432 Class 16 Slides

github.com/THOMASELOVE/2020-432

2020-03-24

Today's Topic

Regression Models for Count Outcomes

- Six modeling approaches are illustrated in these slides.
 - Poisson Regression
 - Negative Binomial Regression
 - Two types of Zero-inflated model
 - ZIP (Zero-inflated Poisson)
 - ZINB (Zero-inflated Neg. Binomial)
 - Two types of Hurdle model
 - using a Poisson approach
 - using a Negative Binomial approach

Chapter 19 of the Course Notes describes this material.

Setup

We've previously installed the countreg package from R-Forge.

```
library(magrittr); library(here); library(janitor)
library(knitr)
library(caret)
library(MASS)
library(pscl)
library(VGAM)
library(broom)
library(tidyverse)
```

An Overview

Generalized Linear Models for Count Outcomes

We want to build a generalized linear model to predict count data using one or more predictors.

In count data, the observations are non-negative integers (0, 1, 2, 3, ...)

- the number of COVID-19 hospitalizations in Ohio yesterday
- the number of mutations within a particular search grid
- the number of days in the past 30 where your mental health was poor

The Poisson and the Negative Binomial probability distributions will be useful.

The Poisson Probability Distribution

The Poisson probability model describes the probability of a given number of events occurring in a fixed interval of time or space.

- If events occur with a constant mean rate, and independently of the time since the last event, the Poisson model is appropriate.
- The probability mass function for a discrete random variable with Poisson distribution follows.

$$Pr(Y = k) = \frac{\lambda^k e^{-\lambda}}{k!}$$

- *k* is the number of times an event occurs in an interval, and *k* can take the values 0, 1, 2, 3, ...
- The parameter λ (lambda) is equal to the expected value (mean) of Y and is also equal to the variance of Y.

The Negative Binomial Probability Distribution

The Negative Binomial distribution models the number of failures in a sequence of independent and identically distributed Bernoulli trials before a specified number of successes occurs.

 The probability mass function for a discrete random variable with a negative binomial distribution follows.

$$Pr(Y = k) = \binom{k+r-1}{k} p^r (1-p)^k$$

- k is the number of failures (units of time) before the rth event occurs, and k can take the values 0, 1, 2, 3, ...
- The mean of the random variable Y which follows a negative binomial distribution is rp/(1-p) and the variance is $rp/(1-p)^2$.

Poisson Regression and the possibility of overdispersion

- Poisson regression assumes that the outcome Y follows a Poisson distribution, and that the logarithm of the expected value of Y (its mean) can be modeled by a linear combination of a set of predictors.
 - A Poisson regression makes the strong assumption that the variance of Y is equal to its mean.
 - A Poisson model might fit poorly due to overdispersion, where the variance of Y is larger than we'd expect based on the mean of Y.
 - Quasipoisson models are available which estimate an overdispersion parameter, but we'll skip those.

We will show the use of glm to fit Poisson models, by using family = "Poisson".

Negative Binomial Regression to generalize the Poisson

• Negative binomial regression is a generalization of Poisson regression which loosens the assumption that the variance of Y is equal to its mean, and thus produces models which fit a broader class of data.

We will demonstrate the use of glm.nb from the MASS package to fit negative binomial regression models.

Zero-inflated approaches

- Both the Poisson and Negative Binomial regression approaches may under-estimate the number of zeros compared to the data.
- To better match up the counts of zero, zero-inflated models fit:
 - a logistic regression to predict the extra zeros, along with
 - a Poisson or Negative Binomial model to predict the counts, including some zeros.

We will demonstrate the use of zeroinfl from the pscl package to fit zero-inflated Poisson (or ZIP) and zero-inflated negative binomial (or ZINB) regressions.

Hurdle models

A hurdle model predicts the count outcome by making an assumption that there are two processes at work:

- a process that determines whether the count is zero or not zero (usually using logistic regression), and
- a process that determines the count when we know the subject has a positive count (usually using a truncated Poisson or Negative Binomial model where no zeros are predicted)

We'll use the hurdle function from the pscl package to fit these models.

Comparing Models

- A key tool will be a graphical representation of the fit of the models to the count outcome, called a rootogram. We'll use the rootograms produced by the countreg package to help us.
- We'll also demonstrate a Vuong hypothesis testing approach (from the lmtest package) to help us make decisions between various types of Poisson models or various types of Negative Binomial models on the basis of improvement in fit of things like bias-corrected AIC or BIC.
- We'll also demonstrate the calculation of pseudo-R square statistics for comparing models, which can be compared in a validation sample as well as in the original modeling sample.

The medicare data

The medicare example

The data we will use come from the NMES1988 data set in R's AER package, although I have built a cleaner version for you in the medicare.csv file on our web site. These are essentially the same data as are used in my main resource from the University of Virginia for hurdle models.

These data are a cross-section originating from the US National Medical Expenditure Survey (NMES) conducted in 1987 and 1988. The NMES is based upon a representative, national probability sample of the civilian non-institutionalized population and individuals admitted to long-term care facilities during 1987. The data are a subsample of individuals ages 66 and over all of whom are covered by Medicare (a public insurance program providing substantial protection against health-care costs), and some of whom also have private supplemental insurance.

medicare <- read.csv(here("data/medicare.csv")) %>% tbl_df

The medicare code book

Variable	Description
subject	subject number (code)
visits	outcome of interest: number of physician office visits
hospital	number of hospital stays
health	self-perceived health status (poor, average, excellent)
chronic	number of chronic conditions
sex	male or female
school	number of years of education
insurance	is the subject (also) covered by private insurance? (yes or no)

Today's Goal

Predict visits using main effects of the 6 predictors (excluding subject)

Skimming the medicare tibble

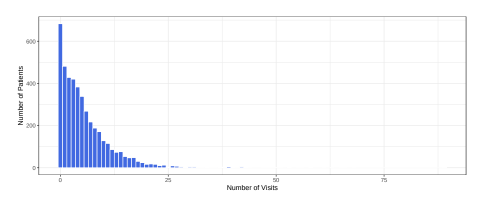
```
skimr::skim(medicare)
-- Data Summary -----
                                                     Values
                                                     medicare
Name
Number of rows
                                                    4406
Number of columns
Column type frequency:
  factor
  numeric
Group variables
                                                     None
-- Variable type: factor -----
# A tibble: 3 x 6
  skim variable n missing complete rate ordered n unique top counts

    x chr>
    x of the late

    x chr>
    x of the late

    x of the late
    x of the late</
 health
3 insurance
-- Variable type: numeric ------
# A tibble: 5 x 11
  skim variable n_missing_complete_rate mean sd_ p0 p25 p50 p75 p100 hist
  0 1 2204. 1272. 1 1102. 2204. 3305. 4406
 . subiect
                                                                    1 5.77 6.76 0 1 4 8 89
1 0.296 0.746 0 0 0 0 8
1 1.54 1.35 0 1 1 2 8
1 10.3 3.74 0 8 11 12 18
2 visits
                                         0 1 5.77
  hospital
  chronic
                                            0
  school
```

Our outcome, visits



```
mosaic::favstats(~ visits, data = medicare)
```

```
min Q1 median Q3 max mean sd n missing 0 1 4 8 89 5.774399 6.759225 4406 0
```

visits numerical summaries

```
medicare %$% Hmisc::describe(visits)
visits
        missing distinct
                        Info
                                 Mean
                                          Gmd
                    60 0.992 5.774 6.227
   4406
    . 05
        .10 .25
                           .50 .75
                                          .90
                                           13
    . 95
     17
lowest: 0 1 2 3 4, highest: 63 65 66 68 89
```

Reiterating the Goal

Predict visits using some combination of these 6 predictors...

Predictor	Description
hospital	number of hospital stays
health	self-perceived health status (poor, average, excellent)
chronic	number of chronic conditions
sex	male or female
school	number of years of education
insurance	is the subject (also) covered by private insurance? (yes or no)

We'll build separate training and test samples to help us validate.

Partitioning the Data into Training vs. Test Samples

```
set.seed(432)
validation samples <- medicare$visits %>%
  createDataPartition(p = 0.75, list = FALSE)
med train = medicare[validation samples,]
med test = medicare[-validation samples,]
I've held out 25% of the medicare data for the test sample.
dim(med_train)
[1] 3306
dim(med test)
[1] 1100
```

mod_1: A Poisson Regression

Poisson Regression

Assume our count data (visits) follows a Poisson distribution with a mean conditional on our predictors.

The Poisson model uses a logarithm as its link function, so the model is actually predicting log(visits).

Note that we're fitting the model here using the training sample alone.

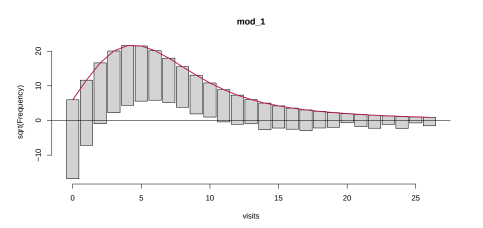
mod_1 (Poisson) model coefficients

term	estimate	std.error	statistic	p.value
(Intercept)	0.990	0.028	35.8	0
hospital	0.165	0.007	24.7	0
healthexcellent	-0.384	0.035	-11.0	0
healthpoor	0.290	0.021	14.1	0
chronic	0.143	0.005	27.2	0
sexmale	-0.085	0.015	-5.7	0
school	0.032	0.002	15.2	0
insuranceyes	0.153	0.019	7.9	0

If Harry and Larry have the same values for all other predictors but only Harry has private insurance, the model predicts Harry to have a 0.153 point larger value of log(visits) than Larry.

Visualize fit with a (Hanging) Rootogram

countreg::rootogram(mod_1)



See the next slide for details on how to interpret this. . .

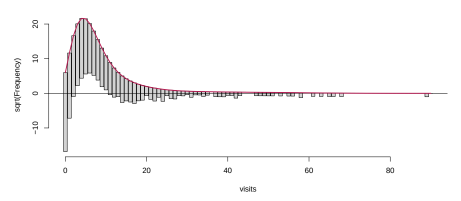
Interpreting the Rootogram

- The red curved line is the theoretical Poisson fit.
- "Hanging" from each point on the red line is a bar, the height of which represents the observed counts.
 - A bar hanging below 0 indicates that the model under-predicts that value. (Model predicts fewer values than the data show.)
 - A bar hanging above 0 indicates over-prediction of that value. (Model predicts more values than the data show.)
- The counts have been transformed with a square root transformation to prevent smaller counts from getting obscured and overwhelmed by larger counts.

For more information on rootograms, check out https://arxiv.org/pdf/1605.01311.

The Complete Rootogram for mod_1

Rootogram for Poisson mod_1



This shows what happens with the subject with 89 visits.

Interpreting the Rootogram for mod_1

In mod_1, we see a great deal of underfitting for counts of 0 and 1, then overfitting for visit counts in the 3-10 range, with some underfitting again at more than a dozen or so visits.

• Our Poisson model (mod_1) doesn't fit enough zeros or ones, and fits too many 3-12 values, then not enough of the higher values.

Store Training Sample mod_1 Predictions

We'll use the augment function to store the predictions within our training sample. Note the use of "response" to predict visits, not log(visits).

Summarizing Training Sample mod_1 Fit

Within our training sample, mod_1_aug now contains both the actual counts (visits) and the predicted counts (in .fitted) from mod_1. We'll summarize the fit...

```
mod_1_summary <- tibble(
  model = "mod_1 (Poisson)",
  R2 = R2(mod_1_aug$.fitted, mod_1_aug$visits),
  RMSE = RMSE(mod_1_aug$.fitted, mod_1_aug$visits),
  MAE = MAE(mod_1_aug$.fitted, mod_1_aug$visits))

mod_1_summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_1 (Poisson)	0.102	6.522	4.124

These will become interesting as we build additional models.

mod_2: A Negative Binomial Regression

Fitting the Negative Binomial Model

The negative binomial model requires the estimation of an additional parameter, called θ (theta). The default link for this generalized linear model is also a logarithm, like the Poisson.

The estimated dispersion parameter value θ is. . .

```
summary(mod_2)$theta
```

```
[1] 1.212527
```

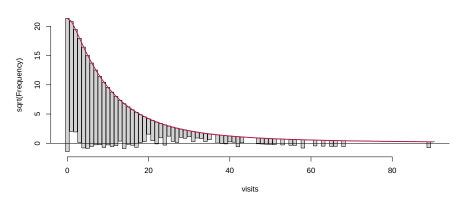
The Poisson model is essentially the negative binomial model assuming a known $\theta=1$.

mod_2 (Negative Binomial) coefficients

term	estimate	std.error	statistic	p.value
(Intercept)	0.873	0.064	13.7	0.000
hospital	0.220	0.023	9.6	0.000
healthexcellent	-0.369	0.070	-5.3	0.000
healthpoor	0.341	0.056	6.1	0.000
chronic	0.176	0.014	12.7	0.000
sexmale	-0.103	0.036	-2.9	0.004
school	0.033	0.005	6.4	0.000
insuranceyes	0.196	0.046	4.3	0.000

Rootogram for Negative Binomial Model

Rootogram for mod_2



Does this look better than the Poisson rootogram?

Store Training Sample mod_2 Predictions

Summarizing Training Sample mod_2 Fit

As before, mod_2_aug now has actual (visits) and predicted counts (in .fitted) from mod_2.

```
mod_2_summary <- tibble(
  model = "mod_2 (Neg. Binomial)",
  R2 = R2(mod_2_aug$.fitted, mod_2_aug$visits),
  RMSE = RMSE(mod_2_aug$.fitted, mod_2_aug$visits),
  MAE = MAE(mod_2_aug$.fitted, mod_2_aug$visits))

mod_2_summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_2 (Neg. Binomial)	0.084	6.834	4.179

So Far in our Training Sample

The reasonable things to summarize in sample look like the impressions from the rootograms and the summaries we've prepared so far.

```
bind_rows(mod_1_summary, mod_2_summary) %>%
  kable(digits = 3)
```

model	R2	RMSE	MAE
mod_1 (Poisson)	0.102	6.522	4.124
mod_2 (Neg. Binomial)	0.084	6.834	4.179

Model	Rootogram impressions
mod_1	Many problems. Data appear overdispersed.
mod_2	Still not enough zeros; some big predictions.

mod_3: Zero-Inflated Poisson (ZIP) Model

Zero-Inflated Poisson (ZIP) model

The zero-inflated Poisson model describes count data with an excess of zero counts.

The model posits that there are two processes involved:

- a logistic regression model is used to predict excess zeros
- while a Poisson model is used to predict the counts

We'll use the pscl package to fit zero-inflated models.

mod_3 ZIP coefficients

Sadly, there's no broom tidying functions for these zero-inflated models.

summary(mod_3)

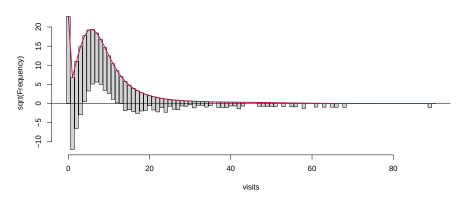
Screenshot on next slide...

```
Call:
  pscl::zeroinfl(formula = visits ~ hospital + health +
     chronic + sex + school + insurance, data = med train)
  Pearson residuals:
     Min
              10 Median
                          30
                                   Max
  -5.3815 -1.1514 -0.4617 0.5647 24.8808
 Count model coefficients (poisson with log link):
                 Estimate Std. Error z value Pr(>|z|)
  (Intercept)
                 1.382706 0.028070 49.258
                                              <2e-16 ***
 hospital
                 0.159542 0.006780 23.533
                                              <2e-16 ***
 healthexcellent -0.307477 0.036003 -8.540
                                              <2e-16 ***
 healthpoor
             0.289720
                            0.020457 14.162
                                              <2e-16 ***
 chronic
                 0.096085 0.005388 17.832
                                              <2e-16 ***
 sexmale
                 -0.038362
                            0.014985 -2.560
                                              0.0105 *
 school
               0.023862 0.002155 11.072
                                              <2e-16 ***
  insuranceyes
                 0.041155
                            0.019668 2.092
                                              0.0364 *
  Zero-inflation model coefficients (binomial with logit link):
                Estimate Std. Error z value Pr(>|z|)
  (Intercept) 0.05980
                            0.16579 0.361 0.718304
                            0.11068 -3.205 0.001349 **
 hospital
            -0.35477
 healthexcellent 0.33687
                            0.16798 2.005 0.044916 *
                -0.05106
                            0.18994 -0.269 0.788064
 healthpoor
  chronic
                -0.53042
                            0.05272 - 10.061 < 2e - 16 ***
 sexmale
               0.38577
                            0.10289 3.749 0.000177 ***
 school
                -0.06615
                            0.01421 -4.655 3.23e-06 ***
                            0.11909 -6.395 1.60e-10 ***
  insuranceyes
                -0.76161
 Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
  Number of iterations in BFGS optimization: 20
 Log-likelihood: -1.21e+04 on 16 Df
github.com/THOMASELOVE/2020-432
                                      432 Class 16 Slides
```

> summarv(mod 3)

Rootogram for ZIP model

ZIP model Rootogram: mod_3

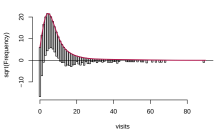


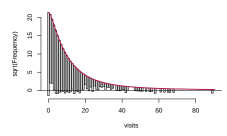
What do you think? Next slide shows all models so far.

First Three Rootograms - Which Looks Best?

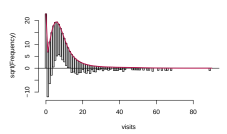


Negative Binomial mod_2





ZIP mod_3



Store Training Sample mod_3 Predictions

We have no augment or other broom functions available for zero-inflated models, so . . .

Summarizing Training Sample mod_3 Fit

mod_3_aug now has actual (visits) and predicted counts (in .fitted) from mod_3, just as we set up for the previous two models.

```
mod_3_summary <- tibble(
  model = "mod_3 (ZIP)",
  R2 = R2(mod_3_aug$.fitted, mod_3_aug$visits),
  RMSE = RMSE(mod_3_aug$.fitted, mod_3_aug$visits),
  MAE = MAE(mod_3_aug$.fitted, mod_3_aug$visits))

mod_3_summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_3 (ZIP)	0.113	6.481	4.093

Training Sample Results through mod_3

```
bind_rows(mod_1_summary, mod_2_summary, mod_3_summary) %>%
  kable(digits = 3)
```

model	R2	RMSE	MAE
mod_1 (Poisson) mod_2 (Neg. Binomial) mod_3 (ZIP)	0.084	6.522 6.834 6.481	4.179

Remember we want a larger R^2 and smaller values of RMSE and MAE.

Comparing models with Vuong's procedure

Vuong's test compares predicted probabilities (for each count) in two non-nested models. How about Poisson vs. ZIP?

```
vuong(mod_1, mod_3)
```

Vuong Non-Nested Hypothesis Test-Statistic: (test-statistic is asymptotically distributed N(0,1) under the null that the models are indistinguishible)

	Vuong z-statistic	H_A	p-value
Raw	-14.93727 model2	> model1	< 2.22e-16
AIC-corrected	-14.85194 model2	> model1	< 2.22e-16
BIC-corrected	-14.59155 model2	> model1	< 2.22e-16

The large negative z-statistic indicates mod_3 (ZIP) fits detectably better than mod_1 (Poisson) in our training sample.

Reference: Vuong, QH (1989) Likelihood ratio tests for model selection and non-nested hypotheses. *Econometrica*, 57:307-333.

mod_4: Zero-Inflated Negative Binomial (ZINB) Model

Zero-Inflated Negative Binomial (ZINB) model

As in the ZIP, we assume there are two processes involved:

- a logistic regression model is used to predict excess zeros
- while a negative binomial model is used to predict the counts

We'll use the pscl package again and the zeroinfl function.

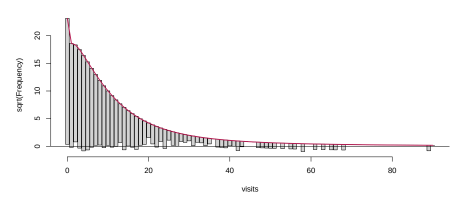
summary(mod_4) results on next slide...

```
Call:
zeroinfl(formula = visits ~ hospital + health + chronic + sex +
school + insurance, data = med_train, dist = "negbin")
Pearson residuals:
            10 Median
   Min
                            30
                                   Max
-1.2029 -0.7074 -0.2836 0.3333 17.9865
Count model coefficients (neabin with log link):
                Estimate Std. Error z value Pr(>|z|)
(Intercept)
                1.158032
                           0.066360 17.451 < 2e-16 ***
hospital
                0.197864
                           0.022812
                                     8.674 < 2e-16 ***
healthexcellent -0.319314
                           0.072892 -4.381 1.18e-05 ***
                           0.053100 6.096 1.09e-09 ***
healthpoor
                0.323694
                                    9.397 < 2e-16 ***
chronic
                0.128298
                           0.013653
                           0.035702 -1.612
sexmale
               -0.057554
                                             0.1069
school
                0.026456
                           0.005064 5.224 1.75e-07 ***
insuranceyes
                0.088092
                           0.049034 1.797
                                             0.0724
Log(theta)
                0.406043
                           0.040469 10.033 < 2e-16 ***
Zero-inflation model coefficients (binomial with logit link):
               Estimate Std. Error z value Pr(>|z|)
(Intercept)
                0.11575
                           0.31399
                                   0.369 0.71240
hospital
               -1.90282
                          1.46412 -1.300 0.19373
healthexcellent 0.30081
                           0.33370
                                   0.901
                                           0.36736
healthpoor
                0.08589
                           0.50858
                                   0.169
                                          0.86589
chronic
               -1.19776
                           0.19684 -6.085 1.16e-09 ***
sexmale
                0.64754
                           0.22609
                                   2.864 0.00418 **
school
               -0.09115
                           0.03149 -2.895 0.00379 **
                           0.27484 -4.542 5.57e-06 ***
              -1.24829
insuranceves
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Theta = 1.5009
Number of iterations in BFGS optimization: 30
Log-likelihood: -9057 on 17 Df
                                         432 Class 16 Slides
```

summary(mod_4)

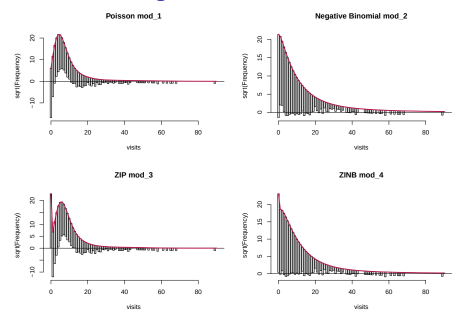
Rootogram for ZIP model

ZINB model Rootogram: mod_4



Again, next slide shows all models so far.

First Four Rootograms - Which Looks Best?



Store Training Sample mod_4 Predictions

Again, there is no augment or other broom functions available for zero-inflated models, so ...

Summarizing Training Sample mod_4 Fit

mod_4_aug now has actual (visits) and predicted counts (in .fitted)
from mod_4.

```
mod_4_summary <- tibble(
  model = "mod_4 (ZINB)",
  R2 = R2(mod_4_aug$.fitted, mod_4_aug$visits),
  RMSE = RMSE(mod_4_aug$.fitted, mod_4_aug$visits),
  MAE = MAE(mod_4_aug$.fitted, mod_4_aug$visits))

mod_4_summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_4 (ZINB)	0.101	6.592	4.111

Training Sample Results through mod_4

model	R2	RMSE	MAE
mod_1 (Poisson)	0.102	6.522	4.124
mod_2 (Neg. Binomial)	0.084	6.834	4.179
mod_3 (ZIP)	0.113	6.481	4.093
mod_4 (ZINB)	0.101	6.592	4.111

What do you think?

Comparing models with Vuong's procedure

Vuong's test compares predicted probabilities (for each count) in two non-nested models. How about Negative Binomial vs. ZINB?

```
vuong(mod_4, mod_2)
```

```
Vuong Non-Nested Hypothesis Test-Statistic:
(test-statistic is asymptotically distributed N(0,1) under the
null that the models are indistinguishible)
```

						_
	Vuong z-statistic			H_A	p-value	
Raw	5.414748	model1	>	model2	3.0688e-08	
AIC-corrected	4.777133	model1	>	model2	8.8906e-07	
BIC-corrected	2.831291	model1	>	model2	0.002318	

The large positive z-statistics indicate mod_4 (ZINB) fits detectably better than mod_2 (Negative Binomial) in our training sample.

mod_5: Poisson-Logistic Hurdle Model

The Hurdle Model

The hurdle model is a two-part model that specifies one process for zero counts and another process for positive counts. The idea is that positive counts occur once a threshold is crossed, or put another way, a hurdle is cleared. If the hurdle is not cleared, then we have a count of 0.

- The first part of the model is typically a binary logistic regression model. This models whether an observation takes a positive count or not.
- The second part of the model is usually a truncated Poisson or Negative Binomial model. Truncated means we're only fitting positive counts, and not zeros.

Fitting a Hurdle Model / Poisson-Logistic

In fitting a hurdle model to our medicare training data, the interpretation would be that one process governs whether a patient visits a doctor or not, and another process governs how many visits are made.

summary(mod_5) results follow...

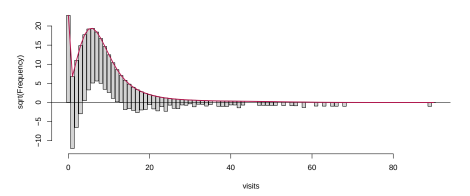
```
hurdle(formula = visits ~ hospital + health + chronic + sex + school +
  insurance, data = med train, dist = "poisson".
     zero.dist = "binomial")
 Pearson residuals:
     Min
             10 Median
                           30
                                  Max
  -5.3847 -1.1511 -0.4635 0.5644 24.8580
 Count model coefficients (truncated poisson with log link):
                 Estimate Std. Error z value Pr(>|z|)
                1.383357
                          0.028078 49.269
                                            <2e-16 ***
 (Intercept)
 hospital
                healthexcellent -0.306767 0.035988 -8.524
                                            <2e-16 ***
 healthpoor
             0.289780 0.020458 14.165 <2e-16 ***
                                            <2e-16 ***
 chronic
                0.095996 0.005386 17.822
 sexmale
                school.
                0.023794 0.002153 11.049
                                            <2e-16 ***
 insuranceves 0.041452
                          0.019658 2.109
                                            0.0350 *
 Zero hurdle model coefficients (binomial with logit link):
                Estimate Std. Error z value Pr(>|z|)
 (Intercept)
               -0.09874
                           0.16296 -0.606 0.544584
 hospital
           0.36432
                          0.11052 3.296 0.000979 ***
 healthexcellent -0.38506
                          0.16052 -2.399 0.016450 *
 healthpoor
                0.06522
                          0.18916 0.345 0.730252
 chronic
                0.53343
                          0.05195 \quad 10.268 \quad < 2e-16 \quad ***
 sexmale
                -0.38411
                          0.10111 -3.799 0.000145 ***
               0.06835
 school
                          0.01392 4.911 9.08e-07 ***
 insuranceves
               0.75178
                           0.11723
                                    6.413 1.43e-10 ***
 Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
 Number of iterations in BFGS optimization: 13
 Log-likelihood: -1.21e+04 on 16 Df
github.com/THOMASELOVE/2020-432
                                     432 Class 16 Slides
```

> summarv(mod 5)

Call:

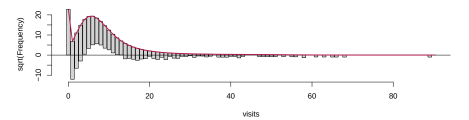
Rootogram for Poisson-Logistic Hurdle model

Poisson-Logistic Hurdle: mod_5

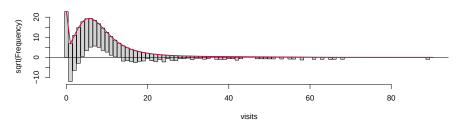


Poisson-Based Rootograms - Which Looks Best?

ZIP mod 3



Poisson-Logistic Hurdle mod_5



Store Training Sample mod_5 Predictions

No augment or other broom functions for hurdle models, so ...

Summarizing Training Sample mod_5 Fit

```
mod_5_aug has actual (visits) and mod_5 predicted (in .fitted) counts.
mod_5_summary <- tibble(
   model = "mod_5 (Poisson Hurdle)",
   R2 = R2(mod_5_aug$.fitted, mod_5_aug$visits),
   RMSE = RMSE(mod_5_aug$.fitted, mod_5_aug$visits),
   MAE = MAE(mod_5_aug$.fitted, mod_5_aug$visits))
mod 5 summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_5 (Poisson Hurdle)	0.113	6.481	4.093

Training Sample Results through mod_5

model	R2	RMSE	MAE
mod_1 (Poisson)	0.102	6.522	4.124
mod_2 (Neg. Binomial)	0.084	6.834	4.179
mod_3 (ZIP)	0.113	6.481	4.093
mod_4 (ZINB)	0.101	6.592	4.111
mod_5 (Poisson Hurdle)	0.113	6.481	4.093

What do you think?

Are ZIP and Poisson-Logistic Hurdle the Same?

```
temp_check <- tibble(</pre>
  subject = mod_3_aug$subject,
  visits = mod_3_aug$visits,
  pred_zip = mod_3_aug$.fitted,
  pred_hur = mod_5_aug$.fitted,
  diff = pred_hur - pred_zip)
mosaic::favstats(~ diff, data = temp_check)
         min
                         01
                                  median
                                                    03
 -0.02810685 -0.0005787973 0.0002147671 0.0008459066
```

-0.02810685 -0.0005787973 0.0002147671 0.00084590 max mean sd n missing 0.04125959 0.0003326582 0.003323227 3306 0

Vuong test: Comparing mod_3 and mod_5

```
vuong(mod_3, mod_5)
```

Vuong Non-Nested Hypothesis Test-Statistic: (test-statistic is asymptotically distributed N(0,1) under the null that the models are indistinguishable)

```
Vuong z-statistic H_A p-value
Raw 1.950967 model1 > model2 0.02553
AIC-corrected 1.950967 model1 > model2 0.02553
BIC-corrected 1.950967 model1 > model2 0.02553
```

There's some evidence mod_3 (ZIP) fits better than mod_5 (Hurdle) in our training sample.

mod_6: Negative Binomial-Logistic Hurdle Model

Fitting a Hurdle Model / NB-Logistic

summary(mod_6) results follow...

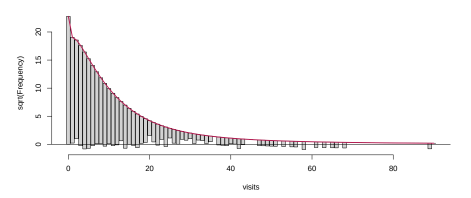
```
hurdle(formula = visits ~ hospital + health + chronic + sex + school +
 insurance, data = med_train, dist = "negbin".
    zero.dist = "binomial")
Pearson residuals:
   Min
            10 Median
                            30
                                  Max
-1.1856 -0.7139 -0.2712 0.3350 18.2426
Count model coefficients (truncated negbin with log link):
                Estimate Std. Error z value Pr(>|z|)
(Intercept)
                1.172949
                           0.068461 17.133 < 2e-16 ***
                                     8.847 < 2e-16 ***
hospital
                0.211206
                           0.023874
healthexcellent -0.337065
                           0.075872 -4.443 8.89e-06 ***
healthpoor
               0.345674
                           0.054985 6.287 3.24e-10 ***
chronic
                0.124764
                           0.014155 8.814 < 2e-16 ***
sexmale
               -0.045048
                           0.037047 -1.216
                                              0.224
school
               0.025236
                           0.005231 4.825 1.40e-06 ***
                           0.049025 1.261
                                              0.207
insuranceves
               0.061841
                0.358188
                           0.049009
                                     7.309 2.70e-13 ***
Log(theta)
Zero hurdle model coefficients (binomial with logit link):
               Estimate Std. Error z value Pr(>|z|)
(Intercept)
              -0.09874
                          0.16296 -0.606 0.544584
hospital
               0.36432
                           0.11052 3.296 0.000979 ***
healthexcellent -0.38506
                           0.16052 -2.399 0.016450 *
healthpoor
                           0.18916 0.345 0.730252
                0.06522
chronic
                0.53343
                           0.05195 10.268 < 2e-16 ***
sexmale
               -0.38411
                           0.10111 -3.799 0.000145 ***
                0.06835
                           0.01392 4.911 9.08e-07 ***
school.
                0.75178
                                    6.413 1.43e-10 ***
insuranceyes
                           0.11723
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Theta: count = 1.4307
Number of iterations in BFGS optimization: 15
Log-likelihood: -9058 on 17 Df
                                        432 Class 16 Slides
```

> summary(mod_6)

Call:

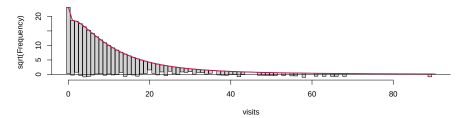
Rootogram for NB-Logistic Hurdle model

NB-Logistic Hurdle: mod_6

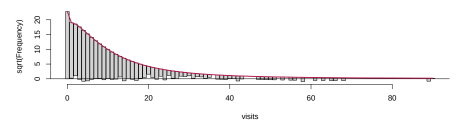


NB-Based Rootograms - Which Looks Best?

ZINB mod 4



NB-Logistic Hurdle mod_6



Store Training Sample mod_6 Predictions

Summarizing Training Sample mod_6 Fit

```
mod_6_aug has actual (visits) and mod_6 predicted (in .fitted) counts.
mod_6_summary <- tibble(
   model = "mod_6 (NB Hurdle)",
   R2 = R2(mod_6_aug$.fitted, mod_6_aug$visits),
   RMSE = RMSE(mod_6_aug$.fitted, mod_6_aug$visits),
   MAE = MAE(mod_6_aug$.fitted, mod_6_aug$visits))
mod 6 summary %>% kable(digits = 3)
```

model	R2	RMSE	MAE
mod_6 (NB Hurdle)	0.096	6.648	4.129

Training Sample Results through mod_6

model	R2	RMSE	MAE
mod_1 (Poisson)	0.102	6.522	4.124
mod_2 (Neg. Binomial)	0.084	6.834	4.179
mod_3 (ZIP)	0.113	6.481	4.093
mod_4 (ZINB)	0.101	6.592	4.111
mod_5 (Poisson Hurdle)	0.113	6.481	4.093
mod_6 (NB Hurdle)	0.096	6.648	4.129

Vuong test: Comparing mod_4 and mod_6

```
vuong(mod_4, mod_6)
```

Vuong Non-Nested Hypothesis Test-Statistic: (test-statistic is asymptotically distributed N(0,1) under the null that the models are indistinguishable)

	Vuong z-statistic		H_A	p-value
Raw	0.1589962	model1 >	model2	0.43684
AIC-corrected	0.1589962	model1 >	model2	0.43684
BIC-corrected	0.1589962	model1 >	model2	0.43684

There's some evidence mod_4 (ZINB) fits better than mod_6 (NB Hurdle) in our training sample.

Cross-Validation

Validation: Test Sample Predictions

Predict the visit counts for each subject in our test sample.

- Use mod_1 as a model for mod_2.
- Use mod_3 as a model for mod_4, mod_5 and mod_6.
- The other models are included with echo = FALSE.

Validation: Test Sample Fit Summaries

I'll show mod_1 and mod_2. The others are in the code with echo = FALSE.

```
mod 1 val <- tibble(</pre>
  model = "mod_1 (Poisson)",
  R2 = R2(test_1_aug$.fitted, test_1_aug$visits),
  RMSE = RMSE(test_1_aug\subseteq.fitted, test_1_aug\subseteq.in),
  MAE = MAE(test_1_aug\$.fitted, test_1_aug\$visits))
mod 2 val <- tibble(</pre>
  model = "mod 2 (Negative Binomial)",
  R2 = R2(test 2 aug$.fitted, test 2 aug$visits),
  RMSE = RMSE(test_2_aug$.fitted, test_2_aug$visits),
  MAE = MAE(test 2 aug$.fitted, test 2 aug$visits))
```

Results on the Next Slide

Validation Results in Test Sample: All Models

model	R2	RMSE	MAE
mod_1 (Poisson)	0.085	6.156	4.159
mod_2 (Negative Binomial)	0.075	6.332	4.221
mod_3 (ZIP)	0.089	6.143	4.143
mod_4 (ZINB)	0.083	6.218	4.171
mod_5 (Poisson Hurdle)	0.089	6.143	4.143
mod_6 (NB Hurdle)	0.081	6.241	4.179

Now which model would you choose based on test sample performance?

Next Time

Modeling Multi-Categorical Outcomes