

Displays for Statistics 5401

Lecture 37

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Class Web Page

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## Evaluation of an estimated rule

When you find a classification rule, whether it's based on non-parametric or parametric estimation of densities from a training sample, you end up with an estimated rule  $\hat{\pi}(\mathbf{x})$ .

The estimated minimum TPM rule is

- $\hat{\pi}(\mathbf{x})$ : Select  $\pi_j$  to maximize  $p_i \hat{f}_i(\mathbf{x})$

The double "hats" signify that the rule is only an estimate of the optimal  $\hat{\pi}$ .

How good is  $\hat{\pi}$ ? That is, how small are

$$\text{TPM} = \text{TPM}(\hat{\pi}) \equiv \sum_{1 \leq i \leq g} p_i \{ \sum_{j \neq i} P(j | i; \hat{\pi}) \}$$

and/or

$$\text{ECM} = \text{ECM}(\hat{\pi}) = \sum_{1 \leq i \leq g} p_i \{ \sum_{j \neq i} P(j | i; \hat{\pi}) C(j | i) \}$$

Both TPM and ECM depend on  $\{p_i\}$  and  $P(j | i; \hat{\pi})$ ,  $i \neq j$ .

It is presumed the  $\{C(j | i)\}$  are known quantities.

Usually prior probabilities  $p_i$  are also treated as known.

Sometimes you can estimate the  $p_i$  from **sample proportions**  $\hat{p}_i = n_i/N$ ,  $n_i$  = number of cases from  $\pi_i$  in the training sample.

This is possible when you obtain the training sample by "mixture sampling"

- randomly select  $\pi_i$  with probability  $p_i$ ,
- then select of  $\mathbf{x}$  using density  $f_i(\mathbf{x})$

But you still need estimates of misclassification probabilities  $P(j | i; \hat{\pi})$  to estimate either  $TPM(\hat{\pi})$  or  $ECM(\hat{\pi})$ .

$P(j | i; \hat{\pi})$  depends on the densities  $f_i(\mathbf{x})$  and is **not** known.

There are two different questions that might be of interest:

**Q<sub>1</sub>** How do you estimate  $P(j | i; \hat{\pi})$  = classification probabilities for the actual best  $\hat{\pi}$ ?

**Q<sub>2</sub>** How do you estimate  $P(j | i; \hat{\pi})$  = classification probabilities for  $\hat{\pi}$  the estimated best  $\hat{\pi}$ ?

An answer to  $Q_1$  might interest a mathematical statistician.

An answer to  $Q_2$  is more useful because, in practice, you will use  $\hat{\pi}$  to classify future cases, not  $\hat{\pi}$ .

In parametric situations (densities of known form with parameter vectors  $\theta_i$ ),

$$P(i | j; \hat{\pi}) = g_{ij}(\theta_i, \theta_j),$$

where  $g_{ij}$  depends of the  $\theta_i$  and  $\theta_j$  that characterize  $\pi_i$  and  $\pi_j$ .

When you can find a mathematical formula for  $g_{ij}(\theta_i, \theta_j) = P(i | j; \hat{\pi})$  and can estimate  $\theta_i$  by  $\hat{\theta}_i$ ,  $i = 1, \dots, g$ , you can estimate  $P(i | j; \hat{\pi})$  consistently by

$$\hat{P}(i | j; \hat{\pi}) = g_{ij}(\hat{\theta}_i, \hat{\theta}_j), \text{ all } i, j.$$

### Example of formula

- $g = 2, p = 2$
- $\mathbf{x}$  either  $N_2(\boldsymbol{\mu}_1, \boldsymbol{\Sigma}) (\pi_1)$  or  $N_2(\boldsymbol{\mu}_2, \boldsymbol{\Sigma}) (\pi_2)$ .
- prior probabilities  $p_1 = p_2 = .5$

Then

$$P(1 | 2) = P(2 | 1) = g_{12}(\boldsymbol{\mu}_1, \boldsymbol{\mu}_2, \boldsymbol{\Sigma}) = g_{21}(\boldsymbol{\mu}_1, \boldsymbol{\mu}_2, \boldsymbol{\Sigma}) = \Phi(-\Delta/2),$$

$$\Phi(x) = \text{cumnor}(x) = \frac{1}{\sqrt{\{2\pi\}}} \int_{-\infty}^x e^{-z^2/2} dz$$

$$\Delta^2 = (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_2)' \boldsymbol{\Sigma}^{-1} (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_2)$$

Therefore,

$$\text{TPM} = .5 * P(1 | 2) + .5 * P(2 | 1) = \Phi(-\Delta/2)$$

You estimate TPM and  $P(i | j)$  by

$$\widehat{\text{TPM}} = \hat{P}(1 | 2) = \hat{P}(2 | 1) = \Phi(-\hat{\Delta}/2)$$

where

$$\hat{\Delta} = \sqrt{\{(\bar{\mathbf{x}}_1 - \bar{\mathbf{x}}_2)' \mathbf{S}^{-1} (\bar{\mathbf{x}}_1 - \bar{\mathbf{x}}_2)\}}, \mathbf{S} = \mathbf{S}_{\text{pooled}}$$

When you don't have a formula for  $g_{ij}(\theta_i, \theta_j)$ , you may be able to estimate  $g_{ij}(\hat{\theta}_i, \hat{\theta}_j)$  by **simulation**:

1. Generate large samples from densities  $f_j(\mathbf{x}; \hat{\theta}_j)$
2. Use  $\hat{\pi}$  to classify them;
3. estimate  $\hat{P}(i | j; \hat{\pi}) = g_{ij}(\hat{\theta}_i, \hat{\theta}_j)$  by relative frequencies.

**Warning:**  $g_{ij}(\hat{\theta}_i, \hat{\theta}_j)$  is useful only when the parametric model is correct. Even when you assume multivariate normality to *derive*  $\hat{\pi}$ , you probably shouldn't believe it accurately describes the populations.

It is better to have a way to estimate  $P(i | j; \hat{\pi})$  regardless of the model, that is, see how well it classifies actual data from known populations.

**Best situation** (validation sample):

You have two *independent* data sets

- the training sample  $\mathbf{x}_1, \mathbf{x}_2, \mathbf{x}_3, \dots, \mathbf{x}_N$  which you use to find rule  $\hat{\pi}(\mathbf{x})$
- a validation sample  $\mathbf{x}_1^*, \mathbf{x}_2^*, \mathbf{x}_3^*, \dots, \mathbf{x}_M^*$  which is independent which you use to estimate  $\text{TPM}(\hat{\pi})$  or  $\text{ECM}(\hat{\pi})$ .

In both samples you know the population each  $\mathbf{x}_i$  or  $\mathbf{x}_i^*$  belongs to.

Suppose there are  $M_j$  cases from  $\pi_j$  in the validation sample.

- Use  $\hat{\pi}(\mathbf{x}_k^*)$  to classify case  $k$  in the *validation* sample,  $k = 1, \dots, M$ .
- Find  $m_{ij}$  = number of  $\mathbf{x}^*$  values from  $\pi_j$  with  $\hat{\pi}(\mathbf{x}^*) = \pi_i$
- Then  $\hat{P}(i | j; \hat{\pi}) = m_{ij} / M_j =$  *sample proportion*.

Because  $E[m_{ij}] = M_j \times P(i | j; \hat{\pi})$ ,  $\hat{P}$  is unbiased, that is

$$E[\hat{P}(i | j; \hat{\pi})] = P(i | j; \hat{\pi}).$$

The estimated TPM and ECM are

$$\begin{aligned} \widehat{\text{TPM}} &= \sum_{1 \leq j \leq g} p_j \{ \sum_{i \neq j} m_{ij} / M_j \} \\ &= 1 - \sum_{1 \leq j \leq g} p_j m_{jj} / M_j \end{aligned}$$

$$\widehat{\text{ECM}} = \sum_{1 \leq j \leq g} p_j \{ \sum_{i \neq j} C(i | j) m_{ij} / M_j \}$$

Because  $\hat{P}(i | j; \hat{\pi})$  is unbiased,  $\widehat{\text{TPM}}$  and  $\widehat{\text{ECM}}$  are also unbiased.

**Disadvantage:**

- You could probably get a better estimate of the optimal rule using *both* the training and validation samples to estimate the rule.

This is not an important issue when the training sample is sufficiently large to provide an acceptable classification rule.

### Re-use the training sample

What if you were to re-use the training sample as if it were a validation sample?

That is, use  $\hat{\pi}$  to classify cases in the training sample and determine the relative frequencies of errors.

This will have optimistic bias, that is, it will make  $\hat{\pi}$  appear to be too good.

- Let  $n_{ij}$  = number of  $\mathbf{x}$ 's in the training sample that belong in  $\pi_j$  but  $\hat{\pi}(\mathbf{x}) = \pi_i$ . Then the estimate based on the training sample is  $\hat{P}(i | j; \hat{\pi}) \equiv n_{ij} / N_j$ .

Estimated TPM and ECM are

- $\widehat{TPM} = \sum_{1 \leq j \leq g} p_j \{ \sum_{i \neq j} n_{ij} / N_j \} = 1 - \sum_{1 \leq j \leq g} p_j n_{jj} / N_j$
- $\widehat{ECM} = \sum_{1 \leq j \leq g} p_j \{ \sum_{i \neq j} C(i | j) n_{ij} / N_j \}$

But  $E[\widehat{TPM}] < TPM$  and  $E[\widehat{ECM}] < ECM$ .

If you estimate  $p_j$  by  $\hat{p}_j = N_j / N$ ,  $\widehat{TPM}$  is

$$\begin{aligned} \widehat{TPM} &= \sum_{j \neq i} n_{ij} / N \\ &= (\text{total number of errors}) / N \end{aligned}$$

This is the **APER** = *apparent error rate*.

APER is directly applicable only when  $\hat{p}_j = N_j / N$  estimates  $p_j$ . Even in that case, the APER is biased *downward* and is thus "optimistic".

$\hat{p}_j = N_j / N$  is a sensible estimate of the prior probabilities  $p_j$  only when

- $\{p_j\}$  are objective probabilities describing the prevalence of the populations,
- The training sample was collected randomly, with the probability of each case belonging to population  $\pi_j$  being  $p_j$ .

## Leave-one-out (Jackknife method)

This widely used method reuses the training sample to estimate classification probabilities of an estimated rule. It is also known as **Lachenbruch's holdout** procedure or the **Lachenbruch-Mickey** method.

You classify each case in the training sample using a rule based on all the other cases, "holding out" the data for the case you are classifying. Specifically:

- For case  $k$ ,  $k = 1, \dots, N$ , estimate a rule  $\hat{\pi}_{(-k)}(\mathbf{x})$  using all the data except  $\mathbf{x}_k$ .
- Then classify each  $\mathbf{x}_k$  as coming from population  $\hat{\pi}_{(-k)}(\mathbf{x}_k)$ .
- Estimate  $P(i | j)$  by  $\hat{P}(i | j) = n_{ij}^* / N_j$  where  $n_{ij}^* =$  number of  $\pi_j$  cases classified as  $\pi_i$ .

$$\widehat{TPM}_{JK} = \sum_{1 \leq j \leq g} p_j \sum_{i \neq j} \hat{P}(i | j) = \sum_{1 \leq j \leq g} p_j (\sum_{i \neq j} n_{ij}^*) / N_j$$

When you estimate  $p_j$  by  $\hat{p}_j = N_j / N$ ,

$$\widehat{TPM}_{JK} = (\sum_{1 \leq j \leq g} \sum_{i \neq j} n_{ij}^*) / N = 1 - \sum_{1 \leq j \leq g} n_{jj}^* / N$$

This might be called  $APER_{JK} =$  JackKnife Apparent Error Rate. It estimates TPM only when you obtained the training sample by "mixture" sampling.

## Example

```

Cmd> data <- read("", "dayfisher")
dayfisher      29      8 format
) Data on seed of Pantago maritima (a Scottish plantain) was
) collected from 29 Scottish localities. The localities could
) be broadly classified in three regions - Coastal, Inland and
) Island. The Coastal localities were further subdivided into
) 4 types (a) waterlogged mud, (b) typical salt marsh, (c)
) drained mud) and (d) coastal meadow above highest tide mark.
)
) Seed from each locality was grown under comparable
) conditions in an experimental garden and various
) measurements made on 100 plants grown from seed from each
) locality. In particular sample means and standard
) deviations were computed for leaf length (L), breadth (B)
) and thickness (T).
)
) This data consists of 1000*log10(means) and 1000*log10(stdev)
) Col. 1: Region (1 = Coastal, 2 = Inland, 3 = Island)
) Col. 2: Locality within Coastal region 1 (1=type a,2=type b,
)          3=type c,4=type d); also 1 for all of regions 2 and 3
) Col. 3: 1000*log10(stdev(length))
) Col. 4: 1000*log10(stdev(breadth))
) Col. 5: 1000*log10(stdev(thickness))
) Col. 6: 1000*log10(mean(length))
) Col. 7: 1000*log10(mean(breadth))
) Col. 8: 1000*log10(mean(thickness))
Read from file "TP1:Stat5401:dayfisher.mat"
    
```

```
Cmd> region <- factor(data[,1])
Cmd> locality <- factor(data[,2])
Cmd> y <- data[,-run(2)]
```

From the description in the header, locality is nested within region. I combined these in a single factor place with 6 levels.

```
Cmd> unique(10*region + locality) # 6 different values
(1)      11      12      13      14      21      31
Cmd> place <- makefactor(10*region + locality)
Cmd> list(place)
place      REAL      29      1      FACTOR with 6 levels
```

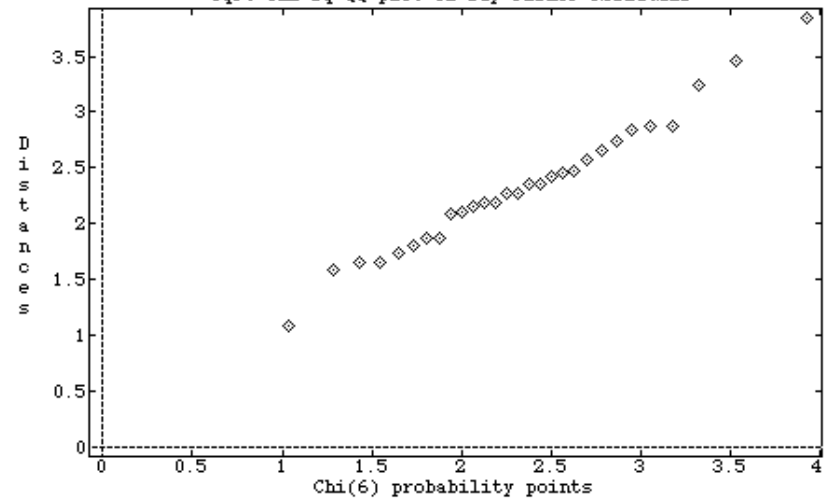
This works because 10\*region+locality consists of the 6 numbers - 11, 12, 13, 14, 21, and 31 which makefactor() turns into a factor with 6 levels.

```
Cmd> manova("y = place")# do MANOVA computates to get residuals
Model used is y = place
WARNING: summaries are sequential
NOTE: SS/SP matrices suppressed because of size; use
'manova(,sssp:T)'
```

	DF	SS and SP Matrices
CONSTANT	1	Type 'SS[1,,]' to see SS/SP matrix
place	5	Type 'SS[2,,]' to see SS/SP matrix
ERROR1	23	Type 'SS[3,,]' to see SS/SP matrix

Check that things aren't too far from normal.

```
Cmd> chiqqplot(RESIDUALS,sqrt:T,\
title:"Sqrt chi-sq QQ plot of Day-Fisher Residuals")
Sqrt chi-sq QQ plot of Day-Fisher Residuals
```



It looks pretty straight.

```
Cmd> N <- nrows(y) # size of training sample (29)
Cmd> setoptions(format:"9.5g") # fit more on line
Cmd> discrimfun <- discrim(place,y) # find linear discrim fun
Cmd> discrimfun # coefficients in estimated min TPM rule
component: coefs      Coefficients of linear part
      place1  place2  place3  place4  place5  place6
(1)  0.83379  0.84765  0.8636  0.88501  0.81145  0.84027
(2) -0.24068 -0.24764 -0.27343 -0.27526 -0.27973 -0.27775
(3) -0.449   -0.46187 -0.47554 -0.50837 -0.45276 -0.47627
(4) -1.0685  -1.0682  -1.0908  -1.1082  -0.9932  -1.0706
(5)  1.0617  1.0838  1.1484  1.1717  1.0317  1.0769
(6)  1.3695  1.3815  1.4215  1.4205  1.283   1.3668
component: addcon  constant to be added
      place1  place2  place3  place4  place5  place6
(1)   -1145  -1178.9 -1270.1 -1293.7 -1035.1 -1142.5
```



```
Cmd> scores <- y %*% discrimfun$coefs + discrimfun$addcon
Cmd> list(scores)
scores          REAL    29    6      (labels)
```

Each row of `scores` contains the linear discriminant scores for the 6 places for that case.

Assuming equal prior probabilities  $p_1 = p_2 = \dots = p_6 = 1/6$ , you classify a case in the population with the largest score.

```
Cmd> scores_1 <- vector(scores[1,]); scores_1 # case 1 scores
(1)  1103.7  1100.5  1086.3  1079.3  1095.5  1100.5
```

You would classify case 1 as place 1 since place 1 has the highest score.

Now find estimated posterior probabilities. Start with case 1.

```
Cmd> exp(scores_1)/sum(exp(scores_1))
WARNING: exp(x) with result too large set to MISSING
WARNING: exp(x) with result too large set to MISSING
WARNING: MISSING values found by sum()
WARNING: arithmetic with missing value(s); operation is /
(1)  MISSING  MISSING  MISSING  MISSING  MISSING  MISSING
```

**Ooops!** This is a case where I should have subtracted some function  $K(\mathbf{x})$  from the scores.

```
Cmd> kx <- scores[,1] # scores for place 1
Cmd> probs <- exp(scores - kx)/vector(sum(exp(scores - kx)'))
Cmd> probs[1,] # repeat of posterior probs for case 1
      place1  place2  place3  place4  place5  place6
(1)  0.92492  0.036058  2.6112e-08  2.301e-11  0.00026554  0.038755
```

You would classify case 1 in crop 1 when  $\hat{\pi}(\mathbf{x})$  computed from all the data.

Now I find  $\hat{\pi}(\mathbf{x}_i)$  for all cases at once.

```
Cmd> daplace <- vector(grade(probs',down:T)[1,])
```

`daplace` contains guessed classifications of all 29 training sample cases.

```
Cmd> print(paste(daplace))
1 1 1 2 2 2 2 2 2 2 3 3 4 4 4 4 5 5 5 5 5 5 2 5 6 6 6 5 6
```

Posterior probabilities and guesses:

```
Cmd> print(format:"5.3f",hconcat(probs, place, daplace))
MATRIX::
      Probabilities      True Guess
(1,1) 0.925 0.036 0.000 0.000 0.000 0.039 1.000 1.000
(2,1) 0.893 0.107 0.000 0.000 0.000 0.000 1.000 1.000
(3,1) 0.826 0.060 0.000 0.000 0.000 0.113 1.000 1.000
(4,1) 0.253 0.396 0.001 0.000 0.005 0.345 1.000 2.000 *
(5,1) 0.354 0.633 0.000 0.000 0.000 0.012 1.000 2.000 *
(6,1) 0.286 0.700 0.001 0.000 0.000 0.013 2.000 2.000
(7,1) 0.268 0.730 0.001 0.000 0.000 0.000 2.000 2.000
(8,1) 0.219 0.471 0.001 0.000 0.005 0.303 2.000 2.000
(9,1) 0.056 0.808 0.119 0.005 0.000 0.012 2.000 2.000
(10,1) 0.082 0.417 0.005 0.000 0.391 0.104 2.000 2.000
(11,1) 0.004 0.059 0.937 0.000 0.000 0.000 3.000 3.000
(12,1) 0.000 0.021 0.968 0.011 0.000 0.000 3.000 3.000
(13,1) 0.000 0.000 0.131 0.869 0.000 0.000 3.000 4.000 *
(14,1) 0.000 0.000 0.032 0.968 0.000 0.000 4.000 4.000
(15,1) 0.000 0.000 0.020 0.980 0.000 0.000 4.000 4.000
(16,1) 0.000 0.000 0.022 0.978 0.000 0.000 4.000 4.000
(17,1) 0.001 0.000 0.000 0.000 0.992 0.007 5.000 5.000
(18,1) 0.013 0.021 0.000 0.000 0.676 0.291 5.000 5.000
(19,1) 0.013 0.005 0.000 0.000 0.956 0.026 5.000 5.000
(20,1) 0.000 0.000 0.000 0.000 0.990 0.010 5.000 5.000
(21,1) 0.000 0.000 0.000 0.000 1.000 0.000 5.000 5.000
(22,1) 0.000 0.000 0.000 0.000 0.999 0.001 5.000 5.000
(23,1) 0.040 0.472 0.009 0.000 0.396 0.083 5.000 2.000 *
(24,1) 0.000 0.000 0.000 0.000 0.993 0.007 5.000 5.000
(25,1) 0.004 0.000 0.000 0.000 0.000 0.996 6.000 6.000
(26,1) 0.145 0.074 0.000 0.000 0.103 0.679 6.000 6.000
(27,1) 0.074 0.126 0.001 0.000 0.001 0.798 6.000 6.000
(28,1) 0.009 0.007 0.000 0.000 0.831 0.153 6.000 5.000 *
(29,1) 0.012 0.168 0.008 0.006 0.016 0.790 6.000 6.000
```

\* means place  $\neq$  guessed place

```
Cmd> sum(daplace != place)
(1,1)      5 misclassified
```

```
Cmd> 5/N
(1) 0.17241 (17% APER)
```

The APER = 5/29 = 17.2%. This estimates TPM only when places were selected by "mixture sampling" so that the prior probabilities can be estimated by  $\hat{p}_i = N_i/N$ . This is unlikely to be true.

```
Cmd> print(tabs(,place,daplace),format:"6.0f")
MATRIX:
(1,1) 3 2 0 0 0 0
(2,1) 0 5 0 0 0 0
(3,1) 0 0 2 1 0 0
(4,1) 0 0 0 3 0 0
(5,1) 0 1 0 0 7 0
(6,1) 0 0 0 0 1 4
```

This is a "confusion matrix, a cross tabulation of the data with

- Rows corresponds to the actual locations cases came from.
- Columns correspond to the guesses  $\hat{\pi}(x_i)$  made for cases.

The diagonal elements are the numbers correctly classified.

Assume equal prior probabilities ( $p_1 = \dots = p_6 = 1/6$ ),

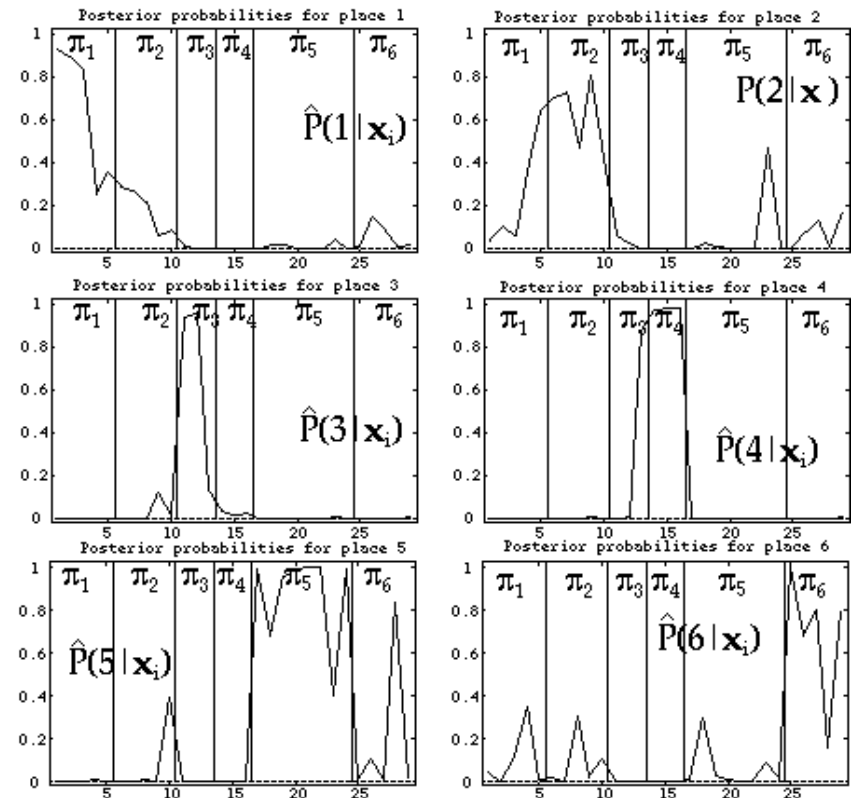
```
Cmd> prior <- rep(1/6,6) # assumed prior probabilities
Cmd> n <- tabs(,place); n # sample sizes
(1)      5      5      3      3      8      5
Cmd> P_ii <- diag(tabs(,place,daplace))/n; P_ii
(1)      0.6      1 0.66667      1 0.875      0.8
```

These are  $\hat{P}(i | i)$ .

```
Cmd> 1 - sum(prior*P_ii)
(1) 0.17639
```

This is  $\widehat{TPM}$ , biased downward because it is computed by treating the training sample as a validation sample.

Plots against case number of the naively estimated posterior probabilities



You can tell from this that  $\hat{\pi}$  does a reasonably good job of classifying the training sample. It would probably not do as well with a true validation sample.

## Jackknife estimation of error probabilities

Illustrate with case 1.

```
Cmd> temp <- discrim(place[-1],y[-1,])
```

temp contains constants for rule computed from all the data except case 1.

```
Cmd> scores_1 <- y[1,] %** temp$coefs + temp$addcon
```

```
Cmd> scores_1 <- vector(scores_1'); scores_1 # case 1 scores
Group 1   Group 2   Group 3   Group 4   Group 5   Group 6
  1100   1097.4   1081.8   1075.5   1093.2   1097.9
```

These are the linear scores for case 1.

Compute estimated posterior probabilities  $\hat{P}(j | \mathbf{x}_1)$  for case 1.

```
Cmd> kx1 <- max(scores_1); exp(scores_1-kx1)/sum(exp(scores_1-
kx1))
Group 1   Group 2   Group 3   Group 4   Group 5   Group 6
0.83416  0.061132  9.9361e-09  1.8822e-11  0.00088082  0.10383
```

This classifies case 1 as coming from  $\pi_1$  as before. Because the maximum  $\hat{P}(j | \mathbf{x}_1)$  is  $\hat{P}(1 | \mathbf{x}_1) = 0.834$ , you would classify with reasonable but not great confidence. Note that  $0.834 < 0.925 =$  "naive" estimate of  $P(1 | \mathbf{x}_1)$ .

jackknife(groups,x) automates leave-one-out computations for linear classification.

```
Cmd> pjk <- jackknife(place, y)
```

```
Cmd> list(pjk)#Cols 1-6 are posterior probs
pjk          REAL    29    7
```

The number of columns is  $g + 1 = 6 + 1$

- $pjk[i, j]$  contains  $\hat{P}(j | \mathbf{x}_i)$  for case  $i$ ,  $j = 1, \dots, g = 6$
- $pjk[i, g+1] = pjk[i, 7]$  holds the group number  $j_{\max}$  with
 
$$\max_j \hat{P}(j | \mathbf{x}_i) = \hat{P}(j_{\max} | \mathbf{x}_i).$$

Case  $i$  would be classified as coming from  $\pi_{j_{\max}}$ .

```

Cmd> print(format:"5.3f",hconcat(p,place))
MATRIX:      Posterior Probabilities      Guess  True
(1,1) 0.834 0.061 0.000 0.000 0.001 0.104 1.000 1.000
(2,1) 0.054 0.909 0.037 0.000 0.000 0.000 2.000 1.000 *
(3,1) 0.539 0.107 0.000 0.000 0.000 0.353 1.000 1.000
(4,1) 0.059 0.467 0.001 0.000 0.009 0.464 2.000 1.000 *
(5,1) 0.054 0.932 0.001 0.000 0.001 0.012 2.000 1.000 *
(6,1) 0.523 0.453 0.001 0.000 0.000 0.023 1.000 2.000
(7,1) 0.684 0.313 0.003 0.000 0.000 0.000 1.000 2.000
(8,1) 0.287 0.296 0.002 0.000 0.009 0.406 6.000 2.000 *
(9,1) 0.084 0.602 0.280 0.011 0.000 0.022 2.000 2.000
(10,1) 0.049 0.026 0.006 0.000 0.773 0.146 5.000 2.000 *
(11,1) 0.166 0.829 0.002 0.000 0.000 0.003 2.000 3.000 *
(12,1) 0.001 0.143 0.799 0.057 0.000 0.000 3.000 3.000
(13,1) 0.000 0.000 0.000 1.000 0.000 0.000 4.000 3.000 *
(14,1) 0.000 0.000 0.162 0.838 0.000 0.000 4.000 4.000
(15,1) 0.000 0.007 0.505 0.488 0.000 0.000 3.000 4.000 *
(16,1) 0.000 0.000 0.047 0.953 0.000 0.000 4.000 4.000
(17,1) 0.004 0.001 0.000 0.000 0.976 0.020 5.000 5.000
(18,1) 0.029 0.047 0.000 0.000 0.348 0.576 6.000 5.000 *
(19,1) 0.183 0.052 0.000 0.000 0.630 0.136 5.000 5.000
(20,1) 0.000 0.000 0.000 0.000 0.956 0.044 5.000 5.000
(21,1) 0.000 0.000 0.000 0.000 1.000 0.000 5.000 5.000
(22,1) 0.000 0.000 0.000 0.000 0.998 0.002 5.000 5.000
(23,1) 0.077 0.787 0.005 0.000 0.035 0.097 2.000 5.000 *
(24,1) 0.000 0.000 0.000 0.000 0.984 0.016 5.000 5.000
(25,1) 0.437 0.000 0.000 0.000 0.000 0.563 6.000 6.000
(26,1) 0.323 0.157 0.000 0.000 0.258 0.262 1.000 6.000 *
(27,1) 0.139 0.240 0.001 0.000 0.002 0.618 6.000 6.000
(28,1) 0.001 0.001 0.000 0.000 0.999 0.000 5.000 6.000 *
(29,1) 0.022 0.465 0.019 0.006 0.040 0.447 2.000 6.000 *

Cmd> confus <- tabs(,place,pjk[,7]);print(confus,format:"4.0F")
confus:
(1,1)  2   3   0   0   0   0
(2,1)  2   1   0   0   1   1
(3,1)  0   1   1   1   0   0
(4,1)  0   0   1   2   0   0
(5,1)  0   1   0   0   6   1
(6,1)  1   1   0   0   1   2
    
```

```

Cmd> N - sum(diag(confus))
(1)      15  Number of misclassifications
    
```

APER<sub>JK</sub> is 15/29 = .517

```

Cmd> 1 - sum(prior*diag(confus)/n)
(1)  0.54167.
    
```

Assuming equal p<sub>j</sub>,

$$\widehat{TPM}_{JK} = 0.54167$$

As you might expect, these are larger than the naive estimates. Usually there is not that big a difference, but here no sample sizes is large (max{N<sub>j</sub>} = 10) and there are two samples of size 3.

Incidentally, you cannot estimate quadratic discriminant functions from these data since N<sub>j</sub> ≤ p for several j. You need S<sub>j</sub><sup>-1</sup> to compute the quadratic discriminant function for group j, but S<sub>j</sub> is singular and has no inverse when N<sub>j</sub> ≤ p = 7 as is the case for every place except place 5.