Discrimination and classification

Thore Egeland/ Raju Rimal

2018

Separating and predicting

Discrimination

- Separating objects in an existing data set
- Find which variables that best separate the g groups

Classification

- Predict group membership for a new observation
- Allocate new objects to known groups

Example 1: Iris Dataset





Iris virginica



This famous (Fisher's or Anderson's) iris data set gives the measurements in centimeters of the variables:

- Sepal Length (X1)
- Sepal Width (X2)
- Petal Length (X3)
- Sepal Width (X4)



Example 1: Iris Dataset

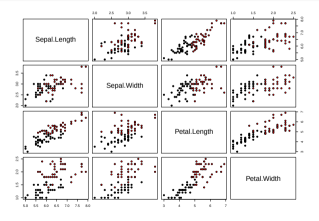
```
load("_data/iris.train.Rdata")
head(iris.train)
```

```
Sepal.Length Sepal.Width Petal.Length Petal.Width
                                                          Species
61
            5.0
                         2.0
                                      3.5
                                                   1.0 versicolor
62
            5.9
                         3.0
                                      4.2
                                                   1.5 versicolor
            6.0
                         2.2
                                      4.0
                                                   1.0 versicolor
63
            6.1
                        2.9
                                      4.7
                                                   1.4 versicolor
64
            5.6
                        2.9
                                      3.6
                                                   1.3 versicolor
65
66
            6.7
                         3.1
                                      4.4
                                                   1.4 versicolor
```

- Can we classify versicolor or virginica based on X-variables?
- What's the best method?

Iris Dataset: Pairs plot

pairs(iris.train[, -5], bg = iris.train[, "Species"], pch = 21)



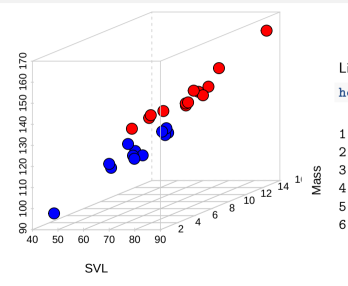
Example 2: Lizards

```
load(" data/Lizard.Rdata")
pairs(Lizard[, -4], bg = Lizard$Sex,
      pch = 21, cex = 1.5
               50 60 70 80
      Mass
                 SVL
                            HLS
         10
                            120 140
```

variable	Description
'Mass' 'SVL'	Weight of Lizard (in grams) Snout-vent length (in mm)
'HLS'	Hind limb span (in mm)

- In addition to sex we have measured
 k = 3 more variables on lizards as above.
- Can we discriminate Sex on the basis of the body measures?

Example 2: Lizards



Lizard Dataset

head(Lizard)

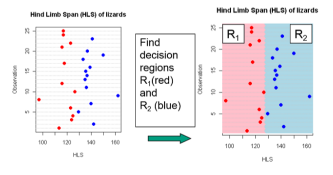
	Mass	SVL	HLS	Sex
1	5.526	59.0	113.5	f
2	10.401	75.0	142.0	m
3	9.213	69.0	124.0	f
4	8.953	67.5	125.0	f
5	7.063	62.0	129.5	m
6	6.610	62.0	123.0	f

Finding decision regions

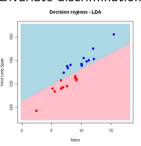
In descriminant analysis we search for decision regions for the g known groups (classes).

Regions R_1, R_2, \ldots, R_g

Univariate Descrimination

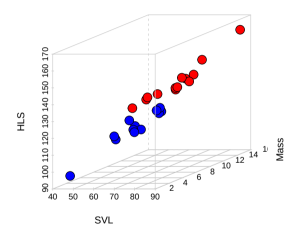


Bivariate discrimination



Using HLS and Mass as predictors (k = 2), we have used a linear decision border. This may also be non-linear.

Finding decision regions



Tri-variate discrimination

- Using all variables (k = 3),
- A linear discriminator is a flat plane in the 3D-space

Most probable group

- Perfect discrimination/classification may not be possible.
- The idea is to find decision regions R_1, R_2, \dots, R_g that minimizes the probability of doing misclassification.
- Intuitively we would allocate an observation to the group which is most probable given the observed value of the predictors
- If, for example, the HLS of a new lizard is 130, it seems most probable based on previous data that the lizard is a male.

Bayes rule

Let $P(c_i|x_1,\ldots,x_p)$ be the probability of class i given the observed variables.

Then by Bayes rule:

$$P(c_i|x_1,\ldots,x_p)\propto f(x_1,\ldots,x_p|c_i)p_i$$

where p_i is the a priori probability of observing an object from class c_i and...

 $f(x_1, \ldots, x_p | c_i)$ is the likelihood ("probability") of the observed data given class i.

A simple classification scheme

Allocate a new observation with observed variables $x_1^*, x_2^*, \dots x_p^*$ to the class which maximizes

$$f(x_1^*,\ldots,x_p^*)p_i$$

We need to estimate $f(x_1^*, \dots, x_p^*)$ from the training data and determine p_i .

In linear discriminant analysis (LDA) we assume that $f(x_1^*, \ldots, x_p^*)$ is a normal distribution, hence we must estimate the mean and (a common) variance for observations from each class.

Often we assume uniform prior: $p_1=p_2=\cdots=p_g=1/g$.

Lizards example

Assume equal probabilities for females (c_1) and males (c_2) a priori: $p_1 = p_2 = 0.5$

Assume $x_1 = HLS$ is the only predictor.

From the data we estimate μ_1 , μ_2 and common σ^2 for both classes. Let f_1 and f_2 denote these two normal distributions.

Classification rule: Classify a new lizard as female if

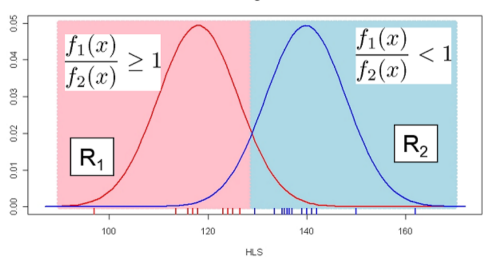
$$f_1(x_1^*)p_1 \geq f_2(x_1^*)p_2$$

or, alternatively if

$$\frac{f_1(x_1^*)p_1}{f_2(x_1^*)p_2} \geq 1$$

Visual representation

Decision regions - LDA



Linear Discreminent Analysis (LDA) in R

The fit is presented as a *confusion matrix* and *accuracy* of classification

```
library (MASS)
DAModel.1 <- lda(
  formula = Sex ~ HLS,
  data = Lizard.
  prior = rep(0.5, 2))
pred mdl <- predict(DAModel.1)</pre>
    Mass SVL HLS Sex
   5.526 59.0 113.5
  10.401 75.0 142.0
   9.213 69.0 124.0
   8.953 67.5 125.0
```

R also returns the predicted classes (fitted) and $P(c_i|x_1^*)$ also known as the *posterior* probability.

```
head(pred_mdl$posterior, 4)
1 0.00590689 0.99409311
2 0.98750273 0.01249727
3 0 16418501 0 83581499
4 0.21513565 0.78486435
head(pred mdl$class)
\lceil 1 \rceil f m f f m f
Levels: m f
```

Confusion Matrix and classification error

From the observed and predicted class of category Sex, we can construct a confusion matrix and calculate classification error.

Remember the classification rule is.

$$f_1(x_1^*)p_1 \geq f_2(x_1^*)p_2$$

Can you answer?

- Can it classify with same accuracy for new observation?
- Since this also depends on prior probability, how will it classify with different priors.

Confusion Matrix

confusion(Lizard\$Sex, pred_mdl\$class)

	True	9
Predicted	m	f
m	13	0
f	0	12
Total	13	12
Correct	13	12
Proportion	. 1	1

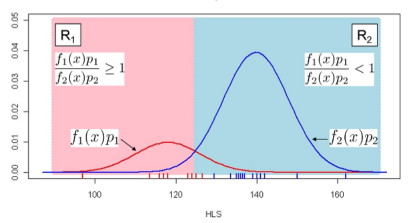
Accuracy

N correct/N total = 25/25 = 1

Using different priors

If we assume males are more abundant: $p_1 = 0.2$ and $p_2 = 0.8$

Decision regions - LDA



LDA in R using different priors

```
DAModel.2 <- lda(
  formula = Sex ~ HLS.
  data = Lizard.
  prior = c(0.8, 0.2)
pred_mdl <- predict(DAModel.2)</pre>
confusion(Lizard$Sex, pred_mdl$class)
```

7	Γrue	e
redicted	m	f
m	13	2.0000000
f	0	10.0000000
Total	13	12.0000000
Correct	13	10.0000000
Proportion	1	0.8333333

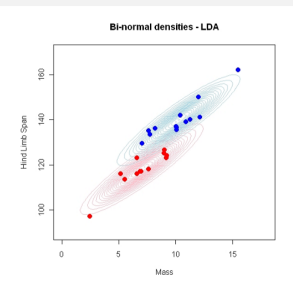
The default choice prior In R the default choice of prior is "empirical" which means that the priors are set equal to the group proportions in the data set Hence, if there are n_i observations in group iand $N = \sum_{i=1}^{g} n_i$ is the total observation number, then

$$\hat{p}_i = \frac{n_i}{N}$$

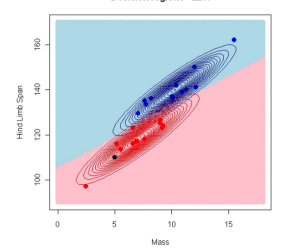
This may be reasonable if one assumes that the sample sizes reflect population sizes.

18 / 25

Bivariate LDA



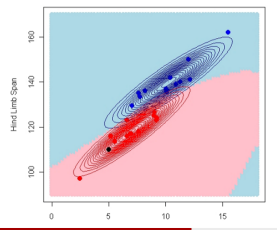
Decision regions - LDA



Quadratic discriminant analysis (QDA)

A Bivariate QDA - Decision regions

Decision regions - QDA



QDA differs from LDA in two ways:

- 1. Class specific variances
- 2. Non-linear decision borders

QDA in R

```
library(MASS)
QDA_model <- qda(
  formula = Sex ~ HLS,
  data = Lizard,
  prior = c(0.8,0.2))
pred_mdl <- predict(QDA_model)</pre>
```

Model selection/evaluation

- We often want to compare the performance of different classifiers.
- Many available statistics for this purpose, but we will consider the accuracy or the apparent error rate (APER)
- The performance of a classifier may as we've seen be summarized in a confusion matrix
- From the error rate, we can also compute the classification accuracy

Apparent Error Rate (APER)

Consider the confusion matrix as.

	True	
	Male	Female
Predicted		
Male	n_m/m	n_m/f
Female	n_f/m	n_f/f
Total	n_m	n_f

$$\mathsf{APER} = \frac{n_m/f + n_f/m}{n_m + n_f} = \frac{\mathsf{Total\ Incorrect}}{\mathsf{Total\ Observation}}$$

Classification accuracy will be 1 - APER.

Example with R: SVL as predictor

```
DAModel.4 <- lda(
  formula = Sex ~ SVL,
  data = Lizard,
 prior = rep(0.5, 2))
pred mdl <- predict(DAModel.4)</pre>
confusion(Lizard$Sex, pred_mdl$class)
           True
Predicted
            10.000 2.000
 m
    3.000 10.000
  Total 13.000 12.000
  Correct 10.000 10.000
  Proportion 0.769 0.833
```

Probabilities of misclassification - APER

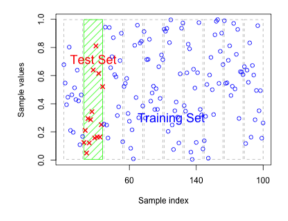
$$P(\text{female } | \text{ male}) = \frac{3}{13} = 0.231$$

 $P(\text{male } | \text{ female}) = \frac{2}{12} = 0.167$

Apparent error rate (APER)
$$= \frac{3+2}{25} = 0.2$$

$$\mathsf{Accuracy} = 1 - \mathsf{APER} = 0.8$$

Validation (Test-data or CV)



- As for regression models the errors based on fitted values may reflect over-fitting
- Should have **new** test data to predict to get an honest classification error.
- Alternatively we can perform cross-validation

Types of Cross-Validation

- K-fold cross-validation and
- Leave-one-out (LOO) Cross-validation

LOO cross-validation: LDA with SVL as predictor

```
da_model <- lda(</pre>
  formula = Sex ~ SVL,
  data = Lizard,
  prior = rep(0.5,2),
  CV = TRUE
confusion(Lizard$Sex, da_model$class)
            True
Predicted
             10.000
                    3.00
 m
             3.000 9.00
  Total 13.000 12.00
  Correct 10.000 9.00
  Proportion 0.769 0.75
```

Typically, as seen here, the accuracy goes down, and the APER increases when new samples are classified.

APER (Error rate)

N incorrect/N total = 6/25 = 0.24 Accuracy

N correct/N total = 19/25 = 0.76

Thank You

