STA2201H Methods of Applied Statistics II

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Week 2: Generalized Linear Models

Overview

Over two weeks

Lecture:

- General Linear Models
- Generalized Linear Models
- Exponential family
- ► Likelihood-based estimation and inference
- Poisson
- Binomial, Multinomial
- Survival analysis

Lab:

- ► EDA
- ► GLM in R

The model fitting process

What are we actually trying to achieve? From last week, applied statistics is:

Using statistical methods to answer questions and draw reasonable conclusions from data that have uncertainty and randomness.

The model fitting process

Overview of process

- 1. Look at the data (EDA, today's lab)
- 2. Decide on a model
 - ▶ Probability distribution for response Y e.g. $Y \sim N(\mu, \sigma^2)$
 - Equation involving explanatory variables (we are trying to explain E[Y])
- 3. Estimate the parameters
- 4. Check the model and residuals
- 5. Inference, interpretation
- 6. Communication

Motivating examples

Outcomes we may be interested in investigating (in relation to other explanatory variables):

- Police stop and frisks in NYC
- Infant deaths in the US
- Number of Canadians who voted for the Liberal party
- Number of Canadians who voted Liberal, Conservatives, LDP
- Concentration of drug at particular times after ingestion

General linear models

We observe $y_1, y_2, \dots y_n$ which are realizations of the random variables Y_1, Y_2, \dots, Y_n

In linear models the y_i 's have two pieces:

1. A **systematic part**, with the form

$$E(\mathbf{Y}) = \mu = \mathbf{X}\beta$$

2. A **random part**, where errors are assumed to be i.i.d such that $E[\epsilon] = 0$ and $var[\epsilon] = \sigma^2$. We usually further assume that errors are Normal with constant variance σ^2 .

General linear models

$$Y_i \sim N(\mu_i, \sigma^2)$$

 $\mu_i = \mathbf{x}_i^T \beta$

General linear models are not appropriate when

- ▶ The range of *Y* is restricted
- ▶ The variance of Y depends on the mean

Generalized Linear Models extend the classical set-up to allow for a wider range of distributions. Introduced by Nelder and Wedderburn (1972) [Later, GAMs in 1990].

Generalized linear models

GLMs have an additional piece on top of the classical linear models:

- 1. **random component**: $Y_i \sim \text{some distribution with } E[Y_i] = \mu_i$
- 2. systematic component: $\mathbf{x}_{i}^{T} \boldsymbol{\beta}$
- 3. The **link function** that links the random and systematic components $g(u_i) = \mathbf{x}_i^T \boldsymbol{\beta}$
- ▶ Set-up is almost the same, particularly in terms of specifying a good linear predictor $\mathbf{x}_i^T \boldsymbol{\beta}$
- Just need to think about the link and the distribution of the outcome

GLMs

$$Y_i \sim G(\mu_i, \phi)$$

$$E[Y_i] = \mu_i$$

$$g(\mu_i) = \mathbf{x}_i^T \beta$$

 $ightharpoonup \phi$ is the scale parameter.

What can Y be distributed as? In principle, anything. In practice (and original formulation), distributions come from the **exponential** family.



Exponential Family

The random variable Y belongs to the exponential family of distributions if its support does not depend upon any unknown parameters and its density or probability mass function takes the form

$$p(y|\theta,\phi) = \exp\left(\frac{y\theta - b(\theta)}{\phi} + c(y,\phi)\right)$$

- $\theta = h(\mu)$ depends on the expected value of y and is the canonical parameter
- $ightharpoonup \phi$ is the scale parameter (if known: one-parameter family)
- b and c are arbitrary functions

Example: Poisson distribution

$$p(y|\theta,\phi) = \exp\left(\frac{y\theta - b(\theta)}{\phi} + c(y,\phi)\right)$$

Poisson:

$$p(y|\mu) = \frac{\mu^y e^{-\mu}}{y!}$$

Write as

$$p(y|\mu) = \exp\{y \log \mu - \mu - \log y!\}$$

- \bullet $\theta = \log \mu$
- \blacktriangleright $b(\theta) = e^{\theta}$
- $c(y, \phi) = -\log y!$
- \blacktriangleright Note that the scale parameter $\phi=1$ so the variance is entirely determined by the mean

Example: Normal distribution

$$p(y|\theta,\phi) = \exp\left(\frac{y\theta - b(\theta)}{\phi} + c(y,\phi)\right)$$

Normal:

$$p(y|\mu,\sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left\{-\frac{(y-\mu)^2}{2\sigma^2}\right\}$$

Write as

$$f(y|\mu,\sigma^2) = \exp\left\{rac{y\mu - rac{1}{2}\mu^2}{\sigma^2} - rac{1}{2}\left[rac{y^2}{\sigma^2} + \log(2\pi\sigma^2)
ight]
ight\}$$

$$\bullet$$
 $\theta = \mu$

$$b(\theta) = \frac{1}{2}\theta^2$$

$$\phi = \sigma^2$$

$$c(y,\phi) = -\frac{1}{2} \left[\frac{y^2}{\sigma^2} + \log(2\pi\sigma^2) \right]$$

Other examples

Other common examples:

- ► Binomial
- Gamma
- Negative binomial
- ► Inverse Gaussian

Propertites of exponential families

Mean and variance for exponential families

$$p(y|\theta,\phi) = \exp\left(\frac{y\theta - b(\theta)}{\phi} + c(y,\phi)\right)$$

It can be shown that

$$E(Y|\theta,\phi) = b'(\theta) = \mu$$

and

$$Var(Y|\theta,\phi) = \phi b''(\theta) = \phi V(\mu)$$

Note the variance of Y depends not only on the scale parameter but also on a function of the mean.

Examples:

$$E(Y|\theta,\phi) = b'(\theta)$$

and

$$Var(Y|\theta,\phi) = \phi b''(\theta)$$

- ▶ Poisson: $E(Y|\theta,\phi) = e^{\theta} = \mu$, $Var(Y|\theta,\phi) = 1 \times e^{\theta} = \mu$
- Normal: $E(Y|\theta,\phi) = \theta = \mu$, $Var(Y|\theta,\phi) = \sigma^2 \times 1 = \sigma^2$

The canonical link

The link function $\eta_i = g(\mu)$ could in theory be any function linking the linear predictor to the distribution of the outcome variable, which is also is **monotonic** and **smooth**.

Recall $\theta = h(\mu)$. If we choose g = h, then

$$\theta_i = h(\mu_i) = h(h^{-1}(\eta_i)) = \eta_i = \mathbf{x}_i^T \beta$$

In other words, it ensures that the systematic component of our model is modeling the parameter of interest.

Canonical links

- ▶ Normal: identity $\theta = h(\mu) = \mu$
- Poisson: $\theta = h(\mu) = \log \mu$
- ▶ Binomial: $\theta = h(\mu) = \log(\frac{\mu}{1-\mu})$

Likelihood-based estimation and inference

Estimation

- Inference is based on MLE, but cannot derive closed form solutions for regression coefficients
- Note we are assuming independence $cov(Y_i, Y_j | \theta_i, \theta_j, \phi) = 0$ for $i \neq j$. (more on dependence later)

The log-likelihood function is:

$$\ell(\theta) = \sum_{i} \ell(\theta_{i}) = \sum_{i} \frac{y_{i}\theta_{i} - b(\theta_{i})}{\phi} + c(y_{i}, \phi)$$

What's our usual approach here?

Score function and Information matrix

$$\mathsf{S}(\beta) = \mathsf{D}^\mathsf{T} \mathsf{V}^{-1} \frac{\mathsf{Y} - \mu(\beta)}{\phi}$$

where D^T is a matrix of the $\partial \mu_i/\partial \beta_j$ and **V** is diagonal with *i*th element $b''(\theta_i)$.

$$I(\beta) = x^t W(\beta) x$$

where **W** is diagonal with $w_i = (\frac{\partial \mu_i}{\partial \eta_i})^2/\phi b''(\theta_i)$.

What about ϕ ?

When ϕ is unknown, can estimate it using

$$\hat{\phi} = \frac{1}{n-k-1} \sum_{i} \frac{(Y_i - \hat{\mu}_i)^2}{V(\hat{\mu})}$$

where $\hat{\mu} = \hat{\mu}(\hat{\beta})$.

Newton-Raphson

Want to find roots such that $S(\beta) = 0$. First order TS approximation:

$$\mathsf{S}(\beta) \approx \mathsf{S}(\beta^{(0)}) + (\beta - \beta^{(0)})^{\mathsf{T}} \mathsf{S}'(\beta^{(0)})$$

Newton-Raphson iterates the step:

$$\beta^{(t+1)} = \beta^{(t)} - S'(\beta^{(t)})^{-1}S(\beta^{(t)})$$

Method of scoring replaces observed information with its expectation $\mathbf{E}[\mathbf{S}'(\beta)] = -\mathbf{I}(\beta)$.

$$\beta^{(t+1)} = \beta^{(t)} + I(\beta^{(t)})^{-1}S(\beta^{(t)})$$

Estimation

Can be rewritten in the form:

$$\widehat{\beta}^{(t+1)} = (\mathbf{x}^\mathsf{T} \mathbf{W} \mathbf{x})^{-1} \mathbf{x}^\mathsf{T} \mathbf{W} \mathbf{z}$$

- **W** and **z** change depending on $\hat{\beta}$ and vice versa
- Use iteratively weighted least squares (IWLS)
 - 1. Choose initial value $\hat{\beta}^{(0)}$
 - 2. Calculate W and z
 - 3. Repeat until convergence

Inference

We know that for the MLE

$$\hat{\beta} \sim \mathcal{N}(\beta, (\mathbf{X}^\mathsf{T}\mathbf{W}\mathbf{X})^{-1}\phi)$$

$$\hat{\beta} \sim N(\beta, I(\hat{\beta})^{-1})$$

Standard errors are the square roots of the inverse of the information matrix.

▶ Use this for the classic Wald Tests e.g. $\sqrt{W} = \frac{\hat{\beta} - \beta_0}{\text{se}(\hat{\beta})}$ follows z distribution.

Likelihood ratio test

Testing nested models, ω_1 and ω_2 , $\omega_1 \in \omega_2$ and number of parameters $p_2 > p1$

$$2[\log L(\widehat{\beta}_1|\mathbf{y}) - \log L(\widehat{\beta}_2|\mathbf{y})] \sim \chi_{\rho_1-\rho_2}$$

Compared to Wald: doesn't assume symmetry in confidence intervals, but requires you to run two models.

GLM in R

- ▶ glm()
- ▶ same set up as lm(); additional family argument with a link
- ► e.g. glm(y~x, family = binomial(link = 'logit')

Poisson regression

Review

- ► mean ?
- variance ?
- ▶ link: ?

What's a problem with just looking at counts?

Offsets

$$Y_i \sim \mathsf{Poisson}(\lambda_i)$$
or $Y_i \sim \mathsf{Poisson}(\mu_i O_i)$
 $\implies \log \mu_i = \mathbf{x_i}^T \beta$

Offset controls for exposure to risk/making inferences to some baseline. e.g.

- population size
- age
- time since exposed

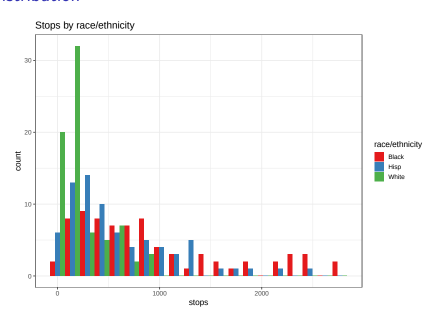
Example: Police stops

Police stop and frisks in NYC (Gelman Hill Chapter 6). Is there a difference in the number of stops by race/ethnicity?

The data look like:

precinct	stops	arrests	race_eth
1	202	980	Black
1	102	295	Hisp
1	81	381	White
2	132	753	Black
2	144	557	Hisp
2	71	431	White
3	752	2188	Black
3	441	627	Hisp
3	410	1238	White
4	385	471	Black

Distribution



GLM

Use arrests as exposure

```
mod1 <- glm(stops-race_eth,family=poisson,offset=log(arrests),data=d)
summary(mod1)</pre>
```

```
##
## Call:
## glm(formula = stops ~ race_eth, family = poisson, data = d, offset = log(arrests))
##
## Deviance Residuals:
      Min
                10 Median 30
                                        Max
## -47.327 -7.740 -0.182 10.241 39.140
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -0.588086 0.003784 -155.40 <2e-16 ***
## race ethHisp 0.070208 0.006061 11.58 <2e-16 ***
## race ethWhite -0.161581 0.008558 -18.88 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
      Null deviance: 46120 on 224 degrees of freedom
## Residual deviance: 45437 on 222 degrees of freedom
## ATC: 47150
##
## Number of Fisher Scoring iterations: 5
```

GLM

Add in factors for precinct

```
mod2 <- glm(stops-race_eth + factor(precinct), family=poisson,offset=log(arrests),data=d)
summary(mod2)[["coefficients"]][1:10,]</pre>
```

```
##
                       Estimate Std. Error
                                               z value
                                                            Pr(>|z|)
## (Intercept)
                    -1.37886803 0.051019006 -27.026556 7.205634e-161
## race_ethHisp
                                              1.497782 1.341899e-01
                     0.01018798 0.006802045
## race ethWhite
                    -0.41900122 0.009434996 -44.409261 0.000000e+00
## factor(precinct)2 -0.14904964 0.074030344
                                             -2.013359 4.407691e-02
## factor(precinct)3
                     0.55995498 0.056758425
                                             9.865583 5.869222e-23
## factor(precinct)4
                     1.21063605 0.057548994
                                             21.036615 3.032678e-98
## factor(precinct)5 0.28286532 0.056794015
                                              4.980548 6.340447e-07
## factor(precinct)6
                    1.14420375 0.058047383 19.711547 1.716374e-86
## factor(precinct)7 0.21817307 0.064335032
                                              3.391202 6.958688e-04
## factor(precinct)8 -0.39056473 0.056867814
                                             -6.867940 6.513564e-12
```

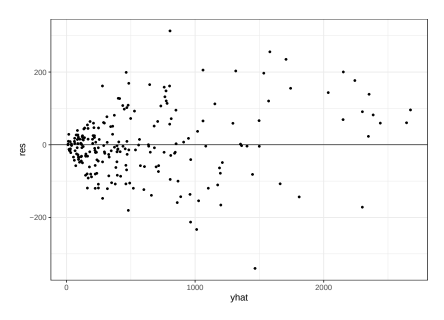
Coefficient interpretation

- ▶ e.g. after controlling for precinct, compared to blacks, whites have 1 exp(-0.42) = 34% less chance of being stopped.
- be wary of exposure variable: stops are compared to the number of arrests in the previous year
- so that the coefficient 'whites' will be less than 1 if the people in that group are stopped disproportionately less than their rates of arrest, as compared to blacks.
- would be different if we had population as exposure variable

Is this a reasonable model?

Look at predicted values versus residuals $(y_i - \hat{y}_i)$. What do we expect?

Predicted values versus residuals



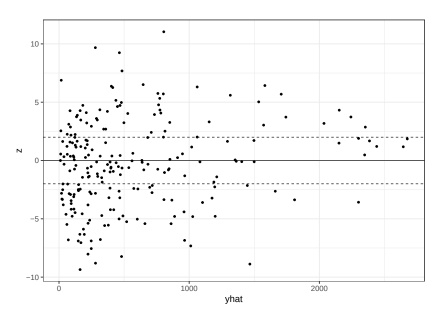
Is this a reasonable model?

Consider standardized residuals

$$z_i = \frac{y_i - \hat{y}_i}{sd(\hat{y}_i)}$$

If Poisson is a good model then these should have mean 0 and sd 1.

Predicted values versus standardized residuals



Overdispersion

- Extra variation in the data beyond what is allowed for in statistical model
- Poisson does not have independent variance parameter

Test for overdispersion: compare sum of squares of standardized residuals to χ^2_{n-k} distribution.

Estimated overdispersion factor is

$$\frac{1}{n-k}\sum_{i}z_{i}^{2}$$

Overdispersion

overdispersion factor is

```
sum(res_df\$z^2)/(n-k)
```

```
## [1] 21.88505
```

P-value of test is

```
pchisq(sum(res_df$z^2), n-k)
```

```
## [1] 1
```

But what's a problem here?

Fit overdispersed Poisson

- ightharpoonup General form includes extra dispersion parameter heta
- Assume variance is proportion to the mean, rather than equal to the mean $E[Y] = \mu\theta$

```
mod3 <- glm(stops-race_eth + factor(precinct), family=quasipoisson,offset=log(arrests),data=d)
summary(mod3)[["coefficients"]][1:10,]</pre>
```

```
##
                       Estimate Std. Error
                                              t value
                                                          Pr(>|t|)
## (Intercept)
                    -1.37886803 0.23867441 -5.7771925 4.326149e-08
## race ethHisp
                     0.01018798 0.03182097 0.3201657 7.492943e-01
## race_ethWhite
                    -0.41900122 0.04413830 -9.4929170 5.489337e-17
## factor(precinct)2 -0.14904964 0.34632483 -0.4303753 6.675488e-01
## factor(precinct)3 0.55995498 0.26552425 2.1088656 3.664011e-02
## factor(precinct)4 1.21063605 0.26922265 4.4967837 1.384310e-05
## factor(precinct)5 0.28286532 0.26569075 1.0646412 2.887722e-01
## factor(precinct)6
                    1.14420375 0.27155419 4.2135374 4.352372e-05
## factor(precinct)7 0.21817307 0.30096874 0.7249028 4.696562e-01
## factor(precinct)8 -0.39056473 0.26603599 -1.4680898 1.442019e-01
```

Notice

```
summary(mod3)[["dispersion"]]
```

```
## [1] 21.88506
```

... and the SEs are inflated $\sim \sqrt{21.9}$.

Overdisperson

Downside to quasi-Poisson it's not true MLE so you don't get likelihood etc to compare models.

Alternative:

- ightharpoonup Could also add a multiplicative random effect θ to represent unobserved heterogeneity.
- ▶ Conditional distribution is Poisson $E[Y|\theta] \sim Pois(\mu\theta)$
- Leads to unconditional distribution being Negative Binomial distribution
- ► Can choose parameters so $E(Y) = \mu$ and $Var(Y) = \mu(1 + \sigma^2 \mu)$

Overdispersion

Fit Negative Binomial

```
library(MASS)
mod4 <- glm.nb(stops-race_eth + factor(precinct), data = d)
summary(mod3)[["coefficients"]][1:10,]</pre>
```

```
##
                        Estimate Std. Error
                                              t value
                                                           Pr(>|t|)
## (Intercept)
                     -1.37886803 0.23867441 -5.7771925 4.326149e-08
## race_ethHisp
                     0.01018798 0.03182097 0.3201657 7.492943e-01
## race_ethWhite
                     -0.41900122 0.04413830 -9.4929170 5.489337e-17
## factor(precinct)2 -0.14904964 0.34632483 -0.4303753 6.675488e-01
## factor(precinct)3 0.55995498 0.26552425 2.1088656 3.664011e-02
## factor(precinct)4
                     1.21063605 0.26922265 4.4967837 1.384310e-05
## factor(precinct)5 0.28286532 0.26569075 1.0646412 2.887722e-01
## factor(precinct)6
                     1.14420375 0.27155419 4.2135374 4.352372e-05
## factor(precinct)7
                     0.21817307 0.30096874 0.7249028 4.696562e-01
## factor(precinct)8 -0.39056473 0.26603599 -1.4680898 1.442019e-01
```

Lab

- Using data from Open Data Portal in Toronto
 - opendatatoronto package by Sharla Gelfand
- ► EDA
- Questions at end need to be handed in via GitHub