mentor effect plot


012345678910111213141516171819 Number of Articles



## GLMs for Count Data



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## Topics

- Generalized linear models
- GLMs for count data
- Example: PhD publications
- Model diagnostics
- Interactions
- Nonlinearity
- Outliers, leverage \& influence
- Overdispersion
- Quasi-poisson models
- Negative binomial models
- Excess zeros
- Zero-inflated models
- Hurdle models


## Count data models: Overview

- Count data models arise when the basic observation is a frequency, $y=0,1,2, \ldots$ of some event and we have some predictors, $x_{1}, x_{2}, \ldots$ to help explain them.
" Typically, these counts ~ Poisson() $\rightarrow$ "poisson regression"
- Examples:
- Number of articles published by PhD candidates
- Predictors: Married?, Female?, Kids < 5?, pubs by mentor
- Number of parasites in blood samples of Norwegian cod
- Predictors: Catch area, Year, length of fish
- Female horseshoe crabs: Number of "satellite" males
- Predictors: Female weight, color, spine condition, shell width
- Special circumstances
- Overdispersion: when the variance > mean
- Zero-counts: When excess 0 counts require an extra model


## Generalized linear models

We have used generalized linear models fit with gIm() in two contexts so far

## Loglinear models

- the outcome variable is the vector of frequencies $\boldsymbol{y}$ in a table cross-classified by factors in a design matrix $\boldsymbol{X}$
- The model is expressed as a linear model for $\log \boldsymbol{y}$

$$
\log (\boldsymbol{y})=\boldsymbol{X} \boldsymbol{\beta}
$$

- The random (or unexplained) variation is expressed as a Poisson distribution for $\mathcal{E}(\boldsymbol{y} \mid \boldsymbol{X})$


## Generalized linear models

## Logistic regression

- the outcome variable is a categorical response $\boldsymbol{y}$, with predictors $\boldsymbol{X}$
- The model is expressed as a linear model for the $\log$ odds that $y=1 \mathrm{vs}$. $y=0$.

$$
\operatorname{logit}(\boldsymbol{y}) \equiv \log \left[\frac{\operatorname{Pr}(y=1)}{\operatorname{Pr}(y=0)}\right]=\boldsymbol{x} \boldsymbol{\beta}
$$

- The random (or unexplained) variation is expressed as a Binomial distribution for $\mathcal{E}(\boldsymbol{y} \mid \boldsymbol{X})$

Hey, aren't these both very like the familiar, classical linear model,

$$
\boldsymbol{y}=\boldsymbol{X} \boldsymbol{\beta}+\boldsymbol{\epsilon}, \quad \epsilon \sim \mathcal{N}\left(0, \sigma^{2} \boldsymbol{I}\right) \quad ?
$$

Yes, for some transformation, $g(\boldsymbol{y})$, and with different distributions!

## Generalized linear models

Nelder \& Wedderburn (1972) said, "Let there be light!", a generalized linear model, encompassing them all, and many more. This has 3 components:

- A random component, specifying the conditional distribution of $\boldsymbol{y}$ given the explanatory variables in $\boldsymbol{X}$, with mean $\mathcal{E}\left(y_{i} \mid \boldsymbol{X}_{i}\right)=\mu_{i}$
- The normal (Gaussian), binomial, and Poisson are already familiar
- But, these are all members of an exponential family
- GLMs now include an even wider family: negative-binomial and others
- The systematic component, a linear function of the predictors called the linear predictor

$$
\boldsymbol{\eta}=\boldsymbol{X} \beta \quad \text { or } \quad \eta_{i}=\beta_{0}+\beta_{1} X_{i 1}+\cdots+\beta_{p} X_{i p}
$$

- An invertible link function, $g\left(\mu_{i}\right)=\eta_{i}=\boldsymbol{X}_{i}^{\top} \boldsymbol{\beta}$ that transforms the expected value of the response to the linear predictor
- The link function is invertable, so we can go back to the mean function $g^{-1}\left(\eta_{i}\right)=\mu_{i}$


## Link functions for the mean

Standard GLM link functions and their inverses:
Table 11.1: Common link functions and their inverses used in generalized linear models
$\left\{\begin{array}{lll}\hline \text { Link name } & \text { Function: } \eta_{i}=g\left(\mu_{i}\right) & \text { Inverse: } \mu_{i}=g^{-1}\left(\eta_{i}\right) \\ \hline \text { identity } & \mu_{i} & \eta_{i} \\ \text { square-root } & \sqrt{\mu_{i}} & \eta_{i}^{2} \\ \log & \log _{e}\left(\mu_{i}\right) & \exp \left(\eta_{i}\right) \\ \text { inverse } & \mu_{i}^{-1} & \eta_{i}^{-1} \\ \text { inverse-square } & \mu_{i}^{-2} & \eta_{i}^{-1 / 2} \\ \hline \text { logit } & \log _{e} \frac{\mu_{i}}{1-\mu_{i}} & \frac{1}{1+\exp \left(-\eta_{i}\right)} \\ \text { probit } & \Phi^{-1}\left(\mu_{i}\right) & \Phi\left(\eta_{i}\right) \\ \operatorname{log-log} & -\log _{e}\left[-\log _{e}\left(\mu_{i}\right)\right] & \exp \left[-\exp \left(-\eta_{i}\right)\right] \\ \text { comp. } \log -\log & \log _{e}\left[-\log _{e}\left(1-\mu_{i}\right)\right] & 1-\exp \left[-\exp \left(\eta_{i}\right)\right] \\ \hline\end{array}\right.$

- The top section recognizes standard transformations of $y_{i}$ often used with classical linear models
- The bottom section is for binomial data, where $y_{\mathrm{i}}$ represents an observed count in $n_{i}$ trials


## Canonical links and variance functions

- For every distribution family, there is a default, canonical link function
- Each one also specifies the expected relation between the mean and variance

Table 11.2: Common distributions in the exponential family used with generalized linear models and their canonical link and variance functions

| Family | Notation | Canonical link | Range of $y$ | Variance function, $\mathcal{V}(\mu \mid \eta)$ |
| :--- | :--- | :--- | :--- | :--- |
| Gaussian | $N\left(\mu, \sigma^{2}\right)$ | identity: $\mu$ | $(-\infty,+\infty)$ | $\phi$ |
| Poisson | $\operatorname{Pois}(\mu)$ | $\log _{e}(\mu)$ | $0,1, \ldots, \infty$ | $\mu$ |
| Negative-Binomial | $\operatorname{NBin}(\mu, \theta)$ | $\log _{e}(\mu)$ | $0,1, \ldots, \infty$ | $\mu+\mu^{2} / \theta$ |
| Binomial | $\operatorname{Bin}(n, \mu) / n$ | $\operatorname{logit}^{( }(\mu)$ | $\{0,1, \ldots, n\} / n$ | $\mu(1-\mu) / n$ |
| Gamma | $G(\mu, \nu)$ | $\mu^{-1}$ | $(0,+\infty)$ | $\phi \mu^{2}$ |
| Inverse-Gaussian | $I G(\mu, \nu)$ | $\mu^{2}$ | $(0,+\infty)$ | $\phi \mu^{3}$ |

## Variance functions \& overdispersion

- In the classical Gaussian linear model, the conditional variance is constant, $\phi=\sigma_{\epsilon}^{2}$.
- For binomial data, the variance function is $\mathcal{V}\left(\mu_{i}\right)=\mu_{i}\left(1-\mu_{i}\right) / n_{i}$, with $\phi$ fixed at 1
- In the Poisson family, $\mathcal{V}\left(\mu_{i}\right)=\mu_{i}$ and the dispersion parameter is fixed at $\phi=1$.
- In practice, it is common for count data to exhibit overdispersion, meaning that $\mathcal{V}\left(\mu_{i}\right)>\mu_{i}$.
- One way to correct for this is to allow the dispersion parameter to be estimated from the data, giving what is called the quasi-Poisson family, with $\mathcal{V}\left(\mu_{i}\right)=\widehat{\phi} \mu_{i}$.


## What is overdispersion?

Overdispersion often results from failures of assumptions of the model

- Supposedly independent observations may be correlated
- The probability of an event may not be constant, or
- it may vary with unmeasured or unmodeled variables


## Maximum likelihood estimation

- GLMs are fit by the method of maximum likelihood
- Likelihood $(\mathcal{L})=\operatorname{Pr}$ (data | model), as function of model parameters
- For the Poisson distribution with mean $\mu$, the probability that the random variable $Y$ takes the values $y=0,1,2, \ldots$ is


## Maximum likelihood estimation

- GLMs are fit by the method of maximum likelihood.
- For the Poisson distribution with mean $\mu$, the probability that the random variable $Y$ takes values $y=0,1,2, \ldots$ is

$$
\operatorname{Pr}(Y=y)=\frac{e^{-\mu} \mu^{y}}{y!}
$$

- In the GLM with a log link, the mean, $\mu_{i}$ depends on the predictors in $\boldsymbol{x}$ through

$$
\log _{e}\left(\mu_{i}\right)=\boldsymbol{x}_{i}^{\top} \boldsymbol{\beta}
$$

- The log-likelihood function (ignoring a constant) for $n$ independent observations has the form

$$
\log _{e} \mathcal{L}(\boldsymbol{\beta})=\sum^{n}\left\{y_{i} \log _{e}\left(\mu_{i}\right)-\mu_{i}\right\}
$$

- It can be shown that the maximum likelihood estimators are solutions to the estimating equations,

$$
\boldsymbol{X}^{\top} \boldsymbol{y}=\boldsymbol{X}^{\top} \mu
$$

- The solutions are found by iteratively re-weighted least squares.


## Goodness of fit

- The residual deviance defined as twice the difference between the maximum log-likelihood for the saturated model that fits perfectly and maximized log-likelihood for the fitted model.

$$
D(\boldsymbol{y}, \widehat{\boldsymbol{\mu}}) \equiv 2\left[\log _{e} \mathcal{L}(\boldsymbol{y} ; \boldsymbol{y})-\log _{e} \mathcal{L}(\boldsymbol{y} ; \widehat{\boldsymbol{\mu}})\right]
$$

- For classical (Gaussian) linear models, this is just the residual sum of squares
- For Poisson models with a log link giving $\boldsymbol{\mu}=\exp \left(\boldsymbol{x}^{\top} \boldsymbol{\beta}\right)$, the deviance takes the form

$$
D(\boldsymbol{y}, \widehat{\boldsymbol{\mu}})=2 \sum_{i=1}^{n}\left[y_{i} \log _{e}\left(\frac{y_{i}}{\widehat{\mu}_{i}}\right)-\left(y_{i}-\widehat{\mu}_{i}\right)\right]
$$

- For a GLM with $p$ parameters, both the Pearson and residual deviance statistics follow approximate $\chi_{n-p}^{2}$ distributions with $n-p$ degrees of freedom.


## GLMs for count data

- Typically, these are fit using
glm(y ~ x1 + x2 + ..., family=poisson, data=mydata)
- As in other linear models, the predictors, $\mathrm{x}_{\mathrm{i}}$, can be discrete factors, quantitative variables, interactions, etc.
- This fixes the dispersion parameter, $\phi$ to 1 , assuming the count variable $\mathrm{y} \mid$ $\mathrm{x} 1, \mathrm{x} 2, \ldots$ is Poisson distributed
- It is possible to relax this, and fit a quasi-Poisson model, allowing $\phi$ to be estimated from the data
- Specify family=quasipoisson. This allows variance to be proportional to the mean

$$
\mathcal{V}\left(y_{i} \mid \eta_{i}\right)=\phi \mu_{i}
$$

- Another possibility is the negative-binomial model, which has

$$
\mathcal{V}\left(y_{i} \mid \eta_{i}\right)=\mu_{i}+\mu_{i}^{2} / \theta
$$

## Example: Publications of PhD candiates

Example 3.24 in DDAR gives data on the number of publications by PhD candidates in biochemistry in the last 3 years of study

```
> data("PhdPubs", package = "vcdExtra")
```

> table(PhdPubs\$articles)

| 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 16 | 19 |
| ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| 275 | 246 | 178 | 84 | 67 | 27 | 17 | 12 | 1 | 2 | 1 | 1 | 2 | 1 | 1 |

Predictors are:

- gender, marital status
- number of young children
- prestige of the doctoral department
- number of publications by the student's mentor


## Example: Publications of PhD candidates

Initially, ignore the predictors
This is equivalent to an intercept-only Poisson model

```
glm(articles ~ 1, family=poisson, data = PhdPubs)
```

As a check on the Poisson assumption, calculate the mean and variance

```
> with(PhdPubs, c(mean=mean(articles),
    var=var(articles),
    ratio=var(articles)/mean(articles)))
    mean var ratio
    1.69 3.71 2.19
```

The assumption that mean = variance could be met when we add predictors

First, look at rootograms:

```
```

plot(goodfit(PhdPubs\$articles), xlab = "Number of Articles",

```
```

plot(goodfit(PhdPubs$articles), xlab = "Number of Articles",
    main = "Poisson")
    main = "Poisson")
plot(goodfit(PhdPubs$articles, type = "nbinomial"),
plot(goodfit(PhdPubs\$articles, type = "nbinomial"),
xlab = "Number of Articles", main = "Negative binomial")

```
```

    xlab = "Number of Articles", main = "Negative binomial")
    ```
```

Poisson

$\begin{array}{lllllllllllll}0 & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 & 10 & 111213141516171819\end{array}$ Number of Articles

Negative binomial

$\begin{array}{llllllllllll}0 & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 & 10111213141516171819\end{array}$ Number of Articles

One reason the Poisson doesn't fit: excess 0s (some never published?)

## Fitting the Poisso model

Fit the model with all main effects; note the ${ }^{\sim}$. notation for this

```
> phd.pois <- glm(articles ~ ., data=PhdPubs, family=poisson)
> Anova(phd.pois)
Analysis of Deviance Table (Type II tests)
Response: articles
    LR Chisq Df Pr(>Chisq)
female
married
kid5
    17.1 1 1 3.6e-05 ***
    22.1 1 2.6e-06 ***
phdprestige
    1.0 1 0.32
mentor
    126.8 1 < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 `*' 0.05 '.' 0.1 '' 1
```

Only phdprestige is NS; it does no harm to keep it, for now

## Interpreting coefficients

$\beta_{j}$ is the increment in log (articles) for a 1 unit change in $x_{j} ; \exp \left(\beta_{j}\right)$ is the multiple of articles:

```
round(cbind(beta = coef(phd.pois),
    expbeta = exp(coef(phd.pois)),
    pct = 100 * (exp(coef(phd.pois)) - 1)), 3)
```

| \#\# | beta | expbeta | pct |  |
| :--- | :--- | ---: | ---: | ---: |
| \#\# | (Intercept) | 0.266 | 1.304 | 30.425 |
| \#\# female1 | -0.224 | 0.799 | -20.102 |  |
| \#\# married1 | 0.157 | 1.170 | 17.037 |  |
| \#\# kid5 | -0.185 | 0.831 | -16.882 |  |
| \#\# phdprestige | 0.025 | 1.026 | 2.570 |  |
| \#\# mentor | 0.025 | 1.026 | 2.555 |  |

Thus:

- females publish -0.224 fewer log (articles), or $0.8 \times$ that of males
- married publish 0.157 more log (articles); or $1.17 \times$ unmarried ( $17 \%$ increase)
- each additional young child decreases this by 0.185 ; or $0.831 \times$ articles (16.9\% decrease)
- each mentor pub multiplies student pub by 1.026, a $2.6 \%$ increase


## Effect plots

As usual, we can understand the fitted model from predicted values for the model effects:
library (effects); plot(allEffects(phd.pois))


These are better visual summaries for a model than a table of coefficients.

## Model diagnostics

Diagnostic methods for count data GLMs are similar to those used for classical linear models

- Test for presence of interactions
- Fit model(s) with some or all two-way interactions
- Non-linear effects of quantitative predictors"
- Component-plus-residual plots- car::crPlot() is useful here
- Outliers? Influential observations?
- car::influencePlot() is your friend
- For count data models we should also check for overdispersion
- Similar to homogeneity of variance checks in Im()


## Checking for interactions

## As a quick check for interactions, fit a model with all two-way terms, . ~ .^2

```
> phd.pois1 <- update(phd.pois, . ~ .^2)
> Anova(phd.poisl)
Analysis of Deviance Table (Type II tests)
```

Response: articles
LR Chisq Df Pr(>Chisq)
female
married
kid5
phdprestige
mentor
female:married
female:kid5
14.510 .00014 ***
6.210 .01277 *
$19.519 .8 \mathrm{e}-06$ ***
1.010 .32655
$128.11<2 e-16$ ***
0.310 .60995
0.110 .72929
female:phdprestige
0.210 .63574
female:mentor
$0.01 \quad 0.91260$
married:kid5
0
married:phdprestige
married:mentor
kid5:phdprestige
1.210 .28203
$0.21 \quad 0.68523$
kid5:mentor
2.810 .09290 .
phdprestige:mentor
3.810 .05094
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

## Compare models

The all main effects and all two-way models are nested, so we can compare them with anova()

```
> anova(phd.pois, phd.poisl, test="Chisq")
Analysis of Deviance Table
Model 1: articles ~ female + married + kid5 + phdprestige + mentor
Model 2: articles ~ female + married + kid5 + phdprestige + mentor +
female:married +
        female:kid5 + female:phdprestige + female:mentor + married:kid5 +
        married:phdprestige + married:mentor + kid5:phdprestige +
        kid5:mentor + phdprestige:mentor
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
1 909 1634
2 900 1618 9 15.2 0.086 .
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 '.' 0.1 ' ' 1
```

$\rightarrow$ No evidence that the two-way terms result in a significantly better model

## Compare models

We can also compare using AIC/BIC with vcdExtra: :Lrstats()

```
> LRstats(phd.pois, phd.poisl)
Likelihood summary table:
    AIC BIC LR Chisq Df Pr(>Chisq)
phd.pois 3313 3342 1634 909 <2e-16 ***
phd.pois1 3316 3388 1618 900 <2e-16 ***
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 '.' 0.1 ' ' 1
```

- There seems to be no reason to include interactions in this model
- Interactions increase AIC \& BIC
- We might want to revisit this, after examining other models for the basic count distribution (quasi-poisson, negative-binomial)


## Basic model plots

Only two of the standard model plots are informative for count data models

```
plot(phd.pois, which=c (1,5))
```




## Nonlinearity diagnostics

- Nonlinear relations are difficult to assess in marginal plots, because they don't control (or adjust) for other predictors
- Component-plus-residual plots (also called: partial residual plots) can show nonlinear relations for numeric predictors
- These graph the value of $\hat{\beta}_{\mathrm{i}} \mathrm{xi}+$ residual $_{\mathrm{i}}$ vs. the predictor $\mathrm{x}_{\mathrm{i}}$
- In this plot, the slope of the points is the coefficient $\hat{\beta}_{\mathrm{i}}$ in the full model
- The residual is $y_{i}-\widehat{y}_{i}$ in the full model
- A non-parametric (e.g., loess()) smooth facilitates detecting nonlinearity


## Nonlinearity diagnostics: crPlot()

Is the relation between article published by the student and by the mentor adequately represented as linear?

```
crPlot(phd.pois, "mentor", pch=16, lwd=4, id = list(n=2))
```



The smoothed curve doesn't differ much from the fitted line

A couple of points stand out: 328, 803, 911, 913

## Residuals

Residuals contain all the information about how a model doesn't fit, and maybe why

For GLMs, there are several types, based on the Pearson and deviance goodness-of-fit statistics

- the Pearson residual is the case-wise contribution to Pearson $\chi^{2}$

$$
r_{i}^{P}=\frac{y_{i}-\widehat{\mu}_{i}}{\sqrt{\widehat{\mathcal{V}}\left(y_{i}\right)}}
$$

- the deviance residual is the signed square root of the contribution to the deviance $G^{2}$

$$
r_{i}^{D}=\operatorname{sign}\left(y_{i}-\widehat{\mu}_{i}\right) \sqrt{d_{i}}
$$

These are raw residuals, on the scale of the counts themselves

## Residuals

- Both of these have standardized forms that correct for conditional variance and leverage, and have approx. $\mathcal{N}(0,1)$ distributions.

$$
\begin{aligned}
\tilde{r}_{i}^{P} & =\frac{r_{i}^{P}}{\sqrt{\widehat{\phi}\left(1-h_{i}\right)}} \\
\tilde{r}_{i}^{D} & =\frac{r_{i}^{D}}{\sqrt{\widehat{\phi}\left(1-h_{i}\right)}}
\end{aligned}
$$

- The most useful is the studentized residual (or deletion residual), rstudent () in R. This estimates the standardized residual resulting from omitting each observation in turn. An approximation is:

$$
\widetilde{r}_{i}^{S}=\operatorname{sign}\left(y_{i}-\widehat{\mu}_{i}\right) \sqrt{\left(1-h_{i}\right)\left(\widetilde{r}_{i}^{D}\right)^{2}+h_{i}\left(\widetilde{r}_{i}^{P}\right)^{2}} .
$$

Don't worry about the formulas, but do know the difference among raw, standardized and studentized residuals

## Outliers, leverage \& influence

influencePlot(phd.pois, id = list( $\mathrm{n}=2$ ) )


Influence $($ CookD $)=$
$\quad$ Leverage (Hat) $\times \mid$ Residual $\mid$
Several cases (913-915) stand out with large + residuals

One observation (328) has a large leverage

Why are they unusual? Do they affect conclusions?

Examine data \& decide what to do

## Who is influential \& why?

At the very least, you should examine these flagged observations in the data

| $>$ | PhdPubs $[\mathrm{c}(328$, | 803, | $913: 915)]$, |  |  |  |
| ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| articles | female married | kid5 | phdprestige | mentor |  |  |
| 328 | 1 | 0 | 1 | 1 | 2 | 77 |
| 803 | 4 | 0 | 1 | 2 | 5 | 66 |
| 913 | 12 | 0 | 1 | 1 | 2 | 5 |
| 914 | 16 | 0 | 1 | 0 | 2 | 21 |
| 915 | 19 | 0 | 1 | 0 | 2 | 42 |

case 328: Mentor published 77 papers! Student, only 1
803: High prestige school, mentor published 66; published a bit less than predicted 913-915: Wow! all published >> than predicted

## Outlier test

- A formal test for outliers can be based on the studentized residuals, rstudent (model), using the standard normal distribution for $p$-values
- A Bonferroni correction should be applied, because interest focuses on the largest $n$ absolute residuals.

For this Poisson model, 4 observations are flagged as large + residuals

```
> outlierTest(phd.pois, cutoff = 0.001)
    rstudent unadjusted p-value Bonferroni p
914 5.54 2.99e-08 2.73e-05
913 5.38 7.36e-08 6.74e-05
911 5.21 1.92e-07 1.75e-04
915 5.15 2.60e-07 2.38e-04
```

What to do?

- Delete them \& refit?
- Keep them, but report as unusual?
- Fit a better model, hope these will go away?


## Overdispersion

- The Poisson model for counts assumes $\mathcal{V}\left(\mu_{i}\right)=\mu_{i}$, i.e., the dispersion parameter $\phi=1$
- But often, the counts exhibit greater variance than the Poisson distribution allows, $\mathcal{V}\left(\mu_{i}\right)>\mu_{i}$ or $\phi>1$
- The observations (counts) may not be independent (clustering)
- The probability of an "event" may not be constant
- There may be unmeasured influences, not accounted for in the model
- These effects are sometimes called "unmodeled heterogeneity"
- The consequences are:
- Standard errors of the coefficients, $\operatorname{se}\left(\widehat{\beta}_{j}\right)$ are optimistically small
- Wald tests, $z_{j}=\widehat{\beta}_{j} / \operatorname{se}\left(\widehat{\beta}_{j}\right)$, are too large, and thus overly liberal.


## Testing overdispersion

- Statistical tests for overdispersion test $\mathrm{H}_{0}: \operatorname{Var}(\mathrm{y})=\mu$ vs. the alternative

$$
H_{1}: \operatorname{Var}(y)=\mu+\text { P } \times f(\mu)
$$

- Implemented in AER: : dispersiontest()
- If significant, overdispersion should not be ignored
- You can try fitting a more general model
- Quasi-poisson
- Negative-binomial


## Quasi-poisson models

- The quasi-poisson model allows the dispersion, $\phi$, to be a free parameter, estimates with other coefficients
- The conditional variance is allowed to be a multiple of the mean

$$
\operatorname{Var}\left(y_{i} \mid \eta_{i}\right)=\phi \mu_{i}
$$

- This model is fit with glm() using family=quasipoisson
- The estimated coefficients $\widehat{\beta}$ are unchanged
- The standard errors are multiplied by $\phi^{1 / 2}$
- Peace, order \& good government is restored!


## Quasi-poisson models

- A simple estimate of the dispersion parameter is the residual deviance divided by degrees of freedom $\phi=D(y, \mu) / d f$
- A Pearson $\chi^{2}$ statistic has better statistical properties $\&$ is more commonly used

$$
\widehat{\phi}=\frac{X_{P}^{2}}{n-p}=\sum_{i=1}^{n} \frac{\left(y_{i}-\widehat{\mu}_{i}\right)^{2}}{\widehat{\mu}_{i}} /(n-p)
$$

For the PhdPubs data, these estimates are quite similar: about $80 \%$ overdispersion

```
> with(phd.pois, deviance/df.residual)
[1] 1.8
> sum(residuals(phd.pois, type = "pearson")^2)/phd.pois$df.residual
[1] 1.83
```


## Fitting the quasi-poisson model

You can fit the quasi-poisson model using glm()
$>$ phd. qpois <- glm(articles ~ ., data = PhdPubs, family = quasipoisson)

The estimate of the dispersion parameter is calculated by the summary() method. You can get it as follows:
> (phi <- summary(phd.qpois) \$dispersion)
[1] 1.83

This is much better than variance/mean ratio of 2.91 calculated for the marginal distribution ignoring the predictors.

Coefficients unchanged; std. errors multiplied by $\widehat{\phi}^{1 / 2}=\sqrt{1.83}=1.35$.

```
> summary(phd.qpois)
```

Call:
glm(formula = articles ~ ., family = quasipoisson, data = PhdPubs)
Deviance Residuals:

| Min | $1 Q$ | Median | $3 Q$ | Max |
| ---: | ---: | ---: | ---: | ---: |
| -3.488 | -1.538 | -0.365 | 0.577 | 5.483 |

Coefficients:

|  | Estimate | Std. Error | t value | $\operatorname{Pr}(>\|t\|)$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| (Intercept) | 0.26562 | 0.13478 | 1.97 | 0.04906 | * |
| female1 | -0.22442 | 0.07384 | -3.04 | 0.00244 | ** |
| marriedl | 0.15732 | 0.08287 | 1.90 | 0.05795 | . |
| kid5 | -0.18491 | 0.05427 | -3.41 | 0.00069 | * |
| phdprestige | 0.02538 | 0.03419 | 0.74 | 0.45815 |  |
| mentor | 0.02523 | 0.00275 | 9.19 | < 2e-16 | *** |
| Signif. code | : 0 \** | ' 0.001 '* | **' 0.01 | ** 0.05 |  |

(Dispersion parameter for quasipoisson family taken to be 1.83)

Null deviance: 1817.4 on 914 degrees of freedom
Residual deviance: 1633.6 on 909 degrees of freedom
AIC: NA

## The negative-binomial model

- The negative-binomial model is a different generalization of the Poisson that allows for over-dispersion
- Mathematically, it allows the mean $\mu \mid \boldsymbol{x}_{i}$ to vary across observations as a gamma distribution with a shape parameter $\theta$.
- The variance function, $\mathcal{V}\left(y_{i}\right)=\mu_{i}+\mu_{i}^{2} / \theta$, allows the variance of $y$ to increase more rapidly than the mean.
- Another parameterization uses $\alpha=1 / \theta$

$$
\mathcal{V}\left(y_{i}\right)=\mu_{i}+\mu_{i}^{2} / \theta=\mu_{i}+\alpha \mu_{i}^{2},
$$

- As $\alpha \rightarrow 0, \mathcal{V}\left(y_{i}\right) \rightarrow \mu_{i}$ and the negative-binomial converges to the Poisson.


## The negative-binomial model



Negative-binomial distributions for varying p \& $\theta$

Overdispersion decreases as $\theta$ increases

## Fitting the negative-binomial

- For fixed $\theta$, the negative-binomial is another special case of the GLM
- This is handled in the MASS package, with

```
family=negative.binomial(theta)
```

- But most often, $\theta$ is unknown, and must be estimated from the data
- This is implemented in glm.nb () in the MASS package.

```
> library(MASS)
> unlist(summary(phd.nbin)[c("theta", "SE.theta")])
    theta SE.theta
    2.267 0.272
```


## Visualizing goodness-of-fit

The countreg package extends rootogram() to work with fitted models:

```
countreg::rootogram(phd.pois, main="PhDPubs: Poisson")
countreg::rootogram(phd.nbin, main="PhDPubs: Negative-Binomial")
```

PhDPubs: Poisson


PhDPubs: Negative-Binomial


The Poisson model shows a systematic, wave-like pattern with excess zeros, too few observed frequencies for counts of 1--3.

## Comparing models: What difference does it make?

The NB is certainly a better fit than the Poisson; the QP cannot be distinguished by standard tests

```
> LRstats(phd.pois, phd.qpois, phd.nbin)
Likelihood summary table:
        AIC BIC LR Chisq Df Pr(>Chisq)
phd.pois 3313 3342 1634 909 <2e-16 ***
phd.qpois
phd.nbin 3135 3169 1004 909 0.015 *
Signif. codes: 0 `***' 0.001 `**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

We can also compare coefficients and their standard errors for these models

|  | pois | qpois | nbin | pois qpois | nbin |  |  |
| :--- | ---: | ---: | ---: | :--- | :--- | :--- | :--- | :--- |
| (Intercept) | 0.266 | 0.266 | 0.213 | (Intercept) | 0.0996 | 0.1348 | 0.1327 |
| female1 | -0.224 | -0.224 | -0.216 | female1 | 0.0546 | 0.0738 | 0.0726 |
| married1 | 0.157 | 0.157 | 0.153 | married1 | 0.0613 | 0.0829 | 0.0819 |
| kid5 | -0.185 | -0.185 | -0.176 | kid5 | 0.0401 | 0.0543 | 0.0528 |
| phdprestige | 0.025 | 0.025 | 0.029 | phdprestige | 0.0253 | 0.0342 | 0.0343 |
| mentor | 0.025 | 0.025 | 0.029 | mentor | 0.0020 | 0.0027 | 0.0032 |

## Visualizing the mean-variance relation

One way to see the difference among models is to plot the variance vs. mean for grouped values of the fitted linear predictor.

- The smoothed (loess) curve gives the empirical mean-variance relationship
- Also plot the theoretical mean-variance from different models
- For PhdPubs, the data is most similar to the negative-binomial
- The models differ most for those with > 3 articles


## What have we learned?

## A summary to this point should use the result of the negative-binomial model

```
> lmtest::coeftest(phd.nbin)
z test of coefficients:
    Estimate Std. Error z value Pr(>|z|)
\begin{tabular}{lrrrrr} 
(Intercept) & 0.21295 & 0.13274 & 1.60 & 0.10866 & \\
female1 & -0.21625 & 0.07259 & -2.98 & 0.00289 & \(* *\) \\
married1 & 0.15279 & 0.08194 & 1.86 & 0.06224 & . \\
kid5 & -0.17634 & 0.05279 & -3.34 & 0.00084 & \(* * *\) \\
phdprestige & 0.02934 & 0.03427 & 0.86 & 0.39192 & \\
mentor & 0.02868 & 0.00324 & 8.86 & \(<2 e-16\) & ***
\end{tabular}
```

For interpretation, examine the coefficients, $\beta$, $\mathrm{e}^{\beta}$ and $\%$ change

```
> round(cbind(beta = coef(phd.nbin),
    expbeta = exp(coef(phd.nbin)),
    pct = 100 * (exp(coef(phd.nbin)) - 1)), 3)
    beta expbeta pct
(Intercept) 0.213 1.237 23.73
female1 -0.216 0.806 -19.45
married1 0.153 1.165 16.51
kid5 -0.176 0.838 -16.17
phdprestige 0.029 1.030 2.98
mentor 0.029 1.029 2.91
```


## What have we learned?

## The number of articles published by PhD candidates:

- Most strongly predicted by mentor pubs, but with a modest effect. On average, each mentor pub increases PhD articles by 2.9\%
- Next, increasing young children (kids5) results in fewer publications. On average, each additional kid reduces PhD articles by 16\%
- Being married is marginally NS, but intriguing. Our estimate shows married candidates publish $16.5 \%$ more articles than non-married.
- Perhaps surprisingly, the prestige of the PhD institution has no significant effect in this purely main-effect model. Yet, a unit change in phdprestige is estimated as a 3\% increase in PhD articles
- Yet, we still have doubts:
- Several cases (328, 913-915) appeared unusual in diagnostic plots. Should we refit w/o them to see if conclusions change?
- The NB model might not be the best way to account for the zero counts - students who never published
- Is there a better way?


## Excess zero counts

- A common problem in count data models is that many sets of data have more observed zero counts than the (quasi) Poisson or NB models can handle.
- In the PhdPubs data, 275 of 915 (30\%) candidates published zilch, bupkis
- The expected count of 0 articles in the Poisson model is only 191 (21\%)
- Maybe there are two types of students giving zero counts:
- Those who never intend to publish (non-academic career path?)
- The rest, who do intend to publish, but have not yet done so
- This suggests the idea of zero inflation
- An alternative idea is that there is some hurdle to overcome before attaining a positive count, e.g., external pressure from the mentor.

Beyond simply identifying this as a problem of lack-of-fit, understanding the reasons for excess zero counts can contribute to a more complete explanation of the phenomenon of interest.

## Models for excess zeros

Two types of models, with different mechanisms for zero counts

- zero-inflated models: The responses with $y_{i}=0$ arise from a mixture of structural, always 0 values, with $\operatorname{Pr}\left(y_{i}=0\right)=\pi_{i}$ and the rest, which are random 0 s , with $\operatorname{Pr}\left(y_{i}=0\right)=1-\pi_{i}$
- hurdle models: One process determines whether $y_{i}=0$ with $\operatorname{Pr}\left(y_{i}=0\right)=\pi_{i}$. A second process determines the distribution of values of positive counts, $\operatorname{Pr}\left(y_{i} \mid y_{i}>0\right)$

Zero-inflated


Hurdle


## Zero-inflated models

The zero-inflated Poisson (ZIP) model has two components:

- A logistic regression model for membership in the unobserved (latent) class of those for whom $y_{i}$ is necessarily zero

$$
\operatorname{logit}\left(\pi_{i}\right)=\boldsymbol{z}_{i}^{\top} \gamma=\gamma_{0}+\gamma_{1} z_{i 1}+\gamma_{2} z_{i 2}+\cdots+\gamma_{q} z_{i q} .
$$

- A Poisson model for the other class (e.g., "publishers"), for whom $y_{i}$ may be 0 or positive.

$$
\log _{e} \mu\left(y_{i} \mid \boldsymbol{x}_{i}\right)=\boldsymbol{x}_{i}^{\top} \boldsymbol{\beta}=\beta_{0}+\beta_{1} x_{i 1}+\beta_{2} x_{i 2}+\cdots+\beta_{q} x_{i p} .
$$

In application, the same predictors can be (and often are) used in both models ( $\mathbf{x}=\mathbf{z}$ )

## Zero-inflated models

In the ZIP model, the probabilities of observing counts of $y_{i}=0$ and $y_{i}>0$ are:

$$
\begin{aligned}
\operatorname{Pr}\left(y_{i}=0 \mid \boldsymbol{x}, \boldsymbol{z}\right) & =\pi_{i} \times\left(1-\pi_{i}\right) e^{-\mu_{i}} \\
\operatorname{Pr}\left(y_{i} \mid \boldsymbol{x}, \boldsymbol{z}\right) & =\left(1-\pi_{i}\right) \times\left[\frac{\mu_{i} y_{i} e^{-\mu_{i}}}{y_{i}!}\right], \quad y_{i} \geq 0 .
\end{aligned}
$$

The conditional expectation and variance of $y_{i}$ then are:

$$
\begin{aligned}
\mathcal{E}\left(y_{i}\right) & =\left(1-\pi_{i}\right) \mu_{i} \\
\mathcal{V}\left(y_{i}\right) & =\left(1-\pi_{i}\right) \mu_{i}\left(1+\mu_{i} \pi_{i}\right) .
\end{aligned}
$$

When $\pi_{i}>0$, the mean of $y$ is always less than $\mu_{i}$; the variance of $y$ is greater than its mean by a dispersion factor of $\left(1+\mu_{i} \pi_{i}\right)$.
The model for the count variable could also be negative-binomial, giving a zero-inflated negative-binomial (ZINB) model using NBin $(\mu, \theta)$

## Exploring zero-inflated data

A little insight can be gained by generating random data from Poisson \& zero-inflated analog. The example uses VGAM::rzipois()
$\operatorname{Pois}(\mu=3)=\operatorname{ZIP}(\mu=3, \pi=0)$
vs. $\quad \operatorname{ZIP}(\mu=3, \pi=.3)$
$>$ set.seed (1234)
> datal <- VGAM::rzipois (200, 3, 0)
> data 2 <- VGAM::rzipois(200, 3, .3)

The tables of counts show far more zeros in data2

```
> table(datal)
data1
\begin{tabular}{rrrrrrrrrr}
0 & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 \\
10 & 31 & 46 & 54 & 24 & 20 & 7 & 3 & 4 & 1
\end{tabular}
> table(data2)
data2
\begin{tabular}{rrrrrrrrr}
0 & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 9 \\
62 & 26 & 33 & 31 & 22 & 9 & 8 & 8 & 1
\end{tabular}
```


## Exploring zero-inflated data

Bar plots of the counts:


The 30\% extra zeros decrease the mean and inflate the variance

## Hurdle models

The Hurdle model has also has two components:

- A logistic regression model, for the probability that $y_{i}=0$ vs. $y_{i}>0$

$$
\operatorname{logit}\left[\frac{\operatorname{Pr}\left(y_{i}=0\right)}{\operatorname{Pr}\left(y_{i}>0\right)}\right]=\boldsymbol{z}_{i}^{\top} \gamma=\gamma_{0}+\gamma_{1} z_{i 1}+\gamma_{2} z_{i 2}+\cdots+\gamma_{q} z_{i q} .
$$

- A model for the positive counts, taken as a left-truncated Poisson or negative-binomial, excluding the zero counts
- Comparing the ZIP and Hurdle models:
- In ZIP models, the first (latent) process generates extra zeros (with probability $\pi_{i}$ ).
- In Hurdle models, $y_{i}=0$ and $y_{i}>0$ are fully observed. The first process generates all the zeros.


## Fitting ZIP \& Hurdle models

In R, these models can be fit using the pscl and countreg packages.
countreg is more mature, but is only available on R-Forge, not on CRAN. Use:
install.packages("countreg", repos="http://R-Forge.R-project.org")
The functions have the following arguments:

```
zeroinfl(formula, data, subset, na.action, weights, offset,
    dist = c("poisson", "negbin", "geometric", "binomial"),
        ...)
hurdle(formula, data, subset, na.action, weights, offset,
    dist = c("poisson", "negbin", "geometric", "binomial"),
        ...)
```

The formula, $y \sim x 1+x 2+\ldots$ uses the same predictors for both models.
Using $y \sim x 1+x 2+\ldots \mid z 1+z 2+\ldots$ allows separate predictors for the 0 submodel.

## Visualizing zero counts

It is often useful to plot the data for the binary distinction between $y_{i}=0$ vs. $y_{i}>0$ as in logistic regression models.

```
plot(factor(articles==0) ~ mentor, data=PhdPubs,
ylevels=1:2, ylab="Zero articles",
breaks=quantile(mentor, probs=seq(0,1,.2)))
```



As expected, zero counts decrease with mentor pubs

NB: this gives a spineplot

## Fitting models

To illustrate, we fit all four models, the combinations of (ZI, hurdle) $\times$ (poisson, nbin) to the phdpubs data.

For simplicity, we use all predictors for both the zero model and the non-zero model.

```
phd.zip <- zeroinfl(articles ~ ., data=PhdPubs, dist="poisson")
phd.znb <- zeroinfl(articles ~ ., data=PhdPubs, dist="negbin")
phd.hp <- hurdle(articles ~ ., data=PhdPubs, dist="poisson")
phd.hnb <- hurdle(articles ~ ., data=PhdPubs, dist="negbin")
```


## Comparing models

Compare the models, sorting by BIC

```
> LRstats(phd.pois, phd.nbin, phd.zip, phd.znb, phd.hp, phd.hnb,
    sortby="BIC")
Likelihood summary table:
    AIC BIC LR Chisq Df Pr(>Chisq)
phd.pois 3313.3 3342.3 3301.3 909< < 2.2e-16 ***
phd.hp 3234.5 3292.4 3210.5 903< 2.2e-16 ***
phd.zip 3233.5 3291.3 3209.5 903< 2.2e-16 ***
phd.hnb 3130.9 3193.5 3104.9 902 < 2.2e-16 ***
phd.znb 3125.8 3188.4 3099.8 902< < . 2e-16 ***
phd.nbin 3135.4 3169.1 3121.4 909 < 2.2e-16 ***
Signif. codes: 0 `***'0.001 `**' 0.01 `*' 0.05 '.' 0.1 ' ' 1
```

The standard negative binomial model looks best by BIC. Why do you think this is? (Hint: look at the residual df)

## Nevertheless, it is useful to examine the coefficients in the ZIP model

```
> lmtest::coeftest(phd.zip)
t test of coefficients:
Estimate Std. Error t value Pr(>|t|)
count_(Intercept) 0.59918 0.11861 5.05 5.3e-07 ***
count_female1 -0.20879 0.06353 -3.29 0.0011 **
count_married1 0.10623 0.07097 1.50 0.1348
count_kid5 -0.14271 0.04744 -3.01 0.0027 **
count phdprestige 0.00700 0.02981 0.23 0.8145
count_mentor 0.01785 0.00233 7.65 5.3e-14 ***
zero_(Intercept) -0.56332 0.49405 -1.14 0.2545
zero_female1 0.10816 0.28173 0.38
zero married1 -0.35558 0.31796 -1.12 0.2637
zero_kid5 0.21974 0.19658 1.12 0.2639
zero_phdprestige -0.00537 0.14118 -0.04 0.9697
zero_mentor -0.13313 0.04643 -2.87 0.0042 **
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 '.' 0.1 ' ' 1
```

Only mentor is significant in the ZIP model

## Let's refit the ZIP and ZNB models using only mentor for the zero models

```
phd.zipl <- zeroinfl(articles ~ .| mentor, data=PhdPubs, dist="poisson")
phd.znbl <- zeroinfl(articles ~ .| mentor, data=PhdPubs, dist="negbin")
```


## Compare models again

```
> LRstats(phd.pois, phd.nbin, phd.zip, phd.znb, phd.hp, phd.hnb,
+ phd.zip1, phd.znb1, sortby="BIC")
Likelihood summary table:
        AIC BIC LR Chisq Df Pr(>Chisq)
phd.pois 3313 3342 3301 909 <2e-16 ***
phd.hp 3235 3292 3211 903 <2e-16 ***
phd.zip 3234 3291 3210 903 <2e-16 ***
phd.zip1 3227 3266 3211 907 <2e-16 ***
phd.hnb 3131 3194 3105 902 <2e-16 ***
phd.znb 3126 3188 3100 902 <2e-16 ***
phd.nbin 3135 3169 3121 909 <2e-16 ***
phd.znb1 3124 3168 3106 906 <2e-16 ***
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ' ' 1
```

Now, the phd. zn.b1 model looks best by BIC. Let's stick with this.

## Model interpretation: Coefficients

Ignoring the NS coefficients in the revised ZNB model (phd. znb1):

```
> coef(phd.znb1)[c(1,2,4,6,7,8)]
count_(Intercept) 
```

- Count model:

$$
\log (\text { articles })=0.357-0.21 \text { female }-0.17 \text { kids } 5+0.024 \text { mentor }
$$

- Zero model:

$$
\operatorname{logit}(\text { articles }=0)=-0.817-0.608 \text { mentor }
$$

Can you describe these in words?

## Model interpretation: Coefficients

Often easier to interpret $\exp (\beta)$

```
> exp(coef(phd.znb1) [c(1,2,4,6,7,8)])
count_(Intercept) count_female1 rount_kid5 % count_mentor
zero_(Intercept) zero_mentor
    0.442 0.544
```

Female: Women publish . 21 fewer log articles, .81 times that of men ( $20 \%$ decrease) Kids5: Each additional kid<5 $\rightarrow .17$ fewer log articles, a 15\% decrease Mentor: Each additional mentor article $\rightarrow .024$ more PhD log pubs (2.4\% increase)

Count model: Each additional mentor article decreases log odds PhDpubs $=0$ by 0.608, a 45\% decrease

## Model interpretation: Effect plots

- The effects package cannot yet handle zero-inflated or hurdle models.
- But the fitted values don't differ very much among these models
- Here, I use the phd. nbin model, and just show the effects for the important terms

```
plot(allEffects(phd.nbin)[c(1,3,5)], rows=1, cols=3)
```

female effect plot

kid5 effect plot

mentor effect plot


The ZIP sub-model for the zero counts ("did not publish") can also be interpreted visually

- As an approximation, fit a separate logistic model for articles==0
- The effect plot for that gives an interpretation of the zero model.

```
phd.zero <- glm((articles==0) ~ mentor, data=PhdPubs, family=binomial)
plot(allEffects(phd.zero), main="Mentor effect on not publishing")
```

Mentor effect on not publishing


## What have we learned?

- The simple Poisson regression model fits very badly
- Standard errors do not reflect overdispersion
- Inference about model effects is compromised by overly liberal tests
- The quasi-poisson model corrects for overdispersion.
- But doesn't account for excess 0s
- The negative-binomial model provides valid tests and fits the 0 counts well.
- But it doesn't provide any insight into why there are so many 0s
- The ZIP and ZNB models fit well, and account for the Os.
- But they lose here on BIC (and AIC) measures, because they have $2 \times$ the number of parameters.
- For simplicity, I have slighted the analogous hurdle models


## What have we learned?

- The revised ZNB model (phd. znb1), with only mentor predicting 0s, wins on parsimony, and has a simple interpretation.
- The log odds that a student does not publish decrease by 0.61 for every article published by the mentor
- Each mentor pub increases student publications by about 2.5\%
- $\Rightarrow$ Encourage or help your supervisor to publish!
- (Or, choose a high publishing one.)
- For this data set, the main substantive interpretation and predicted effects are similar across models. But details matter!
- In data sets where there are substantive reasons for excess 0s, the ZI and hurdle models provide different explanations.
- It is not always just a matter of model fit!
- Hurdle models make the distinction between 0 and $>0$ more explicit
- In ZI models, the interpretation of the mean count is clearer.


## What have we forgotten?

"All models are wrong, but some are useful" --- GEP Box

- Model building and model criticism go hand in hand
- But they don't form a linear series of steps you can put into a flow chart



## What have we forgotten?

- Sometimes, you have to go back and revisit decisions made earlier:

$$
\text { Re-think } \rightarrow \text { Re-fit } \rightarrow \text { Re-interpret }
$$



## What I missed

- In the initial model, phdprestige was NS. I decided to keep it
- In the check for two-way interactions, the interaction
phdprestige:mentor was borderline ( $p=0.051$ )
- I did a global test for all interactions together
- This was NS $(p=0.08)$, so I decided to dismiss them all
- (I wanted to keep he model simple, to go on to other topics: overdispersion, models for excess zeros)


## Back to square TWO

- A question in a former class made me reconsider the phdprestige:mentor interaction
- Perhaps, the effect of mentor varied with phdprestige?

Try this, starting with the negative-binomial, phd.nbin (update () is your friend)

```
> phd.nbin2 <- update(phd.nbin, . ~ . + phdprestige:mentor)
> Anova(phd.nbin2)
Analysis of Deviance Table (Type II tests)
Response: articles
            LR Chisq Df Pr(>Chisq)
female 9.1 1 0.0026 **
married
    3.1 1 0.0762.
kid5 10.7 1 0.0011 **
phdprestige
    0.7 1 0.3921
mentor
    72.8 1 <2e-16 ***
phdprestige:mentor
    5.6 1 0.0179 *
---
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 '.' 0.1 ' ' 1
```


## Visualize the interaction

phd.effnb2 <- allEffects(phd.nbin2)
plot(phd.effnb2[4], x.var="mentor", multiline=TRUE, ci.style="bands", ...)
phdprestige*mentor effect plot


- An effect plot for phdprestige*mentor shows the average over other predictors
- This plot, with mentor on the X-axis shows that the slope for mentor increases with higher prestige of the student's university


## Visualize the interaction- The other way

phd.effnb2 <- allEffects(phd.nbin2)
plot(phd.effnb2[4], multiline=TRUE, ci.style="bands", ...)
phdprestige*mentor effect plot


- This plot, with phdprestige on the X -axis shows that the slopes change sign depending on the value of mentor.
- It explains why the main effect of phdprestige is near 0 .
- The widths of the confidence bands indicate model uncertainty- they get wider as mentor pubs increase, and phdprestige differs from average.


## Back to square ONE

Aren't we done yet?
"All data are wrong, but some are useful - Sitsofe Tsagbey et al. TAS, 2017

- A nagging doubt: what is the coding for phdprestige?
- Email from Scott Long: "the higher the number, the more prestigious the program"
- "PS: The data I used did not categorize the continuous phd scale into discrete categories"
- Found the original Stata data set:

```
library(foreign)
PhdPubs2 <-
    read.dta("http://www.stata-press.com/data/lf2/couart2.dta")
```


## Compare distributions

Histograms with smoothed density estimate of the two versions of phdprestige They are very different!



## What to do?

Re-run the analysis with the new data set, PhdPubs2

- This could be called a sensitivity analysis - does the new data alter conclusions?
- Q: Are the results of the phd.nbin2 and phd.znb2 models about the same. A: YES!
- Q : Is the interaction of phdprestige:mentor about the same. A: YES!
- Q: Does the effect plot look about the same? A: YES!


## What else is there?

The PhdPubs example was rather simple

- There were only a few predictors
- Model selection methods could be based on simple Anova(), coeftest(), LRstats()
- No need for more complex model selection methods or crossvalidation
- Of the quantitative predictors, only mentor \& kid5 had important effects
- The effects of these were sufficiently linear
- No need to try non-linear effects (poly(mentor,2), ns(mentor,2))
- There turned out to be one important interaction
- In Psychology, these are called "moderator" effects
- Interpretation often based on post-hoc tests of simple slopes
- Interpretation is usually simplified in effect plots


## Other methods: Recursive partitioning

- Recursive partitioning, or regression trees are often an attractive alternative to linear models
- Interactions are handled by partitioning the ranges of variables
- Or, models can be fit to subsets of the data defined by recursive partitioning


Logistic regression tree fit to the Titanic data with partykit::glmtree()

## Other methods: Recursive partitioning

Could there be a simpler or different model for the PhdPubs data?

```
library(partykit)
phd.tree <- glmtree(articles ~ mentor| female+married+kid5+phdprestige,
                                    data=PhdPubs, family=poisson)
plot(phd.tree)
```



## Summary

- GLMs introduce a wide class of models for count data, starting from $\log (\boldsymbol{\mu})=\mathbf{X} \boldsymbol{\beta}, \boldsymbol{\mu} \mid \mathbf{X} \sim$ Poisson
- Overdispersion $\rightarrow$ quasi-poisson, negative binomial
- Excess zero counts introduce new ideas \& methods
- ZIP model: structural model for the Os
- Hurdle model: random model for $0 s, 2^{\text {nd }}$ model for $Y>0$
- In all this, we rely on data \& model plots for understanding

