SOSC 4300/5500: Prediction Evaluation

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Outline

Logistics

Evaluation

Train/Test Split

Bias-Variance Trade-off

Regularization

Summary

Logistics

- Say you want to predict something, like flu trends
- How should we choose between different algorithms?
- Two views:
- Structural approach: You believe there is something intrinsically better, and always use that algorithm
- Agnostic approach:
 - there is no model that is intrinsically better or worse
 - choose the method that "optimizes performance for their particular research task" (GRS21)

Prediction evaluations for continuous outcomes

- It's common to use \hat{Y} to denote the predicted value of Y
- For continuous outcomes:
- R²: for linear regression
 - The larger the R², the better the model
- MSE (mean squared error): $1/n\sum_{i=1}^{n}(Y_i-\hat{Y}_i)^2$
 - The most widely used metric
 - The smaller the MSE, the better the model
 - Sometimes we also use RMSE = \sqrt{MSE}
 - For regression: $R^2 = 1 MSE/var(Y)$
- MAE (mean absolute error): $1/n \sum_{i=1}^{n} |Y_i \hat{Y}_i|$

Evaluation

Prediction evaluations for categorical outcomes

- For categorical outcomes, evaluation is more complex
- Cross-entropy loss is a common choice (used by some decision tree algorithms and some deep learning)
 - Also called log-loss or entropy loss
- For binary classification:

$$-\sum_{i=1}^{N} y_i \cdot \log P(\hat{y}_i = 1) + (1 - y_i) \cdot \log (1 - P(\hat{y}_i = 1))$$

Prediction evaluation for categorical outcomes

- Another set of evaluation is based on tabulating predictions and actual values and is more intuitive
- Let us assume that there are 10,000 students/employees at HKUST, and there are 10 infected cases
- ullet We has an algorithm predicting COVID infection (positive =1 vs. negative =0)
- We found that 99% of our predictions are correct. Yeah!
- But wait, is that good enough?

Prediction evaluations for categorical outcomes

- In fact, for any classification task, one of the simplest baseline is to predict every data point as belonging to the majority class
- Here, we know most people are not positive
- So the simplest baseline just predict that every one is negative
- What's the accuracy for this simplest baseline prediction?
- Accuracy = 9990 / 10000
- If class is imbalanced, it is very easy to achieve a high accuracy by predicting the majority class all the time
 - But it is misleading

Evaluation

Prediction evaluations for categorical outcomes

Actual

		1/positive	0/negative
Prediction	1/positive	True Positive (TP)	False Positive (FP)
	0/ negative	False Negative (FN)	True Negative (TN)

- It's better to use confusion matrix
- Each cell is the number of observations fall into the corresponding category
- accuracy = $\frac{TP+TN}{TP+TN+FP+FN}$ precision = $\frac{TP}{TP+FP}$
- - Interpretation: what proportion of predicted positives are
- actual positive? recall = $\frac{TP}{TP+FN}$
 - interpretation: what proportion true positives are identified by predictions?

Simplest baseline: majority class

Actual

		1/positive	0/negative
Prediction	1/positive	True Positive (n = 0)	False Positive (n = 0)
	0/negative	False Negative (n = 10)	True Negative (n = 9900)

- Accuracy: $\frac{TP+TN}{TP+TN+FP+FN}$
 - $\frac{9990}{10000} = 99.9\%$
- precision = $\frac{TP}{TD + FD}$
 - $\frac{0}{0+0}$ = not defined
- recall = $\frac{TP}{TP+FN}$

Evaluation

- $\frac{0}{0.110} = 0\%$
- From precision/recall, we see that this prediction is very bad, which suggests that precision/recall can recognize this majority guess as a bad prediction

Case 1: high precision/ low recall

Actual

1/positive 0/negative True Positive False Positive 1/positive (n = 5)(n = 0)Prediction False Negative True Negative 0/negative (n = 5)(n = 9990)

- Accuracy: $\frac{TP+TN}{TP+TN+FP+FN}$ $\frac{5+9990}{10000} = 99.95\%$
- precision = $\frac{TP}{TP+FP}$ • $\frac{5}{5+0} = 100\%$
- recall = $\frac{TP}{TP+FN}$ • $\frac{5}{5+5} = 50\%$

Evaluation

- So every predicted infected case is indeed infected
- But we missed 50% of actual infected cases

Evaluation

Case 2: high recall/low precision

Actual

		1/positive	0/negative
Prediction	1/positive	True Positive (n = 9)	False Positive (n = 4)
	0/negative	False Negative (n = 1)	True Negative (n = 9986)

- We lower the threshold to be considered as infection case
- Accuracy: $\frac{TP+TN}{TP+TN+FP+FN}$
 - $\frac{9+9986}{10000} = 99.95\%$; the same
- precision = $\frac{TP}{TP+FP}$
 - $\frac{9}{9+4} = 69.23\%$
- recall = $\frac{TP}{TP+FN}$
 - $\frac{9}{911} = 90\%$
- Our prediction captures 90% of actual infected cases
- But less than 70% predicted cases are actually infected

Precision-recall trade-off

- In evaluting perdiction performances for categorical outcome, do not use accuracy
- Instead, use precision and recall
- Depending on tasks, we may emphasize one or the other
- An ideal algorithm will have both high precision and recall
- In real life, one always comes at the cost at the other
- This is called precision-recall trade-off
- [in class activities]: can you think of cases when high precision is more important? Cases when high recall is more important?

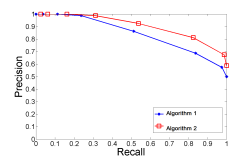
- Many ML algorithms give you predicted probability, and then transform this probability into a binary prediction
- If $P(Y=1) > \phi$; predicted probability is larger than a threshold
 - Predicted value $\hat{Y} = 1$
- If $P(Y = 1) <= \phi$;
 - $\hat{Y} = 0$

Evaluation

- Large threshold ϕ -> high precision
- Small threshold ϕ -> high recall
- Often software will choose threshold to be 0.5 by default
- You can generate new predictions based on whether you want high precision or high recall

Precision-recall curve

- Precision-recall curve is a way to visualize the trade-off
- Imagine you choose many different thresholds
- For each thresholds, obtain binary predictions, and calculate precision/recall
- Then plot the precision against recall



Algorithm 2 is better than 1

Precision-recall and F1 score

- If you do not have a specific reason preferring one or the other
- F1 score is a single-number measure of how good your predictions are
 - 2 * precision*recall precision+recall

False positive and false negative rates

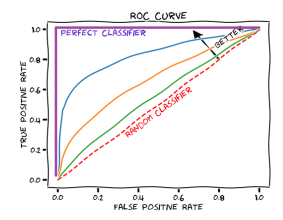
Actual

		1/positive	0/negative
Prediction	1/positive	True Positive (TP)	False Positive (FP)
	0/ negative	False Negative (FN)	True Negative (TN)

- True positive rate = recall: $\frac{TP}{TP+FN}$
- False negative rate: 1 true positive rate $=\frac{FN}{TP+FN}$
 - For COVID example, what percentage of people were infected but predicted as not
- False positive rate: $\frac{FP}{FP+TN}$
 - For COVID example: what percentage of people were not infected but predicted as infected

ROC curve

- Similar to precision-recall curve case, vary decision thresholds and obtain false positive and false negative rates
- Then plot TPR = 1 FNR against FPR



AUC of ROC

- Similar to F1 score is a single-number measure based on precision-recall curve
- Area under the curve (AUC) is a single-number measure based on ROC curve
- Larger AUC -> better prediction performance

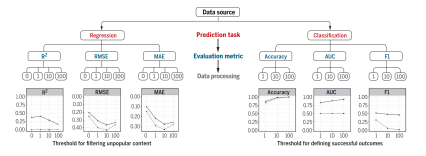
ROC vs Precision/Recall

- Precision-recall curve and ROC curve are much better measure of algorithm performances than ROC curve
- Use ROC curve, if you care both positive and negatives
- Use precision/recall curve, if you care one class more than the other
 - e.g., you care about positive class more than negative class
- Use precision/recall curve, if your data is highly imbalanced, and you only care about one class
 - Often in text analysis, this is the case: you want to extract some information from documents

From binary to categorical

- If you have more than 2 categories
- Calculate precision/recall; FPR/FPR for each category
- Macro-average: treat each category as the same; take the average precision of each category
 - can be problematic if you have imbalanced data.
- Micro-average: take into consideration of the size of each category; preferred

 Jake M. Hofman, Amit Sharma, and Duncan J. Watts, Prediction and explanation in social systems, Science 355 (2017), no. 6324, 486-488



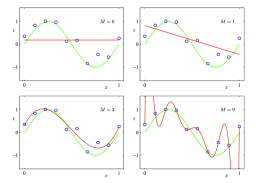
- How social scientists have been using regressions
 - Use the entire dataset to fit a regression model
 - Calculate differences between prediction and the true
- This is going to be problematic, since what works now may not work on new data

- We used Y = sin(X) (the green curve) to generate some data (in blue dots)
- We use a polynomial regression with only one variable X for prediction:
 - Polynomial regression: keep adding higher order terms to independent variables
- When M is larger, we get models that fit the data better and better

$$Y = \beta_0 + \beta_1 X + \beta_2 X^2 + \dots + X^M$$

Example

- M = 1, prediction bad
- M=9, prediction is too good on current data, but is not very well on new data
- M=3, not as great on current data, but well on new data



Principle

- Principle: choose a model that works well on new data
 - If a model predicts well on existing data, it may not work well on new data
- If you have time-series data, easier
 - use past data to train the model; see how well it can predict future data
- If you do not have time-series data, use train/test split

time-series data example

- Training data: CDC counts and search queries from 2003 to 2007
 - Training Procedure:
 - 1. Select a model: e.g., linear regression, CDC counts $\sim \beta \times$ (Google search queries)
 - 2. Training (fitting) model: obtain the value of regression coefficient β
- New data: CDC counts and search queries in 2008
 - Evaluating procedures:
 - 1. calculate predicted CDC counts, based on β and search queries in 2018
 - 2. compare actual CDC counts in 2008, and predicted CDC counts from the above
- Evidence-based way of choosing the best model:
 - if you have another algorithm, just test it on the exact same new data
 - If the new algorithms gives better performance, then use the new algorithm

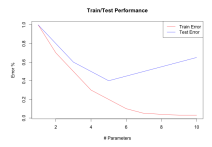
Train/test split

- Time-series data allow very natural way to choose new data
- If you do not have time-series data, use train/test split
- Split your data into two parts (typically 80%/20% or 90%/10%)
 - training data
 - (out-of-sample) test data
- Train your model only with training data
 - ML algorithm must not see the test data
- Evaluate model performances (e.g., precision/recall, false) positive/false negatives)
- Select a model that has the best performance

- Select a model that has the best performance on test data
- We mean both across or within algorithms
 - Across: should you select deep learning or SVM?
 - select the one that optimizes performance on out-of-sample test data
 - Within: should I let the decision tree to be 4 layers? Or 5 layers?
 - select the one that optimizes performance on out-of-sample test data
 - From last lecture, we know that this is also called selecting tuning parameters (or hyperparameters)

rrain/ test spin

- Test error (e.g., MSE or FPR/FNR) typically is larger than training error
- When test error begins to increase, overfitting occurs
- Error/performance metrics based on test data gives more faithful evaluation of how the algorithm will perform in real-world, unseen new data

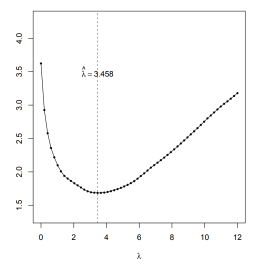


Example: Use train/test split to choose a LASSO model

- Assume we have a LASSO model, and we think our λ can take values 0,1,2,3, \cdots , 10
- For each possible value of λ :
 - Use the training data to fit your statistical model (here: LASSO)
 - ullet Use the estimated parameters to predict Y for the test data
 - Calculate MSE on the test data

$$1/n\sum_{i\in\mathsf{test}}\left(Y_i-\hat{\beta}X\right)$$

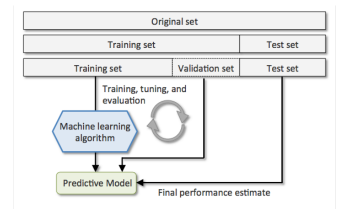
• We should select λ that yields the smallest MSE on the test data



Example: Use train/test split to prune a decision tree

- We can use train/test split to prune a tree
- Leave out some data as test data
- Let the algorithm select the next split for you (grow one branch)
- then predict use the new and old tree for test data
 - if the new MSE is smaller, keep it
 - Otherwise, go back and delete the most recent split (pruning)

- You can see that train/test split allow us to do two things:
 - select the best model
 - evaluate the final model's performance.
- If you have a lot of data, sometimes you will see people do a more complex three-way split
- Further split your training data into:
 - Training data
 - Validation data: select tuning parameters; select a final model
 - test data: report final model's performance on test data

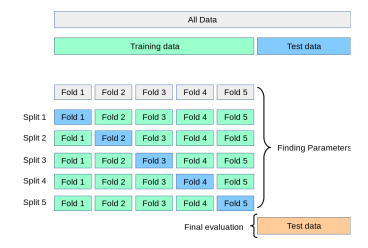


Cross-validation

- Imagine you have chosen a particular three-way split:
 - 60% as training
 - 20% as validation
 - 20% as test
- One concern is that you the particular 80% may be different from the entire sample
- Cross-validation makes every part of data used as training data once

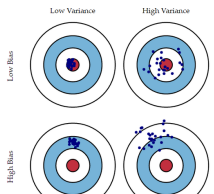
Cross-validation

- K-fold cross validation (below example shows K=5)
- The model performance is the average over 5 evaluations



Bias Variance Trade-Off

- We have seen that more complex models work very well on current data, but may not work well on new data
- A formal reasoning of this intuition is call bias-variance tradeoff
- Typically, more complex prediction algorithm gives less biased predictions, but their variance is also huge



Bias Variance Trade-off

- One implication of high prediction algorithm variance is that it does not generalize well on new data that have not been seen by the ML algorithm
 - When this occurs, people call this overfitting
 - It means that your algorithm learn training data very well, but it is highly unstable on new data and can make a lot of errors
 - Intuition: you remember every thing taught in a class so well, but professor gives you some new exercises you have not seen and your knowledge does not handle new questions

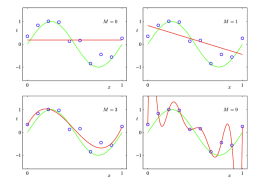
Bias Variance Trade-off (example)

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$$Y = \beta_0 + \beta_1 X + \beta_2 X^2 + \dots + X^M$$

Bias Variance Trade-off (cont'd)

- M = 1, OLS gives lots of estimation bias
- M = 9, the model fits the data so well, but it is highly sensitive to small changes in observations
- M = 3, it achieves a good balance between estimator bias and estimator variance



- There are typically two ways to achieve bias variance trade-off
 - One focus on algorithm side, and the other focus on data side (train/test split)
- Regularization: explicitly make the algorithm less complex to balance bias and variance
 - Note that this is not commonly used in linear regression: people often tell you to add as much variables as possible
 - Each algorithm usually has its own way of doing regularization
- Use train-test split:
 - data-driven way of choosing a model that works the best on new data

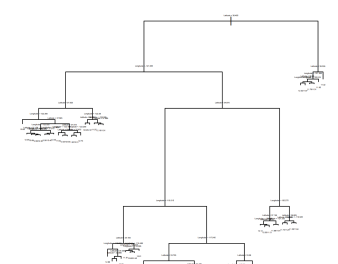
- When there is more regressiors than independent variables
- LASSO: force some variables to be zero
- Ridge: force some variable to be closer to zero (weaker form or regularization)

Decision Tree: Bias vs Variance

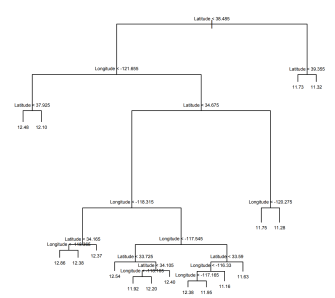
- A decision tree, without any restrictions, can be quite complex
- We can always make a very complex tree by:
 - Try your best to make every single leaf contains only one Y
- Bias-variance trade-off:
 - Complex trees fit the data nearly perfect (low predictor MSE)
- We need to regularize to make tree simpler
 - This is the same principle of LASSO/Ridge
 - To make better predictions, we sometimes have to make the algorithm less complicated

Decision Tree: un-pruned

- In decision tree, regularization is called pruning
- It's like cutting leaves and making the tree smaller



Decision Tree: pruned



- Bagging: simply fit many trees over the entire data
- The key innovation of random forests:
- For each sample from the original training data, randomly select m
 - A common choice: $m = \sqrt{p}$
- In other words, we just force p m predictors to be non-relevant each time

- ML algorithm evaluations:
 - Continuous outcome: MSE
 - Binary: precision/recall; false positive/false negative rates; ROC curve
 - No more "this is the best, magic, computer algorithm that does blah blah"
 - Instead, present empirical evidence
- Bias-variance trade-off
 - Definition; overfitting
 - Regularization
 - Train-test split
 - Allows more faithful evaluation of prediction performance
 - Allow you to select tuning parameters
 - Allow you to select the best algorithm