting of each classifier

Random Forests





- Random forests average a set of deep decision trees
 - Tend to be one of the best *out of the box* classifiers
 - Often close to the best performance of any method on the first run
 - And predictions are very fast
- Does averaging work if you use trees with the same parameters?
- Do deep decision trees make independent errors?
 - No: with the same training data you'll get the same decision tree
- Two key ingredients in random forests:
 - Bootstrapping
 - Random trees





- Start with a standard deck of 52 cards
 - 1. Sample a random card, put it back and re-shuffle
 - 2. Sample a random card, put it back and re-shuffle
 - Sample a random card, put it back and re-shuffle
 - 52 Sample a random card, put it back and re-shuffle
- Make a new deck of the 52 samples

Bootstrap Sampling



- The new 52-card deck is called a **bootstrap sample**
- Some cards will be missing, and some cards will be duplicated
 - So calculations on the bootstrap sample will give different results than original data
- However, the bootstrap sample roughly maintains trends:
 - Roughly 25% of the cards will be diamonds
 - Roughly 3/13 of the cards will be *face* cards
 - There will be roughly four 10 cards
- Common use: compute a statistic based on several bootstrap samples
 - Gives you an idea of how the statistic varies as you vary the data





- Bootstrap sample of a list of n examples
 - A new set of size n chosen independently with replacement

```
forall i \in 1, \ldots, n do
    j = rand(1:n); // pick a random number from \{1, 2, ..., n\}
X_{bootstrap}[i,:] = X[j,:] // use the random sample
end
```

- Gives new dataset of n examples, with some duplicated and some missing
 - For large n, approximately 63% of original examples are included (see next slide)
- Bagging: using bootstrap samples for ensemble learning Bootstrap Aggregating
 - Generate several bootstrap samples of the examples (x_i, y_i)
 - Fit a classifier to each bootstrap sample
 - At test time, average the predictions





■ Probability of an arbitrary x_i being selected in a bootstrap sample

```
p(\text{selected at least once in } n \text{ trials})
= 1 - p(\text{not selected in any of } n \text{ trials})
= 1 - (p(\text{not selected in one trial})^n //\text{trials are independent}
= 1 - (1 - \frac{1}{n})^n //prob = \frac{n-1}{n} \text{ for choosing any of the } n-1 \text{ other samples}
\approx 1 - \frac{1}{e} //((1 - \frac{1}{n})^n \to e^{-1} \text{ as } n \to \inf
\approx 0.632
```

Random Forest Ingredient 2: Random Trees



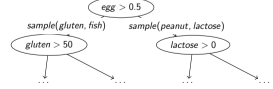
- For each split in a random tree model
 - Randomly sample a small number of possible features (typically \sqrt{d})
 - Only consider these random features when searching for the optimal rule
- Random tree 1: sample(milk, oranges)

 $\begin{array}{c} \textit{milk} > 0.5 \\ \textit{sample(egg, peanut)} & \textit{sample(lactose, gluten)} \\ \textit{egg} > 1 & \textit{lactose} > 0 \\ \end{array}$



Random tree 2:

sample(egg, lactose)







- Splits will tend to use different features in different trees
 - They will still overfit, but hopefully errors will be more independent
- So the average tends to have a much lower test error
- Empirically, random forests are one of the best classifiers
- Fernandez-Delgado et al. [2014]
 - Compared 179 classifiers on 121 datasets
 - Random forests are most likely to be the best classifier

AdaBoost





- A classic boosting algorithm for binary classification is AdaBoost
- AdaBoost assumes we have a base binary classifier that
 - Is simple enough that it does not overfit much
 - Can obtain > 50% weighted accuracy on any dataset.
- Example: decision stumps or low-depth decision trees
 - Easy to modify stumps/trees to use weighted accuracy as score

AdaBoost: Classic Boosting Algorithm



- Overview of AdaBoost:
 - 1. Fit a classifier on the training data
 - 2. Give a higher weight to examples that the classifier got wrong
 - 3. Fit a classifier on the weighted training data
 - 4. Go back to 2
 - Weight gets exponentially larger each time you are wrong.
- Final prediction: weighted vote of individual classifier predictions
 - Trees with higher (weighted) accuracy get higher weight

AdaBoost: Classic Boosting Algorithm



- Are decision stumps a good base classifier?
 - They tend not to overfit
 - Easy to get > 50% weighted accuracy
- Base classifiers that don't work:
 - Deep decision trees (no errors to boost)
 - Decision stumps with infogain (does not guarantee > 50% weighted accuracy)
 - lacktriangle Weighted logistic regression (does not guarantee >50% weighted accuracy)





- AdaBoost with shallow decision trees gives fast/accurate classifiers
 - Classically viewed as one of the best off the shelf classifiers
 - Procedure originally came from ideas in learning theory
- Many attempts to extend theory beyond binary case
 - Led to gradient boosting, which is like gradient descent with trees
- Modern boosting methods:
 - Look like AdaBoost, but don't necessarily have it as a special case

XGBoost



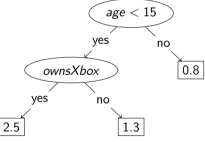


- Boosting has seen a recent resurgence, partially due to XGBoost:
 - A boosting implementation that allows huge datasets
 - Has been part of many recent winners of Kaggle competitions
- As base classifier, XGBoost uses regularized regression trees





- Regression trees used in XGBoost
 - Each split is based on 1 feature
 - Each leaf gives a real-valued prediction
 - Example: How many hours of video games per day?



(Sorry for this example!)

■ We would predict 2.5 hours for a 14-year-old who owns an Xbox

Regression Trees



- How can we fit a regression tree?
- Simple approach:
 - Predict: at each leaf, predict mean of the training y_i assigned to the leaf
 - Weight w_L at leaf L is set to $mean(y_i)$ among y_i at the leaf node
 - Train: set the w_L values by minimizing the squared error

$$f(w_1, w_2, \ldots) = \sum_{i=1}^n (w_{L_i} - y_i)^2$$

- Same speed as fitting decision trees from earlier in the semester
 - Use mean instead of mode, and use squared error instead of accuracy / infogain
- Use greedy strategy for growing tree, as earlier





- Consider an ensemble of regression trees
 - For an example i, they each make a continuous prediction







$$\hat{y}_{i_3}=0.1$$

■ In XGBoost, final prediction is the sum of individual predictions

$$\hat{y}_i = \hat{y}_{i_1} + \hat{y}_{i_2} + \hat{y}_{i_3} + \ldots + \hat{y}_{i_k}$$

= 0.2 + (-0.1) + 0.1 + \ldots + (-0.01)







- Notice we are **not using the mean** as we would with random forests
 - What we do instead?
 - In boosting, each tree is **not individually trying to predict the true** y_i value (we assume they underfit)
 - Instead, each new tree tries to fix the prediction made by the old trees, so that sum is y_i





■ Consider the following *gradient tree boosting* procedure:

```
    Tree[1] = fit(X, y)
    ŷ = Tree[1].predict(X)
    Tree[2] = fit(X, y − ŷ)
    ŷ = ŷ + Tree[2].predict(X)
    Tree[3] = fit(X, y − ŷ)
    ŷ = ŷ + Tree[3].predict(X)
    Tree[4] = fit(X, y − ŷ)
    ŷ = ŷ + Tree[4].predict(X)
```

- **Each** tree is trying to predict *residuals* $(\hat{y}_i y_i)$ of current prediction
 - "True label is 0.9, old prediction is 0.8, so I can improve \hat{y}_i by predicting 0.1"





- Procedure monotonically decreases the training error
 - As long as not all $w_L = 0$, each tree decreases training error
- Can it overfit?
 - It can overfit if trees are too deep or you have too many trees
 - To restrict depth, add L0-regularization (stop splitting if $w_L = 0$)

$$f(w_1, w_1, \ldots) = \sum_{i=1}^n (w_{L_i} - r_i) + \lambda_0 ||w||_0$$

- Only split if you decrease squared error by λ_0 .
- To further fight overfitting, XGBoost also adds L2-regularization of w

$$f(w_1, w_1, \ldots) = \sum_{i=1}^n (w_{L_i} - r_i) + \lambda_0 ||w||_0 + \lambda_2 ||w||^2$$





- Instead of pruning trees if score does not improve, grows full trees
 - And then prunes parts that don't improve score with L0-regularizer added
- Cost of fitting trees in XGBoostis same as usual decision tree cost
 - XGBoost includes a lot of tricks to make this efficient
 - But cannot be done in parallel like random forest Why?
- In XGBoost, it's the residuals that act like the weights in AdaBoost
 - Focuses on decreasing error in examples with large residuals
- How do you maintain efficiency if not using squared error?
 - For non-quadratic losses like logistic, there is no closed-form solution
 - Approximates non-quadratic losses with second-order Taylor expansion
 - Maintains least squares efficiency for other losses (by approximating with quadratic)

Summary



- Ensembles combine predictions of base classifiers
- Averaging
 - Improves predictions of multiple classifiers if errors are independent
- Bagging
 - Ensemble method where we apply same classifier to bootstrap samples
- Boosting
 - Ensemble methods that improve training error
- XGBoost: modern boosting method based on regression trees
 - Each tree modifies the prediction made by the previous trees

Discussion



- In which cases would you **not** use Ensembles?
- How would you handle preprocessing in ensembles?
 - How about feature selection / importance?

Literature



- Chapter 14 in Bishop
- Chapter 11 in Peter Flach's *Machine Learning* (short)



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Thank you for your attention!

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