Binary Response and Logistic Regression

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- 1 Model
- 2 Implementation in R
- 3 Model Evaluation
- 4 Outlook

Model •0000

Model

Binary Response Data

- General setting: $x_i \in \mathbb{R}^n$, $y_i \in \{0,1\}$
- Examples:
 - Stimulus discrimination tasks
 - Shooter paradigm
 - Just noticeable difference in color perception
 - Mortality
 - Is regular physical exercise life extending?
 - What are risk factors for severe COVID-19 symptoms?

Model Evaluation

- Performance assessment
 - Student assessment tests
 - Exam design: modeling task difficulty

Estimation

Model

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Basic idea: predict probability for class 1 as

$$P(Y=1) = \frac{\exp(\eta)}{1 + \exp(\eta)}$$
 with $\eta = \beta_0 + \beta_1 x_1 + \ldots + \beta_k x_k$

• Underlying linear model $\eta_i = \beta_0 + \sum_{j=1}^k \beta_j x_j$ that is plugged into the logistic function to obtain $P(Y = 1) \in [0, 1]$ and $P(Y = 1) + P(Y \neq 1) = 1.$

Likelihood

Likelihood $\mathcal{L}(\beta|x_1,\ldots,x_n) = \prod_{i=1}^n \mathcal{L}_i(\beta|x_i)$

$$I(\beta|x_1,...,x_n) = \sum_{i=1}^n l_i(\beta|x_i)$$

$$= \sum (1 - y_i) \log(1 - p(x_i;\beta)) + y_i \log p(x_i;\beta)$$

$$= \sum y_i \log \frac{p(x_i;\beta)}{1 - p(x_i;\beta)} + \log(1 - p(x_i;\beta))$$

$$= \sum y_i x_i \beta - \log(1 + \exp(x_i\beta))$$

$$= \sum_{i=1}^n y_i \eta_i - \log(1 + \exp(\eta))$$

■ Maximize log-likelihood $I = \sum_{i=1}^{n} y_i \eta_i - \log(1 + \exp(\eta)) =: LL$

Inference

Model

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Data format

- The criterion must be coded as 0/1 or as factor.
- Predictors must be metric or dummy-coded

Interpretation

- Odds ratio for predictor x_j is equal to $\exp(\beta_j)$
- Odds ratio OR_j quantifies how the odds $\frac{P(Y=1)}{P(Y=0)}$ change when x_j increases by 1 unit.

Implementation in R

Syntax

Model

General syntax

- Command glm() (generalized linear model)
- Formula syntax:
 - y~x1+x2+x3 (no interactions)
 - y~x1*x2 (interactions)
 - y~x1+x2+x3+x1:x2 (selected interactions)
- Must provide a link function to glm() through the family parameter (cf. section Other link functions)
 - For logistic regression, we use family=binomial('logit').

Examples

```
m1 = glm(correct_response - iq + math_skill, data=df, family=binomial('logit'))
m2 = glm(fatal_accident - bmi + risk_seeking + gender, family=binomial('logit'))
m3 = glm(vaccination_skeptic - iq * income, family=binomial('logit'))
```

Toy Example

Model

The dataset df^1 contains the yearly sick leave hours, number of tweets on Twitter, IQ, and hair length of N=100 employees along with their gender (binary: male/female) and whether they have ever suffered from depression (binary: yes/no):

```
df %>% slice(sample(nrow(df))) %>% head(5)
     ID depr gender avg_sickhours n_tweets iq hairlength
                                          3 90
## 1 75
          nο
               male
                              13.0
                                                      19.5
## 2 91
               male
                              12.7
                                          3 84
                                                       4.8
## 3 23 ves female
                              11.5
                                         12 90
                                                      46.3
## 4 33
          no female
                              5.4
                                          3 97
                                                      40.4
               male
                              8.6
                                                       4.4
## 5 86
                                          1 83
table(df$gender)
```

```
##
## female male
## 50 50
```

¹www.github.com/marvinschmitt/talk-binary-response

We define gender as criterion and avg_sickhours as predictor. The logit link function leads to a logistic regression. The output's coefficients correspond to β_0, \ldots, β_k .

```
m = glm(gender ~ avg_sickhours, data = df, family = binomial('logit'))
m$coefficients
### (Intercept) avg_sickhours
```

(Intercept) avg_sickhours ## -4.2693393 0.4440937

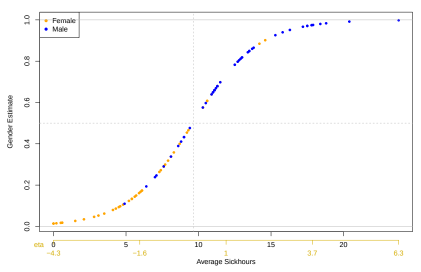
Model

- We can calculate η from the underlying linear model: $\eta = -4.27 + 0.44x_1$
- Thus, the criterion estimate is:

$$\hat{P}(Y = \mathsf{male}) = \frac{\exp(-4.27 + 0.44x_1)}{1 + \exp(-4.27 + 0.44x_1)}$$

■ The odds for gender=m are increased by the factor exp(0.44) = 1.55 per additional sick leave hour.

Gender Estimate by Average Sickhours

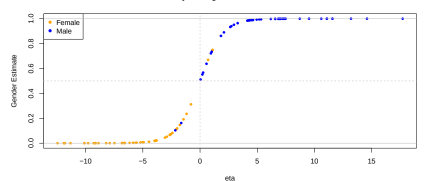


Predictors: Avg. sick hours (x_1) , Number of tweets (x_2)

```
m = glm(gender ~ avg_sickhours + n_tweets, data = df, family = binomial('logit'))
m$coefficients
```

$$\eta = -10.11 + 1.62x_1 + -1.2x_2$$

Gender Estimate by Average Sickhours and number of tweets



```
Predictors: Avg. sick hours (x_1), Number of tweets (x_2), IQ (x_3)
```

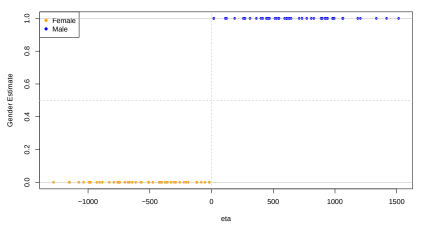
Model Evaluation

```
m = glm(gender ~ avg sickhours + n tweets + ig, data = df, family = binomial('logit'))
## Warning: glm.fit: algorithm did not converge
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
m$coefficients
```

```
##
    (Intercept) avg sickhours
                             n tweets
     1998.03388
                 150.38876 -79.29431
                                        -33,13080
##
```

- $\eta = 1998.03 + 150.39x_1 + -79.29x_2 + -33.13x_3$
- Note the output Warning: glm.fit: algorithm did not converge
 - Issue: Data is linearly separable (cf. section Issue: Linear Separability)
 - See the plot (next slide)

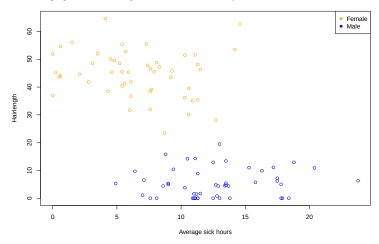
Gender Estimate by Average Sickhours, number of tweets, and IQ



Predictors: Avg. sick hours (x_1) , Hairlength (x_2)

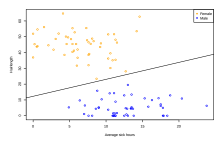
```
m = glm(gender ~ avg_sickhours + hairlength, data = df, family = binomial('logit'))
```

- ## Warning: glm.fit: algorithm did not converge
- ## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred



Issue: Linear Separability

Linear separability of the data causes convergence issues.



- Unstable estimates of the parameters and their standard errors.
- Alternative: Exact logistic regression

Model Evaluation •000000000

Logarithmic Scoring

Model

- Basic evaluation of predicted probabilities:
 - For $Y_{true} = 1$, the predicted probability $\hat{P}(Y = 1)$ should be close to 1.
 - For $Y_{true} = 0$, the predicted probability $\hat{P}(Y = 1)$ should be close to 0.
- Compute logScore = $Y_i \ln(\hat{Y}_i) + (1 Y_i) \ln(1 \hat{Y}_i)$
 - $\blacksquare = \ln(\hat{Y}_i)$ if $Y_i = 1$
 - $= \ln(1 \hat{Y}_i)$ if $Y_i = 0$

Model Selection

Model

- \blacksquare Model selection aims at selecting a model M_i from a set of candidate models M_1, \ldots, M_m .
- The choice depends on the selection criterion and the search method.
- \blacksquare For a model M with k parameters, the Akaike-Information-Criterion is defined as

$$AIC_M = -2\log L_M + 2k$$

- R provides the step() function for stepwise selection from a set of models²
- The step() function uses the AIC as selection criterion for logistic regression models.

²stepwise selection is not an ideal selection technique. Contemporary alternative: regularized methods

```
m0 = glm(chd~1, data=chd data, family=binomial('logit'))
m1 = step(m0, direction='both', trace=0.
         scope='~cigs+chol+weight+age')
summary(m1)
##
## Call:
## glm(formula = chd ~ chol + age + cigs, family = binomial("logit"),
      data = chd_data)
##
## Deviance Residuals:
              1Q Median
      Min
                                        Max
## -1.1307 -0.4596 -0.3550 -0.2587 2.6391
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -8.655131 1.660169 -5.213 1.85e-07 ***
## chol 0.014092 0.003422 4.118 3.82e-05 ***
## age 0.056783 0.029257 1.941 0.0523 .
## cigs 0.017611 0.010368 1.699 0.0894 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 302.35 on 498 degrees of freedom
## Residual deviance: 276.65 on 495 degrees of freedom
## ATC: 284.65
##
## Number of Fisher Scoring iterations: 5
```

Likelihood Ratio Test

- We can test competing nested models with the Likelihood Ratio (LR) Test
- Small model M_1 with k_1 parameters, Large model M_2 with k_2 parameters
- The difference of log-likelihoods is χ^2 distributed:
 - Test statistic $G^2 = -2LL_{M_1} (-2LL_{M_2}) \sim \chi^2(df = k_2 k_1)$
 - If p < .05, the larger model improves model fit.
- Implementation in R for example with anova([models], test='Chisq')

Model Evaluation 0000000000

```
## Analysis of Deviance Table
##
## Model 1: chd ~ 1
## Model 2: chd ~ age
## Model 3: chd ~ age + cigs
## Model 4: chd ~ age + cigs + chol
## Model 5: chd ~ age + cigs + chol + height
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
         498
                 302.35
## 2
         497
                 296.76 1 5.5866 0.01810 *
## 3
         496
               293.78 1
                            2.9815 0.08422 .
## 4
         495
               276.65 1 17.1342 3.483e-05 ***
                 275.52 1 1.1310 0.28755
## 5
         494
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Hosmer and Lemeshow Goodness-of-fit test

- Approach of the HL test:
 - Partition the model population space into bins (risk decentiles)
 - Compare observed relative bin counts with predicted relative bin counts
 - Test statistic follows a χ^2 distribution
- Interpretation

Model

- p < .05 indicates a systematic deviance between observed and predicted bin counts.
- Reasons for a bad fit.
 - Nonlinear influence of predictors on η
 - Solution: Polynomial logistic regression
 - e.g. $\eta = \beta_0 + \beta_{11}x_1 + \beta_{12}x_1^2 + \ldots + \beta_{1p}x_1^p + \beta_{21}x_2 + \ldots + \beta_{kp}x_k^p$
 - Interaction between predictors
 - Solution: Allow and analyze interactions (y~x1*x2).

data: df\$gender01, v_hat

X-squared = 0.57315, df = 3, p-value = 0.9026

```
df$gender01 = as.numeric(df$gender)-1  # recode to 0/1 for HL-test
m = glm(gender01 - avg_sickhours, data=df, family=binomial('logit'))
y_hat = predict(m, type="response")
hoslem.test(df$gender01, y_hat)  # default: g=10 bins

##
## Hosmer and Lemeshow goodness of fit (GOF) test
##
## data: df$gender01, y_hat
## X-squared = 3.6825, df = 8, p-value = 0.8846
hoslem.test(df$gender01, y_hat, g=5)  # g=5 bins

##
## Hosmer and Lemeshow goodness of fit (GOF) test
##
## Hosmer and Lemeshow goodness of fit (GOF) test
##
```

Effect size

Model

McFadden's ρ^2

- **Problem:** We cannot calculate the explained variance R^2
- **Approach:** Calculate McFadden's ρ^2 with the log-likelihood of the full model m1 and the log-likelihood of the null model m0 without predictors.
- **Optional:** Include a correction for the number of predictors k. correction
- Formula: $\rho^2 = 1 \frac{\ln(L_1) k}{\ln(L_1)}$

```
m0 = glm(gender ~ 1,
                            data=df, family=binomial('logit'))
m1 = glm(gender ~ avg sickhours, data=df, family=binomial('logit'))
K = length(m1$coefficients) - 1 # number of predictors
as.numeric(1 - logLik(m1) /logLik(m0)) %>% round(3)
```

```
## [1] 0.36
as.numeric(1 - (logLik(m1)-K)/logLik(m0)) %% round(3)
```

 \blacksquare Another approach to compute an analogon to R^2 is Nagelkerke's (pseudo-) R^2 :

```
m = glm(gender ~ avg_sickhours, data=df, family=binomial('logit'))
N = nrow(df)
(1-exp((m$dev-m$null)/N))/(1-exp(-m$null/N))
```

[1] 0.5238511

Outlook

Predictions

Model

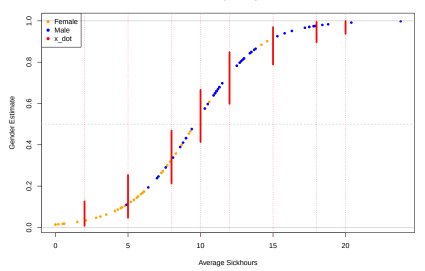
- Given a new input \dot{x} , the prediction on the linear predictor is $\hat{\eta} = \dot{x}\hat{\beta}.$
- This prediction η can be equipped with a confidence interval.
- To obtain a probability confidence interval, $\hat{\eta}$ can be transformed with the well-known inverse link function:

$$\hat{p} = \frac{\exp(\eta)}{1 + \exp(\eta)}$$

```
glm(gender~avg_sickhours, data=df, family=binomial('logit'))
pred = predict(m,newdata=data.frame(avg_sickhours=10),se=T)
(pred_ci = c(pred\fit-1.96*pred\fit, pred\fit+1.96*pred\fit) \%% ilogit())
```

```
## 0 4141107 0 6660099
```

Gender Estimate by Average Sickhours



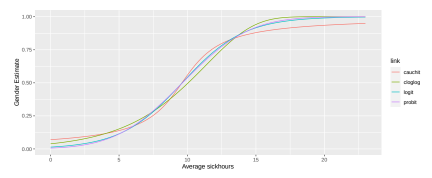
Model

Model Evaluation

- 95% CI: $[\exp(\hat{\beta} 1.96\hat{\sigma}_{\beta}), \exp(\hat{\beta} + 1.96\hat{\sigma}_{\beta})]$
- Invariant to the value of \dot{x}
- Typically reported in clinical research papers.

Other link functions

```
mlogit
          = glm(gender ~ avg_sickhours, data = df, family = binomial(link='logit'))
                         avg_sickhours, data = df, family = binomial(link='probit'))
mprobit
          = glm(gender ~
          = glm(gender ~ avg_sickhours, data = df, family = binomial(link='cloglog'))
mcloglog
mcauchit
          = glm(gender ~ avg_sickhours, data = df, family = binomial(link='cauchit'))
```



Model

- Usually, most observed data lies in the center of the distribution
- Different link functions are typically similar in the center but differ in the tails

Model Evaluation

Approach: Select the link function based on *theoretical* assumptions, experience, and domain expertise.

Questions?