Applied Logistic Regression – Assignment 4

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1. Residuals in Logistic Regression

We'll go back to the dataset on Sentiments Toward Racial Integration in Public Housing. That dataset is currently in so-called Bernoulli format, in which one row corresponds to one binary observation. To ease the following computation of residuals, we will convert it into binomial format, which is better for assessing the accuracy of the model predictions for each combination of predictor values.

```
##
     Contact
                            Y Obs.
                    Norms
## 1
           0
                Favorable
                                 96 0.4270833
## 2
           1
               Favorable 120
                               172 0.6976744
## 3
           0 Unfavorable
                           56
                               201 0.2786070
           1 Unfavorable 66
                              139 0.4748201
summary(Model<-glm(Y/Obs. ~ Contact + Norms, weights = Obs.,</pre>
  family=binomial(link=logit), data=Contact))
```

```
##
## Call:
  glm(formula = Y/Obs. ~ Contact + Norms, family = binomial(link = logit),
##
       data = Contact, weights = Obs.)
##
##
  Deviance Residuals:
##
  -0.4652
             0.3729
                      0.3592 - 0.3842
##
##
  Coefficients:
##
                    Estimate Std. Error z value Pr(>|z|)
                                        -1.189
## (Intercept)
                     -0.1980
                                 0.1664
                                                   0.234
                      0.9726
## Contact
                                 0.1742
                                          5.583 2.37e-08 ***
## NormsUnfavorable
                     -0.8102
                                 0.1747
                                        -4.638 3.52e-06 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 67.93867
                                on 3 degrees of freedom
## Residual deviance: 0.63214
                                on 1 degrees of freedom
## AIC: 27.992
##
## Number of Fisher Scoring iterations: 3
```

In logistic regression, the residual of a single observation is of limited interest due to its dichotomous nature – with the response limited to 0 or 1, its residual is also limited to theoretical maximum of 1. Therefore,

rather than examining residuals of individual observations, it is more illuminating to group the data by each distinct $covariate\ pattern\ -$ also known as explanatory variable pattern or EVP – i.e., each distinct combination of predictor values observed in the data. Each group sharing an EVP also necessarily shares the same predicted probability. Multiplying each group's n by the predicted probability yields the expected number of success outcomes for that group, which can then be compared to the observed success count. Assuming n is sufficiently large for a given covariate pattern, the difference between the observed and expected success counts is asymptotically standard normal.

Grouping by EVP requires that the explanatory variables be categorical. The presence of a truly continuous covariate in the model has the consequence that the number of EVPs approaches or equals the total number of observations in the dataset. In such scenarios, we need the Hosmer-Lemeshow procedure, which groups the data by deciles of predicted probability instead of covariate patterns. Fortunately, both predictors in our racial integration model are dichotomous. We will now calculate both the Pearson residuals and deviance residuals for each covariate pattern:

```
Contact$phat<-fitted.values(Model)
Contact$yhat<-with(Contact,Obs.*phat)
Contact$PearsonRes<-with(Contact,(Y-yhat)/sqrt(yhat*(1-phat)))
Contact$DevianceRes<-with(Contact,
    sign(Y-yhat)*sqrt(2)*sqrt(Y*log(Y/yhat)+(Obs.-Y)*log((Obs.-Y)/(Obs.-yhat))))
Contact[,c(1:2,5:6,3,7:9)]</pre>
```

```
##
     Contact
                   Norms
                                        phat
                                               Y
                                                       yhat PearsonRes
                                 р
               Favorable 0.4270833 0.4506658 41
## 1
                                                  43.26392 -0.4643859
## 2
               Favorable 0.6976744 0.6845121 120 117.73608
                                                             0.3714618
## 3
           0 Unfavorable 0.2786070 0.2673437 56
                                                   53.73608 0.3608088
## 4
           1 Unfavorable 0.4748201 0.4911073 66
                                                  68.26392 -0.3841064
##
    DevianceRes
## 1
     -0.4652041
## 2
       0.3728929
## 3
       0.3592364
     -0.3842147
```

The group-specific residuals suggest a good fit. None of them is anywhere near 1.96 standard deviations in absolute value. We'll also calculate a summary statistic of the fit by adding up the squared Pearson residuals. This statistic follows a chi-squared distribution, with degrees of freedom equal to the number of covariate patterns (groups) minus the number of parameters (intercept plus predictors):

```
(OverallPearson <- sum(Contact$PearsonRes^2))

## [1] 0.6313589

1-pchisq(OverallPearson, df = 1)

## [1] 0.4268573
```

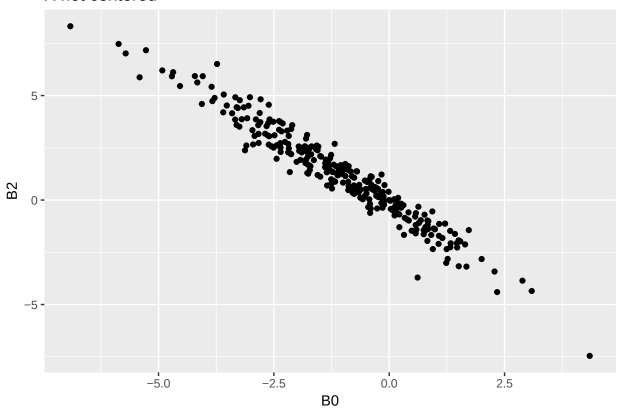
The residual degrees of freedom equal the number of binomial observations (4) minus the number of parameters in the model (3) = 1. The p-value is .43, indicating a statistically non-significant lack of fit. We can thereby conclude that the model fits quite well.

2. Effects of Variable Centering: Simulation

```
n <- 50
G <- c(rep(1,20),rep(0,30))
x <- vector(); for(i in 1:n) {xi <- i/n; x <- c(x,xi)}; rm(xi)
X <- cbind(rep(1,n),G,x)</pre>
```

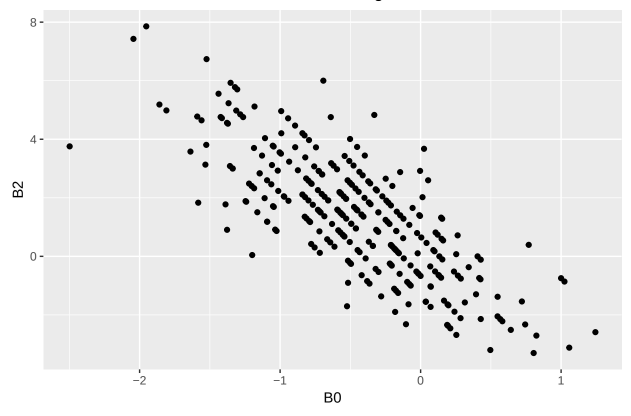
```
B \leftarrow c(-0.9, 0.3, 1)
logodds \leftarrow X[,1]*B[1] + X[,2]*B[2] + X[,3]*B[3]
p <- exp(logodds)/(1+exp(logodds))</pre>
N < -300
# x NOT centered:
MLEs <- data.frame()</pre>
for (i in 1:N){
  U <- runif(n)
  y <- as.numeric(p > U)
  model <- glm(y ~ G+x, family=binomial(link=logit))</pre>
  MLEs <- rbind(MLEs, coef(model))</pre>
};rm(model,y,U); names(MLEs)<-c("B0","B1","B2")</pre>
#Look at the MLEs:
sapply(MLEs, mean);cat("\n B2B0 correlation:\n");with(MLEs,cor(B2,B0))
                       B1
                                   B2
## -1.0680775 0.3666033 1.2101537
##
## B2B0 correlation:
## [1] -0.9712115
#x CENTERED:
x2 <- x-mean(x)
MLEs.2 <- data.frame()</pre>
for (i in 1:N){
  U <- runif(n)
  y <- as.numeric(p > U)
  model <- glm(y ~ G+x2, family=binomial(link=logit))</pre>
 MLEs.2 <- rbind(MLEs.2, coef(model))</pre>
};rm(model,y,U); names(MLEs.2)<-c("B0","B1","B2")</pre>
#Look at the MLEs:
sapply(MLEs.2, mean);cat("\n B2B0 correlation:\n");with(MLEs.2,cor(B2,B0))
##
           B0
                       В1
                                   B2
## -0.4631212 0.3841451 1.3523382
## B2B0 correlation:
## [1] -0.7762432
library(ggplot2)
(scatter1<-ggplot(data=MLEs, aes(x=B0, y=B2)) +
  geom_point(aes(x=B0,y=B2)) +
  ggtitle("X not centered"))
```

X not centered



```
(scatter2<-ggplot(data=MLEs.2, aes(x=B0, y=B2)) +
  geom_point(aes(x=B0,y=B2)) +
  ggtitle("X centered around its mean. Note the change in scale!"))</pre>
```

X centered around its mean. Note the change in scale!



The constant term, or intercept, is the estimated logit of the outcome when all continuous covariates equal 0 and all categorical covariates are at their reference level. This means that the coding of continuous covariates ought to be given careful thought – else the MLE for the intercept becomes uninteresting. In general, centering continuous covariates around their means and contrasting qualitative covariates with their most common category simplifies the interpretation of any regression model – in particular, it makes the constant term immediately informative. This is well illustrated by continuous covariates such as a subject's height and weight, where zero values are impossible in real data. Without centering, the intercept will be the logit-scale probability of the outcome for an individual who weighs 0 pounds and stands 0 feet tall, which is much less useful than an intercept representing the logit-scale probability for a person of average weight and height.

In our simulation data above, the true mean probability of the outcome is .43. The uncentered continuous covariate x takes on positive values from 0 to 1. True B2 is 1, while "true" B0 (intercept) is -0.9. If we center x around a smaller value after the outcomes are generated, its MLE remains the same because it is not based on the x values themselves but on the relationship between the observed variation in x and the observed variation in y. B0, by contrast, represents not the relationship between some variable and the outcome but the estimated logit-scale probability of the outcome before the covariates have had an effect – in other words, when they equal zero. Therefore, when the values of x are reduced, i.e., brought closer to zero by centering it while the data is kept fixed, x0 must necessarily increase, because the effect of x at "zero" now represents the effect of x at its mean.

All of the above entails that it is possible to manipulate the intercept of a model in almost any way we want by adjusting the origins and/or scales of the continuous covariates. Such manipulation is sometimes advisable if it can make the interpretation of the model coefficients simpler and more intuitive.

3. Conditional Logistic Regression for Matched-Pairs Case Control Data

We'll now investigate what factors might be associated with low birth weight in newborns. The data comes in matched-pairs case-control format, so we'll use conditional logistic regression to remove the unmeasured, pair-specific effects from the equation. The measured covariates available are:

- 1. Race (nominal: 1=white, 2=black, 3=other)
- 2. Smoking during pregnancy (dichotomous)
- 3. PTD: history of premature labor (dichotomous)
- 4. HT: History of hypertension (dichotomous)
- 5. UI: Intrauterine Irritability (dichotomous)

I began the analysis by entering the aforementioned 5 variables plus every conceivable two-way interaction term into the model. It was messy and therefore isn't shown here. Many of the interactions could not be analyzed because the data was too sparse, or there were singularities. Out of the interactions that could be analyzed, however, none was statistically significant. We can thus proceed to fitting a model with main effects only:

```
library(survival)
lowbw <- read.table(file="lowbwtm11CSV.csv",sep=";",header=T)</pre>
names(lowbw)
## [1] "i..PAIR" "LOW"
                             "AGE"
                                       "LWT"
                                                  "RACE"
                                                             "SMOKE"
                                                                        "PTD"
## [8] "HT"
                  "IJT"
names(lowbw)[1]<-"PAIR"</pre>
lowbw$RACE <- as.factor(lowbw$RACE)</pre>
levels(lowbw$RACE) <- c("WHITE", "BLACK", "OTHER")</pre>
summary(bw1 <- clogit(LOW ~ RACE + SMOKE + PTD + HT + UI + strata(PAIR), data = lowbw))</pre>
## Call:
  coxph(formula = Surv(rep(1, 112L), LOW) ~ RACE + SMOKE + PTD +
##
##
       HT + UI + strata(PAIR), data = lowbw, method = "exact")
##
##
     n= 112, number of events= 56
##
                coef exp(coef) se(coef)
##
                                             z Pr(>|z|)
## RACEBLACK 0.4340
                        1.5434
                                                 0.52260
                                  0.6788 0.639
## RACEOTHER 0.5308
                        1.7003
                                  0.5803 0.915
                                                 0.36031
## SMOKE
             1.5939
                        4.9230
                                  0.5947 2.680
                                                 0.00736 **
## PTD
                        5.3795
                                  0.7209 2.334
             1.6826
                                                 0.01960 *
             1.5011
## HT
                        4.4865
                                  0.8322 1.804
                                                 0.07126 .
## UI
             1.3320
                        3.7887
                                  0.7034 1.894
                                                0.05826 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
             exp(coef) exp(-coef) lower .95 upper .95
## RACEBLACK
                  1.543
                            0.6479
                                       0.4080
                                                   5.838
                  1.700
                            0.5881
                                                   5.302
## RACEOTHER
                                       0.5453
## SMOKE
                  4.923
                            0.2031
                                       1.5347
                                                  15.792
## PTD
                  5.379
                            0.1859
                                       1.3095
                                                  22.099
## HT
                  4.487
                            0.2229
                                       0.8782
                                                  22.921
## UI
                  3.789
                            0.2639
                                       0.9545
                                                  15.039
##
## Rsquare= 0.18
                    (max possible= 0.5)
```

```
## Likelihood ratio test= 22.25 on 6 df, p=0.001089

## Wald test = 12.42 on 6 df, p=0.05325

## Score (logrank) test = 17.83 on 6 df, p=0.00666
```

Race is about to be dropped from the model. We'll verify this through a comparison of nested models:

```
bw2 <- clogit(LOW ~ SMOKE + PTD + HT + UI + strata(PAIR), data = lowbw)
anova(bw1,bw2,test = "Chisq")</pre>
```

```
## Analysis of Deviance Table
## Cox model: response is Surv(rep(1, 112L), LOW)
## Model 1: ~ RACE + SMOKE + PTD + HT + UI + strata(PAIR)
## Model 2: ~ SMOKE + PTD + HT + UI + strata(PAIR)
## loglik Chisq Df P(>|Chi|)
## 1 -27.689
## 2 -28.149 0.9197 2 0.6314
```

Dropping RACE did not significantly worsen the fit, and it made the model more parsimonious:

summary(bw2)

```
## Call:
   coxph(formula = Surv(rep(1, 112L), LOW) ~ SMOKE + PTD + HT +
##
##
       UI + strata(PAIR), data = lowbw, method = "exact")
##
##
     n= 112, number of events= 56
##
##
           coef exp(coef) se(coef)
                                         z Pr(>|z|)
## SMOKE 1.3808
                    3.9779
                             0.5256 2.627
                                            0.00861 **
         1.6411
                    5.1611
                             0.6891 2.382
## PTD
                                            0.01723
## HT
         1.6042
                    4.9739
                             0.7918 2.026
                                            0.04275 *
## UI
         1.3592
                    3.8929
                             0.7017 1.937
                                            0.05277 .
##
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
         exp(coef) exp(-coef) lower .95 upper .95
## SMOKE
             3.978
                        0.2514
                                   1.4199
                                              11.14
## PTD
             5.161
                        0.1938
                                   1.3372
                                              19.92
## HT
             4.974
                        0.2010
                                   1.0538
                                              23.48
## UI
             3.893
                        0.2569
                                   0.9839
                                              15.40
##
## Rsquare= 0.173
                     (max possible= 0.5)
## Likelihood ratio test= 21.33
                                  on 4 df,
                                              p=0.0002719
## Wald test
                         = 12.15
                                  on 4 df,
                                              p=0.01629
## Score (logrank) test = 17.15
                                  on 4 df,
                                              p=0.001804
```

All the remaining variables are significant. Intrauterine irritability has a p-value on the borderline of significance, but we'll keep it in the model. Newborn babies' health is at stake here, so it's prudent to be cautious and attentive of any suspicious factors, even if they fall 2.7 per mille short of the conventional significance threshold. The model output shows the logit-scale MLEs converted into odds ratios as well as the 95% confidence intervals of these odds ratios: smoking during pregnancy increases the odds of the baby having a low birth weight by a factor of 1.4 to 11. For those having a history of premature labor, the odds of delivering a baby with a low birth weight are from 1.3 to 20 times as high as the odds for those without such a history. Hypertension increases the odds by a factor ranging from 1.05 to 23. Intrauterine irritability multiplies the odds by factor of 1 to 15.

4. Predicting Membership in the Hilmo Drug User Database

We'll now use logistic regression to model the probability of double registration for criminal drug users. Unfortunately the data file is not in a format immediately usable by R, so we have to edit it a little before importing:

```
orig.file<-scan(what="char",file="drug-users.txt",sep="\n")
head(orig.file,10)
    [1] "Data on Drug Users"
##
##
    [2] "hilmo = 1, if registered in hilmo, otherwise = 0"
##
    [3] "riki = 1, if registered in riki, otherwise = 0"
    [4] "age (in years)"
##
   [5] "male = 1, if male, = 0 if female"
    [6] "sta = family status: 1 = married or cohabiting, 2 = single, 3 = widowed, 4 = divorced, 5 = not
    [7] "hilmo riki age male
                                   sta\t"
##
##
    [8] " 1
                 0
                       41
                             0
   [9] " 1
##
                       32
                             1
                                   2"
## [10] " 0
                                   2"
                 1
                       23
                             0
library(stringr)
drug_orig<-str_replace_all(orig.file[7:length(orig.file)],"^ +","")</pre>
drug_orig<-str_replace_all(drug_orig," +","\t")</pre>
drug_orig<-str_replace_all(drug_orig,"\t(?=$)",""); cat(drug_orig,file="drug_orig.txt",sep="\n")</pre>
drug_orig<-read.table(file="drug_orig.txt",sep="\t",header=T)</pre>
We also have to convert the relationship status variable from numeric to nominal:
drug_orig$sta<-as.factor(drug_orig$sta); levels(drug_orig$sta)</pre>
## [1] "1" "2" "3" "4" "5"
(levels(drug_orig$sta)<-c("HasPartner", "Single", "Widowed", "Divorced", NA))
## [1] "HasPartner" "Single"
                                    "Widowed"
                                                  "Divorced"
drug_orig[which(is.na(drug_orig$sta)),]
##
      hilmo riki age male sta
## 98
                   46
                          O <NA>
                1
Unfortunately, this lone NA value in the sta varuabke will cause problems with our probability calculations
later. A NA value will prevent the automated calculation of a fitted probability for that observation, which
will consequently prevent the calculation of means and other statistical parameters for the fitted probabilities.
We will therefore impute a likely value for the missing field. And since the sta variable is categorical, what
```

better way to impute it than logistic regression? We'll just need to use the polytomous variety:

```
library(nnet)
drug <- drug_orig</pre>
drug$sta[98] <- predict(</pre>
  multinom(sta ~ riki + hilmo + age + male, data=drug), newdata=drug[98,], type="class")
## # weights: 24 (15 variable)
## initial value 3593.274984
## iter 10 value 1774.095050
## iter 20 value 1479.097664
## iter 30 value 1478.872792
## iter 30 value 1478.872779
```

```
## iter 30 value 1478.872779
## final value 1478.872779
## converged
drug[98,]
```

```
## hilmo riki age male sta
## 98     0     1     46     0 Divorced
```

Much better. Now we can proceed.

There are two databases that keep track of drug users in Finland – the Hospital Discharge Register hilmo and the Criminal Report Register riki. Our aggregate dataset drug lists everyone who was registered as a drug user in one or both of these databases around the year 2000. The ultimate goal of these exercises is to estimate the total number of drug users in the Uusimaa province of Southern Finland at the turn of the millennium. First we will model the probability that subjects registered in riki are also found in hilmo. The covariates at our disposal are age, sex, and relationship status. We'll start by analyzing the main effects plus all potential two-way interactions. Also, having learned from exercise 2, we'll center the age variable around its mean:

```
d0<-drug[drug$riki==1,]
d0$ageCentered<-d0$age-mean(d0$age)
summary(drugmodel0<-glm(hilmo ~ ageCentered*male + male*sta + ageCentered*sta,
    family=binomial(link=logit), data=d0))</pre>
```

```
##
## Call:
  glm(formula = hilmo ~ ageCentered * male + male * sta + ageCentered *
##
       sta, family = binomial(link = logit), data = d0)
##
## Deviance Residuals:
##
       Min
                 10
                      Median
                                   30
                                           Max
   -0.6927
           -0.4091 -0.3752 -0.3265
                                        2.5018
##
##
## Coefficients:
##
                             Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                           -2.443e+00
                                       6.346e-01
                                                  -3.849 0.000119 ***
## ageCentered
                           -1.261e-01
                                       7.705e-02
                                                  -1.636 0.101809
## male
                           -6.202e-01
                                       8.659e-01
                                                  -0.716 0.473838
## staSingle
                            1.837e-01
                                       6.936e-01
                                                   0.265 0.791151
## staWidowed
                            1.803e+01
                                       1.145e+03
                                                   0.016 0.987439
## staDivorced
                           -1.228e-01
                                       9.436e-01
                                                  -0.130 0.896430
## ageCentered:male
                           -3.288e-03
                                       4.302e-02
                                                  -0.076 0.939080
## male:staSingle
                            1.883e-01
                                       9.152e-01
                                                   0.206 0.836987
## male:staWidowed
                           -3.053e+01
                                       1.913e+03
                                                  -0.016 0.987269
## male:staDivorced
                            6.291e-01
                                       1.163e+00
                                                    0.541 0.588636
## ageCentered:staSingle
                            9.346e-02
                                       7.607e-02
                                                    1.229 0.219208
## ageCentered:staWidowed
                            1.293e-01
                                       9.077e+01
                                                    0.001 0.998863
## ageCentered:staDivorced 1.024e-01
                                       8.361e-02
                                                    1.225 0.220499
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 813.71 on 1578
##
                                       degrees of freedom
## Residual deviance: 787.62 on 1566
                                       degrees of freedom
```

```
## AIC: 813.62
##
## Number of Fisher Scoring iterations: 14
None of the interactions is significant, so we'll drop them. Next, we'll analyze main effects only:
summary(drugmodel1<-glm(hilmo ~ ageCentered + male + sta, family=binomial(link=logit), data=d0))</pre>
##
## glm(formula = hilmo ~ ageCentered + male + sta, family = binomial(link = logit),
##
       data = d0)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -0.9888
           -0.4137 -0.3731
                              -0.3157
                                         2.6095
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
                            0.44110
                                     -5.860 4.63e-09 ***
## (Intercept) -2.58483
## ageCentered -0.04297
                            0.01677
                                     -2.562
                                              0.0104 *
               -0.42248
                            0.23168
                                     -1.824
                                              0.0682
## male
## staSingle
                0.29022
                            0.44789
                                      0.648
                                              0.5170
## staWidowed
                2.54505
                            1.03910
                                      2.449
                                              0.0143 *
## staDivorced 0.45167
                            0.52100
                                      0.867
                                              0.3860
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 813.71 on 1578
                                       degrees of freedom
## Residual deviance: 794.79 on 1573
                                        degrees of freedom
## AIC: 806.79
##
## Number of Fisher Scoring iterations: 5
```

Thanks to our centering of age, the intercept is immediately relevant. It tells us that, according to this model, a drug user who is aged 28, female, and in a relationship, has a probability of

$$e^{-2.58388/(1+e^{-2.58388})} = .07$$

of inclusion in *hilmo*. Age is negatively associated with inclusion in the database, while being Widowed has a very strong positive association. Maleness is negatively associated – a phenomenon which is intuitively believable since common sense suggests that men are less likely than women to seek medical help in distress. The effect falls just short of statistical significance, however.

Within the categorical variable describing relationship status, only the Widowed category has a significant effect. It is therefore worthwhile experimenting with a dummy variable that indicates simply whether the person is Widowed (1) or not (0). This effectively collapses the three remaining levels together into one large reference category representing the vast majority of the subjects in the dataset:

```
d0$Widowed<-as.numeric(d0$sta=="Widowed")
summary(drugmodel2<-glm(hilmo~ageCentered+male+Widowed,family=binomial(link=logit),data=d0))
##
## Call:
## glm(formula = hilmo ~ ageCentered + male + Widowed, family = binomial(link = logit),
## data = d0)</pre>
```

```
##
## Deviance Residuals:
##
       Min
                  1Q
                       Median
                                             Max
  -0.9886 -0.4122 -0.3730 -0.3174
                                          2.6582
##
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                            0.20219 -11.364
## (Intercept) -2.29755
                                              < 2e-16 ***
## ageCentered -0.04157
                            0.01444
                                      -2.879
                                             0.00399 **
## male
               -0.41708
                            0.22765
                                     -1.832
                                              0.06694
## Widowed
                2.25190
                            0.95705
                                       2.353
                                              0.01863 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 813.71 on 1578
                                         degrees of freedom
## Residual deviance: 795.58
                               on 1575
                                         degrees of freedom
## AIC: 803.58
##
## Number of Fisher Scoring iterations: 5
anova(drugmodel1,drugmodel2,test="LRT")
## Analysis of Deviance Table
##
## Model 1: hilmo ~ ageCentered + male + sta
## Model 2: hilmo ~ ageCentered + male + Widowed
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1573
                    794.79
## 2
          1575
                    795.58 -2 -0.78917
                                           0.674
This is a clear improvement. Residual deviance increased only negligibly (p = .67), while df improved by 2
points and AIC by 5 points. We'll keep this change. Next, let's try breaking age into cohorts in order to
examine the exact nature of its effect. We'll contrast the other age brackets with the one containing the mean.
summary(d0$age)
##
      Min. 1st Qu.
                     Median
                               Mean 3rd Qu.
                                                Max.
```

##

15.00

hist(d0\$age)

22.00

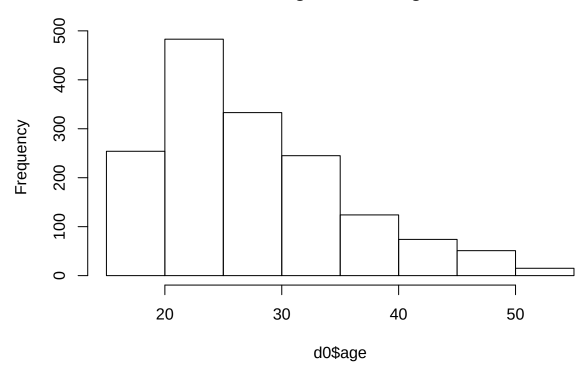
26.00

28.02

33.00

55.00

Histogram of d0\$age



```
d0\$agegroup < -cut(d0\$age, breaks = c(min(d0\$age), 19, 24, 29, 34, 39, 44, 49, max(d0\$age)), include.lowest = TRUE)
addmargins(table(d0$agegroup)); nrow(d0)
## [15,19] (19,24] (24,29] (29,34] (34,39] (39,44] (44,49] (49,55]
                                                                           Sum
       162
                483
                        365
                                272
                                                                          1579
##
                                         138
                                                   83
                                                           55
                                                                    21
## [1] 1579
d0$agegroup<-relevel(d0$agegroup,ref="(24,29]")
summary(drugmodel3<-glm(hilmo~agegroup+male+Widowed,family=binomial(link=logit),data=d0))</pre>
##
   glm(formula = hilmo ~ agegroup + male + Widowed, family = binomial(link = logit),
       data = d0)
##
##
## Deviance Residuals:
##
       Min
                  1Q
                       Median
                                     3Q
                                             Max
   -1.2114
            -0.4368
                     -0.3808
                              -0.3340
                                          2.8437
##
## Coefficients:
##
                    Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                     -1.8041
                                 0.2594 -6.956 3.51e-12 ***
                    -0.4075
                                          -1.154
## agegroup[15,19]
                                 0.3531
                                                    0.2485
                    -0.2858
## agegroup(19,24]
                                 0.2457
                                          -1.163
                                                    0.2448
## agegroup(29,34]
                     -0.5567
                                 0.3127
                                          -1.781
                                                    0.0750
```

0.0755 .

-1.778

0.4296

agegroup(34,39]

-0.7636

```
## agegroup(39,44]
                    -2.2214
                                         -2.174
                                                   0.0297 *
                                 1.0220
                                         -1.419
                                                   0.1560
## agegroup(44,49]
                    -1.0527
                                 0.7421
## agegroup(49,55] -14.5978
                               505.4343
                                         -0.029
                                                   0.9770
                    -0.4978
                                 0.2304
                                         -2.160
                                                   0.0308 *
## male
## Widowed
                      2.3814
                                 1.0301
                                          2.312
                                                   0.0208 *
##
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 813.71
                               on 1578
                                        degrees of freedom
## Residual deviance: 787.33
                              on 1569
                                        degrees of freedom
   AIC: 807.33
##
## Number of Fisher Scoring iterations: 15
```

The effect of age not linear. Instead, it seems to have a similar effect as it does on athletic performance, which is generally high in youth, peaks in the late 20s and then starts to deteriorate as a function of time. The reference category (25-to-29-year-olds) has by far the highest risk of inclusion.

One unexpected consequence of dividing age into cohorts is that maleness becomes statistically significant, which it wasn't in the model where age was continuous. How can this be explained? One possibility is that one or more of the cohorts may, purely by chance, have a high proportion of males avoiding hospitalization on the top end of the age bracket. With age as categorical, the model would attribute such within-bracket associations between age and the outcome solely to maleness rather than dividing it between age and maleness.

Which is the better model between the one with continuous age and the one where it's bracketed? We cannot compare them directly using a LRT because the models are not nested – one treats age as continuous, the other as nominal, so this is a matter of different variables rather than extra variables.

Judging by the smaller AIC, we should perhaps prefer the simpler model with continuous age to the one with age brackets. Though the effect of age is not exactly linear, the overall trend is still that old people are less likely to be registered than young people. In fact, a binary age division between the old and the young might further improve the model. The output of the bracketed-age model already gives us clues as to what the best threshold value might be. We'll try to find one now.

After experimenting with different age cutoffs (not shown), I learned that a binary age distinction between those under 33 and the rest results in a better fit than having age as continuous, and the model is equally parsimonious.

```
d0$AgeUnder33<-as.numeric(d0$age<33)
summary(drugmodel4<-glm(hilmo~AgeUnder33+Widowed+male,family=binomial(link=logit),data=d0))</pre>
##
## Call:
   glm(formula = hilmo ~ AgeUnder33 + Widowed + male, family = binomial(link = logit),
##
       data = d0)
##
## Deviance Residuals:
##
       Min
                  10
                       Median
                                     30
                                             Max
## -1.0666
            -0.3955
                      -0.3955
                               -0.2527
                                          2.6307
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                -2.9655
                             0.3274
                                      -9.058
                                             < 2e-16 ***
                             0.2935
                                              0.00173 **
## AgeUnder33
                  0.9193
                                       3.132
## Widowed
                  2.2429
                             0.9618
                                       2.332
                                              0.01970 *
```

```
-0.4630
                            0.2263 -2.046 0.04075 *
## male
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 813.71 on 1578 degrees of freedom
                                      degrees of freedom
## Residual deviance: 792.83 on 1575
## AIC: 800.83
##
## Number of Fisher Scoring iterations: 6
anova(drugmodel2, drugmodel4)
## Analysis of Deviance Table
##
## Model 1: hilmo ~ ageCentered + male + Widowed
## Model 2: hilmo ~ AgeUnder33 + Widowed + male
     Resid. Df Resid. Dev Df Deviance
## 1
         1575
                  795.58
## 2
          1575
                   792.83 0
                               2.7572
```

The age-bracketed model above has lower residual deviance, but its higher number of parameters results in diminished parsimony and a higher AIC (807.33 as opposed to the current model's 800.83). Therefore, drugmodel4 is our choice. It's got three binary predictors, each of which has high predictive power and is simple to interpret. We'll finish by predicting the probabilities of belonging to *hilmo* for every subject in the entire drug dataset, and viewing 10 of them at random:

```
drug$Widowed<-as.numeric(drug$sta=="Widowed")
drug$AgeUnder33<-as.numeric(drug$age<33)
drug$FittedHilmoProbs <- predict(drugmodel4,newdata=drug,type="response")
sample(drug$FittedHilmoProbs, size=10)

## [1] 0.03141681 0.07521518 0.03141681 0.07521518 0.07521518 0.03141681
## [7] 0.07521518 0.07521518 0.03141681 0.03141681</pre>
```

5. Predicting Membership in Criminal Report Rekister Riki

Now we'll perform a similar analysis in the other direction, trying to predict whether those registered as drug users in hilmo have been caught committing crimes and listed in riki as drug users. We'll begin with a model containing all the main effects and potential two-variable interactions. We will also center age around its mean again:

```
d1<-drug[drug$hilmo==1,]; row.names(d1)<-1:nrow(d1)
d1$ageCentered<-d1$age-mean(d1$age)
summary(DrugModel0<-glm(riki ~ ageCentered*male + male*sta +ageCentered*sta, family=binomial(link=logit
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
##
## Call:
## glm(formula = riki ~ ageCentered * male + male * sta + ageCentered *</pre>
```

Max

sta, family = binomial(link = logit), data = d1)

3Q

##

##

Deviance Residuals:

1Q

Median

Min

```
## -0.6647 -0.5146 -0.4518 -0.3492
                                       2.6432
##
## Coefficients:
                            Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                          -3.002e+00 6.513e-01 -4.609 4.05e-06 ***
## ageCentered
                          -1.359e-01 7.922e-02 -1.716
                                                          0.0862 .
                                                 0.368
                                                          0.7132
## male
                           3.222e-01 8.764e-01
## staSingle
                           5.382e-01 7.049e-01
                                                  0.764
                                                          0.4452
## staWidowed
                           2.429e+00 5.770e+02
                                                  0.004
                                                          0.9966
## staDivorced
                           3.587e-01 9.379e-01
                                                  0.382
                                                          0.7022
## ageCentered:male
                           1.141e-03 3.935e-02
                                                  0.029
                                                          0.9769
## male:staSingle
                          -2.495e-02 9.214e-01 -0.027
                                                          0.9784
## male:staWidowed
                          -2.769e+01 2.059e+03 -0.013
                                                          0.9893
## male:staDivorced
                                                          0.5979
                           6.150e-01 1.166e+00
                                                  0.527
## ageCentered:staSingle
                           8.848e-02 7.974e-02
                                                          0.2672
                                                  1.110
## ageCentered:staWidowed -3.142e+00 1.285e+02 -0.024
                                                          0.9805
## ageCentered:staDivorced 8.817e-02 8.733e-02
                                                 1.010
                                                          0.3127
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 734.05 on 1126 degrees of freedom
## Residual deviance: 702.04 on 1114 degrees of freedom
## AIC: 728.04
## Number of Fisher Scoring iterations: 16
None of the interaction terms is significant, and the whole model is quite confusing to interpret. We'll drop
the interactions:
summary(DrugModel1<-glm(riki ~ ageCentered + male + sta, family=binomial(link=logit), data=d1))</pre>
##
## Call:
  glm(formula = riki ~ ageCentered + male + sta, family = binomial(link = logit),
##
       data = d1)
##
## Deviance Residuals:
      Min
                10
                     Median
                                  30
                                          Max
## -0.9205 -0.5166 -0.4427 -0.3387
                                        2.5389
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.98793
                          0.43928 -6.802 1.03e-11 ***
## ageCentered -0.05742
                          0.01659 -3.460 0.00054 ***
## male
               0.36228
                          0.23368
                                    1.550 0.12107
## staSingle
               0.43188
                          0.44917
                                    0.962 0.33630
              1.77874
## staWidowed
                                     1.943 0.05203 .
                          0.91552
## staDivorced 0.81244
                          0.52622
                                    1.544 0.12260
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
```

```
## Null deviance: 734.05 on 1126 degrees of freedom
## Residual deviance: 712.68 on 1121 degrees of freedom
## AIC: 724.68
##
## Number of Fisher Scoring iterations: 5
```

##

15.00

22.00

26.00

28.61

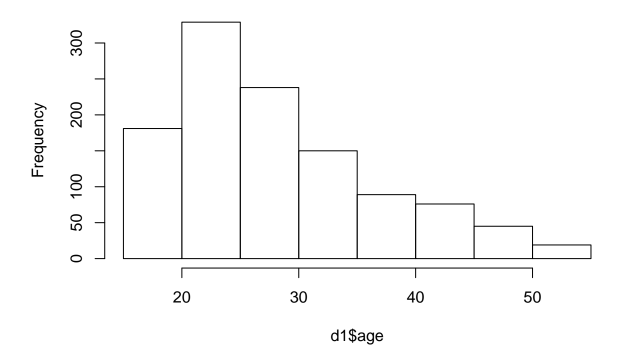
Here, age has an even stronger (disfavoring) effect than in the previous dataset. This makes intuitive sense—the old are less hotheaded than the young and therefore less prone to rash acts of violence and crime. Much like the risk of inclusion in hilmo, being Widowed also increases the risk of inclusion in riki. Maleness increases the risk of inclusion, but at p=.12 the effect falls somewhat short of statistical significance. Let's study the effect of age more closely by dividing the variable into cohorts:

```
summary(d1$age); hist(d1$age)
## Min. 1st Qu. Median Mean 3rd Qu. Max.
```

33.00

Histogram of d1\$age

55.00



```
d1$agegroup<-cut(d1$age,breaks=c(min(d1$age),19,24,29,34,39,44,49,max(d1$age)),include.lowest = TRUE)
addmargins(table(d1$agegroup)); nrow(d1)
##
## [15,19] (19,24] (24,29] (29,34] (34,39] (39,44] (44,49] (49,55] Sum</pre>
```

```
## [15,19] (19,24] (24,29] (29,34] (34,39] (39,44] (44,49] (49,55] Sum
## 111 339 265 157 98 77 57 23 1127

## [1] 1127
d1$agegroup<-relevel(d1$agegroup,ref="(24,29]") #Make 25-30 the reference level
summary(DrugModel2<-glm(riki ~ agegroup + male + sta, family=binomial(link=logit), data=d1))</pre>
```

```
##
## Call:
##
  glm(formula = riki ~ agegroup + male + sta, family = binomial(link = logit),
##
       data = d1)
##
## Deviance Residuals:
       Min
                 10
                      Median
                                    30
                                            Max
  -0.9877 -0.5161 -0.4604 -0.3203
                                         2.8992
##
##
## Coefficients:
##
                   Estimate Std. Error z value Pr(>|z|)
                    -2.5467
                                0.4646
                                        -5.482 4.22e-08 ***
## (Intercept)
## agegroup[15,19]
                    -0.2224
                                 0.3617
                                         -0.615
                                                  0.5387
## agegroup(19,24]
                                         -0.650
                    -0.1635
                                 0.2515
                                                  0.5155
                    -0.3977
                                         -1.218
                                                  0.2233
## agegroup(29,34]
                                 0.3266
## agegroup(34,39]
                    -0.8433
                                 0.4475
                                         -1.885
                                                  0.0595
                    -2.5417
                                         -2.448
## agegroup(39,44]
                                                  0.0144 *
                                 1.0384
## agegroup(44,49]
                    -1.5648
                                 0.7662
                                         -2.042
                                                  0.0411 *
                                         -0.030
## agegroup(49,55] -14.8648
                               487.9613
                                                  0.9757
## male
                     0.2421
                                 0.2367
                                          1.023
                                                  0.3065
## staSingle
                     0.5193
                                 0.4468
                                          1.162
                                                  0.2452
## staWidowed
                     1.8404
                                 0.9395
                                          1.959
                                                  0.0501 .
## staDivorced
                     0.9008
                                          1.691
                                                  0.0909 .
                                 0.5329
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
  (Dispersion parameter for binomial family taken to be 1)
##
##
##
       Null deviance: 734.05 on 1126 degrees of freedom
## Residual deviance: 704.57 on 1115 degrees of freedom
## AIC: 728.57
##
## Number of Fisher Scoring iterations: 15
```

The effect of age seems similarly shaped as in the previous dataset – its favoring effect peaks in the late twenties, then reverses. I'll now see if I can find a threshold value at which age could be conveniently dichotomized to improve the model (experimentation not shown).

It appears that 33 is the best threshold for a binary age classification is this dataset too:

Estimate Std. Error z value Pr(>|z|)

0.5085

0.3295

Coefficients:

-3.8801

1.1715

(Intercept)

AgeUnder33

```
d1$AgeUnder33<-as.numeric(d1$age<33)
summary(DrugModel3<-glm(riki~AgeUnder33+male+sta,family=binomial(link=logit),data=d1))</pre>
##
## Call:
   glm(formula = riki ~ AgeUnder33 + male + sta, family = binomial(link = logit),
##
       data = d1)
##
##
  Deviance Residuals:
##
                 1Q
                      Median
                                     30
                                             Max
##
   -1.0006
           -0.5302 -0.4623
                               -0.3023
                                          2.6032
##
```

-7.630 2.35e-14 ***

3.556 0.000377 ***

```
## male
                 0.2913
                            0.2306
                                     1.263 0.206486
## staSingle
                 0.5261
                            0.4429
                                     1.188 0.234881
## staWidowed
                 1.9862
                            0.9269
                                     2.143 0.032120 *
## staDivorced
                 0.8484
                            0.5280
                                     1.607 0.108067
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 734.05
                              on 1126
                                       degrees of freedom
## Residual deviance: 710.95
                              on 1121
                                       degrees of freedom
  AIC: 722.95
##
##
## Number of Fisher Scoring iterations: 5
anova(DrugModel1,DrugModel3)
## Analysis of Deviance Table
## Model 1: riki ~ ageCentered + male + sta
## Model 2: riki ~ AgeUnder33 + male + sta
     Resid. Df Resid. Dev Df Deviance
## 1
          1121
                   712.68
## 2
          1121
                   710.95 0
                               1.7256
```

This model fits the data better, on the same degrees of freedom, as the one where age was continuous. It seems safe to prefer the binary classification.

A further improvement can be accomplished by removing the non-significant distinction between single and divorced people within the *sta* variable. We will collapse these two categories together into a new reference category that will be contrasted with the more statistically significant classes. Being Widowed is clearly significant, but having a partner might be significant too – we have no direct measure of its effect because it is the variable's current baseline, and it is being contrasted with three different categories. By changing the baseline to Single/Divorced, we will get an overt estimate of the effect of having a partner. The merging the two non-significant cateogries will gain us a degree of freedom at little cost in residual deviance. We will call our new, trimmed version of the variable *sta.3way*:

```
levels(d1$sta)
## [1] "HasPartner" "Single"
                                   "Widowed"
                                                 "Divorced"
d1$sta.3way <- d1$sta
levels(d1$sta.3way) <- c("HasPartner", "Single/Divorced", "Widowed", "Single/Divorced")</pre>
d1$sta.3way <- relevel(d1$sta.3way, "Single/Divorced")</pre>
summary(DrugModel4<-glm(riki~AgeUnder33+male+sta.3way,family=binomial(link=logit),data=d1))</pre>
##
## Call:
   glm(formula = riki ~ AgeUnder33 + male + sta.3way, family = binomial(link = logit),
##
       data = d1)
##
## Deviance Residuals:
##
       Min
                  10
                       Median
                                     3Q
                                             Max
##
   -0.9687 -0.5337
                     -0.4708 -0.3220
                                          2.5458
##
## Coefficients:
```

Estimate Std. Error z value Pr(>|z|)

##

```
## (Intercept)
                       -3.2006
                                   0.3340 -9.584 < 2e-16 ***
                                            3.522 0.000429 ***
                        1.0569
## AgeUnder33
                                   0.3001
                        0.2668
## male
                                   0.2287
                                            1.167 0.243398
## sta.3wayHasPartner -0.5809
                                   0.4384
                                           -1.325 0.185164
## sta.3wayWidowed
                        1.3639
                                   0.8369
                                            1.630 0.103156
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 734.05 on 1126 degrees of freedom
## Residual deviance: 711.72 on 1122 degrees of freedom
## AIC: 721.72
##
## Number of Fisher Scoring iterations: 5
anova(DrugModel3,DrugModel4,test="LRT")
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + male + sta
## Model 2: riki ~ AgeUnder33 + male + sta.3way
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1121
                   710.95
## 2
          1122
                   711.72 -1 -0.76874
                                        0.3806
```

The chi-square statistic for the increase in deviance is only .77 at 1 degree of freedom, which supports our decision to trim the variable.

We now face a decision regarding the gender variable. It is not having the same effect as in the riki dataset. Let's try dropping it:

```
summary(DrugModel5<-glm(riki~AgeUnder33+sta.3way,family=binomial(link=logit),data=d1))</pre>
```

```
##
## Call:
## glm(formula = riki ~ AgeUnder33 + sta.3way, family = binomial(link = logit),
##
       data = d1)
##
## Deviance Residuals:
##
                 1Q
                      Median
                                   3Q
                                           Max
## -0.8909
           -0.5158
                    -0.5158 -0.3114
                                        2.4702
##
## Coefficients:
##
                      Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                       -3.0024
                                   0.2844 -10.556 < 2e-16 ***
## AgeUnder33
                        1.0525
                                   0.3001
                                            3.507 0.000453 ***
## sta.3wayHasPartner -0.6396
                                   0.4355
                                           -1.469 0.141909
## sta.3wayWidowed
                        1.2307
                                   0.8261
                                            1.490 0.136312
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
                                       degrees of freedom
       Null deviance: 734.05 on 1126
## Residual deviance: 713.12 on 1123
                                       degrees of freedom
```

```
## AIC: 721.12
##
## Number of Fisher Scoring iterations: 5
anova(DrugModel4,DrugModel5,test = "LRT")
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + male + sta.3way
## Model 2: riki ~ AgeUnder33 + sta.3way
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1122
                   711.72
## 2
          1123
                   713.12 -1 -1.4042
                                          0.236
```

At p = 0.24, there is relatively robust statistical evidence for removing gender from the model. We will drop the variable. However, since gender is always of scientific interest, we will state for the record that its effect on inclusion in *riki* generally ranged between 1 and 1.6 on the z-scale, varying according to what other predictors were in the model.

The choices get harder from this point on. We will make use of two more statistics in our endeavor to select the best model. In addition to the familiar LRT, ANOVA and AIC, we will be measuring each model candidate's testing error in leave-one-out crossvalidation. Secondly, we will calculate BIC for each model. AIC and BIC are both designed to provide a measure of the model's fit while taking its complexity into account. They accomplish this by adding a penalty to the deviance residual for each additional parameter, and their difference lies in the magnitude of that penalty. No consensus exists on which criterion is better, but AIC is considered relatively lenient towards added complexity, while BIC penalizes it more severely.

The two relationship statuses contrasted with the new baseline have opposite effects of almost identical magnitude. Both also fall somewhat short of the statistical significance threshold. Let's see what happens if we remove the ostensibly least significant remaining variable, i.e., the indicator for having a romantic partner.

summary(DrugModel6<-glm(riki~AgeUnder33+Widowed,family=binomial(link=logit),data=d1))</pre>

```
##
## Call:
  glm(formula = riki ~ AgeUnder33 + Widowed, family = binomial(link = logit),
       data = d1)
##
##
  Deviance Residuals:
##
##
                 10
                      Median
                                   30
                                           Max
           -0.5062 -0.5062
  -0.9031
                             -0.2967
                                        2.5080
##
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                            0.2793 -11.104 < 2e-16 ***
   (Intercept)
                -3.1011
## AgeUnder33
                 1.1109
                            0.2981
                                     3.727 0.000194 ***
## Widowed
                 1.3039
                            0.8270
                                     1.577 0.114859
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 734.05 on 1126
                                       degrees of freedom
  Residual deviance: 715.66
                             on 1124
                                       degrees of freedom
  AIC: 721.66
##
## Number of Fisher Scoring iterations: 5
```

```
anova(DrugModel5,DrugModel6,test = "LRT")
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + sta.3way
## Model 2: riki ~ AgeUnder33 + Widowed
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1123
                   713.12
## 2
          1124
                    715.66 -1 -2.5351
                                         0.1113
if(exists("survival",mode="function")){detach("package:survival",unload=TRUE)}
library(boot)
##
## Attaching package: 'boot'
## The following object is masked from 'package:survival':
##
##
       aml
ErrorRate<-function(Y,prob){mean(abs(Y-prob)>0.5)} #Define how Error Rate is calculated
ModelCandidates <-data.frame(TestingErrorCV=cv.glm(data=d1,glmfit=DrugModel5,
  cost=ErrorRate)$delta[1],AIC=DrugModel5$aic,BIC=BIC(DrugModel5),DevianceRes=DrugModel5$deviance,
  row.names="AgeUnder33+sta.3way")
ModelCandidates <- rbind (ModelCandidates,
  "AgeUnder33+Widowed"=c(TestingErrorCV=cv.glm(data=d1,glmfit=DrugModel6,
  cost=ErrorRate) $delta[1], AIC=DrugModel6$aic, BIC=BIC(DrugModel6), DevianceRes=DrugModel6$deviance))
ModelCandidates
                        TestingErrorCV
##
                                             AIC
                                                      BIC DevianceRes
## AgeUnder33+sta.3way
                             0.1002662 721.1245 741.2338
                                                              713.1245
## AgeUnder33+Widowed
                             0.1002662 721.6596 736.7416
                                                              715.6596
P-value for removing the indicator is .11. Though not statistically significant, I find this unsettlingly low.
AIC also deteriorates. BIC improves, but deviance jumps by 2.5 points. Testing error rates are identical.
This evidence is very inconclusive. Before deciding, let's check what happens if we remove the widowhood
indicator instead, and keep the dummy for having a partner.
d1$HasPartner<-as.numeric(d1$sta.3way=="HasPartner")</pre>
summary(DrugModel6b<-glm(riki~AgeUnder33+HasPartner,family=binomial(link=logit),data=d1))</pre>
##
  glm(formula = riki ~ AgeUnder33 + HasPartner, family = binomial(link = logit),
##
       data = d1)
##
## Deviance Residuals:
##
                      Median
                                    3Q
       Min
                 1Q
                                             Max
## -0.5178 -0.5178 -0.5178 -0.3198
                                          2.4489
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
               -2.9473
                             0.2783 -10.589 < 2e-16 ***
                                      3.402 0.000669 ***
## AgeUnder33
                 1.0055
                             0.2955
## HasPartner
                -0.6603
                             0.4351 -1.518 0.129123
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 734.05
                              on 1126
                                        degrees of freedom
## Residual deviance: 714.90
                              on 1124
                                        degrees of freedom
  AIC: 720.9
## Number of Fisher Scoring iterations: 5
anova(DrugModel5,DrugModel6b,test="LRT")
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + sta.3way
## Model 2: riki ~ AgeUnder33 + HasPartner
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          1123
                   713.12
## 2
          1124
                   714.90 -1 -1.7769
                                         0.1825
ModelCandidates <- rbind (ModelCandidates,
  "AgeUnder33+HasPartner"=c(TestingErrorCV=cv.glm(data=d1,glmfit=DrugModel6b,
  cost=ErrorRate) $delta[1], AIC=DrugModel6b$aic, BIC=BIC(DrugModel6b), DevianceRes=DrugModel6b$deviance))
```

Paradoxically, removing the widowhood indicator has less of an effect on deviance than removing the indicator for having a romantic partner, even though the former variable was estimated to be a more significant predictor. How can this be? The answer lies in different vastly different n's.

```
##
## Single/Divorced HasPartner Widowed
## 993 124 10
```

Widowhood is the better predictor in the few cases where it applies, but it is applicable to less than 1% of the subjects in the dataset. HasPartner is applicable to over 10% of the data. Thus, removing HasPartner worsens the fit considerably more than does removing Widowed, even though the former's predictive power on a single observation is slightly lower. Put another way, the widowhood dummy looks more impressive in the model output but is overall less useful than HasPartner. CV Testing Error remains identical between models 5 and 6b. AIC and BIC, as well as LRT, seem to support the removal of the widowhood dummy, so out it goes. Two predictors remain. Can we remove one?

```
DrugModel7<-glm(riki~AgeUnder33,family=binomial(link=logit),data=d1)
anova(DrugModel6b,DrugModel7,test="LRT")</pre>
```

```
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + HasPartner
## Model 2: riki ~ AgeUnder33
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
##
## 1
          1124
                   714.90
## 2
          1125
                   717.62 -1 -2.7228 0.09892 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(ModelCandidates <- rbind (ModelCandidates,
  "AgeUnder33"=c(TestingErrorCV=cv.glm(data=d1,glmfit=DrugModel7,
  cost=ErrorRate)$delta[1],AIC=DrugModel7$aic,BIC=BIC(DrugModel7),DevianceRes=DrugModel7$deviance)))
```

```
##
                         TestingErrorCV
                                              AIC
                                                       BIC DevianceRes
## AgeUnder33+sta.3way
                                                               713.1245
                               0.1002662 721.1245 741.2338
                               0.1002662 721.6596 736.7416
## AgeUnder33+Widowed
                                                               715.6596
## AgeUnder33+HasPartner
                               0.1002662 720.9015 735.9834
                                                               714.9015
## AgeUnder33
                               0.1002662 721.6243 731.6789
                                                               717.6243
```

No, I don't think this removal is a good idea. The p-value for dropping HasPartner is below .10, and the deviance increase is unsavory. We have a pretty good model now. It is parsimonious with only two predictors, both of which make intuitive sense are simple to interpret. However, we perhaps do still a little better in terms of predictive power. Let's look more closely at the effect of relationship status:

```
table(d1$sta.3way,d1$riki)
```

```
## ## 0 1
## Single/Divorced 888 105
## HasPartner 118 6
## Widowed 8 2
```

Widowed people seem at an elevated risk of inclusion in riki. We knew this. Now let's condition the effect on sex:

```
cat("men\n"); with(d1[d1$male==1,],table(sta.3way,riki)); cat("\n\n")
```

```
## men
##
                     riki
##
  sta.3way
                         0
                             1
##
     Single/Divorced 625
                            80
##
     HasPartner
                       58
                             3
##
     Widowed
                         2
cat("women\n"); with(d1[d1$male==0,],table(sta.3way,riki))
```

```
## riki

## sta.3way 0 1

## Single/Divorced 263 25

## HasPartner 60 3

## Widowed 6 2
```

The odds ratio of being in riki between widowed men and widowed women is infinite. For the bigger picture, let's view the same conditional crosstables from the riki dataset from the previous exercise:

```
cat("men\n"); with(d0[d0$male==1,],table(sta,hilmo)); cat("\n\n")
```

```
## men
##
                hilmo
## sta
                   0
                        1
                        3
##
     HasPartner
                  87
##
     Single
                 968
                       71
                        0
##
     Widowed
                   3
     Divorced
                 148
cat("women\n"); with(d0[d0$male==0,],table(sta,hilmo))
```

```
## women
```

women

hilmo

```
## sta
                    0
##
                         3
     HasPartner
                   35
##
     Single
                  179
                        22
##
     Widowed
                    0
                         2
     Divorced
                   46
```

Again the odds ratio between women and men is infinite. Based on this (very limited) data, widowhood seems much more dangerous to women than to men. The implication for our model selection exercise is that we should consider making a dummy variable for being a widow in the word's exact sense, i.e., a **woman** who has lost her partner to death and remains alone.

```
d1$fem.widow<-as.numeric(d1$male==0 & d1$Widowed)
summary(DrugModel8<-glm(riki~AgeUnder33+HasPartner+fem.widow,family=binomial,data=d1))</pre>
##
## Call:
  glm(formula = riki ~ AgeUnder33 + HasPartner + fem.widow, family = binomial,
##
       data = d1)
##
##
  Deviance Residuals:
##
       Min
                      Median
                                    30
                                            Max
                 10
   -0.6592 -0.5159 -0.5159
                              -0.3088
                                         2.4767
##
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
               -3.0192
                            0.2865 -10.537 < 2e-16 ***
                                      3.539 0.000402 ***
## AgeUnder33
                 1.0695
                             0.3022
                -0.6353
## HasPartner
                            0.4355
                                   -1.459 0.144655
## fem.widow
                 1.6031
                            0.8591
                                      1.866 0.062027 .
##
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 734.05
                                        degrees of freedom
                              on 1126
## Residual deviance: 712.20
                              on 1123
                                        degrees of freedom
## AIC: 720.2
## Number of Fisher Scoring iterations: 5
(ModelCandidates <- rbind (ModelCandidates,
  "AgeUnder33+HasPartner+fem.widow"=c(TestingErrorCV=cv.glm(data=d1,glmfit=DrugModel8,
  cost=ErrorRate)$delta[1],AIC=DrugModel8$aic,BIC=BIC(DrugModel8),DevianceRes=DrugModel8$deviance)))
                                    TestingErrorCV
                                                        AIC
## AgeUnder33+sta.3way
                                         0.1002662 721.1245 741.2338
## AgeUnder33+Widowed
                                         0.1002662 721.6596 736.7416
## AgeUnder33+HasPartner
                                         0.1002662 720.9015 735.9834
## AgeUnder33
                                         0.1002662 721.6243 731.6789
## AgeUnder33+HasPartner+fem.widow
                                         0.1002662 720.2001 740.3093
##
                                    DevianceRes
## AgeUnder33+sta.3way
                                       713.1245
## AgeUnder33+Widowed
                                       715.6596
## AgeUnder33+HasPartner
                                       714.9015
                                       717.6243
## AgeUnder33
```

712.2001

AgeUnder33+HasPartner+fem.widow

anova(DrugModel6b,DrugModel8,test="LRT")

```
## Analysis of Deviance Table
##
## Model 1: riki ~ AgeUnder33 + HasPartner
## Model 2: riki ~ AgeUnder33 + HasPartner + fem.widow
## Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1 1124 714.9
## 2 1123 712.2 1 2.7014 0.1003
```

The improvement in fit that is achieved by adding the dummy for being a widow has a p-value of .10, so there's no clear-cut answer to whether its inclusion is justified. It is risky to include covariates whose predictive power is based on such tiny samples – the reported effect could very well be due to mere chance. To illustrate, let's compare these three predictors by the extent to which they are influenced by single observations:

```
leverage <- dfbetas(DrugModel8); leverage <- as.data.frame(leverage)
(maxlev<-sapply(abs(leverage), max))</pre>
```

```
## (Intercept) AgeUnder33 HasPartner fem.widow
## 0.1892240 0.1999408 0.3110654 0.8549638
```

The dfbeta estimates how much the MLE of the covariate would change if the observation was deleted. We can see that the maximum leverage of an individual observation on the widow dummy is about 3 to 4 times as high as on the other two covariates. We see this difference leverage difference at work when we delete a single observation exerting maximum influence for each predictor and compare the changes in the MLEs:

```
leverage[which(abs(leverage$fem.widow)==maxlev["fem.widow"]),]
```

```
##
       (Intercept) AgeUnder33 HasPartner fem.widow
      -0.1863602 0.1999408 0.02011602 0.8549638
leverage[which(abs(leverage$HasPartner)==maxlev["HasPartner"]),]
       (Intercept) AgeUnder33 HasPartner fem.widow
## 128 -0.05525641 0.05928311 0.3110654 0.01134464
summary(glm(riki~AgeUnder33+HasPartner+fem.widow,family=binomial,data=d1[c(-128,-393),]))
##
## Call:
  glm(formula = riki ~ AgeUnder33 + HasPartner + fem.widow, family = binomial,
##
       data = d1[c(-128, -393), ])
##
##
  Deviance Residuals:
                      Median
##
      Min
                 1Q
                                   30
                                           Max
##
  -0.5140
           -0.5140 -0.5140 -0.3175
                                        2.4547
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
               -2.9624
## (Intercept)
                            0.2835 -10.450 < 2e-16 ***
## AgeUnder33
                 1.0049
                            0.3006
                                     3.343 0.000827 ***
## HasPartner
                -0.8252
                            0.4721
                                    -1.748 0.080475
                 0.9803
                                     0.880 0.379114
## fem.widow
                            1.1145
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
```

```
## Null deviance: 724.82 on 1124 degrees of freedom
## Residual deviance: 704.59 on 1121 degrees of freedom
## AIC: 712.59
##
## Number of Fisher Scoring iterations: 5
```

The MLE for AgeUnder33 stays virtually unchanged. HasPartner becomes more significant as its p-value decreases from .14 to .8 – this is because the most influential observation is one that deviates from the general trend, so getting rid of it makes the trend more clear. The effect of the widow dummy, on the other hand, is based on a mere 2 success outcomes, thus it vanishes completely when we remove one of them.

At the same time, the effect of being a widowed female has the same direction in both datasets – a strong positive association. Our choice is whether to ignore this effect due to the small sample or to assume that the association seen in this meager sample is an indicator of a wider phenomenon of prematurely widowed women having an elevated risk of antisocial and self-destructive behavior.

I choose to "believe my data" here. Losing your husband/boyfriend to death at a young age is undoubtedly a traumatic experience, so it stands to reason that women having experienced it are more susceptible to problem behavior. There seems no particular reason to assume that the effect if due to chance alone. Therefore, the final model chosen to predict riki membership is DrugModel8, containing AgeUnder33 and the dummies HasPartner and fem.widow. We'll now create these dummy variables for the combined drug dataset, and then use the model to predict the probability of inclusion in riki for every subject.

```
drug$HasPartner<-as.numeric(drug$sta=="HasPartner")
drug$fem.widow<-as.numeric(drug$male==0 & drug$sta=="Widowed")
drug$FittedRikiProbs<-predict(DrugModel8, newdata=drug, type="response")</pre>
```

6. The Horvitz-Thompson Estimator of Population Total

If we take a random sample from a population of unknown size, mark each subject in the sample, release them back into the population, and then collect a new random sample of the same population, the degree of overlap between the two random samples is going to be approximately inversely proportional to the total population size. This is the capture-recapture method, which is used to estimate population sizes in fields such as biology. In our example, we are estimating the total population of drug users in Uusimaa, and our two random samples are the hilmo and riki registries, both of which keep independent track of drug users in the area. As per the instructions, we'll rename the hilmo and riki inclusion probabilities calculated for each subject in the druq dataset as p1 and p2, respectively:

```
names(drug) [which(names(drug)=="FittedHilmoProbs")] <- "p1"
names(drug) [which(names(drug)=="FittedRikiProbs")] <- "p2"
#drug <- drug[, c(1,2,4,7,6,9,10,8,11)]</pre>
```

For each of the 2593 subjects, the probability of being registered in either hilmo or riki is:

```
drug$theta <- with(drug, p1+p2 - p2*p2)</pre>
```

Here's a summary of theta:

```
summary(drug$theta)
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.05600 0.09341 0.18430 0.16210 0.18430 0.79170
```

Our two models rely on a combined total of 5 explanatory variables, all of which are dichotomous. In the drug dataset, 12 different combinations of these variables occur. The following table lists the estimated p1, p2, and theta for each combination. I'll leave the fem.widow dummy out of this summary table to save space, because it is already implicitly included as the 0.1 combination of male and widowed:

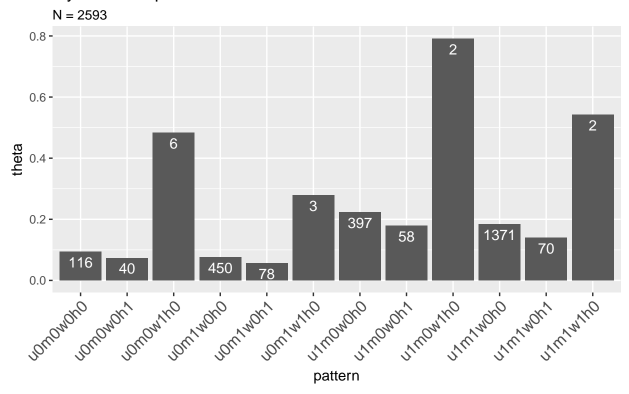
```
EVPs.Bernoulli<-split(drug, list(drug$AgeUnder33,drug$Widowed,drug$male,drug$HasPartner),drop=TRUE)
EVPs <- data.frame(); n<-vector()
for(i in 1:length(EVPs.Bernoulli)){
    EVPs <- rbind(EVPs,EVPs.Bernoulli[[i]][1,]); n<-c(n,nrow(EVPs.Bernoulli[[i]]]))}
EVPs$n<-n; row.names(EVPs)<-1:nrow(EVPs)
EVPs<-EVPs[,c(7,4,6,9,8,11:13)]
round(EVPs,digits=6)</pre>
```

```
##
      AgeUnder33 male Widowed HasPartner
                                                         p2
                                                                theta
                                                р1
                                                                         n
## 1
               0
                    0
                            0
                                        0 0.049011 0.046564 0.093407
                                                                       116
                                                                       397
## 2
               1
                    0
                            0
                                        0 0.114439 0.124581 0.223500
## 3
               0
                    0
                            1
                                        0 0.326838 0.195268 0.483977
                                        0 0.549033 0.414195 0.791670
## 4
                    0
                                                                         2
               1
                            1
## 5
               0
                    1
                            0
                                        0 0.031417 0.046564 0.075813 450
                            0
                                        0 0.075215 0.124581 0.184276 1371
## 6
               1
                    1
## 7
               0
                    1
                            1
                                        0 0.234056 0.046564 0.278452
## 8
               1
                    1
                            1
                                        0 0.433824 0.124581 0.542884
                                                                         2
## 9
               0
                    0
                            0
                                        1 0.049011 0.025221 0.073595
                                                                        40
## 10
                    0
                            0
                                        1 0.114439 0.070106 0.179630
                                                                        58
               1
## 11
                            0
                                        1 0.031417 0.025221 0.056001
               0
                    1
                                                                        78
## 12
                                        1 0.075215 0.070106 0.140406
               1
                    1
                            0
                                                                        70
```

Let u1 = age under 33, m1 = male, w1 = widowed, and h1 = has a partner. The following plot illustrates the variation in *theta* across covariate patterns:

```
EVPs<-cbind(pattern=c("u0m0w0h0","u1m0w0h0","u0m0w1h0","u1m0w1h0","u0m1w0h0",
    "u1m1w0h0","u0m1w1h0","u1m1w1h0","u0m0w0h1","u1m0w0h1","u0m1w0h1","u1m1w0h1"),EVPs)
ggplot(EVPs, aes(x=pattern)) + geom_col(aes(y=theta)) +
    theme(axis.text.x = element_text(angle = 45, hjust = 1, size = 12)) +
    geom_text(aes(y=theta,label=n), vjust=1.5, colour="white") +
    ggtitle("Probability of inclusion in at least one drug user database,\n by covariate pattern",
    subtitle="N = 2593")</pre>
```

Probability of inclusion in at least one drug user database, by covariate pattern



The right side of the plot represents those under 33. The left side are the older people. In both age groups, it is the widowed women followed by the widowed men who have the highest estimated probability of inclusion. We can also see how low n is in each of these four groups.

Finally, we'll calculate the Horvitz-Thompson estimate of the population total of drug users in Uusimaa around 2000. It is the sum over the inverse inclusion probabilities (thetas) of each subject in the full dataset:

(Horvitz.Thompson <- sum(1/drug\$theta))</pre>

[1] 19180.92

Based on the probability estimates yielded by our models, there were approximately 19,000 drug users in Uusimaa at the turn of the century.