Machine Learning Course - CS-433

Classification

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Classification

Similar to regression, classification relates input variables \mathbf{x} to the output variable y, but now y can only take discrete values, i.e., y is a categorical (or nominal) variable.

Binary classification

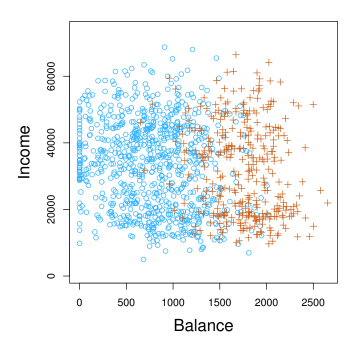
When y can only take two discrete values, it is called binary classification. There are many notations in use. Often we say that $y \in \{C_1, C_2\}$. The values C_i are called class labels or simply classes. Other common notations are $y \in \{-1, +1\}$ or $y \in \{0, 1\}$, although there may not necessarily be any ordering between the two classes.

Multi-class classification

In a multi-class classification, y can take multiple discrete values i.e., $y \in \{C_0, C_1, \dots, C_{K-1}\}$ for a K-class problem. Again, there is no notion of ordering among these classes, but we may ignore this fact and may sometimes use the following notation for convenience: $y \in \{0, 1, 2, \dots, K-1\}$.

Examples of classification problems

An credit card service must be able to determine whether or not a requested transaction is fraudulent or not. They might have at their disposal the users IP address (if the transaction happens on the web), past transaction history, and perhaps some other features. An example is shown below on the *default* dataset from JWHT. We have at our disposal the annual incomes and monthly credit card balances of a number of individuals. The individuals who defaulted on their credit card payments are shown in orange, and those who did default not are shown in blue.



A person arrives at the emergency room with a set of symptoms that could possibly be attributed to one of three medical conditions. Which of the three conditions does the individual have?

Classifier

A classifier will divide the input space into a collection of regions belonging to each class. The boundaries of these regions are calld decision boundaries. A classifier can be linear or nonlinear. This distinction is less strict than it might seem at first. E.g., if you look at the classifier in the right-hand side of "Figure 4.1" you will see that the decision regions are non-linear (not lines). But in fact the classifier that led to these region is linear. It is just that some non-linear features were added to the feature vector (think polynomial basis) before performing the linear classification.

Elements of Statistical Learning (2nd Ed.) © Hastie, Tibshirani & Friedman 2009 Chap 4

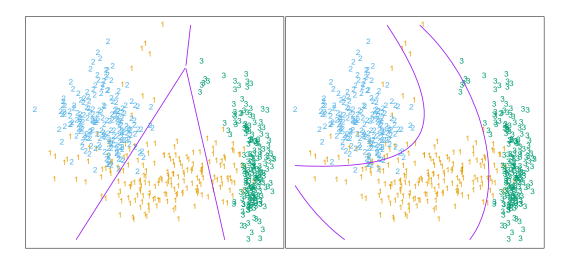


FIGURE 4.1. The left plot shows some data from three classes, with linear decision boundaries found by linear discriminant analysis. The right plot shows quadratic decision boundaries. These were obtained by finding linear boundaries in the five-dimensional space $X_1, X_2, X_1X_2, X_1^2, X_2^2$. Linear inequalities in this space are quadratic inequalities in the original space.

What is the aim of classification?

In some situations we are interested in classification in itself. This means, we are constructing a predictor based on a training set and are interested in applying this predictor to "new" incoming data. E.g., consider the example of a credit card company that wants to predict if a customer will default or not.

But in some instances we are interested in more. We would like to "understand" the cause. E.g., think of a decease prediction. Not only are we interested to know who is at risk but we would like to understand why somebody is at risk. So we are interested in the "interpretation" of the prediction. In particular for this second task it is important to have simple models and to have means of eliminating features that do not contribute significantly to the prediction.

Consider "Figure 4.12". In this data set various risk factors are given for a particular heart disease. These risk factors are blood pressure (sbp), tobacco usage, family history, obesity, alcohol consumption, and age. The figure shows for always pairs of this risk factors how the cases (people who have this disease) separate from the controls (people who do not have the desease). The idea is that rather than predicting the deases from all these risk factors, it would be better to predict it from a very small subset. This might make it easier to interpret the result.

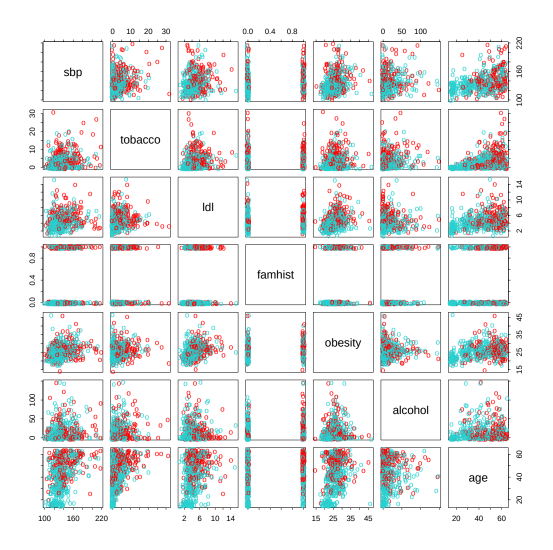


FIGURE 4.12. A scatterplot matrix of the South African heart disease data. Each plot shows a pair of risk factors, and the cases and controls are color coded (red is a case). The variable family history of heart disease (famhist) is binary (yes or no).

Classification as a special case of regression

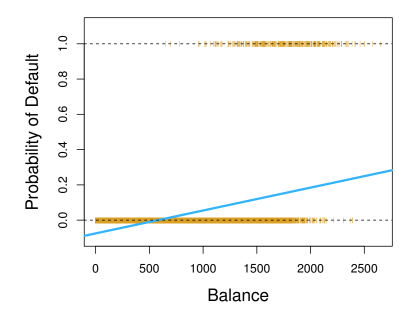
From the very definition we see that the classification problem is a special case of the regression problem. The most obvious difference is that for the classification problem the output is restricted to a small discrete set. So it might appear there is not much to be discussed and we should simply apply our standard regression techniques to this special case. E.g., we can assign y = 0 for C_1 and y = 1 for C_2 and, given a training set S_t , we can use e.g. least-squares to learn a prediction function f_{S_t} for this regression problem. To convert the regression into a classification we could decide that we predict C_1 when $f_{S_t}(\mathbf{x}) < 0.5$ and C_2 when $f_{S_t}(\mathbf{x}) > 0.5$.

In the figure below this approach is applied to the credit-card default problem. To keep things simple we use as feature only the *balance* (as we have seen in a previous plot the second feature we have at out disposal, the *income*, does not seem to contain much information anyway).

We think of y = 0 as "no default" and y = 1 as "default". The dots we see corresponds to the various data points and all dots are either on te line y = 0 or y = 1. The horizontal axis corresponds to the input \mathbf{x} , the balance.

In the figure the output y is labeled as probability. This is just a convenient way of interpretation. Since the label y is either 0 or 1 we can think of y as the probability of a default. So let us now run a regression on the training data S_t . To keep things simple we run a linear regression and learn the linear function $f_{S_t}(\mathbf{x})$. The result is the blue curve that is indicated.

We might want to interpret the value $f_{S_t}(\mathbf{x})$ as the probability of a default and then assign a label depending on whether this "probability" is smaller or larger than 0.5. Of course, this "probability" can be negative or be larger than 1 so such



an interpretation has to be taken with a grain of salt. It is easy to so that the result is questionable. Even if the cases and controls are well separated, the general "position" of the line will depend crucially on how many points are in each class and where these ponts lie. E.g., if we add a few points with y = 1 and a very large balance this will shift the curve significantly even though only a few points changed. Why does this happen? The dominant reason is that the squared loss function is not very well matched to our objective. We would like that the fraction of misclassified cases is small. But the mean squared error is only very loosely related to this objective. In particular, the mean squared error counts positive and negative deviations from the class label equally bad, although only one of them can potentially lead to a misclassification. If we do have a very small mean squared error then indeed we can guarantee a small classification error but the opposite is not true – a regression function can have arbitrarily large mean-squared error even though

the fraction of misclassified cases is arbitrarily small. We therefore might have to work "much harder" than we should in bringing down the mean-squared error and so to have a guarantee on the misclassification error.

Based on the above observation, we see that classification is not just a special form of regression with a simple loss function like the mean-squared error.

Some basic ideas of how to perform classification

Many different approaches have been developed over the years of how to efficiently perform classification. Our aim right now is not to give an exhaustive account of all possible techniques. Rather, let us quickly discuss some basic ideas. We will come back and discuss several of those in much more detail in later lectures.

Nearest Neighbor

In some cases it is reasonable to postulate that inputs that are "close" are also likely have the same label attached. Here "close" might e.g. be measured by the Euclidean distance. If we believe that this assumption is good, then, given an input \mathbf{x} and a training set S_t , we can look for that point \mathbf{x}^* which is closest to \mathbf{x} and an element of S_t and then output y^* , the label attached to \mathbf{x}^* .

The good point about such a classifier is that it might work well even in cases where the decision boundaries are very (see the Figure 2.3 below). But, as we will discuss, such a

scheme fails miserably in high dimensions since in this case the geometry renders the notion of "close by" meaningless.

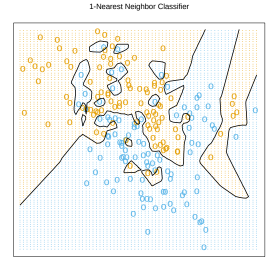


FIGURE 2.3. The same classification example in two dimensions as in Figure 2.1. The classes are coded as a binary variable (BLUE = 0, ORANGE = 1), and then predicted by 1-nearest-neighbor classification.

There are natural generalizations of this concept. Instead of using a single neighbor we can use lets say the k nearest neighbors or we can take a weighted linear combination of elements in our neighborhood. The latter is is called $smoothing\ kernels$.

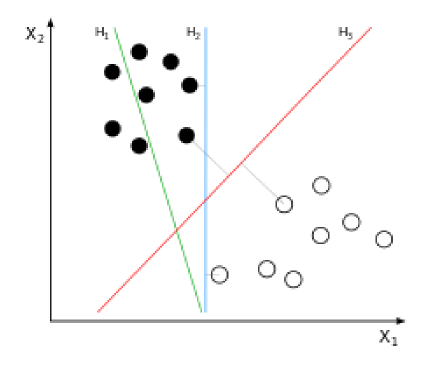
Linear decision boundaries

One starting point is to assume that decision boundaries are linear (hyperplanes). Consider e.g. a binary classification problem and look at schemes where the boundary between the two classes is a hyperplane. We can ask now how to pick this boundary. To keep things simple, assume that there exists a "separating hyperplane" i.e., a hyperplane so that not point in the training set is misclassified.

In general, there might be many hyperplanes that do the trick (assuming there is at least one). So which one should we pick?

One idea is to pick a hyperplane so that the decision has as much "robustness/margin" with respect to the training set as we can have. I.e., even if the training data was slightly changed we would like that the number of misclassifications stays as low as possible.

E.g., in the figure below (taken from Wikipedia) we see that H_1 does not separate the data, but H_2 and H_3 do. But H_3 is preferable, since it has a larger "margin."



This idea will lead us to *support vector machines* (SVM). The technique of SVM is not the only method that leads to linear decision regions. One very popular and powerful technique that leads to linear decision regions is called *logistic regression*. We will spend the whole next class to talk about it.

Non-linear decision boundaries

In many cases linear decision boundaries will not allow us to separate the data and non-linearities are needed. One option is to simply augment the feature vector with some non-linear functions. The *kernel trick* is a method of doing this in an efficient way.

Another way is to find an appropriate non-linear transform of the input so that the transformed input is then linearly separable. This is what is done when we are using *neural* networks.