StatLearn / Homework 02

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Due Sunday, June 17, 2018, 23:00 PM on Moodle

General Instructions

I expect you to upload your solutions on Moodle as a **single running** R Markdown file (.rmd), named with your surnames. In case you decide to work in Python or other programming language, you still have to provide a **running**, **sharable** notebook with detailed instructions on how to use it, **plus** a doc with the final result.

R Markdown Test

To be sure that everything is working fine, start RStudio and create an empty project called HW1. Now open a new R Markdown file (File > New File > R Markdown...); set the output to HTML mode, press OK and then click on Knit HTML. This should produce a web page with the knitting procedure executing the default code blocks. You can now start editing this file to produce your homework submission.

Please Notice

- For more info on R Markdown, check the support webpage that explains the main steps and ingredients: R Markdown from RStudio.
- For more info on how to write math formulas in LaTex: Wikibooks.
- Remember our **policy on collaboration**: Collaboration on homework assignments with fellow students is **encouraged**. However, such collaboration should be clearly acknowledged, by listing the names of the students with whom you have had discussions concerning your solution. You may **not**, however, share written work or code after discussing a problem with others. The solutions should be written by **you**.

Density Estimation Classification & Naïve Bayes

Introduction

The problem of predicting a **discrete** random variable Y from a random vector \mathbf{X} is called **classification**, supervised learning, discrimination, or pattern recognition.

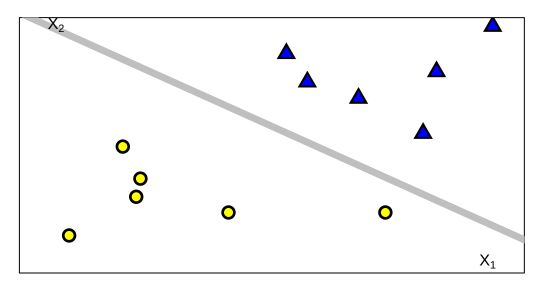
Let's consider the case of a two-class problem where we have IID data $\{(Y_1, \mathbf{X}_1), (Y_2, \mathbf{X}_2), \dots, (Y_n, \mathbf{X}_n)\}$, with $Y_i \in \{0, 1\}$ the (binary) response variable and $\mathbf{X}_i \in \mathcal{X} \subset \mathbb{R}^d$ a vector of explanatory variables (or features, or covariates).

A classification rule or classifier is then <u>any</u> function $\eta: \mathcal{X} \mapsto \{0,1\}$. When we observe a new covariate vector, \mathbf{X}' say, we then predict the response Y' to be $\eta(\mathbf{X}')$.

Just as an example, consider the fake n=12 data-points displayed in the figure below. The covariate $\mathbf{X}=(X_1,X_2)$ is 2-dimensional and the outcome $Y \in \{0,1\} = \{\triangle,\bigcirc\}$. Also shown is a **linear** classification rule represented by the solid line. For suitable values of the parameters (α,β_1,β_2) to be estimated from the data, this rule is of the form

$$\eta(\mathbf{x}) = \begin{cases} 1 & \text{if } \alpha + \beta_1 x_1 + \beta_2 x_2 > 0, \\ 0 & \text{otherwise.} \end{cases}$$

Everything above the line is classified as a 0 and everything below the line is classified as a 1. These two groups are perfectly separated by the <u>linear decision boundary</u> (see the definition below); you probably won't see such a simple pattern in real datasets.



Ok, now that we know what a classifier is, how do we evaluate its performance? Let's start introducing the following (quite natural) loss function and its *empirical* counterpart

- The true error rate of a classifier $\eta(\cdot)$ is: $L(\eta) = \mathbb{P}(\{\eta(\mathbf{X}) \neq Y\})$.
- The empirical error rate or training error rate is: $\widehat{L}_n(\eta) = \frac{1}{n} \sum_{i=1}^n \mathbb{I}(\eta(\mathbf{X}_i) \neq Y_i)$.

To keep going, let's look a bit closer at the structure of the **regression function** (i.e. the conditional expected value of Ygiven X) once specialized to this context

$$r(\mathbf{x}) = \mathbb{E}(Y \mid \mathbf{X}) \underset{Y \text{ binary}}{=} \mathbb{P}(Y = 1 \mid \mathbf{X} = \mathbf{x}) \underset{\text{Bayes'TH}}{=} \frac{f(\mathbf{x} \mid Y = 1) \mathbb{P}(Y = 1)}{f(\mathbf{x} \mid Y = 1) \mathbb{P}(Y = 1) + f(\mathbf{x} \mid Y = 0) \mathbb{P}(Y = 0)} = \frac{\pi_1 \cdot f_1(\mathbf{x})}{\pi_1 \cdot f_1(\mathbf{x}) + (1 - \pi_1) \cdot f_0(\mathbf{x})}$$

where

$$f_0(\mathbf{x}) = f(\mathbf{x} \mid Y = 0), \quad f_1(\mathbf{x}) = f(\mathbf{x} \mid Y = 1), \quad \pi_1 = \mathbb{P}(Y = 1).$$

Nice. Last round of definitions and then the result

- The set $\mathcal{X}(\eta) = \{\mathbf{x} \in \mathcal{X} : \mathbb{P}(Y = 1 \mid \mathbf{X} = \mathbf{x}) = \mathbb{P}(Y = 0 \mid \mathbf{X} = \mathbf{x})\}$ is called the **decision boundary**.
- We define the **Bayes classification rule** $\eta^*(\cdot)$ as

$$\eta^{\star}(\mathbf{x}) = \begin{cases} 1 & \text{if } r(\mathbf{x}) > \frac{1}{2} \\ 0 & \text{otherwise} \end{cases} = \begin{cases} 1 & \text{if } \mathbb{P}(Y = 1 \mid \mathbf{X} = \mathbf{x}) > \mathbb{P}(Y = 0 \mid \mathbf{X} = \mathbf{x}) \\ 0 & \text{otherwise} \end{cases} = \begin{cases} 1 & \text{if } \pi_1 f_1(\mathbf{x}) > (1 - \pi_1) f_0(\mathbf{x}) \\ 0 & \text{otherwise} \end{cases}$$

And here we go with the main result I mentioned before:

• The Bayes rule is **optimal**, that is, if $\eta(\cdot)$ is any other classifier, then $L(\eta^*) \leq L(\eta)$.

The Bayes rule clearly depends on unknown quantities that we need to estimate from data, but please notice: the Bayes rule has **nothing** whatsoever to do with Bayesian inference. We could estimate the Bayes rule using either Bayesian or frequentist methods... and this is actually what you gonna do in this exercise!

So, once we have some data at hand, $\mathcal{D}_n = \{(Y_1, \mathbf{X}_1), (Y_2, \mathbf{X}_2), \dots, (Y_n, \mathbf{X}_n)\}$ say, we may proceed to estimate:

- $f_1(\cdot)$ from the \mathbf{X}_i for which $Y_i = 1$, obtaining $\widehat{f}_1(\cdot)$,
- f₀(·) from the X_i for which Y_i = 0, obtaining f̂₀(·),
 the subpopulation proportion π₁ with π̂₁ = ½ ∑ ∑ i=1 Y_i.

and define

$$\widehat{r}(\mathbf{x}) = \frac{\widehat{\pi}_1 \cdot \widehat{f}_1(\mathbf{x})}{\widehat{\pi}_1 \cdot \widehat{f}_1(\mathbf{x}) + (1 - \widehat{\pi}_1) \cdot \widehat{f}_0(\mathbf{x})} \quad \rightsquigarrow \quad \widehat{\eta}(\mathbf{x}) = \begin{cases} 1 & \text{if } \widehat{r}(\mathbf{x}) > \frac{1}{2} \\ 0 & \text{otherwise} \end{cases}$$

The simplest approach would consist in assuming a **parametric** model for the class-conditional densities $f_1(\cdot)$ and $f_0(\cdot)$. In case we pick a multivariate Gaussian for both, depending on the covariance structure we then get back well-known classifiers known as linear and quadratic discriminant analysis models. But why would you be a dull parametric–guy? Let's go nonparametric and put a kernel density estimator at work here, as simple as that...

Nice and smooth, isn't it? Just a little caveat: in typical classification problems, the dimension d of the covariate vector \mathbf{X} may be quite large and any standard density estimator (histograms, kernels, etc) will miserably fall under the blows of the curse of dimensionality. What now? Well, let's try to be naive (...Bayes...) and assume that, conditionally on the class label, the feature vector \mathbf{X} has independent components, that is

$$f_1(\mathbf{x}) = \prod_{j=1}^d f_{1,j}(x_j)$$
 and $f_0(\mathbf{x}) = \prod_{j=1}^d f_{0,j}(x_j)$,

while this assumption is generally **not** true, it does simplify a lot the estimation process because:

- the individual class-conditional densities $\{f_{1,j}(\cdot)\}_j$ and $\{f_{0,j}(\cdot)\}_j$ can each be estimated separately using 1-dimensional kernel density estimates;
- if a component X_i of **X** is discrete, then an appropriate estimate can be used and seamlessly mixed with the others.

Notice that, in spite of the rather optimistic assumption, naïve Bayes classifiers often outperform far more sophisticated alternatives.

Finally, before we go to the actual exercise, just a few things on how to evaluate a classifier in practice. The keyword is always the same: **data splitting**. There are of course various schemes and options, but the basic one is quite simple and consists in randomly dividing the original dataset \mathcal{D}_n in two subsets of size n_1 and n_2 with $n = n_1 + n_2$: a **training set** $\mathcal{D}_{n_1}^{\text{TR}} = \{(Y_i, \mathbf{X}_i)\}_{i=1}^{n_1}$ and a **test set** $\mathcal{D}_{n_2}^{\text{TE}} = \{(Y_i, \mathbf{X}_i')\}_{i=1}^{n_2}$ in typical proportions of 70% and 30% respectively¹.

- on the **training set** we estimate the classifier as usual. In our case, for example, this step simply consists in getting the optimal bandwidths by cross-validation to obtain $\widehat{\eta}(\cdot)$ as described above.
- on the **test set**, instead, we check the performance of $\widehat{\eta}(\cdot)$ artificially treating $\mathcal{D}_{n_2}^{\text{TE}}$ as if it was a real brand new dataset with no class–label inside. More specifically, we take each feature vector \mathbf{X}_i' in $\mathcal{D}_{n_2}^{\text{TE}}$ and use the classifier $\widehat{\eta}(\cdot)$ to predict its associated outcome $\widehat{Y}_i' = \widehat{\eta}(\mathbf{X}_i')$ for $i \in \{1, \ldots, n_2\}$.

At this point, we take out of their grave the observed responses $\{Y_i'\}_{i=1}^{n_2}$ and compare them with the predictions $\{\hat{Y}_i'\}_{i=1}^{n_2}$ in order to get the **test error rate**:

$$\widehat{L}_{n_2}^{\mathrm{TE}} = \widehat{L}_{n_2} \left(\widehat{\boldsymbol{\eta}} \right) = \frac{1}{n_2} \sum_{i=1}^{n_2} \mathbb{I} \left(\boldsymbol{\eta}(\mathbf{X}_i') \neq Y_i' \right) = \frac{1}{n_2} \sum_{i=1}^{n_2} \mathbb{I} \left(\widehat{Y}_i' \neq Y_i' \right).$$

Since the classifier $\hat{\eta}(\cdot)$ and the test error rate $\hat{L}_{n_2}^{\text{TE}}$ are based on independent samples, the latter surely is more realistic than the (rather optimistic) training error rate

$$\widehat{L}_{n_1}^{\mathtt{TR}} = \widehat{L}_{n_1} \big(\widehat{\boldsymbol{\eta}} \big) = \frac{1}{n_1} \sum_{i=1}^{n_1} \mathbb{I} \big(\boldsymbol{\eta}(\mathbf{X}_i) \neq Y_i \big) = \frac{1}{n_1} \sum_{i=1}^{n_1} \mathbb{I} \left(\widehat{Y}_i \neq Y_i \right),$$

as an estimate of the *true* missclassification error rate $L(\hat{\eta})$.

Remark: Beside basic R, for a unified approach to classification and prediction, I strongly suggest the caret package. There's a lot of documentation available online. You may start from the dedicated webpage and the package vignette, to continue reading a book written in Markdown (sic!) by the package developers, their blog and a not-so-recent jstatsoft paper. Some of the functions you might wanna look at are: createDataPartition(), train(), and confusionMatrix(); together with the long list of available models and a summary of the performance measures suitable for a classification problem – quite useful to interpret the output of confusionMatrix().

Exercise

1. As I said above, in spite of the rather optimistic assumption, naïve Bayes classifiers often outperform far more sophisticated alternatives. Think a bit, look around if you wish – citing all your sources, of course – and write down why, in your opinion, this might actually be the case. In other words, give me your – personal but informed – point of view on the reason why, in a classification context, we don't really need all the gory details of the conditional marginals but only some particular feature that the simplifying hypothesis behind naïve Bayes can actually catch.

¹Notice that in basic R you may simply use the function sample() and setdiff() to create them.

2. Consider the problem of distinguishing human activities performed while wearing inertial and magnetic sensor units. The dataset is publicly available at the UCI Machine Learning Repository and is described in Barshan and Yusek (2013), where it is used to classify 19 activities performed by 8 people that wear sensor units on the chest, arms and legs. For this exercise, I took the results on just 4 activities (walking, stepper, cross trainer, jumping) performed by a single person (#1) and as covariates the measurements taken by all the 9 sensors (x,y,z accelerometers, x,y,z gyroscopes, x,y,z magnetometers) on each of the 5 units on torso (T), right arm (RA), left arm (LA), right leg (RL), left leg (LL) for a total of 45 features.

In this exercise you have to:

- Load the dataset daily-sport. RData into R and take a look with the usual functions (e.g. str()).
- Use the function subset() followed by droplevels() to reduce the dataset to two activities (e.g. running and crosstr) and only 3 sensors on one location (e.g. x,y,z magnetometers on the left leg). Call the resulting dataframe ds.small.
- Take a look at ds.small treating it as a 3D point cloud in the Euclidean space (e.g. plot 500 points at random using any function you like).
- Use the function sample() to split ds.small in ds.train and ds.test containing 70% and 30% of the original dataset respectively.
- Use the function lda() in the MASS package on ds.train to estimate a linear discriminant analysis model, call it lda.out.
- Use the function predict() on lda.out and ds.train to predict the class-labels on the training set, call this vector pred.tr. Use pred.tr to evalute the *training error rate* of the classifier.
- Use the function predict() on lda.out and ds.test to predict the class-labels on the test set, call this vector pred.te.

 Use pred.te to evalute the test error rate of the classifier.
- Compare test and training error rates and make some quick comment.
- Repeat everything using the function naiveBayes() in the package e1071 and briefly comment on its performance.
- Estimate the 1-dimensional class–conditional densities of each of the three covariates using histograms or kernels and build 95% bootstrap confidence bands around them.
- Implement your <u>own version</u> of **naïve Bayes classifiers** and see how it compares with the **naiveBayes()** implementation you used in the previous point.
- 3. ... and finally, do your best on this dataset! Crunch it as you wish to get the best classification accuracy!