Data Mining: Lecture 2 Classification

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Schedule changes

- 01.10.24 Cluster analysis I.
- 08.10.24 Association pattern mining. (10.10 Home assignment I defense.)
- 15.10.24 Cluster analysis II. (EM algorithm)
- 22.10.24 Anomaly and outlier analysis.

Introduction

Introduction

- Classification problem may be seen as learning the structure of a data set of examples, already partitioned into categories or classes (labeled data set).
- Typically, learning leads the model.
- Model is used to estimate labels of the **previously unseen** data.
- Majority of the classification algorithms consist of two phases: Training and Testing
- Usually output of the classification is either Label or Numeric score.

Data

Features or attributes (numeric/categorical/binary) Label to be determined

Decision boundary, geometric interpretation, 2D

- Decision boundary (decision surface) (statistical classification with two classes) is a hypersurface that partitions the data set into two subsets, one for each class.
- Classifier tries to learn (construct) decision boundary that will lead minimal empirical error.

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Decision boundary, 3D

Feature selection

- Filter models: a subset of features is evaluated with the use of a class-sensitive discriminative criterion.
	- \blacktriangleright Gini index.
	- ▶ Entropy.
	- ▶ Fisher score.
	- \blacktriangleright Fisher linear discriminant.
- Wrapper models.
- **Embedded models.**

Gini index

- Measures the discriminative power of a particular feature.
- Typically, it is used for categorical variables, but it can be generalized to numeric attributes by the process of discretization.
- Let v_1, \ldots, v_r are the possible values of the particular categorical attribute.
- Let p_i denotes the fraction of the data points containing attribute value v_i belonging to the class $j \in \{1, \ldots, k\}$ to the data points containing attribute value v_i then Gini index defined as follows:

$$
G(v_i) = 1 - \sum_{j=1}^{k} p_j^2.
$$

- The value $1 1/k$ indicates that the different classes are distributed evenly for a particular attribute value.
- Lower values of the Gini index imply greater discrimination.

Gini index

0

The value-specific Gini index may be converted into an attribute wise Gini index.

$$
G(v_i) = 1 - \sum_{j=1}^{k} p_j^2.
$$

Let n_j denote the number of data points that take the value v_i ,
 $\sum_{i=1}^r n_i = n$. Overall Gini index is defined as $\sum\limits_{i=1}^r n_i=n$. Overall Gini index is defined as

$$
G = \sum_{i=1}^{r} \frac{n_i G(v_i)}{n}.
$$

Lower values of the Gini index imply greater discriminative power.

Entropy

- The class-based entropy measure is related to notions of information gain resulting from fixing a specific attribute value.
- The class- base entropy is defied as follows:

$$
E(v_i) = -\sum_{j=1}^{k} p_i \log_2(p_j))
$$

takes its values in $[0,\log_2(k)]$, whereas greater values indicate greater mixing.

By analogy with Gini index one may define overall Entropy as

$$
E = \sum_{i=1}^{r} \frac{n_i E(v_i)}{n}.
$$

Fisher score

- The Fisher score is naturally designed for numeric attributes to measure the ratio of the average interclass separation to the average intraclass separation.
- The larger the Fisher score, the greater the discriminatory power of the attribute.
- Let μ_i and σ_i denote the mean and the standard deviation of the of the data points belonging to the class j , for a particular feature. And let p_j be the fraction of the points belonging to the class j. Finally let μ define the mean of the entire data set. The Fisher index is defined as follows:

$$
F = \frac{\sum_{j=1}^{k} p_j (\mu_j - \mu)^2}{\sum_{j=1}^{k} p_j \sigma_j^2}
$$

• The attributes with the largest value of the Fisher score may be selected for use with the classification algorithm.

 k -nearest neighbour (k-NN) classification

• Let N be a labeled set of points belonging to c different classes such that

$$
\sum_{i=1}^{c} N_i = N
$$

- Classification of a given point x
	- \blacktriangleright Find k nearest points to the point x.
	- Assign x the majority label of neighbouring (k -nearest) points

Example

Classification model goodness!

- How good is the model?
- What is the goal of modeling?

Classification outcome

- **Consider binary classifier.**
- \bullet In the data set there are two classes: Positive (P) and negative (N)
- Outcomes of the classification: True positive, true negative, false positive (type I error), false negative(type II error).

Context of information retrieval

NB! Observe notions!

- Relevant elements of the data set. One is interested to find (retrieve elements of the certain class).
- **Precision is defined as:**

$$
precision = \frac{|\text{relevant} \cap \text{retrieved}|}{|\text{retrieved}|}
$$

• Recall is defined as:

$$
recall = \frac{|relevant \cap retrieved|}{|relevant|}
$$

Context of classification I

Denote: tp - true positive, tn - true negative, fp - false positive and fn false negative.

Precision:

$$
Precision = \frac{tp}{tp + fp}
$$

Recall:

$$
\text{Recall} = \frac{tp}{tp + fn}
$$

• True negative rate (Specificity):

$$
\text{TNR} = \frac{tn}{tn + fp}
$$

• Accuracy:

$$
Accuracy = \frac{tp + tn}{tp + tn + fp + fn}
$$

• Predicted positive condition rate

Predicted positive condition rate $=$ $\frac{tp + fp}{tp + tn + fp + fn}$

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F-measure not to be confused with similarly named values!!!

Frequently referred as F_1 -score ... is harmonic average of precision and recall.

 \bullet

$$
F = 2 * \frac{\text{precision} \times \text{recall}}{\text{precision} + \text{recall}}
$$

• More general definition:

$$
F_{\beta} = (1 + \beta^2) \frac{\text{precision} \times \text{recall}}{\beta^2 \text{precision} + \text{recall}}
$$

Decision trees

- Non-parametric supervised learning technique.
- Tree-like graph is used to represent the model of decision making and possible consequences of such decisions.
- Internal nodes are conditions (questions). terminal nodes represent labels of classes.
- Questions or conditions play a role of features. Answers to the questions are referred as feature values.
- Training a tree model is referred as tree growing.

Growing a tree 1

Greedy heuristic is the most popular technique. Let F be the possible set of features and S is the subset of data. The idea is to find most useful feature (among remaining) at each node.

$$
j(S) = \arg\min_{j \in F} \text{cost}(\{x_i, y_i : x_i \in S, x_{i,j} = c_k\})
$$

$$
+ \text{cost}(\{x_i, y_i \mid x_i \in S, x_{i,j} \neq c_k\})
$$

Classification cost:

$$
\hat{\pi_c} = \frac{1}{|S|} \sum_{x_i \in S} \mathbb{1}\{y_i = c\}
$$

Misclassification rate:

$$
\frac{1}{|S|} \sum_{x_j} \in S1(y_i \neq \hat{y}) = 1 - \hat{\pi y}
$$

Cost functions

• Entropy:

$$
\mathbb{H}(\hat{\pi}) = -\sum_{c=1}^{C} \hat{\pi_c} \log_2 \hat{\pi_c}
$$

Minimizing entropy is equivalent to maximizing information gain which is $\mathbb{H}(Y) - \mathbb{H}(Y|X_i)$.

• Gini index:

$$
G = \sum_{c=1}^{C} \hat{\pi_c} (1 - \hat{\pi_c})
$$

Growing a tree 3

- Repeat:
	- \triangleright For each feature divide data into corresponding subsets. Evaluate accuracy of such split with respect to response variable.
	- ▶ "Most accurate" feature wins. It will become condition at a given node.
	- ▶ Exclude chosen feature from the feature set.
- **Q.** Until no more features left.

Example: When to play tennis

Information gain

Definition

Information gain G_I of an action is the decrease of the ambiguity achieved as the result of the action.

- In the context of decision tree growing the action is splitting the node.
- If entropy is chosen as the cost function then information gain is defined as follows:

$$
G_I = E - (E_l \cdot p_l + E_r \cdot p_r)
$$

where E is the entropy before splitting E_l is the entropy of left child and E_r is the entropy of the right child. Indexes r and l have the same meaning for the proportions p .

Growing the tree: case of continues features

Denote X the matrix where columns correspond to different features and rows correspond to the different observation points.

- If all the data points are of the same class return the leaf node that predicts this class.
- Among all splitting points for each column find the one giving largest information gain.
- Then chose the column with the maximum gain.
- Perform splitting.
- If stopping criteria is satisfied return the tree.
- If stopping criteria is not satisfied apply tree growing procedure to each child.

Pruning

- In order prevent overfitting stop growing the tree when the decrease is not sufficient to justify adding extra subtree.
- **•** Grow a full tree and then prune the branches giving less decrease in error.

Wrapper Models

- Filter models are agnostic to the particular classification algorithm being used. In some cases, it may be useful to leverage the characteristics of the specific classification algorithm to select features.
- Wrapper models can optimize the feature selection process to the classification algorithm at hand.
- \bullet Let $\mathcal A$ denote a specific classification algorithm. The basic strategy in wrapper models is to iteratively refine a current set of features F by successively adding features to it.
- The algorithm starts with the F set to be empty then two following steps are repeated
	- ▶ Create an augmented set of features F by adding one or more features to the current feature set.
	- ▶ Use a classification algorithm $\mathcal A$ to evaluate the accuracy of the set of features F . Use the accuracy to either accept or reject the augmentation of F .

Rule-Based Classifiers

• Rule-based classifiers use a set of "if–then" rules $\mathcal{R} = \{ \mathcal{R}_1, \ldots, \mathcal{R}_m \}$ to match antecedents to consequents. A rule is typically expressed in the following form:

IF Condition THEN Condition

- The condition on the left-hand side of the rule, also referred to as the antecedent, may contain a variety of logical operators.
- The right-hand side of the rule is referred to as the consequent, and it contains the class variable.
- The training phase of a rule-based algorithm creates a set of rules. The classification phase for a test instance discovers all rules that are triggered by the test instance.
- In some cases, methods are required to resolve the conflicts in class label prediction.

Bayes theorem

- \bullet Let us suppose that there k classes are given.
- The posterior probability of a class C_k for an input x is:

$$
p(C_k | x) = \frac{p(x | C_k)p(C_k)}{p(x)}
$$

- $p(x | C_k)$ is the likelihood, $p(C_k)$ is the prior probability, $p(x)$ is the marginal data likelihood.
- $p(C_k)$ is the probability of a class $p(C_k)$ a priori, before getting about any knowledge about the data.
- $p(C_k | x)$ is the class probability a posteriori, after getting knowledge about the data.
- Bayes theorem updates prior distribution into posterior on the basis of empiric information.

Conditional and unconditional independence

 \bullet If X and Y are unconditionally independent then their joint distribution is the product of the marginal distributions:

$$
X \perp Y \Leftrightarrow p(X, Y) = p(X)p(Y)
$$

If the influence is mediated through a third variable Z, then X and Y are said to be conditionally independent

$$
X \perp Y \mid Z \Leftrightarrow p(X, Y \mid Z) = p(X \mid Z)p(Y \mid Z)
$$

Conditional independence does not imply unconditional independence and vice versa:

$$
X \perp Y \mid Z \nleftrightarrow X \perp Y
$$

Na¨ıve Bayes assumption

• Likelihood is computed as:

$$
p(\boldsymbol{x} \mid y) = \prod_{j=1}^{n} p(x_j \mid y)
$$

- Naïve Bayes assumption: the features are conditionally independent given the class label.
- the word naïve refers to the fact that actually features are not expected to be independent or conditionally independent.
- Model has relatively few parameters and therefore immune to overfilling.

Prediction with naïve Bayes model

- \bullet the goal is to find wether a new element is of class 1 or 0 (in the example of spam filtering wether given e-mail message is spam or not).
- According to Bayes theorem.

$$
p(y = 1 | \mathbf{x}, \boldsymbol{\theta}) \propto p(\mathbf{x} | y, \boldsymbol{\theta}) p(y | \boldsymbol{\theta}) = p(y = 1 | \boldsymbol{\theta}) \prod_{j=1}^{n} p(x_{i,j} | y = 1, \boldsymbol{\theta})
$$

$$
p(y = 0 | \mathbf{x}, \boldsymbol{\theta}) \propto p(\mathbf{x} | y, \boldsymbol{\theta}) p(y | \boldsymbol{\theta}) = p(y = 0 | \boldsymbol{\theta}) \prod_{j=1}^{n} p(x_{i,j} | y = 0, \boldsymbol{\theta})
$$

• Predict the class with highest posterior probability:

$$
y^* = \arg\max_{y \in \{0,1\}} p(y \mid \bm{x, \theta})
$$