Analysis of the minn38 Dataset Using Log-Linear Models and Proportional Odds Models

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I.

1. Analysis

• Initial model

The data frame minn38 in MASS gives a dataset with four factors and a numeric column of frequencies. These are the first 6 rows of the data.

```
> head(minn38)
  hs phs fol sex f
1 L C F1 M 87
2 L C F2 M 72
3 L C F3 M 52
4 L C F4 M 88
5 L C F5 M 32
6 L C F6 M 14
```

I will use surrogate log linear model to proceed the analysis. Our initial model, also our minimal model, may be described as having the conditional probabilities for each of the four phs classes the same for all hs x fol x sex groups. In other words, phs is independent of the other explanatory factors.

```
Call:
glm(formula = f \sim hs * fol * sex + phs, family = poisson, data = minn38)
Deviance Residuals:
    Min 10
                  Median
                              30
                                     Max
-12.3618 -2.5793
                 -0.7388
                          1.8396
                                  15.0211
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 18660.9
                              on 167
                                       dearees of freedom
Residual deviance: 3010.1 on 123
                                       degrees of freedom
AIC: 4011
Number of Fisher Scoring iterations: 5
```

The high residual deviance clearly indicates that this simple model is inadequate, so the probabilities do appear to vary with the explanatory factors. We now consider adding the interaction terms between response and explanatory factors to the model that allow for some variation of this kind.

• Stepwise Selection

After achieving our minimal model, I will choose a candidate model by AIC in a Stepwise Algorithm.

```
mn.step<-step(mn.minimal,scope=list(lower=~.,upper=~.^2))
glm(formula = terms(f ~ hs * fol * sex + phs + fol:phs + hs:phs +
   sex:phs + fol:sex:phs + hs:fol:phs, keep.order = T), family = poisson,
   data = minn38)
Deviance Residuals:
    Min
        10
                 Median
                                    Max
-1.53279 -0.41864 0.00036 0.37007 1.70933
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 18660.923 on 167
                                        degrees of freedom
Residual deviance:
                       52.881 on 42
                                        degrees of freedom
AIC: 1215.8
Number of Fisher Scoring iterations: 4
> 1 - pchisq(deviance(mn.m1), mn.m1$df.resid)
Γ17 0.1211617
```

As we can see the difference between the minimal model and the model by stepwise is that we add five interaction terms phs: (hs+fol+sex), hs:fol:phs and fol:sex:phs. The stepwise procedure greatly reduced the AIC from 4011 to 1215.8. The deviance also indicates that it is satisfactorily fitting model, but we might need check if there are other adjustments to the model are needed.

Note that the first term here is part of the minimum model and hence it may not be removed. Only terms that contain the response factor, phs, are of any interest to us for this analysis. However, deleting them will not help us reduce the deviance or AIC. Now we consider adding possible interaction terms.

```
> add1(mn.m1, ~. + hs:sex:phs)
Single term additions
Model:
f ~ hs * fol * sex + phs + fol:phs + hs:phs + sex:phs + fol:sex:phs
    hs:fol:phs
            Df Deviance
                            AIC
                 52.881 1215.8
hs:sex:phs 6 47.745 1222.7
Likelihood ratio test
Model 1: f ~ hs * fol * sex + phs + fol:phs + hs:phs + sex:phs + fol:sex:phs +
   hs:fol:phs
Model 2: f ~ hs * fol * sex + phs + fol:phs + hs:phs + sex:phs + fol:sex:phs +
   hs:fol:phs + hs:sex:phs
 #Df LogLik Df Chisq Pr(>Chisq)
1 126 -481.90
2 132 -479.33 6 5.1359
                          0.5265
```

I added the interaction term hs: sex: phs, so it is my maximum model. However, as it increases the AIC and the likelihood ratio test showed that there is no significant difference between the model by stepwise and the add1 model, I choose not to include it on the grounds of simplicity, although in some circumstances we might view this decision differently. Therefore, my final model will be the model after stepwise selection.

```
Final model:
```

```
f ~ hs * fol * sex + phs + fol:phs + hs:phs + sex:phs + fol:sex:phs + hs:fol:phs
```

2. Presentation

I present my finial model final fit in the form of estimated cell probabilities.

```
hs fol sex prob.C prob.E prob.N prob.O

1 L F1 F 0.328 0.112 0.054 0.507

2 M F1 F 0.484 0.079 0.085 0.351

3 U F1 F 0.689 0.078 0.038 0.195

4 L F2 F 0.177 0.079 0.083 0.661

5 M F2 F 0.267 0.118 0.100 0.516

6 U F2 F 0.422 0.124 0.090 0.365

7 L F3 F 0.081 0.074 0.052 0.792

8 M F3 F 0.151 0.131 0.057 0.661

9 U F3 F 0.231 0.260 0.072 0.436

10 L F4 F 0.166 0.083 0.061 0.690

11 M F4 F 0.243 0.119 0.085 0.553
```

```
12 U F4 F 0.408 0.114 0.101 0.377
13 L F5 F 0.118 0.104 0.033 0.745
14 M F5 F 0.159 0.137 0.055 0.650
15 U F5 F 0.320 0.108 0.091 0.481
16 L F6 F 0.061 0.080 0.012 0.847
17 M F6 F 0.092 0.112 0.057 0.739
18 U F6 F 0.225 0.102 0.052 0.621
19 L F7 F 0.052 0.047 0.020 0.881
20 M F7 F 0.121 0.088 0.046 0.746
21 U F7 F 0.210 0.107 0.077 0.607
22 L F1 M 0.430 0.063 0.009 0.498
23 M F1 M 0.610 0.043 0.015 0.332
24 U F1 M 0.788 0.039 0.006 0.167
25 L F2 M 0.253 0.050 0.025 0.672
26 M F2 M 0.378 0.073 0.030 0.519
27 U F2 M 0.559 0.072 0.025 0.344
28 L F3 M 0.082 0.024 0.018 0.876
29 M F3 M 0.160 0.045 0.021 0.774
30 U F3 M 0.282 0.102 0.031 0.586
31 L F4 M 0.202 0.044 0.022 0.733
32 M F4 M 0.303 0.065 0.031 0.601
33 U F4 M 0.501 0.060 0.036 0.402
34 L F5 M 0.171 0.051 0.011 0.767
35 M F5 M 0.233 0.068 0.018 0.680
36 U F5 M 0.444 0.051 0.029 0.476
37 L F6 M 0.080 0.045 0.007 0.868
38 M F6 M 0.125 0.065 0.034 0.776
39 U F6 M 0.291 0.056 0.030 0.623
40 L F7 M 0.132 0.040 0.015 0.814
41 M F7 M 0.279 0.067 0.030 0.624
42 U F7 M 0.431 0.073 0.045 0.452
```

3. Discussion

The message of the fitted model is now clear. The factor having most effect on the probabilities is high school rank. With an increase in high school rank increasing the probability of enrolling in college, however, middle and low high school rank students have higher probabilities of doing other stuffs. The next important factor is the sex. Males have higher probability to go to college. Finally, the father's occupational level has a relatively small effect on the post high school status.

1. Analysis

Initial model

Our initial model, also our minimal model, may be described as having the conditional probabilities for each of the twelve phs and hs pairs the same for all fol x sex groups. In other words, phs and hs pair is independent of the other two explanatory factors.

```
glm(formula = f ~ fol * sex + hs + phs, family = poisson, data = minn38)
Deviance Residuals:
    Min    1Q    Median    3Q    Max
-12.707   -2.993   -1.203    1.485    18.029

    Null deviance: 18660.9 on 167 degrees of freedom
Residual deviance: 3638.2 on 149 degrees of freedom
AIC: 4587.1
```

The high residual deviance clearly indicates that this simple model is inadequate, so the probabilities do appear to vary with the explanatory factors. We now consider adding the interaction terms between response and explanatory factors to the model that allow for some variation of this kind.

• Stepwise Selection

After achieving our minimal model, I will choose a candidate model by AIC in a Stepwise Algorithm.

As we can see the difference between the minimal model and the model by stepwise is that we add four interaction terms fol:phs, hs:phs, sex:hs, fol:hs

and fol:sex:phs. The stepwise procedure greatly reduced the AIC from 4587.1 to1227.1. However, the deviance indicates that it is not a satisfactorily fitting model, so we might need check if there are other adjustments we can do to the model.

```
Single term deletions
Model:
f ~ fol * sex + hs + phs + fol:phs + hs:phs + sex:hs + sex:phs +
    fol:hs + fol:sex:phs
           Df Deviance
                          AIC
<none>
                160.17 1227.1
hs:phs
            6 1098.38 2153.3
sex:hs
            2
                562.68 1625.6
fol:hs
           12
                195.68 1238.6
fol:sex:phs 18 220.04 1251.0
```

Note that deleting these interaction terms will increase the AIC and the deviance, so we may not remove them. Now consider adding possible interaction terms.

```
Single term additions
Model:
f ~ fol * sex + hs + phs + fol:phs + hs:phs + sex:hs + sex:phs +
    fol:hs + fol:sex:phs
            Df Deviance
                            AIC
                 160.17 1227.1
<none>
                 136.35 1227.3
fol:sex:hs 12
Likelihood ratio test
Model 1: f \sim fol * sex + hs + phs + fol:phs + hs:phs + sex:hs + sex:phs +
    fol:hs + fol:sex:phs
Model 2: f \sim fol * sex + hs + phs + fol:phs + hs:phs + sex:hs + sex:phs +
    fol:hs + fol:sex:phs + fol:sex:hs
  #Df LogLik Df Chisq Pr(>Chisq)
1 78 -535.54
2 90 -523.63 12 23.819
                          0.02152 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
```

I added the interaction term fol: sex: hs, so it is my maximum model. As it slightly increases the AIC and the likelihood ratio test showed that there is a significant difference between the model by stepwise and the add1 model, I choose to include it to have a lower deviance model, although in some circumstances we might view this decision differently. Therefore, my final model will be the model after adding fol: sex: hs.

Final model:

```
f ~ fol * sex + hs + phs + fol:phs + hs:phs + sex:hs + sex:phs + fol:hs + fol:sex:phs + fol:sex:hs
```

2. Presentation

I present my finial model final fit in the form of estimated cell probabilities.

	fol	sex	prob.C-L	prob.C-M	prob.C-U	prob.E-L	prob.E-M	prob.E-U	prob.N-L	prob.N-M	prob.N-U	prob.O-L	prob.O-M	prob.O-U
1	F1	F	0.065	0.187	0.083	0.128	0.145	0.057	0.012	0.211	0.041	0.011	0.048	0.011
2	F2	F	0.004	0.040	0.009	0.001	0.010	0.001	0.043	0.411	0.057	0.063	0.332	0.028
3	F3	F	0.149	0.235	0.008	0.197	0.107	0.013	0.025	0.209	0.008	0.016	0.028	0.006
4	F4	F	0.015	0.056	0.003	0.003	0.009	0.002	0.086	0.366	0.092	0.085	0.173	0.110
5	F5	F	0.362	0.056	0.028	0.280	0.095	0.029	0.048	0.026	0.025	0.017	0.021	0.012
6	F6	F	0.024	0.019	0.009	0.003	0.009	0.004	0.086	0.199	0.164	0.049	0.303	0.131
7	F7	F	0.056	0.228	0.062	0.113	0.258	0.038	0.025	0.094	0.043	0.023	0.049	0.012
8	F1	М	0.011	0.041	0.008	0.005	0.014	0.002	0.109	0.253	0.092	0.165	0.257	0.043
9	F2	М	0.134	0.266	0.008	0.182	0.177	0.021	0.053	0.086	0.006	0.033	0.026	0.006
10	F3	М	0.037	0.048	0.003	0.011	0.009	0.002	0.206	0.187	0.087	0.208	0.111	0.089
11	F4	М	0.351	0.022	0.034	0.280	0.038	0.060	0.110	0.016	0.024	0.040	0.010	0.014
12	F5	М	0.060	0.007	0.012	0.011	0.003	0.005	0.210	0.121	0.175	0.125	0.151	0.120
13	F6	М	0.121	0.096	0.084	0.140	0.114	0.086	0.156	0.061	0.046	0.053	0.027	0.016
14	F7	М	0.017	0.010	0.007	0.007	0.002	0.002	0.312	0.097	0.062	0.378	0.081	0.025

3. Discussion

Our main interest lies in the conditional probabilities of the pair of the two response factors given the two explanatory factors. However, either sex or the father's occupational level has a relatively small effect on the pair of post high school status and high school rank. It is hard to interpret or get some useful knowledge by just observing the cell probabilities.

III.

By taking hs as response and fol and sex as predictors, I fitted two proportional odds models, the interaction model and the additive model, to the data.

```
Call: polr(formula = hs \sim sex + fol, data = minn38, weights = f, Hess = TRUE)
Call: polr(formula = hs \sim sex * fol, data = minn38, weights = f, Hess = TRUE)
             Value Std. Error t value
                                                                                       sexM -0.6738
                                                                                                                 0.03197 -21.078
                            0.07125 -6.2069
0.07931 -3.7386
0.09901 -4.6798
                                                                                       folF2 -0.1469
                                                                                                                 0.05940 -2.473
             -0.296528
-0.463325
                                                                                       folF3 -0.5109
                                                                                                                 0.05210 -9.806
             -0.709666
-0.477875
                                                                                       folF4 -0.4374
                                                                                                                 0.05696
sexM:folF2 -0.317580
sexM:folF3 -0.130369
sexM:folF4 -0.289398
                                                                                       folF5 -0.5492
                                                                                                                 0.07298
                            0.11879
0.10421
0.11374
                                      -2.6735
-1.2510
-2.5443
                                                                                       folF6 -0.7294
                                                                                                                 0.07364
                                                                                       folF7 -0.5759
                                                                                                                 0.07765 -7.416
 sexM:folF5 -0.173647
sexM:folF6 -0.022173
sexM:folF7 -0.201774
                                                                                       Intercepts:
                                                                                       Value Std. Error t value
LIM -1.7626 0.0495 -35.6116
MIU -0.0094 0.0470 -0.2004
     Value Std. Error t value
-1.6777 0.0637 -26.3550
0.0768 0.0619 1.2393
                                                                                       Residual Deviance: 29913.68
 Residual Deviance: 29901.69
```

```
Likelihood ratio test

Model 1: hs ~ sex * fol

Model 2: hs ~ sex + fol

#Df LogLik Df Chisq Pr(>Chisq)

1  15 -14951

2  9 -14957 -6 11.99  0.0622 .

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

From the above plots, our initial observation is that while the interaction model offers slight improvements in reducing residual deviance and AIC. Also, the likelihood ratio test showed that there is no significant difference between the two models. Therefore, the additive model was ultimately selected as the preferred proportional odds model due to its greater interpretability and similar performance. To evaluate the adequacy of this chosen model, one can simply review the AIC and residual deviance values provided in the model's summary. The resulting residual deviance is 29913.68 and the AIC is 29931.68, both of which

suggest that the model is insufficient in explaining the variability in high school ranking. Since the two models has no significant difference, we can conclude that sex and fol are independent of each other.