# Binary Logistic DrPH (Epidemiology)

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# Contents

Modeling Binomial Data	<b>2</b>
Locate files	2
Set the folder	2
Read data	2
Describe data	2
Explore data	3
Estimate parameters	3
Make inference	4
Make prediction	5
Compare models	5
Linearity in logits	7
Diagnostics for a model with a binomial response	9
Diagnostic plot	12
Interaction	12
Resources	14
Additional materials	14

# Modeling Binomial Data

- 1. Describe data
- 2. Explore data Exploratory Data Analysis
- 3. Estimate parameters
- 4. Make Inference
- 5. Make Prediction
- 6. Interpretation

### Locate files

- Browse your folders.
- Look for the files.
- Check the path to the folder containing the files

### Set the folder

Set our working directory. REMEMBER! your working directory (working folder) is different from my working directory

```
# this is my working directory. You have to specify yours
setwd("E:/Epi_Stat_Matters/LectureNotes2015/binary-logistic/binary-logistic-DrPH-2015/BinaryLogisticDrP.
getwd()
```

## [1] "E:/Epi\_Stat\_Matters/LectureNotes2015/binary-logistic/binary-logistic-DrPH-2015/BinaryLogisticDr

### Read data

- Read our data in the working folder
- Then, save as a csv file in our working directory

```
mydata <- read.csv("http://www.ats.ucla.edu/stat/data/binary.csv")
write.csv2(mydata,'logistic.csv')</pre>
```

### Describe data

```
#observe data
head(mydata,10)
```

Rank is taken as numerical variable which does not make sense. We need to convert it to a categorical (factor) variable

```
summary(mydata)
```

```
##
        admit.
                          gre
                                                          rank
          :0.0000
## Min.
                     Min.
                           :220.0
                                          :2.260
                                                            :1.000
                                     1st Qu.:3.130
## 1st Qu.:0.0000
                     1st Qu.:520.0
                                                     1st Qu.:2.000
## Median :0.0000
                     Median :580.0
                                     Median :3.395
                                                     Median :2.000
```

```
Mean
           :0.3175
                             :587.7
                                              :3.390
                                                               :2.485
                      Mean
                                       Mean
                                                       Mean
   3rd Qu.:1.0000
                      3rd Qu.:660.0
##
                                       3rd Qu.:3.670
                                                       3rd Qu.:3.000
                                              :4.000
## Max.
           :1.0000
                             :800.0
                                       Max.
                                                       Max.
                                                               :4.000
mydata$rank<-factor(mydata$rank)
summary(mydata$rank)
##
     1
         2
             3
   61 151 121 67
```

### Explore data

Use plots like \* Histogram for numerical variables \* and barplot for categorical variables, at least.

### Estimate parameters

coefficients(mylogit)

- we estimate the logit or the log odds.
- We used summary to see the results stored as mylogit
- We used **coefficients** to examine the regression coefficients

```
mylogit <- glm(admit~gre+gpa+rank,family = 'binomial'(link = logit),data=mydata)</pre>
summary(mylogit)
##
## Call:
  glm(formula = admit ~ gre + gpa + rank, family = binomial(link = logit),
##
       data = mydata)
##
## Deviance Residuals:
                 1Q
                      Median
                                   3Q
                                           Max
## -1.6268 -0.8662 -0.6388
                                        2.0790
                               1.1490
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -3.989979
                           1.139951 -3.500 0.000465 ***
                0.002264
                           0.001094
                                      2.070 0.038465 *
## gre
                0.804038
                           0.331819
                                      2.423 0.015388 *
## gpa
## rank2
               -0.675443
                           0.316490 -2.134 0.032829 *
## rank3
               -1.340204
                           0.345306
                                    -3.881 0.000104 ***
## rank4
               -1.551464
                           0.417832 -3.713 0.000205 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 499.98 on 399
                                      degrees of freedom
## Residual deviance: 458.52 on 394 degrees of freedom
## AIC: 470.52
## Number of Fisher Scoring iterations: 4
```

```
## (Intercept) gre gpa rank2 rank3
## -3.989979073 0.002264426 0.804037549 -0.675442928 -1.340203916
## rank4
## -1.551463677
```

To obtain the odds ratios and their 95% CI, we need to exponentiate using **exp** the regression coefficients or the betas

```
exp(coefficients(mylogit))
                                                                        rank4
## (Intercept)
                        gre
                                    gpa
                                               rank2
                                                           rank3
     0.0185001
                 1.0022670
                              2.2345448
                                           0.5089310
                                                       0.2617923
                                                                    0.2119375
exp(confint(mylogit))
## Waiting for profiling to be done...
                      2.5 %
                               97.5 %
##
## (Intercept) 0.001889165 0.1665354
## gre
               1.000137602 1.0044457
               1.173858216 4.3238349
## gpa
               0.272289674 0.9448343
## rank2
## rank3
               0.131641717 0.5115181
## rank4
               0.090715546 0.4706961
```

### Make inference

Here, we examine the p-values (hypothesis testing) and the confidence intervals.

- First, using the method of maximum likelihood
- Next, using the SE method (function **confint.default**)

0.0001202298 0.004408622

0.1536836760 1.454391423

-1.2957512650 -0.055134591

-2.0169920597 -0.663415773

-2.3703986294 -0.732528724

```
confint(mylogit)
```

## gre ## gpa

## rank2

## rank3

## rank4

```
## Waiting for profiling to be done...
                       2.5 %
## (Intercept) -6.2716202334 -1.792547080
## gre
                0.0001375921 0.004435874
                0.1602959439 1.464142727
## gpa
               -1.3008888002 -0.056745722
## rank2
               -2.0276713127 -0.670372346
## rank3
               -2.4000265384 -0.753542605
## rank4
confint.default(mylogit)
                       2.5 %
                                   97.5 %
## (Intercept) -6.2242418514 -1.755716295
```

# Make prediction

We now can:

- 1. Predict the log odds for having the outcome
- 2. Predict the conditional probability for having the outcome

```
pred.logit<-predict(mylogit,type='link')</pre>
head(pred.logit)
##
                           1.0377118 -1.5273305 -2.0081113 -0.5323458
## -1.5671256 -0.8848442
-3.99+0.00226*380+0.8041*3.61-1.34
## [1] -1.568399
Notice, that similarity between predict(x, type='response' and fitted Remember, we can calculate the
conditional probability of having the outcome
pred.prob<-predict(mylogit,type='response')</pre>
head(pred.prob)
## 0.1726265 0.2921750 0.7384082 0.1783846 0.1183539 0.3699699
head(fitted(mylogit))
##
## 0.1726265 0.2921750 0.7384082 0.1783846 0.1183539 0.3699699
\exp(-1.567)/(1+\exp(-1.567))
## [1] 0.1726445
```

# Compare models

We compare a model with vs and without gre. This is done using the deviance

```
summary(mylogit)
```

```
##
## Call:
   glm(formula = admit ~ gre + gpa + rank, family = binomial(link = logit),
       data = mydata)
##
##
## Deviance Residuals:
##
       Min
                 1Q
                                    3Q
                      Median
                                            Max
## -1.6268 -0.8662 -0.6388
                               1.1490
                                         2.0790
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -3.989979
                           1.139951 -3.500 0.000465 ***
## gre
                0.002264
                           0.001094
                                     2.070 0.038465 *
                                      2.423 0.015388 *
## gpa
                0.804038
                           0.331819
## rank2
               -0.675443
                           0.316490 -2.134 0.032829 *
## rank3
               -1.340204
                           0.345306 -3.881 0.000104 ***
## rank4
               -1.551464
                           0.417832 -3.713 0.000205 ***
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 499.98 on 399 degrees of freedom
##
## Residual deviance: 458.52 on 394 degrees of freedom
## AIC: 470.52
##
## Number of Fisher Scoring iterations: 4
mylogit2 <- glm(admit~gpa+rank,family = 'binomial'(link = logit),data=mydata)</pre>
summary(mylogit2)
##
## Call:
  glm(formula = admit ~ gpa + rank, family = binomial(link = logit),
##
      data = mydata)
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                  3Q
                                          Max
## -1.5055 -0.8663 -0.6590
                             1.1505
                                       2.0913
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
                           1.1003 -3.148 0.001645 **
## (Intercept) -3.4636
## gpa
                1.0521
                           0.3102
                                   3.392 0.000694 ***
## rank2
               -0.6810
                           0.3141 -2.168 0.030181 *
## rank3
               -1.3919
                           0.3419 -4.071 4.68e-05 ***
## rank4
               -1.5943
                           0.4152 -3.840 0.000123 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 499.98 on 399 degrees of freedom
## Residual deviance: 462.88 on 395 degrees of freedom
## AIC: 472.88
## Number of Fisher Scoring iterations: 4
anova(mylogit,mylogit2,test = 'Chisq')
## Analysis of Deviance Table
##
## Model 1: admit ~ gre + gpa + rank
## Model 2: admit ~ gpa + rank
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          394
                  458.52
## 2
                  462.88 -1 -4.3578 0.03684 *
          395
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

# Linearity in logits

gre is tested for linearity in logit. gre is linear but it is rescaled to produce less decimals

```
The linearity of logits is tested using library mfp
library(mfp)
## Loading required package: survival
mylogit3 <- mfp(admit~fp(gre)+gpa+rank,family = 'binomial'(link = logit),data=mydata,verbose=T)</pre>
##
                           Power(s)
   Variable
               Deviance
## -----
## Cycle 1
   rank2
##
               463.096
               458.517
##
##
##
##
##
   rank3
               474.043
##
##
               458.517
                           1
##
##
##
##
   rank4
               473.551
##
##
               458.517
                           1
##
##
##
##
   gpa
##
               464.532
##
               458.517
                           1
##
##
##
##
   gre
##
               462.875
##
               458.517
                           1
##
               458.415
               458.366
                           -2 -2
##
##
##
## Tansformation
##
      shift scale
## rank2
           0
## rank3
            0
## rank4
            0
                  1
## gpa
            0
            0 1000
## gre
## Fractional polynomials
```

```
## df.initial select alpha df.final power1 power2
## rank2
           1
                       1 0.05
                                      1
                                             1
## rank3
                1
                       1 0.05
## rank4
                        1 0.05
                1
                                      1
                                             1
## gpa
                 1
                        1 0.05
                                      1
## gre
                 4
                        1 0.05
                                      1
                                             1
##
##
## Transformations of covariates:
##
               formula
## gre I((gre/1000)^1)
## gpa
                   gpa
## rank
                  rank
##
##
## Deviance table:
            Resid. Dev
## Null model
                499.9765
## Linear model 458.5175
## Final model
                458.5175
summary(mylogit3)
##
## glm(formula = admit ~ rank + gpa + I((gre/1000)^1), family = binomial(link = logit),
##
      data = mydata)
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                  3Q
## -1.6268 -0.8662 -0.6388 1.1490
                                       2.0790
## Coefficients:
                  Estimate Std. Error z value Pr(>|z|)
                              1.1400 -3.500 0.000465 ***
## (Intercept)
                   -3.9900
## rank2
                   -0.6754
                               0.3165 -2.134 0.032829 *
                               0.3453 -3.881 0.000104 ***
## rank3
                   -1.3402
## rank4
                   -1.5515
                               0.4178 -3.713 0.000205 ***
## gpa
                   0.8040
                               0.3318 2.423 0.015388 *
## I((gre/1000)^1) 2.2644
                               1.0940
                                      2.070 0.038465 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 499.98 on 399 degrees of freedom
## Residual deviance: 458.52 on 394 degrees of freedom
## AIC: 470.52
## Number of Fisher Scoring iterations: 4
mylogit3$fptable
        df.initial select alpha df.final power1 power2
## rank2
                 1
                       1 0.05
                                      1
```

## Diagnostics for a model with a binomial response

To do these diagnostics, you need to load library('LogisticDx').

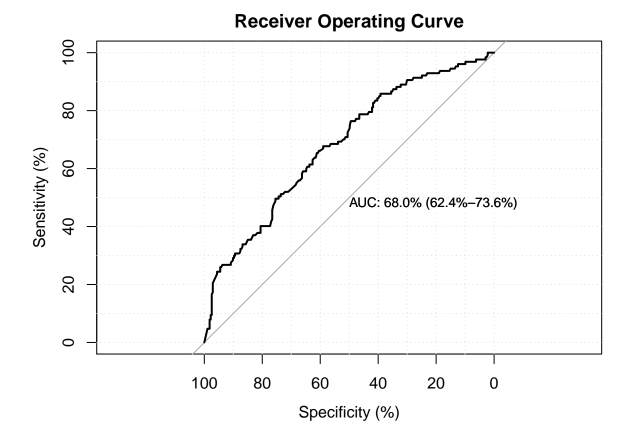
First, we produce the diagnostic measures for a binary regression model by covariate pattern

Next, we produce the Goodness-of-fit for binomial regression. Usually, the number of groups (quantiles) equal 10 to perform the Hosmer-Lemeshow test. At the same time, we plot the ROC curve

Similarly, we can check the auc value

```
library('LogisticDx')
dx(mylogit2,byCov=T)
```

```
gpa rank2 rank3 rank4 y
##
                                                      P n
        (Intercept)
                                                               yhat
##
     1:
                  1 3.94
                             1
                                   0
                                          0 1 0.5001242 2 1.0002483
##
     2:
                  1 3.30
                                          0 1 0.3378663 3 1.0135990
                             1
                                    0
##
     3:
                  1 3.99
                             0
                                          0 1 0.3412259 3 1.0236777
                                   1
##
     4:
                  1 3.35
                             1
                                          0 1 0.3497327 3 1.0491980
##
     5:
                                          0 1 0.3079792 3 0.9239375
                  1 3.17
                             1
                                   0
##
## 256:
                  1 3.52
                             0
                                   0
                                          1 2 0.2051122 3 0.6153366
  257:
                  1 2.68
                             0
                                   1
                                          0 1 0.1154721 1 0.1154721
                  1 3.00
                             0
## 258:
                                   0
                                          1 1 0.1299149 1 0.1299149
## 259:
                  1 2.42
                             0
                                   0
                                          0 1 0.2854368 1 0.2854368
                             0
## 260:
                  1 2.65
                                    1
                                          0 1 0.1122874 1 0.1122874
##
                   Pr
                                 dr
                                               h
                                                           sPr
##
     1: -0.0003511985 -0.0003511985 0.014377039 -0.0003537507 -0.0003537506
##
     2: -0.0165997514 -0.0166181074 0.006902268 -0.0166573376 -0.0166757573
     3: -0.0288329487 -0.0288874508 0.015618469 -0.0290607843 -0.0291157171
##
##
     4: -0.0595625368 -0.0597856465 0.006844794 -0.0597674360 -0.0599913131
##
         0.0954814103
                                                                0.0947908121
##
  256:
         1.9798618855
                       1.7253451406 0.017098521
                                                  1.9970084425
                                                                1.7402874601
  257:
         2.7676886108
                       2.0778480687 0.011720502
                                                  2.7840519420
                                                                2.0901328741
##
##
   258:
         2.5879228606
                       2.0203343433 0.013448875
                                                  2.6055027084
                                                                2.0340585431
##
   259:
         1.5822145401
                       1.5834991249 0.034641074
                                                  1.6103525417
                                                                1.6116599714
##
  260:
         2.8117113691
                       2.0912646219 0.011927131
                                                  2.8286306943
                                                                2.1038486967
##
              dChisq
                             dDev
                                          dBhat
##
     1: 1.251395e-07 1.251395e-07 1.825379e-09
##
     2: 2.774669e-04 2.780809e-04 1.928462e-06
##
     3: 8.445292e-04 8.477250e-04 1.339953e-05
##
     4: 3.572146e-03 3.598958e-03 2.461912e-05
     5: 9.116700e-03 8.985298e-03 6.865333e-05
##
##
## 256: 3.988043e+00 3.028600e+00 6.937586e-02
  257: 7.750945e+00 4.368655e+00 9.192235e-02
  258: 6.788644e+00 4.137394e+00 9.254425e-02
  259: 2.593235e+00 2.597448e+00 9.305602e-02
## 260: 8.001152e+00 4.426179e+00 9.658274e-02
```



#### fit.mylogit2 ## chiSq df pVal ## PrI 400.78 395 0.409645 462.88 395 0.010379 \* ## drI ## PrG 255.36 255 0.481870 ## drG 294.60 255 0.044609 \* ## PrCT 255.36 255 0.481870 ## drCT 294.60 255 0.044609 \* ## ---## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.05 '.' 0.1 ' ' 1 ## val df pVal 8.190458 8 0.415090 ## HL chiSq ## mHL F 1.484491 9 0.153918 ## OsRo Z 0.027903 NA 0.977740 ## SstPgeq0.5 Z 0.217564 NA 0.827769 ## SstP10.5 Z 0.422859 NA 0.672398 ## SstBoth chiSq 0.226144 2 0.893086 ## SllPgeq0.5 chiSq 0.047739 1 0.827045 ## SllPl0.5 chiSq 0.176745 1 0.674186 ## SllBoth chiSq 0.177398 2 0.915121 #area under curve

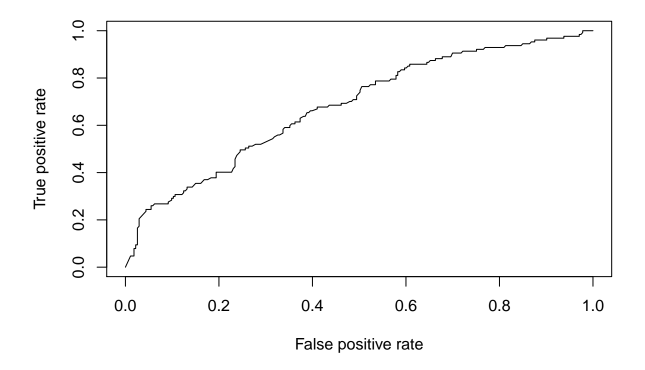
fit.mylogit2\$auc

```
auc lower 95% CI upper 95% CI
##
      68.01073
                   62.39737
                                73.62409
## attr(,"interpret")
## [1] "auc = 0.5
                       --> useless" "0.7 < auc < 0.8 --> good"
## [3] "0.8 < auc < 0.9 --> excellent"
#chi square test for gof
fit.mylogit2$chiSq
##
      test
             chiSq df
                             pVal
## 1: PrI 400.7833 395 0.40964543
## 2: drI 462.8753 395 0.01037867
## 3: PrG 255.3601 255 0.48187033
## 4: drG 294.6050 255 0.04460896
## 5: PrCT 255.3601 255 0.48187033
## 6: drCT 294.6050 255 0.04460896
#contigency table for HL test
fit.mylogit2$ctHL
          P y1
                   y1hat y0
                               yOhat n
## 1: 0.149 6 5.037833 34 34.96217 40 0.1259458
## 2: 0.192 5 6.824095 35 33.17591 40 0.1706024
## 3: 0.219 7 8.260683 33 31.73932 40 0.2065171
## 4: 0.258 11 9.327547 28 29.67245 39 0.2391679
## 5: 0.293 12 11.338710 29 29.66129 41 0.2765539
## 6: 0.34 15 12.556219 25 27.44378 40 0.3139055
## 7: 0.376 14 13.796927 25 25.20307 39 0.3537674
## 8: 0.437 14 16.561867 27 24.43813 41 0.4039480
## 9: 0.511 13 18.253216 25 19.74678 38 0.4803478
## 10: 0.678 30 25.042903 12 16.95710 42 0.5962596
#GOF test
fit.mylogit2$gof
           test stat
                             val df
                                         pVal
## 1:
             HL chiSq 8.19045835 8 0.4150903
## 2:
            mHL
                    F 1.48449073 9 0.1539182
## 3:
           OsRo
                    Z 0.02790263 NA 0.9777398
## 4: SstPgeq0.5
                    Z 0.21756392 NA 0.8277689
## 5:
      SstPl0.5
                    Z 0.42285933 NA 0.6723979
        SstBoth chiSq 0.22614408 2 0.8930863
## 6:
## 7: SllPgeq0.5 chiSq 0.04773921 1 0.8270450
       SllPl0.5 chiSq 0.17674490 1 0.6741857
## 8:
## 9:
        SllBoth chiSq 0.17739848 2 0.9151208
# another package for ROC
library(ROCR)
## Loading required package: gplots
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
##
      lowess
```

```
pred.prob2<-predict(mylogit2, type='response')
head(pred.prob2)

## 1 2 3 4 5 6
## 0.2577653 0.2700256 0.6779993 0.1542276 0.1218149 0.2712217

pred.prob22<-prediction(pred.prob2, mydata$admit )
pred.prob22f<-performance(pred.prob22, measure='tpr', x.measure='fpr')
plot(pred.prob22f)</pre>
```



```
auc2<-performance(pred.prob22, measure='auc')
auc2@y.values[[1]]</pre>
```

## [1] 0.6801073

# Diagnostic plot

Will return many diagnostic plot

```
plot(mylogit2)
```

### Interaction

Let use see how we deal an interaction. First, read data from this text file.

```
Columns (variables) no 2, and from 5 to 10 need to be converted to categorical (factor) variables
data.l<-read.table("LOWBWT.txt",header=T)</pre>
data.1[,c(2,5:10)] < -lapply(data.1[,c(2,5)],factor)
To simulate a binary predictor variable, we now recode LWT to LWD (LWT<110 vs >=110)
data.l$LWD<-findInterval(data.l$LWT,110)</pre>
data.l$LWD<-factor(data.l$LWD,labels = c("less 110",">=110"))
head(data.l$LWD,10)
                                    less 110 less 110 >=110
##
   [1] >=110
                 >=110
                           >=110
                                                                 less 110
## [8] >=110
                 >=110
                           >=110
## Levels: less 110 >=110
head(data.l$LWT,10)
## [1] 120 130 187 105 85 150 97 128 132 165
str(data.1$LWD)
## Factor w/ 2 levels "less 110",">=110": 2 2 2 1 1 2 1 2 2 2 ...
Model the relationship; outcome (LOW=0,1) with predictors of LWD and AGE interact with each other.
You may try with using #
mod.lwd.age<-glm(LOW~LWD*AGE,family = binomial(link =logit ),data=data.l)</pre>
summary(mod.lwd.age)
##
## Call:
## glm(formula = LOW ~ LWD * AGE, family = binomial(link = logit),
##
       data = data.l)
##
## Deviance Residuals:
                      Median
##
                 1Q
                                    3Q
       Min
                                            Max
## -1.4257 -0.8554 -0.6960
                                1.1602
                                         2.0329
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -1.16959
                             1.46515 -0.798
                             1.72481
                                                0.2597
## LWD>=110
                 1.94409
                                       1.127
## AGE
                 0.05262
                             0.06449
                                       0.816
                                                0.4145
## LWD>=110:AGE -0.13220
                             0.07570 -1.746
                                                0.0807 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 234.67 on 188 degrees of freedom
## Residual deviance: 221.14 on 185 degrees of freedom
## AIC: 229.14
## Number of Fisher Scoring iterations: 4
Predict our model using new data. Before doing so, we need to create a dataset containing new data
newdata.2<-data.frame(AGE=c(15,15,20,20),LWD=rep(c("less 110",">=110"),2))
```

Now let us predict the log odds

```
predict(mod.lwd.age,newdata=newdata.2)
                       2
## -0.3802252 -0.4190874 -0.1171023 -0.8169483
newdata.2
##
     AGE
              LWD
## 1 15 less 110
## 2 15
            >=110
## 3 20 less 110
## 4 20
            >=110
Can you prove these?
-1.1696+0+0.0526*15+0
## [1] -0.3806
-1.1696+1.944*1+0.0526*15-0.1322*1*15
## [1] -0.4196
-1.1696+0+0.0526*20+0
## [1] -0.1176
-1.1696+1.944*1+0.0526*20-0.1322*1*20
## [1] -0.8176
```

### Resources

- 1. http://www.ats.ucla.edu/stat/r/dae/logit.htm
- 2. https://cran.r-project.org/web/packages/HSAUR/vignettes/Ch\_logistic\_regression\_glm.pdf

### Additional materials

1. http://www.shizukalab.com/toolkits/plotting-logistic-regression-in-r

First, we'll create a fake dataset of 20 individuals of different body sizes:

```
bodysize<-rnorm(20,30,2) # generates 20 values, with mean of 30 & s.d.=2
bodysize<-sort(bodysize) # sorts these values in ascending order.
survive<-c(0,0,0,0,0,1,0,1,0,0,1,1,0,1,1,1,0,1,1,1) # assign 'survival' to these 20 individuals non-ran dat<-as.data.frame(cbind(bodysize,survive)) # saves dataframe with two columns: body size & survival dat # just shows you what your dataset looks like. It will look something like this:
```

```
##
      bodysize survive
## 1
      28.32004
                     0
## 2
     28.45303
                     0
## 3
     28.47945
                     0
     28.94434
## 4
                     0
## 5 28.96723
                     0
## 6 29.41233
                     1
## 7
      29.44386
                     0
## 8 29.52775
                     1
```

```
29.73249
## 10 29.86694
                      0
## 11 29.96795
## 12 29.99313
                      1
## 13 30.16779
## 14 30.26324
                      1
## 15 32.16110
                      1
## 16 32.75981
## 17 33.04847
                      0
## 18 33.20532
                      1
## 19 33.59674
                      1
## 20 34.59570
                      1
```

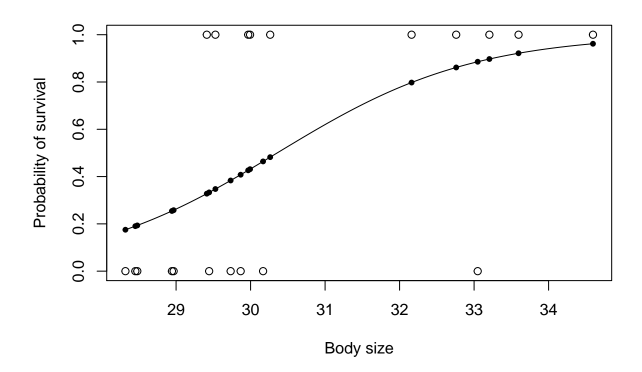
#### Plot

```
#quartz(title="bodysize vs. survival") # creates a quartz window with title

plot(bodysize,survive,xlab="Body size",ylab="Probability of survival") # plot with body size on x-axis
g=glm(survive~bodysize,family=binomial,dat) # run a logistic regression model (in this case, generalize

curve(predict(g,data.frame(bodysize=x),type="resp"),add=TRUE) # draws a curve based on prediction from

points(bodysize,fitted(g),pch=20) # optional: you could skip this draws an invisible set of points of b
```



 $2.\ http://www.cookbook-r.com/Statistical\_analysis/Logistic\_regression/$ 

```
data(mtcars)
dat2 <- subset(mtcars, select=c(mpg, am, vs))</pre>
```

#### dat2

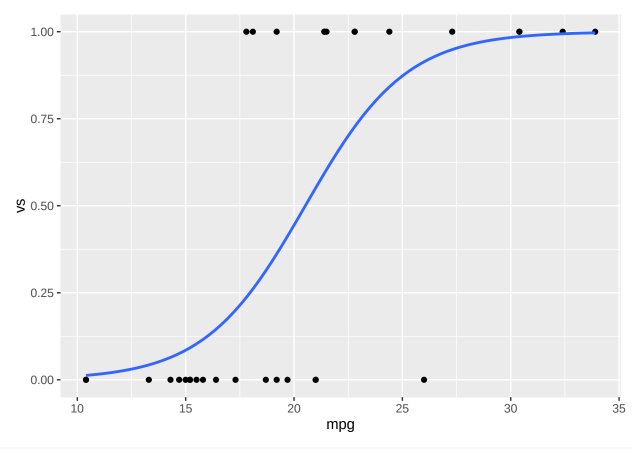
```
##
                     mpg am vs
                    21.0 1
## Mazda RX4
## Mazda RX4 Wag
                    21.0 1 0
## Datsun 710
                    22.8 1 1
## Hornet 4 Drive
                    21.4 0 1
## Hornet Sportabout 18.7 0 0
## Valiant
                    18.1 0 1
## Duster 360
                   14.3 0 0
## Merc 240D
                    24.4 0 1
## Merc 230
                    22.8 0 1
## Merc 280
                    19.2 0 1
## Merc 280C
                    17.8 0 1
## Merc 450SE
                    16.4 0 0
## Merc 450SL
                    17.3 0 0
## Merc 450SLC
                    15.2 0 0
## Cadillac Fleetwood 10.4 0 0
## Lincoln Continental 10.4 0 0
## Chrysler Imperial 14.7 0 0
## Fiat 128
                    32.4 1 1
## Honda Civic
                    30.4 1 1
                   33.9 1 1
## Toyota Corolla
## Toyota Corona
                    21.5 0 1
## Dodge Challenger 15.5 0 0
## AMC Javelin
                 15.2 0 0
## Camaro Z28
                   13.3 0 0
## Pontiac Firebird 19.2 0 0
## Fiat X1-9
                  27.3 1 1
## Porsche 914-2
                   26.0 1 0
                    30.4 1 1
## Lotus Europa
## Ford Pantera L
                    15.8 1 0
                    19.7 1 0
## Ferrari Dino
## Maserati Bora
                    15.0 1 0
                    21.4 1 1
## Volvo 142E
```

### Continuous predictor

```
# Do the logistic regression - both of these have the same effect.
# ("logit" is the default model when family is binomial.)
logr_vm <- glm(vs ~ mpg, data=dat2, family=binomial)
logr_vm <- glm(vs ~ mpg, data=dat2, family=binomial(link="logit"))</pre>
```

Plotting, first using ggplot2 then base graphics

```
library(ggplot2)
ggplot(dat2, aes(x=mpg, y=vs)) + geom_point()+
    stat_smooth(method="glm",method.args ="binomial",se=FALSE)
```



plot(dat2\$mpg, dat2\$vs)
curve(predict(logr\_vm, data.frame(mpg=x), type="response"), add=TRUE)

