randomLCA Examples

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1 Introduction

Following are two examples of using randomLCA for latent class analysis. Some aspects will certainly change but most code should still work. Two things that will change are the use of accessor functions and better labelling of results.

2 Example 1

This example demonstrates the fitting of data from Rindskopf and Rindskopf (1986), where latent class analysis is used to determine diagnostic classifications based on medical tests. Although this example is for medical data, the model is simply standard latent class so the methods can be applied to data from other areas.

A series of latent class models for 1 to 4 classes can be fitted using the commands

```
> myocardial.lca1 <- randomLCA(myocardial[, 1:4],
+    freq = myocardial$freq, nclass = 1)
> myocardial.lca2 <- randomLCA(myocardial[, 1:4],
+    freq = myocardial$freq, nclass = 2, calcSE = TRUE)
> myocardial.lca3 <- randomLCA(myocardial[, 1:4],
+    freq = myocardial$freq, nclass = 3)</pre>
```

The BIC values may be extracted from the fitted objects and are shown in Table 1.

```
> bic.data <- data.frame(classes = 1:3, bic = c(BIC(myocardial.lca1),
+ BIC(myocardial.lca2), BIC(myocardial.lca3)))</pre>
```

classes	bic
1	524.7
2	402.3
3	421.1

Table 1: BIC by class.

Using BIC as a selection method, this selects the 2 class model, indicating a nice beakdown into diseased and nondiseased, which it is assumed represent those with and without myocardial infarction. The true nature of classes is always debateable.

Summary may be used to display the fitted results

> summary(myocardial.1ca2)

```
Classes
               AIC
                        BIC
                                logLik
        2 379.3954 402.2851 -180.6977
Class probabilities
Class 1 Class
  0.5422
           0.4578
Outcome probabilities
            Q.wave History
                                LDH
                                       CPK
Class
       1 1.964e-08
                    0.1951 0.02692 0.1955
      2 7.668e-01
                    0.7914 0.82791 1.0000
Class
```

Individual results may be obtained from summary, for example the outcome probabilities shown in Table 2.

> outcomep.data <- summary(myocardial.1ca2)\$outcomep</pre>

	Q.wave	History	LDH	CPK
Class 1	0.000	0.195	0.027	0.196
Class 2	0.767	0.791	0.828	1.000

Table 2: Outcome Probabilities.

This gives some interesting information. In Class 1, those without myocardial infarction, will have abscence of Q.wave but in those with myocardial infarction it will only be present in 76.7%. The class probabilities can

be obtained as myocardial.lca2\$classp of 0.54 and 0.46 for Class 1 and 2 respectively.

One aspect of latent class is that no subject is uniquely allocated to a given class, although in some cases a subject may have an extremely high probability.

The class probs can be obtained as

```
> classprobs <- cbind(myocardial.lca2$patterns,
+ myocardial.lca2$classprob)
> colnames(classprobs) <- c(names(myocardial)[1:4],
+ "Class 1", "Class 2")</pre>
```

with results shown in Table 3. This shows subjects with 3 or 4 positive tests to be strongly classified as having myocardial infarction, and even some with 2, depending on which to to be well classified. Having only one positive test makes it unlikely that it is myocardial infarction.

Q.wave	History	LDH	СРК	Class 1	Class 2
1	1	1	1	0.000	1.000
0	1	1	1	0.008	0.992
1	0	1	1	0.000	1.000
0	0	1	1	0.111	0.889
1	1	0	1	0.000	1.000
0	1	0	1	0.581	0.419
1	0	0	1	0.000	1.000
0 1 0 1 0	0	0	1	0.956	0.044
	1	1	0	0.662	0.338
	1	1	0	1.000	0.000
	0	1	0	0.968	0.032
	0	1	0	1.000	0.000
1	1	0	0	0.997	0.003
0	1	0	0	1.000	0.000
1	0	0	0	1.000	0.000
0	0	0	0	1.000	0.000

Table 3: Class Probabilities.

Outcome probabilities are shown in Figure 1.

```
> trellis.par.set(col.whitebg())
> print(plot(myocardial.lca2, type = "l", xlab = "Test",
+ ylab = "Outcome Probability", scales = list(x = list(at = 1:4,
+ labels = names(myocardial)[1:4]))))
```

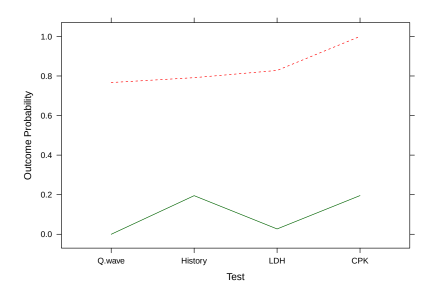


Figure 1: Outcome probabilities for 2 Class Latent Class model.

3 Example 2

This example shows the fitting of the dentistry data from Qu et al. (1996). The data consists of the results of five dentists evaluating x-rays for presence or absence of caries. As there is no gold standard, the latent class method is to assume two classes, diseased and non-diseased which are identified from the data.

3.1 Latent Class

A series of latent class models for 1 to 4 classes can be fitted using the commands

```
> dentistry.lca1 <- randomLCA(dentistry[, 1:5],
+ freq = dentistry$freq, nclass = 1)
> dentistry.lca2 <- randomLCA(dentistry[, 1:5],
+ freq = dentistry$freq, nclass = 2, calcSE = TRUE)
> dentistry.lca3 <- randomLCA(dentistry[, 1:5],
+ freq = dentistry$freq, nclass = 3)
> dentistry.lca4 <- randomLCA(dentistry[, 1:5],
+ freq = dentistry$freq, nclass = 4)</pre>
```

The BIC values may be extracted from the fitted objects and are shown in Table 4. This indicates the presence of 3 classes. A possible interpretation is that there is a class of subjects with moderate disease, or the alternative of heterogeneous disease which will be covered in the next section. Outcome probabilities are shown in Figure 2 and for the 2 class model in Figure 3.

```
> bic.data <- data.frame(classes = 1:4, bic = c(BIC(dentistry.lca1),
+ BIC(dentistry.lca2), BIC(dentistry.lca3),
+ BIC(dentistry.lca4)))</pre>
```

classes	bic
1	17531.1
2	15021.6
3	14962.9
4	15000.0

Table 4: BIC by class.

```
> trellis.par.set(col.whitebg())
> print(plot(dentistry.lca3, type = "1", xlab = "Dentist",
+ ylab = "Outcome Probability"))
```

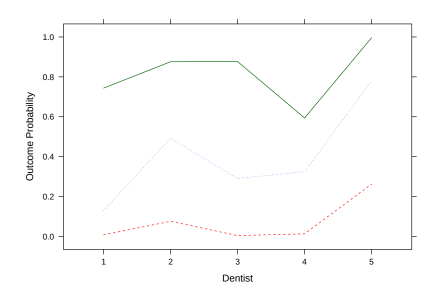


Figure 2: Outcome probabilities for 3 Class Latent Class model.

```
> trellis.par.set(col.whitebg())
> print(plot(dentistry.lca2, type = "1", xlab = "Dentist",
+ ylab = "Outcome Probability"))
```

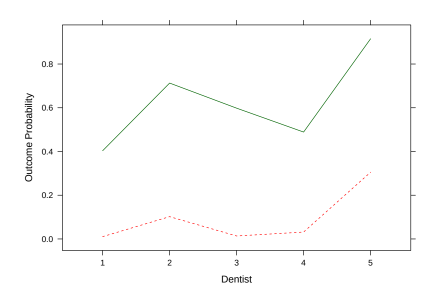


Figure 3: Outcome probabilities for 2 Class Latent Class model.

The 2 Class results can be interpreted as a diagnostic test. Important results for diagnostic testing are the sensitivity and specificity for each test. The sensitivity is the probability of the test correctly identifying the subject as diseased given that the subject is diseased. In classical diagnostic testing the "true" status of a subject is known through use of a "gold standard" which is assumed to, sometiems optimistically, correctly classify the subject. The latent class method constructs a hypothetical standard, which has the disadvantage that this is not known with certainty but it allows correctly for any uncertainty in the underlying disease state. The other measure is specificity which is the probability of correctly identifying a subject as not diseased. The sensitivities are then simply the outcome probabilities for the diseased class, Class 1 and the specificity one minus the outcome probabilities for the non-diseased class, Class 2. These can be obtained with 95% confidence intervals (provided the model is fitted with calcSE=TRUE) using the outcome probs function.

> outcome.probs(dentistry.lca2)

```
Class
       1
   Outcome p
                 2.5 %
                          97.5 %
V1 0.4033505 0.3616407 0.4465060
V2 0.7128811 0.6691318 0.7529812
V3 0.5981280 0.5494253 0.6449681
V4 0.4888445 0.4468912 0.5309555
V5 0.9154705 0.8856533 0.9380563
Class
       2
                    2.5 %
                              97.5 %
    Outcome p
V1 0.01061846 0.006938882 0.01621740
V2 0.10198782 0.089733601 0.11570279
V3 0.01359118 0.008592905 0.02143395
V4 0.03156296 0.024211140 0.04105326
V5 0.30527861 0.287118825 0.32406485
```

The sensitivity and specificity are shown in Table 5. A reasonable conclusion is that the dentists are fairly good at identifying teeth that are not diseased (except for dentist 5), but not too good at identifying teeth that are diseased.

```
> probs <- outcome.probs(dentistry.lca2)
> order <- ifelse(dentistry.lca2$classp[2] > dentistry.lca2$classp[1],
+ 1, 2)
```

```
> spec <- NULL
> sens <- NULL
> for (i in 1:5) {
      sens <- c(sens, sprintf("%3.2f(%3.2f,%3.2f)",
          probs[[order]]$Outcome[i], probs[[order]]$"2.5 %"[i],
          probs[[order]]$"97.5 %"[i]))
      spec <- c(spec, sprintf("%3.2f (%3.2f,%3.2f)",</pre>
          1 - probs[[3 - order]]$Outcome[i], 1 -
              probs[[3 - order]]$"97.5 %"[i], 1 -
              probs[[3 - order]]$"2.5 %"[i]))
+ }
> stable <- data.frame(sens, spec)</pre>
> names(stable) <- c("Sensitivity", "Specificity")</pre>
> row.names(stable) <- paste("V", 1:5, sep = "")
> print(xtable(stable, digits = c(0, 2, 2), caption = "Sensitivity and Specific
      label = "tab:outcomeconfint"), include.rownames = TRUE)
```

	Sensitivity	Specificity
$\overline{V1}$	0.40 (0.36,0.45)	0.99 (0.98,0.99)
V2	$0.71 \ (0.67, 0.75)$	$0.90 \ (0.88, 0.91)$
V3	$0.60 \ (0.55, 0.64)$	0.99 (0.98, 0.99)
V4	$0.49 \ (0.45, 0.53)$	0.97 (0.96, 0.98)
V5	0.92 (0.89,0.94)	0.69 (0.68,0.71)

Table 5: Sensitivity and Specificity

The confidence intervals for the outcome probabilities can be calculated using the parametric bootstrap.

```
>dentistry.lca2.outcomes.boot
<- outcome.probs(dentistry.lca2,boot=TRUE)</pre>
```

These are shown in Table 6 and are in agreement with those from the standard errors.

The true and false positive rates can be plotted for each dentist, and are shown in Figure 4. This gives a better explanation of what is happening. It appears that the difference between dentists is mainly related to the threshold for what they classify as diseased. Dentist 5 is more likely to correctly identify teeth as diseased but at the expense of being more likely to identify non-diseased teeth as diseased.

	Sensitivity	Specificity
V1	$0.40 \ (0.36, 0.44)$	$0.99 \ (0.98, 0.99)$
V2	$0.71 \ (0.68, 0.75)$	$0.90 \ (0.88, 0.91)$
V3	$0.60 \ (0.56, 0.64)$	0.99 (0.98, 0.99)
V4	$0.49 \ (0.45, 0.53)$	0.97 (0.96, 0.97)
V5	$0.92\ (0.89, 0.94)$	$0.69 \ (0.68, 0.71)$

Table 6: Sensitivity and Specificity

```
> itpr <- ifelse(dentistry.lca2$classp[2] > dentistry.lca2$classp[1],
+          1, 2)
> ifpr <- 3 - itpr
> probs <- outcome.probs(dentistry.lca2)
> probs <- data.frame(tpr = probs[[itpr]][, 1],
+          fpr = probs[[ifpr]][, 1])</pre>
```

3.2 Latent Class with Random Effects

The method used in Qu et al. (1996) is to introduce a random effect to model heterogeneity within classes. In their model the probabilities are transformed to the probit scale and then a normal random effect introduced. In practice it usually makes little difference if a probit or logit transform is used.

```
> dentistry.lca2random <- randomLCA(dentistry[,
+ 1:5], freq = dentistry$freq, initmodel = dentistry.lca2,
+ nclass = 2, random = TRUE, quadpoints = 31,
+ probit = TRUE)</pre>
```

The BIC is reduced to 14944.7 showing an improvement over any of the latent class models. An alternative model is to allow the variance of the random effect to vary by outcome (dentist). This can be performed using the blocksize parameter. This allows the structure of the data to be set as a series of blocks, and within each block each outcome has a different loading.

```
> dentistry.lca2random1 <- randomLCA(dentistry[,
+ 1:5], freq = dentistry$freq, initmodel = dentistry.lca2random,
+ nclass = 2, random = TRUE, probit = TRUE,
+ quadpoints = 31, blocksize = 5)</pre>
```

```
> trellis.par.set(col.whitebg())
> print(plot(tpr ~ fpr, type = "p", xlab = "False Positive Rate\n(1-Specificity
+ ylab = "True Positive Rate (Sensitivity)",
+ data = probs))
```

NULL

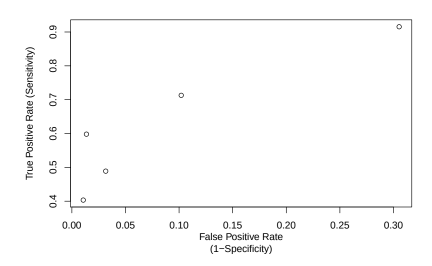


Figure 4: True and False Positive Rates by Dentist.

This increases the BIC to 14949.4, and is the 2LCR model obtained by Qu et al. (1996). It appears that the simpler model is more appropriate.

A further extension is to allow the loading or random effect variance to vary by class.

```
> dentistry.lca2random2 <- randomLCA(dentistry[,
+ 1:5], freq = dentistry$freq, initmodel = dentistry.lca2random1,
+ nclass = 2, random = TRUE, probit = TRUE,
+ blocksize = 5, byclass = TRUE, quadpoints = 41)</pre>
```

The BIC increases to 14987.6. It is not surprising that this model isn't an improvement, there are now 21 parameters fitted to 32 observations. This also may give problems with the fitting algorithm so the number of quadrature points is increase to 41.

The marginal outcome probabilities, obtained by integrating over the random effect can be plotted, as in Figure 5.

Outcome probabilities with confidence intervals may be calculated for models with random effects only using the parametric bootstrap.

```
dentistry.lca2random.outcomes.boot
<- outcome.probs(dentistry.lca2random,boot=TRUE)</pre>
```

These are shown in Table 7.

```
> print(xtable(stable, digits = c(0, 2, 2), caption = "Sensitivity and Specific")
+ label = "tab:outcomeconfintboot2"), include.rownames = TRUE)
```

	Sensitivity	Specificity
V1	0.40 (0.33,0.47)	0.98 (0.97,0.99)
V2	$0.65 \ (0.56, 0.73)$	0.87 (0.85, 0.89)
V3	$0.62 \ (0.53, 0.77)$	0.98 (0.96, 1.00)
V4	$0.42 \ (0.35, 0.49)$	$0.94 \ (0.93, 0.96)$
V5	$0.86 \ (0.79, 0.92)$	$0.67 \ (0.64, 0.69)$

Table 7: Sensitivity and Specificity

The observed and fitted values can be obtained and are shown in Table 8. Differences from the Qu et al paper result from different approximation methods.

```
> trellis.par.set(col.whitebg())
> print(plot(dentistry.lca2random1, graphtype = "marginal",
+ type = "l", xlab = "Dentist", ylab = "Marginal Outcome Probability"))
NULL
```

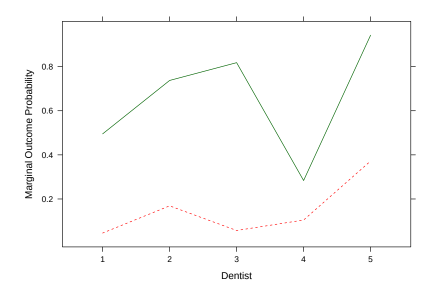


Figure 5: Marginal Outcome Probabilities for 2 Class Latent Class with Random Effect (2LCR) model.

```
> obs.data <- data.frame(dentistry.lca2random1$patterns,
+ dentistry.lca2random1$observed, dentistry.lca2$fitted,
+ dentistry.lca2random1$fitted)
> names(obs.data) <- c("V1", "V2", "V3", "V4", "V5",
+ "Obs", "Exp 2LC", "Exp 2LCR")</pre>
```

References

- Y Qu, M Tan, and MH Kutner. Random effects models in latent class analysis for evaluating accuracy of diagnostic tests. *Biometrics*, 52(3):797–810, 1996.
- D Rindskopf and W Rindskopf. The value of latent class analysis in medical diagnosis. *Statistics in Medicine*, 5:21–27, 1986.

V1	V2	V3	V4	V5	Obs	Exp 2LC	Exp 2LCR
0	0	0	0	0	1880	1836.3	1882.6
0	0	0	0	1	789	830.4	784.7
0	0	0	1	0	43	61.9	38.2
0	0	0	1	1	75	49.6	79.7
0	0	1	0	0	23	28.6	24.2
0	0	1	0	1	63	47.5	63.8
0	0	1	1	0	8	4.0	6.8
0	0	1	1	1	22	35.1	25.8
0	1	0	0	0	188	213.9	184.7
0	1	0	0	1	191	152.2	192.5
0	1	0	1	0	17	12.1	23.1
0	1	0	1	1	67	61.0	67.2
0	1	1	0	0	15	11.2	12.5
0	1	1	0	1	85	91.6	87.4
0	1	1	1	0	8	8.1	7.1
0	1	1	1	1	56	86.4	50.8
1	0	0	0	0	22	21.2	18.5
1	0	0	0	1	26	25.2	27.9
1	0	0	1	0	6	2.1	4.8
1	0	0	1	1	14	16.1	16.0
1	0	1	0	0	1	2.5	2.3
1	0	1	0	1	20	24.7	19.7
1	0	1	1	0	2	2.2	1.8
1	0	1	1	1	17	23.5	14.5
1	1	0	0	0	2	6.0	7.3
1	1	0	0	1	20	42.0	19.8
1	1	0	1	0	6	3.7	4.7
1	1	0	1	1	27	39.3	22.4
1	1	1	0	0	3	5.7	2.7
1	1	1	0	1	72	61.1	69.6
1	1	1	1	0	1	5.4	3.2
1	1	1	1	1	100	58.4	103.0

Table 8: Observed and expected frequencies